Vitamin D Modulation of Mitochondrial Oxidative Metabolism and mTOR Enforces Stress Adaptations and Anticancer Responses

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ABSTRACT
The relationship between the active form of vitamin D₃ (1,25-dihydroxyvitamin D, 1,25(OH)₂D) and reactive oxygen species (ROS), two integral signaling molecules of the cell, is poorly understood. This is striking, given that both factors are involved in cancer cell regulation and metabolism. Mitochondria (mt) dysfunction is one of the main drivers of cancer, producing more mitochondria, higher cellular energy, and ROS that can enhance oxidative stress and stress tolerance responses. To study the effects of 1,25(OH)₂D on metabolic and mt dysfunction, we used the vitamin D receptor (VDR)-sensitive MG-63 osteosarcoma cell model. Using biochemical approaches, 1,25(OH)₂D decreased mt ROS levels, membrane potential (ΔΨₘ), biogenesis, and translation, while enforcing endoplasmic reticulum/mitohormetic stress adaptive responses. Using a mitochondria-focused transcriptomic approach, gene set enrichment and pathway analyses show that 1,25(OH)₂D lowered mt fusion/fission and oxidative phosphorylation (OXPHOS). By contrast, mitophagy, ROS defense, and epigenetic gene regulation were enhanced after 1,25(OH)₂D treatment, as well as key metabolic enzymes that regulate fluxes of substrates for cellular architecture and a shift toward non-oxidative energy metabolism. ATACseq revealed putative oxi-sensitive and tumor-suppressing transcription factors that may regulate important mt functional genes such as the mTORC1 inhibitor, DDIT4/REDD1. DDIT4/REDD1 was predominantly localized to the outer mt membrane in untreated MG-63 cells yet sequestered in the cytoplasm after 1,25(OH)₂D and rotenone treatments, suggesting a level of control by membrane depolarization to facilitate its cytoplasmic mTORC1 inhibitory function. The results show that 1,25(OH)₂D activates distinct adaptive metabolic responses involving mitochondria to regain redox balance and control the growth of osteosarcoma cells. © 2021 The Authors. JBMR Plus published by Wiley Periodicals LLC on behalf of American Society for Bone and Mineral Research.

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1. Introduction

The altered metabolism of cancer cells imparts a large impact on redox (reduction–oxidation) homeostasis, resulting in enhanced production of reactive oxygen species (ROS) that drives and sustains cancer cell proliferation and evasion. One counter measurement to elevated ROS is the increase of antioxidants, which allows cancer cells to persist in a damaging environment. However, as cancer cells persist in the continuing pro-oxidant environment, DNA is further damaged and the genome becomes more unstable with concomitant metabolic reprogramming. Over time, the metabolic reprogramming further distances cancer cells from redox homeostasis, leading to a vicious cycle of accumulating genetic lesions toward cancer progression. Understanding the mechanisms and how to control the dysfunctional and dysregulated metabolism of cancer cells is a major challenge in cancer biology.
Vitamin D consists of two major forms known as vitamin D2 (ergocalciferol) and vitamin D3 (cholecalciferol) that are most known for their roles in the maintenance of mineral homeostasis and growth of the body. In humans, vitamin D3 is synthesized in the skin upon UVB exposure, whereby vitamin D3 is obtained from plant sources in our diets. Both forms of vitamin D are biologically inert and require conversion and hydroxylation to 25(OH)D by vitamin D-25-hydroxylase in the liver, the major measurement of one’s vitamin D status. 25(OH)D is further hydroxylated in the kidneys or within specialized cell types by the 25(OH)-D1-24-25-24-hydroxylase (CYP27B1) to yield the biologically active metabolite of vitamin D, 1-alpha, 25-dihydroxyvitamin D (also called 1,25(OH)2D). Circulating 1,25(OH)2D effects are mediated by the vitamin D receptor (VDR), a member of the intracellular nuclear receptor superfamily. Perturbation of the vitamin D metabolic and signaling systems in humans and animals are associated with several diseases and disorders, including alopecia, diabetes, osteosarcopenia, diabetes, and cancer. A large body of studies have all implicated a suppressive role of vitamin D in cancer development and improved cancer patient and animal survival. And most recently, a large clinical trial involving vitamin D supplementation suggests that cholecalciferol can benefit patients with advanced or lethal cancers by decreasing mortality and prolonging survival in the study population. Despite these encouraging results from the clinical trial, the precise mechanism for anticancer effects of vitamin D and its metabolites remains elusive.

Understanding how what we eat (e.g., fruits and vegetables, which are rich in antioxidants like vitamin C), what we expose ourselves to (e.g., sunlight, pollution), and the metabolism that can lead to energy-reducing equivalents that drive cellular replication and differentiation at the genetic level provides insight toward the circle of life. In contrast, understanding how molecular factors such as ROS and their antioxidants can regulate the cell cycle, differentiation, and adaptive responses (e.g., DNA damage, mutagenesis) at the genetic level provides insight toward the circle of death. Numerous mechanisms exist to help dictate the consequences of cellular oxidative stress. For example, the cytoprotective upregulation of DNA repair transcripts allow for DNA mismatch repair, non-homologous end-joining, and base excision repair in cells to handle the high levels of oxidative damage. Other avenues may consist of oxidation of membrane receptors, signaling molecules, redox-sensitive transcription factors, and epigenetic transcriptional regulators, including histone deacetylase family members to regulate the cell’s response to stress and cancer development. Understanding the metabolic oxidation/reduction reactions and cellular responses to various biological and environmental factors remains a major milestone in cancer biology and therapy.

Thus, in the present study, we postulated that effects on the metabolic oxidation/reduction reactions that are characteristic of cancer cells may be crucial to the anticancer effects of the environmental/nutritional factor 1,25(OH)2D (also called calcitriol). We applied genomewide transcriptomic and epigenomic approaches to provide a comprehensive understanding of how 1,25(OH)2D modulates mitochondrial functions and counteracts tumorigenicity within the MG-63 osteosarcoma cell model, followed by mitochondrial biochemical and ultrastructural investigations. Based on these approaches, we provide evidence of novel regulators of mitochondrial stress, biogenesis, translation, organelar hormesis, and cancer metabolic fates mediated by 1,25(OH)2D, and cooperating factors leading to an overall reduction in oxidative stress levels. In particular, although the tumor suppressor DNA damage-inducible transcript 4 (DDIT4) is induced under stress conditions in normal bone cells to inhibit metabolism activated by the mammalian target of rapamycin (mTOR) in the cytoplasm, MG-63 osteosarcoma cells exhibit high levels of DDIT4 sequestered to the mitochondria as a potential mechanism to regulate mTOR activation and cancer progression. In contrast, 1,25(OH)2D treatment of MG-63 cells increased the expression and cytoplasmic localization of DDIT4 through separation from the outer mitochondrial membrane. Ironically, a meta-analysis of numerous cancer cell types identified DDIT4 as being overexpressed compared with non-cancerous cells and associated with poor survival outcomes despite being a potent mTOR inhibitor. Based on our findings from MG-63 cells, we propose that 1,25(OH)2D may suppress tumor progression of other cancer types that involves mitochondrial-to-cytoplasmic DDIT4/RED1 exchange. Overall, the results herein establish that 1,25(OH)2D can target the deregulation of specific metabolic hubs within osteosarcoma cells to suppress tumorigenicity, thus attempting to maintain the delicate balance between the cycle of life and death.
numbers >8.5. Library preparation and RNA-sequencing were performed at the Oncogenomics Core Facility at the Sylvester Comprehensive Cancer Center (University of Miami). Samples were sequenced using 75 bp paired ends with an Illumina (San Diego, CA, USA) NextSeq 500 generating reads of ~30 million per sample that were trimmed and filtered using Cutadapt. Compressed Fastq.gz files were uploaded to a Galaxy account (https://usegalaxy.org/), and data sets were concatenated tail-to-head (Galaxy Version 0.1.0). A MultiQC analysis was performed on all FastQC raw data files to generate a summarized QC report. HISAT2 was performed for sample alignment to the human genome (Ensembl 87: GRCh38. p7 human transcriptome) to generate BAM files with the most reads (80% to 90%) aligning successfully. Samtools stat was performed for further quality control of BAM files, and HTSeq-count was performed to generate non-normalized gene counts for each sample. Normalization of RNAseq gene counts (counts per million [CPM]), and differential gene expression and visualization analyses were performed with iDEP (http://bioinformatics.sdstate.edu/idep93). Differentially expressed genes were determined using DESeq2 with the false discovery rate (FDR) set to 0.05 and logFC (1) compared with controls. For individual gene expression plots derived from RNAseq, a two-way ANOVA was performed with Bonferroni’s multiple comparisons testing using the CPM values, where the p value summaries were depicted as ***p ≤ 0.0001, **p ≤ 0.001, *p ≤ 0.01, and p ≤ 0.05. Gprofiler was utilized to perform gene name conversion (ENTREZ TO ENTREZ) and basic functional annotation analyses (http://biit.cs.ut.ee/gprofiler/gost). To adjust for the FDR, we only considered terms with a Benjamini–Hochberg adjusted p value of 0.05. iDEP was utilized for the distribution of transformed data and the generation of scatter plots of sample correlations, hierarchical and k-means heatmap generation, and pathway analyses. GSEA and GAGE were performed using statistically significant differentially expressed genes to determine whether a priori defined set of genes were different between the two biological states. For GSEA and GAGE, the molecular Signatures Database v7.3 with the hallmark and canonical (KEGG) gene sets were applied.

2.4 ATAC-Seq and data analysis

Cells from two independent experiments were collected and open chromatin was assessed using an ATACseq kit (Active Motif, Carlsbad, CA, USA; S3150). DNAseq library preparation was completed at the Oncogenomics Core Facility at the Sylvester Comprehensive Cancer Center. Samples were sequenced using 100-bp paired ends with an Illumina NovaSeq 6000. Compressed Fastq.gz files were uploaded to a Galaxy account (https://usegalaxy.org/), concatenated, and subsequent sequencing reads (~40 million per sample) were trimmed off the Nextera adapter sequences and filtered using Cutadapt. Reads were mapped to the reference genome (hg38 Canonical) using Bowtie2 with presets “very sensitive end-to-end (very-sensitive),” “set the maximum fragment length for valid paired-end alignments: 1000,” and allowing mate dovetailing to generate BAM files. We filtered uninformative reads with low mapping quality and were not properly paired using Filter BAM data sets on a variety of attributes (Galaxy Version 2.4.1). Filter on read mapping quality (phred scale) was set to ≥30. ATACseq motif discovery was conducted using HOMER (http://www.homer.ucsd.edu).

2.5 Multi-omics analysis of genes that encode mitochondrial proteins

We examined the differential expression of genes that encode mitochondria-related proteins based on a compendium from MitoCarta (https://www.broadinstitute.org/mitocarta) and mitoXplorer (http://mitoxplorer.ibdm.univ-mrs.fr). We appraised mitochondrial protein-encoding genes using Venn analysis (http://www.interactivevenn.net) between the mitochondrial compendium and the differentially regulated genes derived from our RNAseq studies.

2.6 Quantitative real-time RT-PCR (qPCR) and analysis

RNA was prepared like the RNAseq studies. cDNA was synthesized using 200 ng total RNA with the ProtoScript First Strand cDNA Synthesis kit (New England Biolabs, Ipswich, MA, USA) utilizing random hexamers. All cDNAs were amplified under the following conditions: 95°C for 10 minutes to activate AmpliTaq Gold Polymerase, followed by 40 cycles of 95°C for 15 seconds and 60°C for 1 minute with an internal ROX reference dye. qPCR analysis was performed on a QuantStudio 3 Real-Time instrument (Thermo Fisher Scientific) utilizing the Power SYBR Green PCR Master mix (Thermo Fisher Scientific; Supplemental Table S1). Target genes were normalized to beta actin mRNA expression. For the primer design, the human genome sequence coverage assembly GRCh38.p13 was utilized from the Genome Reference Consortium. Data were presented as fold induction of treatments compared with 0 nM (vehicle) normalized to beta actin mRNA levels (i.e., the comparative CT Livak method). Melting curve analysis was performed for all primer sets to eliminate those that yielded primer-dimers. The p values reflect the log fold-change compared with the vehicle (0 nM) condition (n = 3 experimental samples ± SD). A two-way ANOVA test with Sidak’s multiple comparisons test was performed between vehicle and treatment data sets using Prism (GraphPad, La Jolla, CA, USA) where the p value summaries were depicted as ***p ≤ 0.0001, **p ≤ 0.001, *p ≤ 0.01, and p ≤ 0.05.

2.7 MitoSOX red mitochondrial superoxide indicator and live-cell Apoptome imaging

MitoSOX Red reagent (Thermo Fisher Scientific, M36008) was used to detect mitochondrial superoxide levels in live cells. MG-63 cells were cultured in Millicell EZ chamber slides (EMD Millipore). A 5 mM MitoSOX Red reagent stock solution was made by dilution into dimethyl sulfoxide (DMSO). A 5 μM MitoSOX Red reagent working solution was made by diluting the stock into a culture medium. Cells were loaded with MitoSOX Red reagent by incubating for 10 minutes at 37°C protected from light. Hoechst 33342 (1:2000) live-cell dye was used as a counterstain to detect the nuclei of live cells (Thermo Fisher Scientific, H1399). Cells were washed three times with a warm medium. Intensity measurements were obtained using the Zen Blue software (Zeiss) analyzed using Prism 8 (GraphPad). Rote-none (Sigma, St. Louis, MO, USA; R8875) was applied as a positive control. For each replicate (n = 6 replicates/condition), average ratios were derived from four different fields of views of 5 to 10 individual cells. Data are presented as mean ± SEM error bars; ***p ≤ 0.0001, **p ≤ 0.001, *p ≤ 0.01, and p ≤ 0.05 (one-way ANOVA with Tukey’s multiple comparisons test compared with vehicle).
2.8 Mitochondrial membrane potential (\(\Delta \Psi_M\)) measurements and live-cell Apoptome imaging

A JC-1 (5,5,6,6'-tetrachloro-1',3',3'-tetracyanobenzimidazolyl-carbocyanine iodide) mitochondrial membrane potential detection kit (Biotium, Fremont, CA, USA; 30001) was used to measure mitochondrial membrane potential changes in live cells. MG-63 cells were cultured in Millicell EZ chamber slides (EMD Millipore). All experiments were performed in a low-light setting. A 1× working solution of JC-1 dye was prepared in a cell culture medium, and cells were incubated in a 37°C cell culture incubator for 15 minutes. Cells were washed once with PBS and replenished with a fresh culture medium. Hoechst 33342 live-cell dye was used to detect the nuclei of live cells (Thermo Fisher Scientific, H1399). Cells were observed immediately using a Zeiss Observer 7 ApoTome2 microscope using a dual bandpass filter designed to simultaneously detect fluorescein and rhodamine, or fluorescein and Texas Red. The spectral properties of the JC-1 dye consist of Excitation/Emission (cytoplasm): 510/527 nm (green) and Excitation/Emission (polarized mitochondria): 585/590 nm (red). The ellipsoid spot measurement tool in Imaris was used to select perinuclear JC-1-labeled mitochondria to determine JC-1 aggregate:monomer intensity ratios. The spot intensity tool (Imaris) was used to determine the intensity (y axis) and position (x axis) between spot-to-spot (i.e., a series of mitochondria-to-mitochondria) in treated cells. For each replicate (n = 6–8 replicates/condition), average ratios were derived from four different fields of views of 6 to 8 individual cells. Data are presented as mean ± SEM error bars; ***\(p \leq 0.0001\), **\(p \leq 0.001\), *\(p \leq 0.01\), and \(p \leq 0.05\) (two-way ANOVA with Sidak’s multiple comparisons test compared to vehicle).

2.9 Mitochondrial biogenesis in-cell ELISA assay

Mitochondrial biogenesis and translation were monitored in MG-63 cells using an in-cell ELISA kit (Abcam, Cambridge, MA, USA; Ab110217). In brief, MG-63 cells were cultured in collagen-coated 96-well plates and treated with 1,25(OH)2D at various concentrations and durations in five replications. After the treatment series, cells were fixed with 4% PFA and then quenched for endogenous alkaline phosphatase activity using acetic acid. Primary antibody cocktails containing COX-1 and SDHA recognizing antibodies were added to the wells. Afterward, secondary HRP and AP-conjugated antibodies were applied and detected using a microplate reader at OD 405 nm (for AP detection of COX-1) and 600 nm (for HRP detection of COX-1). Measurements were normalized to the Janus Green staining intensity at OD 595 nm to account for differences in cell seeding.

2.10 Immunofluorescence labeling and analysis of MG-63 cells

MG-63 cells were cultured in Millicell EZ chamber slides (EMD Millipore) and fixed in either 80% methanol or 4% paraformaldehyde (PFA) in 0.1 M phosphate buffer (PBS, pH 7.4) for 10 minutes. PFA fixed cells were permeabilized with 0.2% Triton X-100 in PBS for 5 to 15 minutes at room temperature, followed by washes with PBS. Cells were blocked with normal horse/goat serum for non-specific background and then incubated with primary antibodies at a 1:200 dilution for 1 hour at room temperature. Primary antibodies used in this study included rabbit monoclonal to VDAC1 (Abcam, ab154856), and rabbit monoclonal to REDD1/DDIT4 (Abcam, ab191871). After washing steps in phosphate-buffered saline-Tween-20, the cells were incubated at room temperature for 20 minutes with corresponding species-specific secondary antibodies (Alexa series at 1:2000, Life Technologies). The slides were covered with Vectashield medium containing 4,6-diamidino-2-phenylindole (DAPI; Vector Laboratories, Burlingame, CA, USA; H-1200-10) for nuclei staining and then mounted with a glass coverslip. Negative controls were included that had either no primary or secondary antibodies in the blocking buffer. Immunofluorescence confocal-like microscopy was performed using a Zeiss Observer 7 ApoTome2 system. We carefully selected the wavelength ranges, emission filters, and dichroic mirrors to avoid signal bleed-through. Deconvolution and ApoTome processing (i.e., extended depth of view) of stacked images was performed using the ZenBlue software (Zeiss Microscopy). Image stacks were reconstructed and visualized as three-dimensional (3D) volumes with ZenBlue software (Bitplane, Zurich, Switzerland). The Imaris Spot detection algorithm was used as described by the manufacturer for semiautomatic identification and counting of fluorescently labeled mitochondria and cytoplasmic components. Means of expression intensity of mitochondrial and cytoplasmic regions within individual cells were compared between vehicle and 1,25(OH)2D-treated samples. For analysis, individual experiments (n = 4) were performed whereby each experiment entailed an assessment of 4 to 6 individual sets of cells for technical replication. For some experiments, Imaris (Bitplane) and MATLAB were used to generate 3D rendered models of protein expression and colocalization. Spots were located at the local maxima of the filtered image with background subtraction. Imaris calculated a “spot quality” (minimum of 100) based on intensity differences and shapes for spot rendering and was adjusted to include the signal of interest. Colocalization analysis was performed using the “spot” tool to designate the distance threshold and the mean distance between the “VDAC” and “DDIT4” colocalized spots. A two-way ANOVA test with Sidak’s multiple comparisons test was performed between vehicle and treatment data sets using Prism (GraphPad) where the \(p\) value summaries were depicted as ***\(p \leq 0.0001\), **\(p \leq 0.001\), *\(p \leq 0.01\), and \(p \leq 0.05\). Statistical significance was accepted at \(p \leq 0.05\).

2.11 Transmission electron microscopy (TEM)

TEM was performed at the Transmission Electron Microscopy Core Facility at the Miller School of Medicine, University of Miami. The TEM Core prepared the cells for electron microscopy and performed embedding and semi-thin (1 μm) and thin (100 nm) sectioning of the samples and final imaging with a JEOL JEM-1400 electron microscope. For analysis, 7 to 10 cells were investigated per condition, in which we averaged parameters between 20 to 40 mitochondria per cell.

2.12 Stimulation and measurement of ER stress

Known ER stress inducers tunicamycin (Sigma-Aldrich, T7765) and thapsigargin (Sigma-Aldrich, T9033) were diluted in ethanol and exposed to cells for 6 hours with appropriate vehicle controls. Both endpoint semi-quantitative and quantitative real-time PCR methods were used to assess ER stress based on Yoon Seung-Bin and colleagues25 with adjustments (e.g., the annealing temperature of 62°C was used instead). For the endpoint
PCR reaction, the Phusion DNA polymerase (Thermo Fisher Scientific) was used, and a 2.5% agarose gel was utilized to assess ER stress PCR products. U5/RXB1 primers relative to the 26 bp of XB1 removed by IRE1 were used for real-time PCR reactions (Supplemental Table S1). Housekeeping genes (Gapdh, 18sRNA) and the total amount of XB1 (Supplemental Table S1) were used to normalize gene expression.

3. Results

3.1 Genomewide assessment of 1,25(OH)2D-mediated transcription using RNAseq

Previous studies have shown that 1,25(OH)2D can suppress the growth of MG-63 cells but not of receptor-poor cell lines in standard 2D culture assays within the range of 100 nM (10−8 M or 40 ng/mL) and 10 nM (10−9 M or 4 ng/mL). However, 3D colony formation in soft agar is the gold-standard assay to study anchorage-independent carcinogenesis in vitro and has not been tested on 1,25(OH)2D treated MG-63 cells to date. After 14 days of treatment, both 10 nM and 100 nM of 1,25(OH)2D resulted in a significant reduction in overall colony size (Fig. 1A, B). However, 10 nM of 1,25(OH)2D did not significantly affect the number of colonies after treatment, in contrast to 100 nM (Fig. 1B). This suggests that 1,25(OH)2D may induce cell apoptosis at the higher 100 nM concentration and that 1,25(OH)2D can suppress MG-63 tumor growth at 10 nM via non-apoptotic mechanisms and lower toxicity (see later).

To define the genomewide impact on transcription by 1,25(OH)2D on MG-63 cells under the growth-inhibiting conditions at 10 nM, high throughput RNAseq analysis was carried out. After 24 hours of continuous 1,25(OH)2D treatment, there were 462 and 432 differentially up- and downregulated genes, respectively, compared with vehicle treatment using a false discovery rate (FDR) threshold of 0.05 and log fold change (1) with a high degree of correlation (Pearson correlation coefficient = 0.96–0.99; principal component analysis) (Fig. 1C, Supplemental Fig. S1, and Supplemental Worksheet S1). After 48 hours of 1,25(OH)2D treatment, there were 1015 and 1139 differentially up- and downregulated genes, respectively. To identify subgroups of genes that share expression patterns, we ranked genes by their standard deviation to obtain hierarchical clusters using DESeq2 (Fig. 1D and Supplemental Worksheet S2). An interactive heatmap of the 50 most variable genes shows that 1,25(OH)2D induced genes such as SOD2, IRS2, BIRC3, and DUSP1/5, which are either cytoplasmic or mitochondrial signaling molecules that mediate the effects of growth factors and/or cytokine interactions with known anticancer properties. For example, 1,25(OH)2D strongly induced the expression of the mitochondrial, but not cytosolic, manganese superoxide dismutase, SOD2, which converts the free radical O2− (superoxide) to H2O2 to defend against free radicals. The 50 most variable downregulated genes included the cytochrome P450 family 24 subfamily A member 1 (CYP24A1), the onco-channel TRPV6, and DKK2 (i.e., a Wnt mediator of tumor immune evasion). CYP24A1 functions as a mitochondrial monooxygenase that catalyzes the 24-hydroxylation and catabolism of 1,25(OH)2D, suggesting a negative feedback response to preserve 1,25(OH)2D signaling and its anticancer effects in MG63 cells.

Next, we studied the relationships among the up- and downregulated genes relative to their enriched gene ontology (GO) terms using hierarchical clustering trees (Supplemental Fig. S2A). The downregulated genes were overwhelmingly involved in nucleosome/chromatin assembly and organization, as well as DNA replication. Since nucleosomes assemble and become octameric during DNA replication amassing on daughter DNA strands, these findings suggest 1,25(OH)2D decreases replication, replication stress, and genomic instability associated with cancer. The decrease in nucleosome assembly also suggests that 1,25(OH)2D-treated MG-63 cells may be in interphase of the cell cycle, supporting our previous studies, where DNA is less compact and associated with increased chromatin accessibility (see later). Although chromatin remodeling, per se, was not a feature of the analysis, we did identify chromatin-modifying enzymes (e.g., SIRT1,4) that were differentially regulated after 1,25(OH)2D treatment. Nevertheless, the upregulated genes were associated with programmed cell death, translation, and response to organic substance. Of note, although regulators of apoptotic pathways were found to be enriched, we observed no changes in the early apoptosis marker Annexin V phosphatidylserine in 1,25(OH)2D-treated MG-63 cells at 10 nM (data not shown). We also used the dimension reduction algorithm, t-SNE, to map the top genes, and then identified four clusters of enriched pathways called k-means that were further mapped to GO biological processes (Supplemental Fig. S2B and Supplemental Worksheet S3). Cluster A consisted of genes upregulated after 48 hours of 1,25(OH)2D treatment that was enriched for the defense response to virus pathway. Cluster B consisted of genes upregulated after 1,25(OH)2D treatment for both 24 and 48 hours that were enriched for the stress response pathway. Cluster C consisted of genes downregulated after 48-hour 1,25(OH)2D treatment that enriched for the chromosome organization pathway. Lastly, Cluster D consisted of genes downregulated after both 24 and 48 hours that were enriched for chromatin/nucleosome assembly and cell development pathways. These findings show that 1,25(OH)2D regulates genome architecture and downstream stress response pathways as part of its anticancer response.

3.2 Functional enrichment analysis reveals 1,25(OH)2D-mediated cancer inhibition via mitochondrial OXPHOS and stress regulators

Functional annotation and gene set enrichment analysis (GSEA) were performed using several methods to reflect the heterogeneity of data repositories and statistical approaches. We first used the g:GOst program to map genes to known functional information to determine statistically significant enriched relationships. The data were stratified based on GO molecular functions (MF), biological processes (BP), and cellular components (Supplemental Worksheets S4 and S5). Based on GO-MF subset analysis, genes that regulate fatty acid desaturases were upregulated after 1,25(OH)2D treatment, suggesting a putative role in unsaturated fatty acid biosynthesis and utilization (Fig. 1E). Based on GO-BP, 1,25(OH)2D treatment induced genes that regulate unfolded proteins, programmed cell death, and the detoxification of metal ions. On the other hand, 1,25(OH)2D suppressed growth factors and structural molecule activity-related genes based on GO-MF. Based on GO-BP, 1,25(OH)2D suppressed chromatin assembly, morphogenesis, and oxidative phosphorylation (OXPHOS)-related genes. The OXPHOS genes include COX11, which is a copper-binding subunit of the cytochrome c oxidase enzyme in the electron transfer chain in the mitochondria. Several respiratory
After 48 hours of 1,25(OH)2D treatment, the enriched pathways at https://biit.cs.ut.ee/gpro that maintain the loose mesenchymal phenotype (Fig. 2A). For example, the data suggest that 1,25(OH)2D can reverse EMT to suppress mesenchymal metastasis through a transcription factor (Fig. 1). Significant decreases in O2 levels were observed in 1,25(OH)2D-treated samples revealed gene sets related to inflammation, hypoxia, and epithelial-mesenchymal transition (EMT) pathways that were not discovered using g:GOSt (Fig. 2A). For example, the data suggest that 1,25(OH)2D can reverse EMT to suppress mesenchymal metastasis through downregulation of SNAI2, a key zinc finger transcription factor that maintains the loose mesenchymal phenotype (Fig. 2A, B). After 48 hours of 1,25(OH)2D treatment, the enriched pathways were related to hypoxia, glycolysis, inflammation, unfolded protein response, mTOR pathway, cholesterol homeostasis, apoptosis, xenobiotic metabolism, and p53 signaling (Fig. 2B). Key upregulated genes include DDIT4/REDD1 and sequestosome 1 (SQSTM1), which target the direct inhibition of mTOR or indirect effects through autophagy, respectively. In terms of hypoxia, decreased OXPHOS after 1,25(OH)2D treatment is likely to increase molecular oxygen levels as hypoxia in cancer cells is partly due to increasing O2 consumption and reduction to water that can thereby induce EMT. Hyperoxia is also supported by the increased SOD2 levels after 1,25(OH)2D treatment, as SOD2 metabolizes superoxide radicals into hydrogen peroxide. These findings suggest that 1,25(OH)2D affects major pathways involved in oxygen levels and the growth regulation of tumor cells.

In addition, we applied generally applicable gene-set enrichment (GAGE) analysis that has no limitations on sample size based on a parametric gene randomization method to test the significance of gene sets using log-based fold changes as the per gene statistic. By using the absolute values of fold change in the GAGE analysis combined with the Kyoto Encyclopedia of Genes and Genomes (KEGG) database, 1,25(OH)2D was shown to significantly downregulate a more dynamic OXPHOS gene set at both 24 and 48 hours of treatment (Fig. 2C and Supplemental Worksheet S7). The GAGE output was shared with Pathview to rationalize the OXPHOS genes (Fig. 2D), whereby the analysis shows that 1,25(OH)2D downregulates the second (II) and third (III) large enzyme complexes (i.e., succinate dehydrogenase and cytochrome bc1 complex, respectively) in the respiratory electron transport chain of MG-63 cells. The cytochrome bc1 complex is responsible for the proton gradient as well as for the formation of O2. Disruption of the flow of electrons across the membrane is predicted to alter the transmembrane difference of proton electrochemical potential, which ATP synthases use to generate energy (see later).
Furthermore, 1,25(OH)₂D downregulated ATP synthase components (e.g., ATP5D, ATP5A1, ATP5C1) as another mode to regulate overall mitochondrial activity. Collectively, these findings suggest that 1,25(OH)₂D promotes additional metabolic shifts that involve the suppression of mitochondrial OXPHOS as part of its anticancer strategy.
3.3 1,25(OH)₂D-mediated organellar hormesis enforces stress tolerance and growth inhibition of MG-63 cells

Our genomewide bioinformatics analysis suggests the involvement of the unfolded protein response (UPR) in 1,25(OH)₂D-treated MG-63 cells, which is known to mediate stress tolerance and organismal longevity involving a process called hormesis. Hormesis describes a phenomenon where mild cellular stress caused by unfolded proteins stimulates alternative signaling pathways with beneficial, overcompensating outcomes and organellar connectivity. To better understand how 1,25(OH)₂D modulates hormetic responses in cancer cells, we investigated various ER and mitochondrial hormetic signaling pathways that involve antioxidants and protein-folding chaperones (Fig. 3). The ER transmembrane receptor protein kinases (ER-TRK) IRE1 and PERK and the transcription factor ATF6 govern the expression of factors that protect cells as part of the hormetic UPR by promoting cell cycle arrest, protein translation inhibition, and chaperone production. A proxy for activated IRE1 is cleavage of 26 base pairs from its substrate, XBP1, to generate a spliced form called sXBP1 that functions as a transcription factor for expression of binding immunoglobulin protein (BIP, also called GRP78 or HSPA5), which

![Fig 3. 1,25(OH)₂D and ER/mitochondrial unfolded protein stress regulation. (A) Representative endpoint PCR analysis of IRE1-XBP1 expression after 6 hours of positive control treatments. u (unspliced XBP1; 256 bp), sp (spliced XBP1; 230 bp), 18sRNA (190 bp). (B) Real-time PCR analysis of IRE1-XBP1 expression after 6 hours of positive control treatments. The graph depicts fold change of either uXBP1 (unspliced) or sXBP1 (spliced) normalized to the total XBP1 levels. Data are presented as mean ± SEM error bars (n = 3 samples/condition); ***p < 0.001 (one-way ANOVA with Tukey’s multiple comparisons test compared with respective vehicle). (C) Real-time PCR analysis of BIP/HSPA5 expression in positive controls. Data are presented as mean ± SEM error bars (n = 3 samples/condition); ***p < 0.001 (one-way ANOVA with Tukey’s multiple comparisons test compared with respective vehicle). (D) Representative endpoint PCR analysis of IRE1-XBP1 expression after 24 to 48 hours of 1,25(OH)₂D treatments. u (unspliced XBP1; 256 bp), sp (spliced XBP1; 230 bp), GAPDH (350 bp). (E) Real-time PCR analysis of IRE1-XBP1 expression after 24 to 48 hours of 1,25(OH)₂D treatments. The graph depicts fold change of either uXBP1 (unspliced) or sXBP1 (spliced) normalized to the total XBP1 levels. Data are presented as mean ± SEM error bars (n = 3 samples/condition); ***p < 0.001, **p < 0.01 (one-way ANOVA with Tukey’s multiple comparisons test compared with respective vehicle). (F) Real-time PCR analysis of BIP/HSPA5 and ATF5 expression after 24 to 48 hours of 1,25(OH)₂D treatments. Data are presented as mean ± SEM error bars (n = 3 samples/condition); **p < 0.01 (one-way ANOVA with Tukey’s multiple comparisons test compared with respective vehicle). (G–J) RNAseq analysis of ER/mitochondrial stress and hormetic regulators. A two-way ANOVA was performed with Bonferroni’s multiple comparisons test using the counts per million (CPM) values (n = 2 samples/condition), where the p value summaries were depicted as ****p < 0.0001, ***p < 0.001, and **p < 0.01. ns = not significant; UPR = unfolded protein response. (K) Proposed model: 1,25(OH)₂D enforces stress tolerance in cancer cells via metabolic reprogramming involving ER/mitohormesis.
functions as a major ER stress chaperone. To characterize UPR in the MG-63 cell system, thapsigargin and tunicamycin (i.e., blockers of the ER ATPase/SERCA pump and glycoprotein synthesis, respectively) were first used and found to induce a dose-dependent increase in xBP1 and BIP/HSPA5 (Fig. 3A–C). Interestingly, 1,25(OH)₂D treatment enhanced xBP1 in a time-dependent manner at 10 nM but not at 100 nM (Fig. 3D, E) with no change in BIP mRNA levels across all concentrations, suggesting a hormetic response to insoluble proteins (Fig. 3G, H). As the proxies for ATF6 activation are upregulation of BIP and uXBP1, our findings also suggest that ATF6 plays a minimal role in the 1,25(OH)₂D response (Fig. 3E). Two proxies for PERK activation are ATF4 and CHOP (also called DDIT3 or GADD153), whereby RNAseq analysis showed no changes in both transcripts after 1,25(OH)₂D treatment (Fig. 3F and Suplemental Worksheet S1). Therefore, we investigated potential hormetic antioxidative responses of the alternative ER-TRK, recently described in C. elegans, (31) in the context of 1,25(OH)₂D by appraising the human glutathione S-transferase family of genes. We only observed statistically significant increases in glutathione S-transferase kappa 1 (GSTK1) and glutathione S-transferase Mu 4 (GSTM4) after 1,25(OH)₂D treatment of MG-63 cells (Fig. 3I), whereby lower levels of GSTK1 have been linked to the elevation of mt ROS underlying hypertrophic cardiomyopathy. (32) Lastly, since the bioinformatics analysis also suggests the downregulation of OXPHOS, we assessed mitochondrial UPR by way of activating transcription factor 5 (ATF5) (Fig. 3J). ATF5 is a major mitochondrial stress regulator that can induce proteostasis and chaperonin production, (33) whereby 10 nM of 1,25(OH)₂D treatment significantly downregulated ATF5 in MG-63 cells, the effect of which dissipated at higher concentrations, signifying a hormetic response (Fig. 3J). Overall, the results suggest that 1,25(OH)₂D activates distinct hormetic adaptive responses in the ER and mitochondria to regain control of the growth of cancer cells, which may underly beneficial interorganellar communication to overcome cancer stress (Fig. 3K).

3.4 A multi-omics approach to study mitochondrial anticancer responses to 1,25(OH)₂D

Given that 1,25(OH)₂D suppresses mitochondrial UPR, we performed a more granular multi-omics assessment of mitochondrial transcriptional changes using the annotated databases MitoCarta and mitoXplorer. MitoCarta currently annotates 1136 genes encoding mitochondrial proteins, while mitoXplorer contains 1229 genes. First, we used MitoCarta (version 3.0) to identify differentially regulated mitochondria-related genes from our RNAseq data set. (34) Among the 1477 upregulated 1,25(OH)₂D-mediated differently expressed genes (DEGs) (Fig. 1), we identified 79 genes that encode mitochondria proteins within the combined 24- and 48-hour gene sets (~5%; Fig. 4A and Supplemental Worksheet S8). Among the 1571 downregulated 1,25(OH)₂D-mediated DEGs (Fig. 1), we identified 45 genes encoding mitochondrial proteins in total (~2.8%; Fig. 4A and Supplemental Worksheet S8). However, MitoCarta provides no annotation on the genes, and to understand the biological significance behind these changes, we utilized the annotated mitoXplorer (version 1.0) necessary for pathway analysis. In all, there were 64 and 37 1,25(OH)₂D-mediated up- and downregulated mitochondrial genes, respectively, that were common between the two repositories (Fig. 4B). There were only 15 and 8 up- and downregulated 1,25(OH)₂D-mediated mitochondrial genes, respectively, that were specific to the MitoCarta repository and not included in the mitoXplorer annotative analysis.

Based on the mitoXplorer analysis, the 1,25(OH)₂D-mediated downregulated DEGs after 24 hours included MRPS18B, which encodes a 28S subunit mitoribosomal protein involved in protein translation (Fig. 4C and Supplemental Worksheet S8). In addition, HSPA1A and B, members of the heat shock protein family, were also downregulated by 1,25(OH)₂D, suggesting a lowering of stress aggregation and increased protein stability in mitochondria. In terms of metabolism, dimethylglycine dehydrogenase (DMGDH), a mitochondrial enzyme involved in phosphatidylcholine and lipid metabolism and glycine modifications, was elevated after 1,25(OH)₂D treatment (Fig. 1E). Recently, studies have shown that DMGDH can play a role in antioxidant defense, (35) and low levels can be a diagnostic and prognostic marker for hepatocellular carcinoma metastasis by acting on the Akt pathway. (36) Genes that regulated beta oxidation of fatty acids were also found to be suppressed by 1,25(OH)₂D treatment, suggesting another mean for ROS reduction. (37) Interestingly, mitochondrial amino acid metabolism and detoxification were upregulated after 1,25(OH)₂D treatment by way of glutamate-ammonia ligase (GLUL), which is a mitochondrial enzyme that catalyzes the synthesis of glutamine from the more toxic glutamate and ammonia. In addition, nitrilase omega-amidase (NIT2) was upregulated by 1,25(OH)₂D, which is known to play a role in arresting cells to remove toxic intermediates such as 2-oxoglutaramate. (38) Pyruvate metabolism was also affected after 1,25(OH)₂D treatment by way of upregulation of the mitochondrial pyruvate dehydrogenase kinase 4 (PDK4). PDK4 inhibits the mitochondrial pyruvate dehydrogenase complex to reduce pyruvate conversion from glucose, suggesting that 1,25(OH)₂D may conserve glucose metabolism (i.e., slowing glycolysis), as during hibernation, by decreasing its conversion to acetyl-CoA.

In the 48-hour analysis, the overwhelming effect of 1,25(OH)₂D on mitochondrial protein translation at 24 hours was aborted, suggesting adaptive responses (Fig. 4D). Additional selective pressures toward translation occurred via upregulation of MTERF2, a transcription termination factor that modulates cell growth and the cell cycle. (39) Longer treatments of 1,25(OH)₂D did enhance the ROS defense response (iCAT); however, this was countered by decreased MPV17, which is involved in ROS neutralization and mitochondrial protection. (40) Antioxidant responses closely regulate mitochondrial epigenetic signaling factors such as SIRT4, (41) an enzyme with deacetylase and ADP-ribosylation activities, which was downregulated after 1,25(OH)₂D treatment, suggesting a mode for further fine-tuning of epigenomic regulation. Other mitochondrial metabolic and dynamic effects of 1,25(OH)₂D include the suppression of the heme biosynthesis pathway by way of UROS, which is part of the catalytic steps of porphyrin biosynthesis and associated with cancer when heme production is left unchecked. (42) Furthermore, mitofusion 1 (MFN1) was downregulated after 1,25(OH)₂D treatment that mediates mitochondrial fusion, suggesting reduced mitochondrial networks, ATP production, and OXPHOS. SQSTM1, a protein involved in mitophagy, was upregulated after 1,25(OH)₂D treatment, suggesting a selective and adaptive process to remove dysfunctional mitochondria from cancer cells. The TCA cycle, which provides electrons via the reducing agent NADH for OXPHOS, was enhanced after 48 hours of 1,25(OH)₂D treatment despite the suppression of OXPHOS, raising the possibility of non-redox roles. (43) For example, 1,25(OH)₂D may involve substrate-level phosphorylation as a metabolic reaction to generate energy instead of OXPHOS. SUCGL2, a
GTP-specific beta subunit of succinyl-CoA synthase that forms succinyl-CoA, succinate, and ATP through the coupling of this reaction independent of OXPHOS, was elevated after 1,25(OH)₂D treatment. Also, OGDH, a dehydrogenase that catalyzes the conversion of 2-oxoglutarate to succinyl-CoA and carbon dioxide, was increased after 1,25(OH)₂D treatment, which may further drive energy production via TCA non-redox intermediates.

Lastly, several known mitochondrial genes were not co-curated in the MitoCarta and mitoXplorer repositories, including DDIT4/REDD1 (44) (see later) that were validated by qPCR derived from our RNAseq data sets (Fig. 4). We also observed a consistent downregulation of known mitochondrial chaperonin PPID (cyclophilin D). Although there was an upward trend for the mitophagy marker, P62 (Fig. 4F), qPCR reanalysis showed a statistically significant increase in transcript levels (Fig. 4H), suggesting a possible role in conjunction with SQSTM1 toward mitophagy. In addition, mitochondrial BCL2/adenovirus E1B 19 kDa protein-interacting protein 3 (BNIP3) transcripts were increased after 1,25(OH)₂D treatment and may interact with LC3 to remove damaged ER and mitochondria to recycle cellular content to promote the health of cells (Fig. 4F). Again, in terms of mitochondrial dynamics, 1,25(OH)₂D treatment resulted in the downregulation of mitochondrial fission transcript, FIS1, and of OPA3, a dynamin-related GTPase that regulates the equilibrium between mitochondrial fusion and mitochondrial fission (Fig. 4G). Endothelial PAS domain protein 1 (EPAS1) mRNA, which encodes a protein involved in mitochondrial biogenesis, was decreased after 1,25(OH)₂D treatment (Fig. 4G). Overall, the multi-omics approach revealed novel factors and pathways as part of 1,25(OH)₂D’s mitochondrial-mediated anticancer response.

3.5 1,25(OH)₂D-mediated epigenetic regulation of mitochondrial-related genes in MG-63 osteosarcoma cells

Next, to identify functional chromosomal regions that may govern anticancer responses and may be coregulated by 1,25(OH)₂D and oxidative stress, we used assay for transposase-accessible chromatin using sequencing (ATACseq) and assessment of transcription factor (TF) binding motifs. This method appraises genomewide chromatin accessibility using hyperactive Tn5 transposase that inserts sequencing adapters into open chromatin regions (Fig. 5A). The data show that most peaks were...
located within intronic, intergenic, and promoter regions across samples with 96% to 97% of reads with ≥Q30 scores, satisfying the quality-control requirements (Supplemental Worksheet S9). Globally, there were 97,739 overlapping peaks, 14,210 1,25(OH)2D unique peaks, and 7535 vehicle-unique peaks after 1,25(OH)2D treatment for 24 hours. (C) Heatmap depicting the pattern of peaks outlined in the Venn diagram. There is one column per group used in the comparison. The color bars on the left correspond to Venn diagram grouping of peaks. The heatmap displays the read coverage density (redder means more reads at that location), with each row corresponding to the average peak profile for a single peak that is averaged within each group, across all groups in the comparison. (D) Heatmap for transcriptional start sites (TSS) after 1,25(OH)2D [10 nM] and vehicle treatments. Chromatin accessibility at TSS significantly increased in response to 1,25(OH)2D. The lower half of the 1,25(OH)2D plot harbors the 1,25(OH)2D-unique peaks from (B) and (C). The plots specify a window of ± 3000 bp around the TSS of genes. Read density scores presented as the right index. (E) Genome browser track of ATACseq results for select mitochondrial downregulated genes. Differentially accessible regions identified for select genes. (F) Genome browser track of ATACseq results for select mitochondrial upregulated genes. Of note, a 1,25(OH)2D-unique peak was identified in intron 9–10 of SUCLG2. (G) List of transcription factor (TF) motifs enriched in accessible chromatin upon 1,25(OH)2D treatment.

For upregulated genes (Fig. 5F), we observed enhanced chromatin accessibility at both the proximal and promoter regions of DDIT4. On the contrary, there appeared to be nominal epigenetic regulation of SOD2 by 1,25(OH)2D. This finding suggests either posttranslational and/or posttranscriptional modes of SOD2 mRNA regulation after 1,25(OH)2D treatment. Interestingly, one of the most significantly affected chromosomal regions identified by ATACseq was in intron 9–10 of the SUCLG2 gene, suggesting a regulatory epigenetic mechanism induced by 1,25(OH)2D to control succinyl-CoA synthase expression.

We next used hypergeometric optimization of motif enrichment (HOMER) for transcription factor (TF) motif discovery within 1,25(OH)2D-sensitive open chromatin (Supplemental Worksheet S10). From this analysis, GATA3 was the most highly associated TF identified (Fig. 5G), whereby GATA3 is essential for normal tissue development and is commonly mutated in breast cancers. Not surprisingly, the VDR motif was the second most highly correlated nuclear TF identified from our analysis. Interestingly, we also identified the nuclear respiratory factor 1 (NRF1), a redox-sensitive member of the Cap-N-Collar family of TFs that binds to antioxidant response elements (AREs), as a potential regulator of 1,25(OH)2D-mediated epigenetic responses, suggesting that AREs may cooperate with 1,25(OH)2D response elements (VDREs) via NRF1-VDR binding. 1,25(OH)2D treatment also may promote estrogen receptor binding, suggesting a synergistic effect to help promote the normal bone-forming osteoblast phenotype in osteosarcoma cells. Overall, 1,25(OH)2D
promotes chromatin accessibility in MG-63 cancer cells to enhance the regulatory effects of specific TFs that may play important roles in oxidative stress defense and normal tissue and cellular developmental processes.

3.6 1,25(OH)₂D-mediated decrease in ΔΨₘ inhibits mitochondrial ROS production

The transcriptomic and epigenomic data thus far suggest that 1,25(OH)₂D regulates mitochondrial functions in MG-63 cells to
promote its anticancer effects. Therefore, we investigated the mitochondrial membrane potential ($\Delta \Psi_M$) using the ratiometric JC-1 dye, where the accumulation of cationic J-aggregates (red) in mitochondrial membranes acts as a proxy for polarized mitochondria. On the other hand, cells that have diminished $\Delta \Psi_M$ will contain JC-1 in its monomeric form (green) in either the mitochondria or cytoplasm during transition states.

To validate the JC-1 dye, we treated MG-63 cells with hydrogen peroxide (H$_2$O$_2$), a known oxidant and mitochondrial membrane depolarizer. Within 20 sections of H$_2$O$_2$ treatment, we observed a decrease in the J-aggregate-to-monomer ratio signifying a decrease in $\Delta \Psi_M$ (Fig. 6A, B). In the 1,25(OH)$_2$D studies, we pretreated MG-63 cells for 24 hours and then measured the JC-1 intensity ratios (Fig. 6C, D). Interestingly, while most of the vehicle-treated MG-63 cells were positive for J-aggregates, only ~25% of 1,25(OH)$_2$D-treated cells contained J-aggregates in their mitochondria, suggesting a complete collapse of the $\Delta \Psi_M$ within most cells (Fig. 6D). Among those 1,25(OH)$_2$D-treated cells that exhibited J-aggregates, their JC-1 intensity ratio was significantly reduced compared with vehicle treatment (Fig. 6E). Using the Imaris software (Bitplane) spot intensity tool, the vehicle-treated cells exhibited an overlap in J-aggregates.
aggregate-to-monomer signals across a series of mitochondria (i.e., spot-to-spot) within cells (Fig. 6F). On the contrary, 1,25(OH)2D-treated cells exhibited an increased level of non-overlapping monomer-to-J-aggregate signals, suggesting the depolarization of the mitochondrial membrane and extramitochondrial presence of the monomers.

A key factor in determining the fate of cells with depolarized mitochondria is the level of ROS. To determine the impact of 1,25(OH)2D on ROS production within MG-63 cells, we measured mitochondria-specific ROS using the MitoSOX Red, a mitochondrial O2•− indicator for live cells (Fig. 6G). 1,25(OH)2D treatment for 24 hours significantly reduced the production of O2•− within MG-63 cells compared with vehicle-treated samples (Fig. 6H). Conversely, treatment with the inhibitor of complex I of the respiratory chain, rotenone, significantly increased mt ROS levels. Thus, MG-63 osteosarcomas are accompanied by mechanisms that prevent mitochondrial depolarization, resulting in chronic intracellular mt ROS. Overall, the data suggest that 1,25(OH)2D treatment is associated with the opening of the mitochondrial permeability pores, loss of the electrochemical proton gradient, and reduced ROS as part of its anticancer effects.
3.7 1,25(OH)_2D modulates mitochondrial structure and dynamics in MG-63 cancer cells

We next appraised mitochondria structure and morphology using immunofluorescence (IF) and electron microscopy (EM). Using antibodies against the outer mitochondrial membrane voltage-dependent anion-selective channel 1 (VDAC-1), we observed the classic elongated tubular shape of mitochondria in vehicle-treated MG-63 cells (Fig. 7A). However, 1,25(OH)_2D treatment promoted changes from a tubular to globular “ring” morphology with weak fluorescence in the center of mitochondria (Fig. 7B). This characteristic suggests mitochondrial shortening, swelling, and structural changes triggered by decreased ΔΨm and increased permeability of the inner mitochondrial membrane. 3D-rendered images of 1,25(OH)_2D-treated cells revealed rough, fragmented surfaces (i.e., structural changes) of individual mitochondria treated for 24 hours (Fig. 7B, lower panel). The reversible changes of mitochondria from the tubular to condensed or fragmented conformations is the classic response of loss of ATP synthesis by OXPHOS. In EM studies, vehicle-treated cells exhibited tubular mitochondria; however, individual cristae were hardly discernible, suggestive of deranged mitochondrial respiration in vehicle-treated MG-63 cells (Fig. 7C, red arrows). Furthermore, some mitochondria were in various stages of membrane fusion/fission as marked by “tethered” structures indicative of dynamic remodeling (Fig. 7C, blue arrow). 1,25(OH)_2D treatment increased the size of mitochondria and generated mitochondria with discernible cristae (Fig. 7D, E), which may reflect partial prevention of cristolysis. The mitochondria also contained electron-lucent cavities, not vacuoles, consistent with the ring-shaped structures observed in the IF studies (Fig. 7B, white arrow) and may represent interorganellar connections commonly observed in normal cells. 1,25(OH)_2D regulation of mitochondrial biogenesis mediates DDIT4/REDD1 availability and mTOR function in the cytoplasm

Lastly, given the results of our functional annotation analysis and recent findings that certain cells express DDIT4/REDD1 in the mitochondria, we focused the remainder of our attention on the role that 1,25(OH)_2D and DDIT4 play in cancer prevention. DDIT4 is a known tumor suppressor gene predominantly expressed in the mitochondria under certain stress conditions to function as a potent mTOR inhibitor. However, recent findings show that DDIT4 is highly expressed in malignant cancers, leading to poor cancer-related prognosis in a paradoxical manner, suggesting that for certain genes the expression profiles cannot be functionally generalized (Supplemental Fig. S3). To help rationalize this paradoxical observation, we investigated DDIT4 cellular flux in MG-63 cells before and after 1,25(OH)_2D treatment. First, 1,25(OH)_2D at 10 nM increased DDIT4 mRNA levels in a time- and VDR-dependent manner (Fig. 8A). Next, we performed Apoptome (Zeiss) structured-illumination imaging of DDIT4 and VDAC1 within vehicle-treated MG-63 cells and found that DDIT4 was exclusively associated with VDAC1-positive mitochondria (Fig. 8B, yellow arrow). Imaris 3D-rendering confirmed the proximity of both VDAC1 and DDIT4 within individual mitochondria (Fig. 8C), whereby VDAC1-free DDIT4 expression in the cytoplasm was uncommon (white arrow) (Fig. 8C). A more dynamic colocalization pattern was observed after 1,25(OH)_2D treatment (Fig. 8D, E). DDIT4 was predominantly expressed in the cytoplasm after 1,25(OH)_2D treatment (Fig. 8E). Interestingly, among the VDAC1-DDIT4 colocalized mitochondria in 1,25(OH)_2D-treated cells, the average spot distance (i.e., the shortest distance between VDAC1 and DDIT4) was significantly increased compared with controls, suggesting the translocation or leaking of DDIT4 into the cytoplasm (Fig. 8F, bottom panel). Furthermore, after rotenone treatment, MG-63 cells contained globular VDAC1-positive mitochondria as previously noted, as well as a disassociation of DDIT4 from the mitochondria, suggesting a potential role of mitochondrial depolarization (Fig. 8H). Nonetheless, the level of colocalized VDAC1-DDIT4 protein in 1,25(OH)_2D-treated cells was significantly decreased compared with vehicle treatment, reflecting the excess of DDIT4 in the cytoplasm (Fig. 8G). In contrast, vehicle-treated MG-63 cells contained a statistically significant higher percentage of VDAC1-DDIT4 colocalized mitochondria with lower levels of cytoplasmic DDIT4 (Fig. 8G). Given the apparent decrease in the number of mitochondria after 1,25(OH)_2D treatment, the increase in cytoplasmic DDIT4 protein may occur, in part, due to reduced mitochondrial biogenesis (i.e., mass/content/self-replication). To address this possibility, we used a duplexing in-cell ELISA assay that quantifies both mitochondrial (mt) DNA- and nuclear (n) DNA-encoded proteins, COX-1 and SDHA, respectively, in MG-63 cells. We observed 1,25(OH)_2D-dependent inhibition of mtDNA-encoded COX-1 protein relative to nDNA-encoded SDHA protein by ~20% after 24 hours (Fig. 8I). These data suggest that enhanced mitochondrial localization of DDIT4 may help confer the cancer state and that the enhanced cytoplasmic localization and expression of DDIT4 may be a mechanism by which 1,25(OH)_2D suppresses osteosarcomas.

4. Discussion

4.1 Relationship between 1,25(OH)_2D and the metabolic oxidation/reduction reactions of cancerous and non-cancerous cells

Findings so far in non-cancerous cells suggest that proper 25(OH)D levels maintain and minimize systemic cellular oxidative stress after the day-to-day exposure to damaging agents such as UV sunlight. Furthermore, loss of VDR functional studies in human skin keratinocytes show increased mitochondrial membrane potential due to increased transcription of the respiratory chain subunits II and IV of cytochrome c oxidase. In addition, the potential for vitamin D3 to reduce oxidative damage to DNA has been linked to a clinical trial where vitamin D3 supplementation reduced 8-hydroxy-2'-deoxyguanosine, a marker of oxidative damage, in colorectal epithelial crypt cells. In other studies, 1,25(OH)_2D was shown to modulate the expression of select antioxidative genes via nuclear factor erythroid 2-related factor 2 (NRF2), which is a key transcription factor that can bind to AREs to protect cells against oxidative stress associated with diabetic neuropathy. These findings suggest that vitamin D metabolites can regulate the respiratory chain and to modulate ancillary metabolic pathways depending on the cellular context and requirements within stressed non-cancerous cells.

Our findings in cancer cells show that 1,25(OH)_2D can influence mitochondrial metabolism, structure, and function to dictate its anticancer effects, which may also intimately involve extra-mitochondrial organelles such as the ER (Figs. 3 and 9). Membrane potential is directly related to the activity of mitochondria, with more activity correlated with higher stress levels. Our findings show that there is lower mitochondria activity through the depolarization of the mitochondrial membrane after 1,25(OH)_2D treatment, hence less stress and ROS production. 1,25(OH)_2D decreased the mitochondrial membrane potential to a level sufficient for cells to survive
but insufficient to generate mt ROS, unlike other mitochondrial depolarizers such as hydrogen peroxide. Mitochondrial depolarization aims to reverse the inner membrane potential to generate minimal ATP with lower mt ROS, which is associated with increased longevity as observed in long-lived naked mole rats and bats. By contrast, in the absence of 1,25(OH)2D, MG-63 osteosarcomas are accompanied by mechanisms that prevent mitochondrial depolarization, resulting in chronic intracellular protein and DNA damage by mt ROS. In addition, 1,25(OH)2D treatment resulted in mitochondria with “ring-like” structures, which may represent intracellular organelar connections such as mitochondria-ER-associated membranes where the ER has been pulled into the lumen of the mitochondria to facilitate the direct transfer of phospholipids between the ER and mitochondria to help organize and generate cristae. It is also possible that 1,25(OH)2D may promote the sharing of other interorganellar resources, such as mitochondria-derived antioxidants or SOD2, to support the stressed ER, that has the potential to produce additional ROS itself. In contrast, untreated MG-63 cells exhibit a lack of distinct mitochondrial cristae, suggesting defects in the conductivity of structural proteins and stages of replication with constant flux between fusion and fission where the double-membrane structures are not as well developed. The potential reduction in fusion and fission after 1,25(OH)2D treatment may also result in reduced complementation of mitochondrial gene products that occurs to repair damaged mitochondria in untreated cancer cells. That is to say, the reduced fission/fusion may reflect the limited need for organelar quality control, supported by the downregulation of ATFS. Overall, in MG-63 cancer cells, 1,25(OH)2D functions to decrease mt ROS levels, which may also prevent ROS leakage to other cellular compartments to maintain molecular and organelar integrity.

4.2 1,25(OH)2D and ROS regulation

How 1,25(OH)2D reduces mitochondrial ROS levels is unclear and likely involves multiple factors, including cellular detoxification as well as reduced synthesis in the mitochondria via downregulation of the respiratory chain complex subunits. Although we showed that 1,25(OH)2D can enhance SOD2 levels, there are other ROS scavenging and degradation systems such as glutathione peroxidase, glutathione reductase, thioredoxin, and catalase, some of which were also regulated by 1,25(OH)2D treatment in our studies. Our results also show that 1,25(OH)2D can decrease the beta oxidation of fatty acids as an additional means to reduce ROS levels. 1,25(OH)2D may also regulate ROS production and turnover in other organelles besides mitochondria. ROS are also produced in the ER through the oxidation of proteins. Although our ER stress results show the activation of the early stage of the UPR after 1,25(OH)2D treatment, the more severe consequences of unfolded proteins (e.g., ER-mediated apoptosis) were not evident, suggesting that 1,25(OH)2D may also prevent this from occurring. In addition, 1,25(OH)2D induced heme oxygenase-1 (HMOX1) expression, which is an essential enzyme localized to the plasma membrane and Golgi apparatus for heme catabolism to form biliverdin, carbon monoxide, and ferrous iron. Free heme promotes ROS production by damaging DNA, lipids, and proteins. In conjunction, the generation of carbon monoxide may also regulate key anti-inflammatory

Fig 9. 1,25(OH)2D regulation of mitochondrial hormesis and functions within MG-63 osteosarcoma cells. (A) 1,25(OH)2D suppresses cancer progression by modulating the ROS threshold. 1,25(OH)2D-mediated mitohormesis and ROS production reprograms MG-63 osteosarcoma cells to enforce stress tolerance and growth inhibition to improve health outcomes. (B) 1,25(OH)2D modulates chromatin assembly and mitochondrial metabolism, oxidative stress, and mTOR inhibition via DDIT4/REDD1 localization to dictate its overall anticancer response. Red triangles represent acetylation.
cytokines (IL-10, IL-1RA, NFKBIA) that may play an anticancer role in MG-63 cells after 1,25(OH)2D treatment.2)

4.3 1,25(OH)2D and ROS consequences

The direct consequences of lowered ROS levels after 1,25(OH)2D treatment on intracellular targets and processes in cancer cells are unknown. It is known that at low levels, ROS can act as signaling molecules in various intracellular processes such as migration. At low levels, ROS differentially alters target protein conformation to modulate stability, folding, and activities of enzymes and ensuing phosphorylation cascades. The decrease in ROS production mediated by 1,25(OH)2D is likely to also affect downstream cystine oxidation of proteins to regulate, for example, DNA repair and damage, to facilitate the anticancer response. Overall, our results suggest that after 1,25(OH)2D treatment, the mitochondria produce low levels of ROS as a new set point that are effectively scavenged by the cancer cells’ antioxidant defense system. It is at this new set point that makes mitochondrial ROS an intracellular signaling molecule that does not induce oxidative stress but instead provides a safe “basal” window for redox signaling to alter the cancer cell fate.

Cell studies have shown that ROS can have both inductive and suppressive effects on global transcription as well as epigenetic responses in a highly cell type–specific manner.60) This may be related to the variable epigenetic and ROS-sensitive and/or insensitive transcription factors or factors that control the availability of antioxidant thiols within a cell. The effects of ROS on epigenetic and genetic mechanisms can involve direct effects that entail modifications of DNA bases and histones or indirect effects on DNA and histone-modifying enzymes to control cancer development.60) ROS can also affect class III histone deacetylases (HDACs) called sirtuins (SIRTs). Unlike class I, II, and IV HDACs that use metals as cofactors, SIRTs require NAD+ as a functional cofactor, thus making this enzyme sensitive to metabolic and redox changes to relay cellular stress in the form of histone modifications and changes in gene expression.60) In our study, 1,25(OH)2D treatment resulted in the downregulation of the mitochondrial SIRT1/4 deacetylases in MG-63 cells, which may signify a decoupling of lysine deacetylation with NAD+ hydrolysis and PDK4-acetyl-CoA (histone acetylation) to promote gene expression. Tumor studies have shown that SIRT4 has both oncogenic and tumor-suppressive activities in cancer depending on the experimental conditions.61) In the context of 1,25(OH)2D signaling and concomitant ROS reduction, SIRT1/4 downregulation may help create an epigenomic landscape and balance to facilitate 1,25(OH)2D-specific anticancer transcriptional responses and genomic stability.

4.4 1,25(OH)2D and stress tolerance and metabolic responses

Unchallenged protein misfolding can elicit cell death, while low levels of stress may be beneficial to cells by eliciting an adaptive UPR.30) Furthermore, the beneficial effects of mild stress on aging and longevity have been studied in experimental animals, whereby mild dietary stress by way of dietary restriction without malnutrition delays age-related physiological changes and extends the life span. Importantly, animal studies have also demonstrated that mild dietary stress can prevent or lessen the severity of cancer.72) Recent findings using the model organism, Caenorhabditis elegans, showed that 1,25(OH)2D can promote longevity by enhancing proteostasis,73) which may be akin to our findings of mitochondrial proteostasis and reduced biogenesis in MG-63 cells. These findings suggest that 1,25(OH)2D may mimic a metabolic state induced by dietary restriction and/or mild UPR to improve the life span and anticancer effects. Indeed, our previous studies showed that 1,25(OH)2D treatment was comparable to serum starvation of cultured osteoblasts, where suppression of the mTOR pathway was identified as a common feature and known also to be involved in life span expansion in mice when inhibited with rapamycin.74) Furthermore, our RNAseq and ATACseq motif analysis revealed associations with hypoxia, suggesting that 1,25(OH)2D may promote tumor starvation by inhibiting vascular perfusion less the negative effects of elevated ROS. Also, 1,25(OH)2D can promote mitochondrial depolarization, which is coupled to the availability of glucose or creatine, akin to dietary restriction to support sufficient mitochondrial ATP. These observations can also be metabolically linked to the increase in PDK4 we observed after vitamin D treatment. PDK4 is increased during hibernation/starvation and helps to decrease metabolism and conserve glucose by reducing its conversion to acetyl-CoA for ATP production.75)

Our model suggests that 1,25(OH)2D changes the metabolism of cancer cells from being responsive to stress to that of tolerant of stress that involves ER/mitohormetic processes with overall ROS reduction (Figs. 3 and 9). There is recent precedence for this model in the natural immunometabolism setting involving microbial-macrophage interactions.76) Timblin and colleagues showed that modulation of initial elevated antimicrobial ROS levels within macrophages involves ROS defense strategies as well as metabolic shifts toward non-oxidative energy metabolism, resulting in a reduction of ROS levels for macrophages to survive and function. Our model similarly shows a parallel paradigm enforced by 1,25(OH)2D on the dysregulated metabolism of MG-63 cancer cells. Co-opting this stress tolerance response identified in this study by 1,25(OH)2D may be a future strategy to consider toward cancer therapy. Importantly, we identified key 1,25(OH)2D-mediated metabolic enzymes that regulate fluxes of small compounds to provide the appropriate basal substrates for cell structure and energy production within dysfunctional osteosarcoma cells. For example, 1,25(OH)2D upregulated DMGDH, whereby it acts as an antioxidant when its enzymatic byproduct, dimethylglycine, is used to support the one-carbon (1-C) metabolism toward cytosolic NADPH production.135) Importantly, increased DMGDH levels are linked to hepatocellular carcinoma suppression.30) Furthermore, 1,25(OH)2D also positively regulates succinyl-CoA synthase, which facilitates the coupling of succinyl-CoA synthesis and hydrolysis to substrate level phosphorylation of ADP to ATP.43) The significance of this finding is that despite mitochondrial depolarization and OXPHOS inhibition after 1,25(OH)2D treatment, the cell can generate sufficient ATP via non-redox metabolism independent of mitochondrial electron acceptors to support anticancer biological activities, including survival.

4.5 Linking 1,25(OH)2D regulation of DDIT4/REDD1 to mitochondria and cancer biology

In the physiologic setting, DDIT4 is highly expressed in the cell cytoplasm under stress conditions such as hypoxia, cigarette smoke,77) and UV-induced DNA damage to function as a potent mTOR inhibitor to suppress cell proliferation and growth, while promoting autophagic processes instead. DDIT4 is also highly expressed in malignant cancers,53,44) despite its known mTOR-
inhibiting properties, suggesting that some cancers have evolved mechanisms to resist DDIT4, which may also antagonize antitumor therapies. For example, a meta-analysis of individual cancer data sets using gene expression profiling interactive analysis (GEPIA) shows that DDIT4 mRNA expression is significantly increased in numerous tumor tissues such as cervical squamous cell carcinoma (CESC)(23) (Supplemental Fig. S3); however, no data on osteosarcoma are currently available. We use GEPIA to further determine the overall cancer survival for CESC based on DDIT4 gene expression levels. DDIT4 levels were normalized for relative comparison between a housekeeping gene, ACTB, and the VDR gene. Using the log-rank test (Mantel-Cox test) for hypothesis evaluation, the hazard ratio (HR) and the 95% confidence interval (CI) information associated with both gene normalization comparisons suggest a significant association with decreased survival of patients with elevated DDIT4 levels (p = 0.0019 and 0.039 and HR = 2.1 and 1.6). The VDR relative comparison resulted in a higher p value and lower HR, suggesting direct regulation of DDIT4 levels by vitamin D across individuals. This association of decreased survival for high DDIT4 cohorts was observed for many other cancer types besides CESC presented in GEPIA, suggesting elevated DDIT4 is associated with poor prognosis and a vitamin D component.

In line with the findings from GEPIA, our findings in MG-63 cancer cells show that the mitochondria and their biogenic state can dictate DDIT4 cellular localization pattern and function. In contrast to MG-63 cancer cells, our previous findings using normal primary osteoblasts showed a robust cytoplasmic expression pattern of DDIT4 under basal settings,(23) which suggests a DDIT4 dichotomy between normal and cancer states. Currently, it is unknown if DDIT4 mitochondrial sequestration and biogenesis in the context of 1,25(OH)2D treatment and oxidative stress (GEPIA) shows that DDIT4 mRNA expression is significantly increased in cervical squamous cell carcinoma (CESC) and osteosarcoma. Future studies will focus on better understanding DDIT4 interactions via reactive cysteines with the mitochondria. This suggests that the n-terminus of DDIT4 that contains a cysteine residue in the cleavage domain (Supplemental Fig. S3). This suggests that 1,25(OH)2D, through its effects on ROS production, may regulate DDIT4 interactions via reactive cysteines with the mitochondria. Given the unknown function of DDIT4 in the mitochondria of MG-63 cells, future studies will focus on better understanding its role in the regulation of cell metabolism and mitochondrial biogenesis in the context of 1,25(OH)2D treatment and oxidative signaling.

Disclosures

All authors state that they have no conflicts of interest.

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Data availability statement

All data generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

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