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Personalized Genome-Scale Metabolic Models Identify Targets of Redox Metabolism in Radiation-Resistant Tumors

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Summary:

Redox cofactor production is integral towards antioxidant generation, clearance of reactive oxygen species, and overall tumor response to ionizing radiation treatment. To identify systems-level alterations in redox metabolism which confer resistance to radiation therapy, we developed a bioinformatics pipeline for integrating multi-omics data into personalized genome-scale flux balance analysis models of 716 radiation-sensitive and 199 radiation-resistant tumors. These models collectively predicted that radiation-resistant tumors reroute metabolic flux to increase mitochondrial NADPH stores and ROS scavenging. Simulated genome-wide knockout screens agreed with experimental siRNA gene knockdowns in matched radiation-sensitive and -resistant cancer cell lines, revealing gene targets involved in mitochondrial NADPH production, central carbon metabolism, and folate metabolism that allow for selective inhibition of glutathione production and H₂O₂ clearance in radiation-resistant cancers. This systems approach represents a significant advancement in developing quantitative genome-scale models of redox metabolism and identifying personalized metabolic targets for improving radiation sensitivity in individual cancer patients.

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By integrating multi-omics data into genome-scale metabolic models of individual patient tumors, Lewis et al. determine that radiation-resistant tumors produce elevated levels of reduced redox cofactors, allowing for increased ROS clearance through mitochondrial and folate-dependent pathways. This systems approach advances the identification of personalized gene targets impacting tumor redox metabolism.

**Graphical Abstract**

![Graphical Abstract Image]

**Keywords**

genome-scale; personalized models; flux balance analysis; redox metabolism; radiation resistance; NADPH; glutathione; hydrogen peroxide; reactive oxygen species; The Cancer Genome Atlas

**Introduction:**

Radiation therapy remains a cornerstone of cancer treatment, with more than half of all cancer patients receiving radiation as part of their treatment regimen (Delaney et al., 2005; Miller et al., 2016). Nonetheless, tumor resistance to radiation therapy constitutes a significant obstacle to long-term cancer patient survival. More than one-fifth of patients in The Cancer Genome Atlas (TCGA) database continued to show stable or progressive disease following radiation treatment, and almost all cancer types had some proportion of radiation-resistant patients (Weinstein et al., 2013). To investigate the underlying pathophysiological mechanisms of radiation resistance and discover targets for improving sensitivity to
radiation therapy, correlative studies using single omics modalities such as genomics or transcriptomics have been previously performed (Chen et al., 2015; Lee et al., 2010; Manem and Dhawan, 2019; Skvortsov et al., 2014; Skvortsova et al., 2008; Smith et al., 2009). However, these previous findings fall short by failing to integrate multiple biological data types, analyze differential expression in the context of genome-scale biochemical and regulatory networks, or provide mechanistic insights into how prospective biomarkers impact tumor function and can be exploited to improve radiation sensitivity.

Redox metabolism relies on the oxidation and reduction of electron-carrying molecules such as NADPH, NADH, and glutathione (GSH), which are used as cellular antioxidants and electron donors for metabolic reactions (Lewis et al., 2019; Xiao et al., 2018). These metabolites represent the reduced forms of redox couples with their associated oxidized forms NADP⁺, NAD⁺, and glutathione disulfide (GSSG), respectively; the ratio of reduced to oxidized forms of these redox couples provides an indication of the intracellular redox potential and oxidative state of the cell (Mallikarjun et al., 2012). Ionizing radiation therapy results in the generation of reactive oxygen species (ROS) such as superoxide (O₂⁻) and hydrogen peroxide (H₂O₂), which oxidize the cellular environment and damage cellular structures including DNA (Brady et al., 2013; Cadet and Wagner, 2013; Reisz et al., 2014; Tominaga et al., 2004). Redox cofactors such as NADPH and GSH can be utilized by H₂O₂-scavenging enzymes to lower cellular levels of ROS (Forshaw et al., 2019; Harris et al., 2015). Additionally, these cofactors can directly promote DNA damage repair following oxidative damage, either by reduction of nitrogenous bases or utilization of NAD(P)H for nucleotide synthesis (Alvarez-Idaboy and Galano, 2012; Chatterjee, 2013; Franklin et al., 2016; Turgeon et al., 2018). Since redox metabolism is critical to the response of tumors to ionizing radiation, identifying targets for inhibiting production of these redox cofactors may provide a valuable strategy for sensitizing tumors to radiation therapy (Lewis et al., 2019).

Because redox cofactors are utilized in thousands of reactions throughout the human metabolic network, computational methods are needed to investigate systems-level redox metabolism and its interconnections with other cellular metabolic pathways (Brunk et al., 2018). Flux balance analysis (FBA) is a computational approach for predicting steady-state metabolic fluxes at a genome scale for cells or tissues of interest (Orth et al., 2010). By combining the stoichiometric representation of the human metabolic network, constraints on the fluxes through metabolic reactions, and an objective function to maximize a particular metabolic phenotype, predictions of maximum reaction fluxes or metabolite production rates under physiological constraints are generated (Bordel, 2018; Oberhardt et al., 2010; Supandi and Van Beek, 2018). Recently, FBA models personalized to individual cancer cell lines or tumors have been developed through the integration of transcriptomic data; however, there remain significant methodological shortcomings in the construction of these models that have hindered their ability to yield accurate and quantitative metabolic predictions (Blazier and Papin, 2012). Commonly-used FBA model extraction methods such as GIMME, iMAT, MADE, and CORDA utilize arbitrary gene expression thresholds to constrain metabolic activity, necessitate the comparison of multiple transcriptomic datasets, or completely remove reactions from the metabolic network with low associated gene expression (Becker and Palsson, 2008; Jensen and Papin, 2011; Schultz and Qutub, 2016; Shlomi et al., 2008). Other methods such as E-Flux constrain maximum reaction fluxes in proportion to the
associated enzyme’s gene expression, but arbitrary reference values and proportionality functions are still used to set flux constraints instead of directly estimating enzyme abundances (Colijn et al., 2009). Additionally, most FBA models fail to incorporate any kinetic or thermodynamic constraints, which greatly affect metabolic fluxes and the directionalities of individual reactions (Henry et al., 2007). These shortcomings in model construction have ultimately limited the clinical utility of FBA models for accurately predicting metabolic phenotypes and directly improving cancer diagnosis and treatment (Zhang and Hua, 2015).

We have resolved many of these limitations in FBA model development and utility by integrating transcriptomic, kinetic, and thermodynamic information into quantitative constraints on the maximum flux and directionality of metabolic reactions. In our previously-developed FBA models of radiation-sensitive and radiation-resistant head and neck squamous cell carcinoma (HNSCC) cell lines, we accurately identified oxidoreductase genes that differentially impacted response to treatment with the NADPH-dependent redox-cycling chemotherapeutic β-lapachone between radiation-sensitive and -resistant cells (Lewis et al., 2018). Additionally, our models suggested that radiation-resistant cancer cells re-route NADH-generating metabolic fluxes through NAD salvage and purine salvage pathways involving NAMPT, an enzyme whose activity has previously been associated with radiation resistance and poor survival in cancer patients (Gujar et al., 2016; Lewis et al., 2019). Here, we extend this approach by developing an automated bioinformatics pipeline for integrating multi-omics information from The Cancer Genome Atlas (TCGA) and publically-available repositories into personalized genome-scale FBA models of 915 patient tumors across multiple cancer types. These personalized FBA models are used to investigate differences in redox cofactor production between radiation-sensitive and -resistant tumors, discover gene targets which differentially impact redox metabolism in radiation-resistant tumors, and identify personalized therapeutic strategies for individual radiation-resistant patients.

Results:

An automated bioinformatics pipeline integrates multi-omics data into personalized FBA models of TCGA patient tumors

The framework for building FBA models of tumor metabolism was initiated with the community-curated Recon3D human metabolic reconstruction (8,401 metabolites, 13,547 reactions, and 3,268 genes; Figure 1A) (Brunk et al., 2018). The stoichiometric representation of this reconstruction is combined with minimum and maximum constraints on reaction fluxes to obtain a solution space for steady-state fluxes throughout the metabolic network. An objective function is typically utilized to narrow the possible FBA solution space to physiologically-optimal metabolic fluxes (Garcia Sanchez and Torres Saez, 2014). We hypothesized that tumors are under selective pressure to maximize production of reduced redox cofactors including NADPH, NADH, and GSH to decrease ROS-mediated damage induced by ionizing radiation; thus, we used an objective function of maximizing redox cofactor reduction. FBA is used to compare maximum cofactor reduction between radiation-
sensitive and -resistant tumors, while flux variability analysis (FVA) is used to predict fluxes through individual metabolic reactions involved in cofactor reduction (Figure 1B).

To generate personalized FBA models of radiation-sensitive and -resistant TCGA patient tumors, Michaelis-Menten $V_{\text{max}}$ constraints were set on all 4,367 Recon3D reactions that contain both a gene-protein-reaction (GPR) rule and enzyme commission (EC) number (Figure 1C). This constraint sets the maximum flux for each reaction ($V_{\text{max}}$, units of mmol gDW$^{-1}$ hr$^{-1}$) equal to the kinetic rate constant of the enzyme catalyzing the reaction ($k_{\text{cat}}$, units of hr$^{-1}$), multiplied by the estimated protein abundance of the enzyme ([E], units of mmol gDW$^{-1}$). A custom protein prediction pipeline was developed to convert RNA-seq transcriptomic data from individual TCGA samples into estimated enzyme abundances ([E]) (Figure S1A). This pipeline takes advantage of models relating the mRNA and protein abundances for individual genes/proteins using experimentally-measured transcription, translation, and degradation rates (Schwanhausser et al., 2011). Predicted enzyme abundances from this pipeline had improved correlation with experimental protein expression values from both NCI-60 and TCGA datasets compared to original gene expression values (Figure S1B–I). Additionally, a custom kinetic rate pipeline was developed to extract physiologically-accurate turnover numbers for metabolic enzymes ($k_{\text{cat}}$) by matching experimentally-measured values from the BRENDA database with the correct enzyme, substrate, organism, and environmental conditions as those in the Recon3D network (Figure S2) (Schomburg et al., 2004). Envision database scores were applied to predict the effect of mutations in individual TCGA samples on the catalytic rate of corresponding metabolic enzymes (Gray et al., 2018). Finally, transformed changes in Gibbs free energy ($\Delta G'$) from the Virtual Metabolic Human (VMH) database were used to set thermodynamic constraints, such that only reactions with a negative $\Delta G'$ can carry non-zero net fluxes (Noor et al., 2013). These proteomic, kinetic, and thermodynamic constraints yield quantitative, patient-specific, and physiologically-accurate predictions of metabolic fluxes on a genome scale.

RECIST classification of TCGA samples provided an evaluation metric of radiation sensitivity based on changes in tumor size in response to radiation therapy (Figure 1D). Patients with a complete or partial response to radiation (greater than 30% decrease in tumor size) were classified as radiation-sensitive, and patients with stable or progressive disease (either less than 30% decrease in tumor size, or increase in tumor size) were classified as radiation-resistant. Using this classification, 716 personalized FBA models of radiation-sensitive tumors and 199 personalized models of radiation-resistant tumors were generated. Comparisons of FBA predictions between radiation-sensitive and -resistant tumor models were controlled for patient and tumor covariates including cancer type and tumor staging by calculating propensity scores for each patient tumor using covariate values and performing analysis of covariance (ANCOVA) (Austin, 2011; Kahlert et al., 2017; Kurth et al., 2006).

**Radiation-resistant tumors display compartmental differences in redox metabolic fluxes compared to radiation-sensitive tumors**

Personalized FBA models were first used to compare differences in redox cofactor production between radiation-sensitive and -resistant tumors. Radiation-resistant tumor
models showed significantly elevated production of reduced cofactors NADPH, NADH, and GSH (Figure 2A). To validate model predictions, we performed experimental measurements in matched pairs of radiation-sensitive and -resistant cell lines across three different cancer types (Figure 2B, Table S1). Two of the three radiation-resistant cell lines had significantly more reduced glutathione half-cell potentials (E_{hc} GSH/GSSG) than their matched radiation-sensitive cell line, indicating greater conversion of oxidized GSSG to reduced GSH as predicted in TCGA models (Figure 2C, Figure S3). Although BRCA cell lines showed the opposite trend, the observed E_{hc} difference was much smaller than in the other two cancer types and could be attributed to compensation for NQO1 depletion by increased expression of other antioxidant enzymes involved in glutathione reduction or which bypass the need for GSH (e.g. peroxiredoxin or thioredoxin systems) (Cao et al., 2014). While predicted cytosolic production of NADPH did not differ between tumor classes, increased mitochondrial NADPH production in radiation-resistant tumors accounted for observed differences in total cellular NADPH production (Figure 2D). In agreement with this finding, increased deoxynucleotide production (which relies on mitochondrial NAD(P)H) was seen in radiation-resistant tumor models, while no significant differences in production of fatty acid precursors including palmitate (which relies on cytosolic NAD(P)H) were observed (Figure S4) (Jones, 1980; Lewis et al., 2014; Schnell et al., 2004; Turgeon et al., 2018). Significant differences were observed in NADPH fluxes between radiation-resistant and -sensitive patients regardless if the tumor sample was collected before or after radiation therapy, with no significant differences within the radiation-resistant or -sensitive classes based on collection before versus after therapy (Figure S5).

Flux variability analysis (FVA) was used to compare fluxes through major NADPH-generating metabolic reactions between radiation-sensitive and -resistant tumors (Figure 2E–F). Predicted fluxes through major cytosolic reactions including G6PD and PGD were greater in radiation-sensitive tumors, in agreement with our previously identified flux distributions in radiation-sensitive HNSCC cell lines as well as experimental findings (Lewis et al., 2018; Mims et al., 2015). On the other hand, predicted fluxes through major mitochondrial and folate-dependent reactions including GLUD1/2, ME2/3, IDH2, MTHFD2, and MTHFD1 were greater in radiation-resistant tumors, accounting for net increased NADPH production in these tumors. High-flux NADPH-generating reactions reached fluxes near their V_{max} for almost all samples, whereas low-flux reactions varied significantly based on reaction compartmentalization and sample radiation response (Figure S6, S7). Hierarchical clustering of NADPH-producing fluxes yielded pronounced separation of cytosolic and mitochondrial reactions except for the folate-dependent reactions MTHFD1 and MTHFD2, both of which displayed increased fluxes in radiation-resistant tumor models (Figure 2G). Optimal sample clustering measured by the silhouette coefficient was found to be based on radiation response rather than other clinical factors such as cancer type (Figure S8). Collectively, these results suggest that differences in mitochondrial- and folate-dependent NADPH metabolism can discriminate between radiation-sensitive and -resistant tumors.
**Simulated genome-wide knockout screen identifies targets of redox metabolism in radiation-resistant cancers**

FBA models enable the assessment of changes in metabolic fluxes in response to knockout of a particular metabolic enzyme-encoding gene, providing insight into the gene’s role within genome-scale metabolism (Figure 3A). A simulated genome-wide knockout screen was performed to predict the effect of knocking out each individual gene in Recon3D on total cellular NADPH production across all TCGA models (Figure 3B, Figure S9). Most gene knockouts did not significantly decrease total NADPH production, corroborating our previous findings that metabolic networks redirect flux through alternate, compensatory pathways following perturbation to optimize NADPH production (Lewis et al., 2018). Among the knockouts with largest predicted effects were genes directly involved in NADPH generation (G6PD, GLUD1, ME1), as well as those involved in glycolysis (ENO1, GAPDH, GPI, HK1, PGAM1, PKM), folate metabolism (DHFR), and amino acid metabolism (PYCR2).

A statistical comparison of gene knockout effect on NADPH production between radiation-sensitive and -resistant tumors was performed to identify redox metabolic targets selective for radiation-resistant tumors (Figure 3C, Table S2). 27 gene knockouts caused a significantly greater decrease in NADPH production among radiation-resistant tumors, including those involved in mitochondrial and folate-dependent NADPH metabolism (GLUD1/2, IDH2, ME2, MTHFD1, MTHFR) as well as many aldehyde dehydrogenase genes (ALDH1L1, ALDH1L2, ALDH5A1); many of these targets are consistent with those previously implicated in the response of radiation-resistant HNSCC to NADPH-depleting chemotherapies (Lewis et al., 2018). 22 gene knockouts caused a significantly greater decrease in NADPH production among radiation-sensitive tumors, including those of the pentose phosphate pathway (G6PD, PGD, PGLS, RPE, RPIA, TALDO1, TKT). Differences in gene expression between radiation-sensitive and -resistant tumors show minimal utility alone in predicting differences in FBA gene knockout effects on NADPH production (Figure S10). While many targets with differential effects on NADPH production were also found to differentially impact GSH production between radiation-sensitive and -resistant tumor models, some notable differences were observed (Figure 3D, Figure S11A, Table S3). Whereas pentose phosphate pathway genes were not predicted to significantly affect GSH production, those involved in the TCA cycle (DLST, FH, OGDH, SDHA), oxidation-reduction reactions (AKR1A1, GSR, PRDX6), and glycine metabolism (AMT, GCSH, GLDC) were identified; glycine metabolism has been previously shown to impact GSH/GSSG ratios and ROS levels in cancer (Zhuang et al., 2018). Additionally, the folate metabolism gene MTHFR had a larger impact on NADPH production in radiation-resistant tumor models but impacted GSH production to a greater extent in radiation-sensitive models, possibly attributing to folate and NADPH’s role in other metabolic pathways including nucleotide synthesis (Fan et al., 2014). Finally, de novo GSH production was found to be significantly increased in radiation-resistant tumor models through γ-glutamyl cysteine synthase (γGCS) and glutathione synthetase (Figure S12); this synthesis flux is highly correlated to glutathione reduction flux (Figure S13).
To validate model-predicted targets of glutathione metabolism, the change in glutathione half-cell potential (ΔE_{hc} GSH/GSSG) between siRNA knockdown cells and negative control siRNA-transfected cells was measured in both radiation-sensitive and -resistant cell lines for 12 different gene targets (Figure 3E). 4 out of 5 model-predicted radiation-resistant targets (ALDH4A1, GLUD1, GLUD2, LDHB) had an overall more oxidizing effect in radiation-resistant cell lines, with GLUD1 having the largest differential oxidizing effect across all siRNA’s tested (Figure 3F). Additionally, 4 of the 5 experimental gene targets with more oxidizing effects in radiation-sensitive cell lines (GSR, MTHFR, OGDH, PRDX6) were identified by FBA models as radiation-sensitive targets. These results suggest that FBA models of TCGA tumors can be used to predict gene targets which differentially impact redox cofactor production associated with radiation sensitivity.

Disparities in redox metabolism and H₂O₂-scavenging systems between radiation-sensitive and -resistant tumors determine differential ROS response

By utilizing an FBA objective function maximizing H₂O₂ clearance, radiation-resistant tumor models predicted increased H₂O₂-scavenging potential compared to radiation-sensitive tumors (Figure 4A). To validate model predictions of H₂O₂ response, matched radiation-sensitive and -resistant cell line pairs were treated for 2 hours with 10 mU/mL of the H₂O₂-generating enzyme glucose oxidase, and measured ATP levels were used as an indicator of cell viability (Adimora et al., 2010; Bankar et al., 2009; Daniela et al., 2015). All cell lines showed decreased viability with glucose oxidase treatment, but 2 out of the 3 radiation-resistant cell lines showed a significantly lesser decrease in relative cell viability compared to their matched radiation-sensitive cell lines (Figure 4B). These findings were consistent with experimentally-measured differences in glutathione half-cell potential (Figure 2C). FVA was used to compare fluxes through major H₂O₂-clearing reactions between radiation-sensitive and -resistant tumors, accounting for enzyme isoform differences between cellular compartments (Figure 4C). Mitochondrial flux through catalase (CAT) accounted for the increased H₂O₂-clearing capability of radiation-resistant tumor models; although CAT does not directly use redox cofactors in the reaction mechanism, it relies on NADPH to prevent oxidation of its active site, providing a link between cofactor production and ROS clearance (Kirkman et al., 1999). Superoxide dismutase isoforms were predicted to have significant impacts on H₂O₂ production (Figure S14, S15).

A simulated genome-wide knockout screen was performed to identify gene targets with greater effects on H₂O₂ clearance in radiation-resistant tumors (Figure 4D, Figure S11B, Table S4). While CAT was the only significant gene with direct H₂O₂-scavenging function, many gene targets of NADPH production were predicted to have significant differential effects on H₂O₂ clearance. Radiation-sensitive targets included members of the pentose phosphate pathway (G6PD, PGD, PGLS, RPE, RPIA, TALDO1, TKT) whereas radiation-resistant targets included genes involved in mitochondrial and folate-dependent NADPH production (GLUD1/2, MTHFD1, MTHFD2, MTHFR) and central carbon metabolism (ACO2, ALDOC, PGAM2, PKM). To validate these targets, Bliss independence scores indicating the effect of siRNA gene knockdown on glucose oxidase response were measured in radiation-sensitive and -resistant cell lines for 9 different gene targets (Figure 4E):
Bliss = \log_2 \left( \frac{\text{siRNA: 10 mU/mL G. O.}}{\text{siRNA: 0 mU/mL G. O.}} \right) = \log_2 \left( \frac{\text{Control: 10 mU/mL G. O.}}{\text{Control: 0 mU/mL G. O.}} \right)

Values < 0 indicate synergistic effects between siRNA and glucose oxidase treatments for decreasing cell viability, whereas values > 0 indicate antagonistic effects between siRNA and glucose oxidase treatments. Thus, Bliss\text{Sens} - Bliss\text{Res} > 0 for siRNA gene knockdowns causing a greater decrease in cell viability with glucose oxidase treatment in the radiation-resistant cell line compared to the radiation-sensitive cell line, corresponding to a model-predicted \log_2(\text{Resistant/Sensitive}) > 0. 3 of the model-predicted radiation-resistant targets (\text{GLUD1}, \text{MTHFD1}, \text{and PGAM2}) had the largest differential response in radiation-resistant cell lines across all siRNA’s tested (Figure 4F). Furthermore, a larger differential response to ALDH1L1 knockdown was observed in 2 of the 3 radiation-resistant cell lines, as predicted by FBA models. TCGA models accurately predicted that \text{MTHFD1} knockdown would have a greater effect on GSH production in radiation-sensitive cancers but greater effect on \text{H}_2\text{O}_2\text{ response in radiation-resistant cancers, suggesting that FBA models accurately capture other metabolic systems which impact ROS clearance besides glutathione-dependent pathways (Forshaw et al., 2019). Overall, the observed agreement between FBA model predictions and experimental validation demonstrates the ability to correctly identify targets of redox metabolism which differentially impact radiation-resistant cancers.}

**Personalized metabolic flux profiles highlight heterogeneity in redox metabolism between radiation-resistant tumors**

Although radiation-resistant tumor models displayed overall differences in redox metabolism compared to radiation-sensitive models, individual patient tumors may exemplify divergent metabolic flux profiles and thus differences in optimal therapeutic strategies for improving radiation sensitivity (Kim and DeBerardinis, 2019). To determine if patient clinical information could be used to distinguish radiation-resistant tumor models with differing metabolic phenotypes, we evaluated the correlation between clinical factors and predicted fluxes through major NADPH-generating reactions and other radiation-resistant NADPH targets (Figure 5A, Figure S16). While cancer type, patient age, and tumor grade were highly correlated with predicted fluxes through most reactions, other clinical factors were associated with a few select reactions. Smoking history was highly associated with G6PD and PGD fluxes, in agreement with previous experimental studies showing that exposure to cigarette smoke causes upregulation of G6PD and shifts glucose metabolism towards the pentose phosphate pathway for increased NADPH production (Agarwal et al., 2014; Noronha-Dutra et al., 1993). The association with smoking history is particularly strong for G6PD fluxes in lung adenocarcinoma (LUAD) and for PGD fluxes in cervical squamous cell carcinoma/endocervical adenocarcinoma (CESC) (Figure S17). Additionally, response to cisplatin treatment was highly associated with fluxes through mitochondrial reactions IDH2 and NNT (Figure S18); treatment with cisplatin has been previously reported to induce a mitochondrial-ROS response, which could lead to upregulation of mitochondrial NADPH-generating enzymes (Choi et al., 2015; Marullo et al., 2013).
To investigate whether patients with similar clinical features also displayed similarities in tumor redox metabolism, we evaluated personalized metabolic flux profiles of radiation-resistant patients with head and neck squamous cell carcinoma, a cancer type characterized by high genetic and metabolic heterogeneity (Figure 5B, Figure S19) (Chen et al., 2018). Predicted fluxes through NADPH-generating reactions differed substantially between patients with very similar clinical features; while some tumor models showed increased NADPH generation through single reactions (TCGA-T2-A6WZ: G6PD; TCGA-CN-6998: ME1), others showed increased fluxes through multiple disparate reactions (TCGA-HD-8224: ME1, ME2/3, NNT). These stark differences in redox metabolism among patients with similar clinical and demographic attributes demonstrate the utility of personalized genome-scale metabolic models for identifying therapeutic targets in individual patients.

Discussion:

Identifying redox metabolic targets which impact radiation sensitivity through modulation of antioxidant levels and oxidative DNA damage is an ongoing challenge (Spitz et al., 2004). Flux balance analysis (FBA) is a genome-scale metabolic modeling approach that has gained attraction for building mechanistic models of cancer metabolism and identifying chemotherapeutic targets (Folger et al., 2011; Nilsson and Nielsen, 2017; Yizhak et al., 2014). Because redox cofactors are involved in thousands of reactions throughout the human metabolic network, genome-scale FBA models are well-suited to study redox metabolism and their interconnections with other metabolic processes (Brunk et al., 2018). Nonetheless, few studies have used FBA models for studying cancer redox metabolism. A notable example is that of Benfeitas et al., who developed FBA models of hepatocellular carcinomas (HCC) to characterize heterogeneity in redox metabolism between HCC tumors at differing stages of progression and within different HCC tumor clusters (Benfeitas et al., 2019). Using these models, the authors discovered distinct differences in NADPH production and \( \text{H}_2\text{O}_2 \) clearance between HCC tumors with high \( \text{G6PD} \) expression and those with high \( \text{ALDH2} \) expression. However, their use of the MADE method for converting transcriptomic data into upper flux bounds results in discrete reaction constraints which lack quantitative accuracy compared to using continuous enzyme abundance values. Furthermore, kinetic and thermodynamic effects on reaction constraints, which have significant impacts on metabolic fluxes, were not present in these models (Henry et al., 2007).

Our bioinformatics and metabolic modeling pipeline represents, to our knowledge, the first study to integrate multiple omics datasets into human genome-scale mechanistic models to compare redox metabolism between radiation-sensitive and -resistant tumors, as well as to identify therapeutic biomarkers for improving radiation therapy response. Methodological shortcomings of previous FBA studies are overcome by incorporating transcriptomic and mutational data from individual patient tumors, as well as genome-scale kinetic and thermodynamic parameter values, into quantitative constraints on metabolic fluxes, allowing for more accurate metabolic predictions (Figure 1). Analysis of differential gene expression between radiation-sensitive and -resistant tumors alone does not accurately predict enzyme-catalyzed metabolic fluxes (Figure S20, S21). Additionally, previous FBA modeling approaches such as E-Flux which solely use transcriptomic data to set reaction constraints fail to identify important differences in redox metabolism between radiation-sensitive and -
resistant tumors (Figure S22) (Colijn et al., 2009). Exclusion of genomic or thermodynamic data from FBA models impacts predictions of redox metabolism in an objective function-specific manner (Figures S23–S26). The use of redox-specific objective functions also allowed for the identification of gene targets with significant differential effects on redox metabolism between radiation-sensitive and -resistant tumors. For example, the NADPH objective function recapitulates many of the differences between radiation-sensitive and -resistant tumor models observed from using the biomass objective function, while also allowing for the identification of redox metabolic targets in radiation-sensitive and -resistant tumors (Figure S27). This combination of multi-omics data integration and context-specific objective functions had significant impacts on FBA model outputs, yielding quantitative metabolic predictions with experimentally-validated accuracy.

FBA models of radiation-resistant TCGA tumors showed increased fluxes through mitochondrial NADPH-producing reactions, allowing for elevated mitochondrial stores of reduced redox cofactors as well as increased fluxes through mitochondrial H\textsubscript{2}O\textsubscript{2}-clearing reactions (Figures 2, 4). Mitochondrial NADPH-producing enzymes identified from this study, including GLUD1, IDH2, ME2, and NNT, have been previously implicated in regulation of oxidative stress and tumor proliferation; additionally, mitochondrial-specific compartmentalization of signaling and energy metabolism has been previously identified to impact chemotherapy and radiation response (Ciccarese and Ciminale, 2017; Hsieh et al., 2015; Jin et al., 2015; Porporato et al., 2018; Stein et al., 2014; Yin et al., 2012). Simulated genome-wide knockout screens and validation with experimental siRNA knockdown experiments suggest that enzymes involved in mitochondrial (GLUD1/2) and folate-dependent (MTHFD1) NADPH production and central carbon metabolism (LDHB, PGAM2) may be viable targets for inhibiting GSH production and/or H\textsubscript{2}O\textsubscript{2} clearance in radiation-resistant tumors (Figures 3, 4). On the other hand, G6PD, the most well-characterized NADPH-producing reaction, exemplified larger fluxes in radiation-sensitive tumor models and showed lower experimental effects on targeted H\textsubscript{2}O\textsubscript{2} clearance in radiation-resistant cell lines compared to the aforementioned gene targets. Interestingly, the G6PD cluster identified by Benfeitas et al. in HCC is highly enriched in genes identified from this current study as radiation-sensitive targets (G6PD, PGD, RPE, RPL4), whereas the ALDH2 cluster is highly enriched in genes identified as radiation-resistant targets (ALDH2, ALDH5A1, ALDH6A1, CAT, MTHFD1, SHMT1) (Benfeitas et al., 2019). Despite the fact that the authors were not analyzing redox metabolism in the context of radiation sensitivity, this correspondence between gene sets from separate analyses may suggest that there exist two fundamental tumor subtypes with distinct redox metabolic phenotypes and corresponding radiation sensitivities.

By developing FBA models of individual patient tumors, personalized metabolic flux profiles were generated to identify patient-specific redox biomarkers (Figure 5). These personalized metabolic predictions also demonstrate the significant amount of heterogeneity in redox metabolism between patient tumors. While the majority of initiatives for precision medicine for cancer treatment have focused on mutations and expression differences in signaling pathway proteins, there is currently a greater focus on exploiting metabolic differences between patient tumors (DeBerardinis and Chandel, 2016; Kanarek et al., 2018; Kanarek et al., 2020). For example, heterogeneity in glycolytic metabolism is being used to
identify diagnostic biomarkers and treatment strategies for both pancreatic cancer and acute myeloid leukemia patients (Follia et al., 2019; Stuani et al., 2019). Personalized nutrition may also be a viable strategy for manipulating tumor redox metabolism in individual patients; most NADPH-generation pathways are supplied by glucose and glutamine intake, and folic acid levels may impact folate-dependent reactions (Choi and Park, 2018; Wallace et al., 2019). Continued development of genome-scale metabolic models of individual patient tumors will undoubtedly aid in the identification of metabolic targets or optimal dietary strategies for individual cancer patients.

Although our bioinformatics and modeling approach towards using multi-omics data for predicting metabolic phenotypes in individual patient tumors represents a significant methodological advancement over previous FBA models, additional improvements and integrations with other modeling strategies would further improve its accuracy and applicability. Currently, rates of metabolite transport through specific plasma membrane transporters are much less characterized than turnover rates of intracellular metabolic enzymes, limiting the implementation of quantitative transport constraints (Schomburg et al., 2004). Unless experimentally-measured metabolite uptake rates from samples of interest are obtained, constraints on uptake reactions are commonly set as binary (i.e. if a metabolite is present in the extracellular medium, intracellular uptake is unconstrained; otherwise, uptake is set to zero). A potential approach towards setting quantitative uptake constraints would be to relate membrane transporter expression with experimentally-measured extracellular metabolite concentrations (for example, from cell culture media or patient blood samples) to predict individualized metabolite uptake rates. Regarding the application of FBA models towards studying redox metabolism, integration of important redox signaling and regulatory networks such as the Nrf2/Keap1/ARE pathway through methods such as integrated FBA (iFBA) could improve the accuracy of predicted reaction fluxes which utilize ARE-regulating genes including G6PD, IDH1, ME1, and PGD (Covert et al., 2008; Jaramillo and Zhang, 2013; Kansanen et al., 2013; Lee et al., 2008; Lin et al., 2016). Additionally, the role of H$_2$O$_2$ in redox-regulated signal transduction is not reflected in the constraint-based models discussed in the current study.

Despite the recent increase in experimental metabolomics studies towards identifying altered metabolic phenotypes in cancer, an integrated assessment of the 13,000+ human metabolic reactions and their collective impact on individualized treatment response is currently infeasible using solely experimental approaches (Kaushik and DeBerardinis, 2018). Instead, computational approaches which integrate multi-omics measurements and global reconstructions of human metabolism into predictive models provide tremendous utility in improving our understanding of pathophysiological processes and discovering personalized metabolic targets (Nilsson and Nielsen, 2017). Our analysis indicates that genome-scale metabolic models of individual patient tumors can identify important differences in redox metabolism between radiation-sensitive and -resistant tumors. Specifically, by comparing properties of 716 radiation-sensitive and 199 radiation-resistant personalized tumor models, we have elucidated multiple mechanisms of how tumors can upregulate metabolic flux through mitochondrial NADPH-generating and H$_2$O$_2$-clearing reactions to increase cellular antioxidant stores, decrease ROS levels, and resist the damaging effects of ionizing radiation treatment. These model-identified targets significantly impact antioxidant production and
ROS response, thus serving as putative biomarkers for the \textit{a priori} prediction of radiation sensitivity, as well as therapeutic strategies for sensitizing tumors to radiation therapy. Ultimately, the development of personalized metabolic models has the potential to facilitate the clinical management of cancer patients and improve long-term outcomes.

\textbf{STAR Methods:}

\textbf{RESOURCE AVAILABILITY}

\textbf{Lead Contact—}Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Melissa L. Kemp (melissa.kemp@bme.gatech.edu).

\textbf{Materials Availability—}This study did not generate new unique reagents.

\textbf{Data and Code Availability}

- \textbf{Source Data:} TCGA tumor models developed for this study are available at https://github.com/kemplab/FBA-pipeline. The following datasets are available at https://github.com/kemplab/FBA-pipeline:
  - Dataset 1. TCGA samples included in the analysis, with corresponding radiation response, patient/tumor factors, and propensity scores
  - Dataset 2. NADPH-generating fluxes for TCGA tumor models [mmol/gDW/hr] (Related to Figure 2)
  - Dataset 3. Simulated effect of gene knockout on \textit{NADP}^{+} \rightarrow \textit{NADPH} [KO/WT] (Related to Figure 3)
  - Dataset 4. Simulated effect of gene knockout on \textit{GSSG} \rightarrow \textit{GSH} [KO/WT] (Related to Figure 3)
  - Dataset 5. Comparison of model-predicted and experimentally-validated \textit{GSSG} \rightarrow \textit{GSH} gene targets (Related to Figure 3)
  - Dataset 6. Simulated effect of gene knockout on \textit{H}_{2}\textit{O}_{2} \rightarrow [KO/WT] (Related to Figure 4)
  - Dataset 7. Comparison of model-predicted and experimentally-validated \textit{H}_{2}\textit{O}_{2} \rightarrow gene targets (Related to Figure 4)

- \textbf{Original Code:} Code for the generation and simulation of personalized FBA models is available at https://github.com/kemplab/FBA-pipeline. Personalized models can be developed for any human sample (such as cell lines or patient tumors) with RNA-seq gene expression data, and mutation data if available. Jupyter notebooks are available for 1) processing of sample RNA-seq data to estimate enzyme abundance values, 2) processing of sample mutation data to estimate kinetic rate parameters, and 3) running FBA analysis with user-specified analysis type, model constraints, media constraints, objective function, and samples of interest.
• **Scripts Used to Generate Figures**: The scripts used to generate the figures reported in this paper are available in the Python packages matplotlib (version 3.1.0, matplotlib.org) and seaborn (version 0.10.1, seaborn.pydata.org). The following functions were used to generate figures from the source data provided above:
  – Bar graphs: matplotlib.pyplot.bar()
  – Box plots: seaborn.boxplot() and seaborn.swarmplot()
  – Heatmaps: seaborn.heatmap()
  – Hierarchical clustering: seaborn.clustermap()
  – Volcano plots and spider plots: matplotlib.pyplot.plot()

• Any additional information required to reproduce this work is available from the Lead Contact.

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

**Cell Lines**—Table S1 provides the matched radiation-sensitive and radiation-resistant human cancer cell lines used for experimental validation of model predictions (Bansal et al., 2014; Huang et al., 2016; Zhao et al., 2011). All cell lines were maintained in RPMI-1640 cell culture media (Thermo Fisher Scientific, Cat#11875) with 10% fetal bovine serum (Sigma-Aldrich, Cat#F4135) at 37°C and 5% CO₂, and were free of *Mycoplasma*.

**METHOD DETAILS**

**Flux Balance Analysis (FBA)**—A metabolic network can be represented by a stoichiometric matrix $S$ of size $m \times r$, where $m$ and $r$ are the number of metabolites and reactions in the network, respectively. Entry $S_{ij}$ is equal to the stoichiometric coefficient of metabolite $i$ in reaction $j$ ($S_{ij} < 0$ for reactants, $> 0$ for products, and $= 0$ if metabolite $i$ is not involved in reaction $j$). The relationship between reaction fluxes and metabolite concentrations in the network can be written as:

$$Sv = \frac{dm}{dt}$$

(2)

where $v$ is a $r \times 1$ vector of reaction fluxes, and $m$ is a $m \times 1$ vector of metabolite concentrations. In flux balance analysis, the steady state is assumed (metabolite concentrations do not change with time), changing Equation 2 to:

$$Sv = 0$$

(3)

Each reaction flux $v_j$ is also constrained by lower and upper bounds:

$$v_{min,j} \leq v_j \leq v_{max,j}$$

(4)

The solutions to Equations 3 and 4 that maximize a particular objective function $f(v)$ are chosen. Thus, the flux balance analysis problem can be represented as a Linear Programming (LP) optimization problem:
\[
\begin{align*}
\max f(v) \\
\text{subject to } Sv = 0 \\
\text{and } v_{\min,j} \leq v_j \leq v_{\max,j}
\end{align*}
\]

Solving Equation 5 provides the maximized value \( f_{\max} \) of the objective function. Gurobi 8.0 optimization software was used to solve these LP problems.

**Flux Variability Analysis (FVA)**—Flux variability analysis allows for calculation of the minimum and maximum allowable fluxes through each metabolic reaction while still maintaining the maximum possible objective function value \( f_{\max} \):

\[
\min/\max v_j \text{ such that } f(v) = f_{\max}
\]

The average flux through each metabolic reaction was estimated by taking the mean of the minimum and maximum fluxes calculated from Equation 6.

**Objective Functions**—To maximize the production of a particular metabolite in the metabolic network, an artificial demand reaction can be added to the network, and the flux through this new objective function reaction can be maximized. This in turn maximizes the fluxes through other reactions throughout the metabolic network that produce the metabolite. FBA provides the objective value, i.e. the maximum amount of metabolite that can be produced, and FVA provides the fluxes through metabolic reactions which produce the metabolite. To maximize the reduction of a metabolite from its oxidized to reduced form, the objective function consists of an artificial demand reaction consisting of the oxidation of the metabolite - this in turn maximizes the fluxes through other metabolic reactions that reduce the metabolite.

For example, to maximize the reduction of NADP\(^+\) to NADPH in the cytosol, the objective function would be:

\[
\text{NADPH}[c] \rightarrow \text{NADP}^+[c] + \text{H}^+[c]
\]

To maximize the production of a metabolite in all cellular compartments, separate objective functions for each compartment with the same weight were simultaneously maximized. Table S5 lists the objective functions used in FBA and FVA. To ensure that biologically viable solutions were obtained, all models were checked to ensure that they were capable of producing physiological ATP levels typical of mammalian cells (1.0625 mmol gDW\(^{-1}\) hr\(^{-1}\)) while maximizing the imposed objective functions (Kilburn et al., 1969).

**Metabolic Network**—Recon 3D version 3.01 was used as the core metabolic network (Brunk et al., 2018). While the “full” version (Recon3D_301.mat) produced erroneous energy-generating cycles (e.g. producing ATP without uptake of any metabolic precursors), the “model” version (Recon3DModel_301.mat) was found not to be the largest subset of reactions without erroneous energy-generating cycles (additional reactions from the “full” version could be added). To this end, following the procedure outlined in Fritzemeier et al.,
reactions from the “full” version were sequentially added back into the “model” version as long as they did not result in erroneous energy-generating cycles (Fritzemeier et al., 2017).

To address missing and inaccurate redox-based reaction information within Recon3D, the following changes were made:

1. The reaction “FTHFDH” was split into two separate cytosolic and mitochondrial reactions. The cytosolic reaction is catalyzed by ALDH1L1, and the mitochondrial reaction is catalyzed by ALDH1L2.

   Originally:
   \[
   1 \text{10fthf}[c] + 1 \text{h2o}[c] + 1 \text{nadp}[c] \rightarrow 1 \text{co2}[c] + 1 \text{h}[c] + 1 \text{nadph}[c] + 1 \text{thf}[c]
   \]
   GPR: 10840.1 or 160428.1
   EC: 1.5.1.6

   Updated:
   \[
   1 \text{10fthf}[c] + 1 \text{h2o}[c] + 1 \text{nadp}[c] \rightarrow 1 \text{co2}[c] + 1 \text{h}[c] + 1 \text{nadph}[c] + 1 \text{thf}[c]
   \]
   GPR: 10840.1
   EC: 1.5.1.6

   \[
   1 \text{10fthf}[m] + 1 \text{h2o}[m] + 1 \text{nadp}[m] \rightarrow 1 \text{co2}[m] + 1 \text{h}[m] + 1 \text{nadph}[m] + 1 \text{thf}[m]
   \]
   GPR: 160428.1
   EC: 1.5.1.6

2. MTHFR was removed from the GPR associated with reaction “MTHFD”. MTHFR catalyzes the reaction converting 5,10-methylenetetrahydrofolate to 5-methyltetrahydrofolate, which is already included in Recon3D.

   Originally:
   \[
   1 \text{mlthf}[c] + 1 \text{nadp}[c] \leftrightarrow 1 \text{methf}[c] + 1 \text{nadph}[c]
   \]
   GPR: 4522.1 or 4524.1
   EC: 1.5.1.5

   Updated:
   \[
   1 \text{mlthf}[c] + 1 \text{nadp}[c] \leftrightarrow 1 \text{methf}[c] + 1 \text{nadph}[c]
   \]
   GPR: 4522.1
   EC: 1.5.1.5

3. Reactions catalyzed by isoforms of NADPH oxidase were added.

   Originally: None
Updated:

- \(1 \text{nadh} + 2 \text{o}_2 \rightarrow 2 \text{o}_2s + 1 \text{nadp} + 1 \text{h} \)
  
  GPR: 27035.1 or 1536.1 or 50508.1 or 79400.1
  
  EC: 1.6.3.1

- \(1 \text{nadh} + 2 \text{o}_2 \rightarrow 2 \text{o}_2s + 1 \text{nadp} + 1 \text{h} \)
  
  GPR: 50507.1
  
  EC: 1.6.3.1

4. PRDX1 and PRDX2 were removed from the GPR associated with reaction “GTHP”. Glutathione peroxidase involves the reduction of hydrogen peroxide by glutathione, which does not involve peroxiredoxins. Peroxiredoxin reactions are added to Recon3D (see #7)

Originally:

- \(1 \text{h}_2\text{o}_2 + 2 \text{gthrd} \rightarrow 2 \text{h}_2\text{o} + 1 \text{gthox} \)
  
  GPR: 7001.3 or 5052.3 or 2877.1 or 2876.2 or 5052.2 or 2876.1 or 5052.1 or 7001.1 or 2879.1 or 7001.2
  
  EC: 1.11.1.9

Updated:

- \(1 \text{h}_2\text{o}_2 + 2 \text{gthrd} \rightarrow 2 \text{h}_2\text{o} + 1 \text{gthox} \)
  
  GPR: 2877.1 or 2876.2 or 2876.1 or 2879.1
  
  EC: 1.11.1.9

5. Similarly to #4, PRDX3 was removed from the GPR associated with reaction “GTHPm”.

Originally:

- \(1 \text{h}_2\text{o}_2 + 2 \text{gthrd} \rightarrow 2 \text{h}_2\text{o} + 1 \text{gthox} \)
  
  GPR: 10935.1 or 10935.2 or 2879.1 or 2876.1
  
  EC: 1.11.1.9

Updated:

- \(1 \text{h}_2\text{o}_2 + 2 \text{gthrd} \rightarrow 2 \text{h}_2\text{o} + 1 \text{gthox} \)
  
  GPR: 2879.1 or 2876.1
  
  EC: 1.11.1.9
6. PRDX6 was removed from the GPR’s associated with the following reactions: HMR_0960, HMR_0963, HMR_0988, HMR_2441, HMR_1048, HMR_1049. These are all glutathione peroxidase reactions, which PRDX6 is not involved in.

7. Reactions catalyzed by isoforms of peroxiredoxin were added:

   Originally: None

   Updated:

   • $1 \text{h}_2\text{O}_2[c] + 1 \text{trd}[c] \rightarrow 2 \text{h}_2\text{O}[c] + 1 \text{trdox}[c]$  
     GPR: 5052.1 or 5052.2 or 5052.3 or 7001.1 or 7001.2 or 7001.3  
     EC: 1.11.1.15
   • $1 \text{h}_2\text{O}_2[m] + 1 \text{trd}[m] \rightarrow 2 \text{h}_2\text{O}[m] + 1 \text{trdox}[m]$  
     GPR: 10935.1 or 10935.2  
     EC: 1.11.1.15

8. The oxidation and glutathionylation of protein thiol groups, as well as their reduction by glutaredoxin and thioredoxin, were added:

   Originally: None

   Updated:

   • $1 \text{h}_2\text{O}_2[c] + 1 \text{Pr-SH}[c] + 1 \text{gthrd}[c] \rightarrow 2 \text{h}_2\text{O}[c] + 1 \text{Pr-SSG}[c]$  
     GPR: -  
     EC: -
   • $1 \text{h}_2\text{O}_2[m] + 1 \text{Pr-SH}[m] + 1 \text{gthrd}[m] \rightarrow 2 \text{h}_2\text{O}[m] + 1 \text{Pr-SSG}[m]$  
     GPR: -  
     EC: -
   • $1 \text{Pr-SSG}[c] + 1 \text{gthrd}[c] \rightarrow 1 \text{Pr-SH}[c] + 1 \text{gthox}[c]$  
     GPR: 2745.1  
     EC: 1.20.4.1
   • $1 \text{Pr-SSG}[m] + 1 \text{gthrd}[m] \rightarrow 1 \text{Pr-SH}[m] + 1 \text{gthox}[m]$  
     GPR: 2745.1  
     EC: 1.20.4.1
   • $1 \text{h}_2\text{O}_2[c] + 1 \text{Pr-SH}[c] \rightarrow 2 \text{h}_2\text{O}[c] + 1 \text{Pr-SS}[c]$  
     GPR: -  
     EC: -
   • $1 \text{h}_2\text{O}_2[m] + 1 \text{Pr-SH}[m] \rightarrow 2 \text{h}_2\text{O}[m] + 1 \text{Pr-SS}[m]$  
     GPR: -
9. The reliance of catalase enzyme on NADPH to prevent oxidation of its active site by H$_2$O$_2$ was incorporated (Kirkman et al., 1999). Kirkman et al. measured rates of H$_2$O$_2$ formation and corresponding rates of NADPH oxidation in mixtures of catalase and varying concentrations of glucose oxidase (Kirkman et al., 1987). If the rate of H$_2$O$_2$ formation by glucose oxidase is assumed to equal the rate of H$_2$O$_2$ clearance by catalase, then the slope of NADPH oxidation vs. H$_2$O$_2$ formation gives the amount of NADPH oxidized by catalase per molecule of H$_2$O$_2$ cleared. Linear regression of data from Kirkman et al. shows that for every 1 molecule of H$_2$O$_2$ cleared by catalase, 0.0641 molecules of NADPH are oxidized; thus, the chemical equation for catalase activity is:

\[
2 \text{H}_2\text{O}_2 + 0.1282 \text{NADPH} \rightarrow 2 \text{H}_2\text{O} + 1 \text{O}_2 + 0.1282 \text{NADP}^+ + 0.1282 \text{H}^+ \tag{8}
\]

This change was made for catalase reactions in the cytosol, mitochondria, peroxisome, and endoplasmic reticulum.

10. The neomorphic 2-hydroxyglutarate-producing reaction catalyzed by mutant IDH1 was added. Reactions for 2-hydroxyglutarate export were also added.

*Originally:* None

*Updated:*

- $1 \text{akg}[c] + 1 \text{nadph}[c] \rightarrow 1 \text{M00653}[c] + 1 \text{nadp}[c]
  
  GPR: 3417.1
  EC: 1.1.1.42

- $1 \text{M00653}[c] \rightarrow 1 \text{M00653}[c]
  
  GPR: None
  EC: None

- $1 \text{M00653}[e] \rightarrow
  
  GPR: None
  EC: None
Because transcriptomic and proteomic data from TCGA were available at the individual gene level but not at the individual isoform level, all isoforms for each individual gene within Recon3D were combined.

Other changes made to Recon3D include:

- Removing the following duplicate reactions (when another reaction exists where the stoichiometric vector is an exact multiple of these duplicate reactions): “HMR_7257”, “PEHSFABPe”, “G6PDH2c”, “GNDe”, “PGLc”, “RPeC”
- Removing EC numbers from all transport reactions, since many of these were incorrect.
- Removing the following genes and any associations with them within Recon3D reactions, since information about these genes could not be found: NCBI Gene ID’s: 0, 100507855, 8041
- Removing reactions within the following subsystems, as this was necessary to prevent de novo production of essential amino acids: “Protein assembly”, “Protein degradation”, “Protein modification”, “Protein formation”
- Removing reactions in the “Bile acid synthesis” subsystem, as these are not expected to be active in most tumors
- Removing NQO1 futile cycle reactions, as these unconstrained reactions resulted in erroneous predictions of NADPH production: “NADPQNOXR”, “NADQNOXR”, “HMR_9538”, “HMR_9722”
- Removing oxalosuccinic acid-producing IDH1 reactions, as these overlapped with α-ketoglutarate-producing IDH1 reactions: “r0423”, “r0424”, “r0425”, “r0422”

**Model Flux Constraints**—Upper flux bounds for reactions within Recon3D with both associated GPR rules and EC numbers were constrained using the Michaelis-Menten $V_{\text{max}}$ parameter:

$$v_{\text{max},j} = k_{\text{cat}}[E]$$  \(9\)

where $k_{\text{cat}}$ and [E] are the turnover number (units of hr⁻¹) and abundance (units of mmol gDW⁻¹) of the associated enzyme catalyzing that reaction, respectively. Equation 9 was used to set upper flux bounds $v_{\text{max},j}$ as well as lower flux bounds $v_{\text{min},j}$ for reversible reactions (where $v_{\text{min},j} < 0$). See sections “Enzyme Abundance Calculation” and “Turnover Number Calculation” for more information.

**Enzyme Abundance Calculation**—RNA-Seq gene expression data from TCGA patient tumors was obtained from Rahman et al.’s alternative preprocessing method (GEO: GSE62944) (Rahman et al., 2015). Data from this preprocessing method showed fewer
missing values, more consistent expression between replicates, and improved prediction of biological pathway activity compared to the original TCGA pipeline.

Enzyme abundances within individual samples were predicted from sample gene expression data using the protein prediction pipeline shown in Figure S1A. The approach described below relies only on gene expression data from the individual sample of interest and not data from other samples included within the dataset, improving the reproducibility of this method as well as allowing for new samples to be analyzed post-hoc. The individual steps in the pipeline are explained herein:

1. **Schwanhäusser et al.** – By measuring the transcription, translation, and degradation rates of individual mRNA’s and proteins in NIH3T3 cells, Schwanhäusser et al. developed ODE models relating mRNA and protein abundances per cell that encompass 955/3268 (29.2%) genes in Recon3D (Schwanhausser et al., 2011). Their model also took into account proteins with half-lives longer than the length of the cell cycle, and how this would impact measurements of protein abundance. The authors demonstrated that these rate constants and models could be applied to accurately predict protein abundances in other samples including cancer cell lines. Using this model, gene expression values (transcripts per kilobase-million (TPM)-normalized) were converted to protein abundance values (parts per million (PPM)-normalized) for the 955 Recon3D genes included.

2. **Linear regression** – For each sample, a linear regression of predicted protein abundance values using the Schwanhäusser et al. method versus measured gene expression values of corresponding genes is performed (example is shown in Figure S1B). This regression model is used to predict the abundance of proteins where an associated gene expression value is available but not corresponding model/parameter values in the Schwanhäusser et al. method.

3. **PaxDB** – If an individual gene expression value is missing for a particular sample, the harmonic mean of abundance values for that associated protein across all *Homo sapiens* samples within the PaxDB database is taken (Wang et al., 2015).

If the associated protein is not available in the PaxDB database, the harmonic mean of abundance values for all proteins across all *Homo sapiens* samples is taken. Imposing the same values for all samples with missing gene expression values acts to prevent artificial differences in FBA model predictions between samples due to missing data. The harmonic mean of PaxDB values is taken since (1) the arithmetic mean tends to overestimate representative values of positively-skewed distributions (as seen with protein abundance values); and (2) the smaller protein abundance estimate of the harmonic mean well characterizes the fact that missing RNA-Seq values are usually at least in part due to low actual mRNA expression.

For genes with associated enzymes in multiple cellular compartments (e.g. catalase, which localizes to the cytosol, mitochondria, peroxisome, and endoplasmic reticulum), the
predicted protein expression value was divided into each cellular compartment with weights proportional to the COMPARTMENTS value for the cellular compartment in which the metabolic reaction takes place (Binder et al., 2014). These COMPARTMENTS values provide a score representing the confidence that a particular protein is found within a particular cellular compartment, using both experimental and computational evidence.

Validation of this pipeline was performed to ensure (1) that the predicted enzyme abundance values correlated better with experimental protein expression data compared to gene expression data; and (2) that the predicted enzyme abundance values were at physiological orders of magnitude. Normalized RPPA experimental protein expression values from TCGA samples were available for 30 genes/proteins in Recon3D (Weinstein et al., 2013). The correlation between experimental enzyme abundance and predicted enzyme abundance values ($R^2_{\text{PredictedAbundance}}$) was greater than the correlation between experimental enzyme abundance and gene expression values ($R^2_{\text{Gene}}$) for 18/30 genes (60%; Figure S1C–E). Additionally, the largest change in correlation for genes where $R^2_{\text{PredictedAbundance}} < R^2_{\text{Gene}}$ was less than 0.004, indicating that the $R^2_{\text{PredictedAbundance}}$ was not substantially lower than the $R^2_{\text{Gene}}$ for any of the 30 genes.

Because experimental non-normalized protein abundance values were not available from TCGA samples for validation, experimentally-measured protein abundances from the NCI-60 panel of cancer cell lines were used to analyze the improvement in correlation to experimental data and physiological accuracy of predicted abundances on a genome scale. Experimental NCI-60 abundance values showed greater correlation to mean values of predicted enzyme abundances across TCGA samples ($R^2 = 0.450$) compared to TCGA gene expression values ($R^2 = 0.314$) (Figure S1F–G). Additionally, the order of magnitude of predicted abundance values matched well with experimental values for individual genes/proteins over the majority of orders of magnitude. The match to experimental values was greatest for enzyme abundances that were calculated at the first step of the pipeline (Schwanhäusser et al. method; $R^2 = 0.586$) compared to abundances calculated at later steps ($R^2 = 0.286$), indicating the necessity for future transcription, translation, and degradation rate measurements to be conducted for genes/proteins not currently covered in the Schwanhäusser et al. method (Figure S1H–I).

**Turnover Number Calculation**—Enzyme turnover numbers for associated reactions within Recon3D were determined using available experimental data from the BRENDA database API (Schomburg et al., 2004). Because the BRENDA database does not contain data for all Recon3D enzymes with the exact same enzyme commission (EC) number and substrate measured using human enzymes at 37°C and the correct cellular compartment-specific pH, a pipeline was developed where the most physiologically-accurate turnover numbers for Recon3D metabolic reactions are identified (Figure S2). For an individual enzyme-catalyzed reaction within Recon3D, the associated turnover number was determined as close to the start of the pipeline as data was available within BRENDA. Values determined towards the start of the pipeline are more accurate, but less turnover number data with this greater accuracy is available within BRENDA. In determining turnover number accuracy, priority was given in the order of (1) using the correct EC number (versus using data from EC numbers that match only the first 3, 2, or 1 digits with that of the Recon3D
reaction); (2) using the correct substrate as given in the Recon3D reaction (versus using data from any available substrate); (3) using turnover number data taken from human enzymes (versus using data from any available organism); and (4) using turnover number data taken at physiological temperature and pH (versus using data near but not at physiological temperature and pH). At a particular step in the pipeline, if multiple turnover number values that met the pipeline criteria were available, a weighted average of these values was taken, with weights equal to the Tanimoto coefficient (a measure of molecular similarity; greater coefficient signifies greater similarity between two molecules) between the desired substrate in the Recon3D reaction and the substrate in BRENDA for which the associated turnover number was determined.

**Mutation Data**—The effect of single nucleotide polymorphisms (SNP’s) located in Recon3D genes on the catalytic activity of the associated metabolic enzyme was estimated using the Envision computational platform developed by Gray et al (Gray et al., 2018). Envision leverages large-scale experimental mutagenesis datasets to predict a score representing the effect of single amino acid changes on protein function; a score less than 1 signifies a loss-of-function, while a score greater than 1 signifies a gain-of-function. These quantitative scores were shown to correlate better with experimentally-measured percent changes in protein activity after mutation than previously-developed methods including SNAP2 and EVmutation. For each TCGA patient with mutation data available (752 of the 915 patients analyzed), the turnover number for a particular Recon3D reaction was multiplied by the Envision score for any SNP’s located in the associated gene. Envision scores used to adjust tumor-specific enzyme catalytic rates are likely very conservative estimates of the impact of mutations on enzymatic function, and exclusion of genomic data did not affect the majority of objective value calculations, nor comparisons between radiation-sensitive and -resistant tumor models. Distributions of mutation numbers and Envision scores across the TCGA dataset are provided in Figure S28.

**IDH1 mutations**—Particular mutations in isocitrate dehydrogenase 1 (IDH1), predominantly at amino acid R132, result in both loss of function of its normal NADPH-generating activity, as well as gain of function of a neomorphic NADPH-consuming activity that produces the oncometabolite 2-hydroxyglutarate (Dang et al., 2009). This change in function was implemented by adding the IDH1-catalyzed neomorphic reaction to Recon3D and imposing turnover numbers of the normal and neomorphic reactions measured by Avellaneda Matteo et al. based on the IDH1 mutation status of each tumor (Table S6) (Avellaneda Matteo et al., 2017).

**Thermodynamic Constraints**—Only metabolic reactions with a negative change in Gibbs free energy (ΔG) can carry a non-zero net flux. Implementing thermodynamic constraints prevents fluxes through these thermodynamically infeasible reactions; additionally, this limits the formation of thermodynamically infeasible loops that carry net fluxes around closed cycles in the metabolic network (Schellenberger et al., 2011). To implement thermodynamic constraints on FBA models, the minimum and maximum transformed change in Gibbs free energy (ΔG’) of Recon3D reactions were obtained from the Virtual Metabolic Human (VMH) database (Noor et al., 2013). Based on the component
contribution method, the VMH database predicts the standard change in Gibbs free energy (ΔG°) value for each metabolic reaction. A confidence interval (minimum and maximum values) on the ΔG° value for each reaction is obtained by using a default concentration range of 10^{-7} M to 10^{-2} M for all metabolites. Transformed ΔG° (ΔG') ranges are then obtained by correcting for a temperature of 310.15 K, ionic strength of 0.15 M, and compartment-specific pH and electrical potential values provided in Haraldsdottir et al (Haraldsdottir et al., 2012). For all reactions where both the minimum and maximum ΔG' values were greater than zero (and for all reversible reactions where both the minimum and maximum ΔG' values were less than zero in the reverse direction), the maximum flux v_{max,j} (or v_{min,j} for reversible reactions) through that reaction was set to zero. 67 forward-direction reactions and 498 reverse-direction reactions were blocked by thermodynamic constraints.

**Media Constraints**—To simulate experimental cell culture conditions, the modeled external compartment contained all metabolites found in RPMI-1640 cell culture media as well as fetal bovine serum (Price and Gregory, 1982). The external compartment also contained water, hydrogen ions, hydroxide ions, oxygen, and carbon dioxide. Exchange reactions (metabolite uptake reactions) for metabolites in this external compartment were unconstrained in FBA models. For metabolites not in this external compartment, no influx was allowed through corresponding exchange reactions. Metabolite efflux was unconstrained for all Recon3D metabolites.

**Analysis of Constraints:** Inclusion of protein abundance estimates, genomic data, and thermodynamic constraints were analyzed individually across the TCGA dataset for effects on flux calculations (Figures S20–S26).

**Radiation Response Classification**—Radiation responses for TCGA tumors were obtained from the GDC Data Portal (see “Code + Models/data/clinical/_data_/input/TCGA/manifest.txt” in our “kemplab/FBA-pipeline” GitHub repository for a list of GDC Data Portal files containing radiation response information). TCGA provides the RECIST classification of each patient tumor response to radiation therapy:

- **Complete Response (CR):** Complete resolution of all target lesions
- **Partial Response (PR):** At least 30% decrease in tumor size (measured as the sum of the largest diameter of all target lesions, using the baseline sum of largest diameters as reference)
- **Stable Disease (SD):** Neither sufficient decrease in tumor size to meet criteria of PR, nor sufficient increase to meet criteria of PD
- **Progressive Disease (PD):** At least 20% increase in tumor size (using the smallest sum of largest diameters since treatment started as reference), or the appearance of new target lesions

All TCGA tumor samples with transcriptomic data and a recorded radiation response were included in the analysis. Patient tumors classified as either CR or PR were included in the radiation-sensitive group, and those classified as either SD or PD were included in the
radiation-resistant group. If a patient received multiple treatments of radiation therapy, only the first measured response was used for classification.

**Controlling for Patient/Tumor Covariates**—To control for patient and tumor covariates when comparing FBA predictions between radiation-sensitive and -resistant tumor models, propensity scores were calculated for each patient tumor using covariate values. The following factors were included as covariates:

- **Patient factors**: age, gender, race, Karnofsky performance status
- **Tumor factors**: cancer type, histological subtype, tumor location, tumor grade, clinical stage/TNM, pathological stage/TNM

Propensity scores were calculated by performing a logistic regression classification, predicting radiation response (sensitive or resistant) from the covariates above (Pedregosa et al., 2011). The propensity score for each patient is equal to the predicted probability of radiation resistance from the logistic regression classifier. These propensity scores were then used to correct comparisons of model predictions between radiation-sensitive and -resistant classes by performing analysis of covariance (ANCOVA), with the dependent variable being the model prediction, the between factor being radiation response (sensitive or resistant), and the covariate being propensity score for each patient (Vallat, 2018). ANCOVA results in a p-value for the significance of radiation response in determining the value of the dependent variable (model prediction), while correcting for the propensity score (covariates). These p-values were used to report the statistical significance of FBA-predicted comparisons between radiation-sensitive and -resistant tumor models. The coefficients of the logistic regression model indicate that the major confounding variables for radiation response are the clinical and pathologic stage of the tumor, rather than the cohort (cancer type) or other patient-specific factors (Figure S29).

**siRNA Transfections**—Cells were seeded in 24-well plates at $5 \times 10^4$ cells per well for glutathione half-cell potential measurements, or 96-well plates at $8 \times 10^3$ cells per well for glucose oxidase response measurements. 24 hours after seeding, cells were transfected using the N-TER Nanoparticle siRNA Transfection System (Sigma-Aldrich; Cat#N2913) at a final siRNA concentration of 50 nM with serum-free medium for 4 hours; afterwards, an equal volume of 2x (20%) FBS-containing medium was added to each well. Negative controls consisted of transfection with the MISSION siRNA Universal Negative Control (Sigma-Aldrich; Cat#SIC001; Lot#WDAA1199). Predesigned MISSION siRNA’s targeting individual genes were ordered from Sigma-Aldrich; the top 3 rated siRNA’s based on expected efficacy for each gene target were pooled together and transfected concurrently, except for GLUD1 and GLUD2, where only 2 and 1 siRNA’s were available, respectively (Table S7). Figure S30 shows the knockdown efficiency of siRNA transfections in each cell line with GAPDH siRNA compared to no siRNA (just N-TER transfection reagent) and negative control siRNA by Western blot (GAPDH antibody: Cell Signaling Technology, Cat#2118; Tubulin antibody: Thermo Fisher Scientific, Cat#62204). Experiments were performed 24 hours after transfection.
**E<sub>hc</sub> GSH/GSSG Measurements**—The protocol and reagents used for measurement of glutathione half-cell potential were adapted from Rahman et al. (Rahman et al., 2006). KPE buffer was made by combining 16mL of solution A (6.8g KH<sub>2</sub>PO<sub>4</sub> (Sigma-Aldrich; Cat#P5655; CAS: 7778–77-0) in 500mL dH<sub>2</sub>O), 84mL of solution B (8.5g K<sub>2</sub>HPO<sub>4</sub> (Sigma-Aldrich; Cat#P8281; CAS: 7758–11-4) in 500mL dH<sub>2</sub>O), and 0.327g EDTA sodium salt (Sigma-Aldrich; Cat#E5134; CAS: 6381–92-6). After removal of cell culture media, 150μL of extraction buffer (0.1% Triton-X100 (Sigma-Aldrich; Cat#X100; CAS: 9002–93-1) and 0.6% sulfosalicylic acid (Sigma-Aldrich; Cat#S2130; CAS: 5965–83-3) in KPE) was added to each well of a 24-well plate containing samples of interest. Plates were shaken at 800rpm for 10min to promote cell lysis. For each well, 100μL of lysate was taken for GSSG measurement, and 20μL was taken for GSSG+GSH measurement.

For GSSG+GSH measurement, 20μL samples were placed in a clear 96-well plate. 20μL of GSSG (Sigma-Aldrich; Cat#G6654; CAS: 27025–41-8) and GSH (Sigma-Aldrich; Cat#S4251; CAS: 70–18-8) standards at concentrations of 2e-2, 1e-2, 5e-3, 2.5e-3, 1e-3, 5e-4, 2.5e-4, 1e-4, 5e-5, 2.5e-5, 1e-5, and 0 mg/mL were placed in separate rows of the 96-well plate. 120μL of 1:1 DTNB:GR solution (DTNB: 2mg 5,5'-dithiobis(2-nitrobenzoic acid) (Sigma-Aldrich; Cat#D8130; CAS: 69–78-3) in 3mL KPE; GR: 40μL of glutathione reductase enzyme (Sigma-Aldrich; Cat#G3664; CAS: 9001–48-3) (250 units/mL) in 3mL KPE) was added to each well. 30sec later, 60μL of NADPH solution (2mg of β-NADPH (Sigma-Aldrich; Cat#N7505; CAS: 2646–71-1 (anhydrous)) in 3mL KPE) was added to each well. Immediately after, absorbance at 412nm in each well was measured every 31sec for a period of 5min10sec (11 measurements) using a BioTek Synergy 4 plate reader. All sample and standard values were background subtracted. To determine the concentration of GSSG+GSH in each sample (mg/mL), the slope of absorbance vs. time from each sample was compared to the average slope of the GSSG and GSH standard concentrations.

For GSSG measurement, 100μL samples as well as 100μL of GSSG and GSH standards were placed in a 96-well plate. 10μL of 2VP solution (2-vinylpyridine (Sigma-Aldrich; Cat#132292; CAS: 100–69-6) diluted 1:50 in KPE) was added to each well. 60min later, 10μL of triethanolamine solution (triethanolamine (Sigma-Aldrich; Cat#T58300; CAS: 102–71-6) diluted 1:10 in KPE) was added to each well. 10min later, 20μL from each well was transferred to a new clear 96-well plate. GSSG was then measured analogously to GSSG+GSH. GSSG measurements were multiplied by 1.2 to account for the dilution of the original 100μL sample with 10μL of 2VP solution and 10μL of triethanolamine solution.

To determine the concentration of GSH (mg/mL) for each sample, the measured concentration of GSSG was subtracted from the measured concentration of GSSG+GSH. To convert lysate concentrations (mg/mL) to intracellular concentrations (M), the following conversion was used:

$$[M] = \frac{[mg/mL] \times V_{sample}}{MM \times V_{cell} \times n_{cells}} \quad (10)$$
where \( MM \) is the molar mass of either GSH or GSSG (g/mol), \( V_{\text{sample}} \) is the volume of sample (20μL), \( V_{\text{cell}} \) is the estimated volume of each cell (1 pL), and \( n_{\text{cells}} \) is the number of cells per sample (5 \( \times \) 10^4).

The glutathione half-cell potential \( E_{hc} \) is calculated using the Nernst equation:

\[
E_{hc} = E_{hc}^o - \frac{RT}{ZF} \ln \left( \frac{[\text{GSH}]}{[\text{GSSG}]} \right)
\]  

(11)

where \( E_{hc}^o \) is the standard half-cell potential at pH 7.4 (−264 mV), \( R \) is the universal gas constant (8.314 J/K/mol), \( T \) is the temperature (310.15 K), \( z \) is the number of electrons (2), and \( F \) is the Faraday constant (96485 C/mol = 96.485 J/mV/mol).

**Glucose Oxidase Response Measurements**—Prior to seeding, cells were stained with CellTracker Red CMTPX Dye (Thermo Fisher Scientific; Cat#C34552) for 30 min. 8 \( \times \) 10^3 cells per well were seeded in white-sided clear-bottom 96-well plates. 24 hr after transfection and prior to treatment with glucose oxidase, CellTracker fluorescence measurements in each well were taken at Ex 577nm, Em 602nm using a BioTek Synergy 4 plate reader as a proxy for number of cells per well. An average of 10 fluorescence measurements was taken, and measurements were background subtracted. Samples were then treated with 10 mU/mL glucose oxidase enzyme (Sigma-Aldrich; Cat#G7141) diluted in 100μL cell culture medium (RPMI-1640 + 10% FBS) for 2 hr. The rate of \( H_2O_2 \) production from 10 mU/mL glucose oxidase is calculated to be 10 μM/min:

\[
10 \text{ mU/mL} \left( \frac{1 \times 10^{-6} \text{ mol } H_2O_2}{\text{min}} \right) = 1 \times 10^{-5} \text{ M/min} = 10 \mu\text{M/min}
\]  

(12)

After 2 hr treatment, 100μL of CellTiter-Glo 2.0 Cell Viability Assay (Promega; Cat#G9241), which measures cellular ATP levels, was added to each well. Plates were shaken at 800 rpm for 2 min to promote cell lysis, and then incubated at room temperature for 10 min. Luminescence measurements were then taken using a BioTek Synergy 4 plate reader and background subtracted. Cell viability normalized by cell count was measured by dividing the luminescence measurement by the average CellTracker fluorescence measurement.

The relative effect of glucose oxidase on cell viability between siRNA-treated and negative controls for a particular cell line was calculated using the Bliss independence score (Equation 1) where each of the 4 values (siRNA 10 mU/mL, siRNA 0 mU/mL, control 10 mU/mL, control 0 mU/mL) represents the luminescence cell viability measurement normalized by fluorescence cell count measurement as explained above. Values < 0 indicate synergistic effects between siRNA and glucose oxidase treatments for decreasing cell viability, whereas values > 0 indicate antagonistic effects between siRNA and glucose oxidase treatments. Thus, BlissSens - BlissRes > 0 for siRNA gene knockdowns causing a greater decrease in cell viability with glucose oxidase treatment in the radiation-resistant cell line compared to the radiation-sensitive cell line.
QUANTIFICATION AND STATISTICAL ANALYSIS

- Comparison of gene expression values, as well as FBA-predicted objective values, reactions fluxes, and gene knockout effects between radiation-sensitive and -resistant TCGA samples (Fig. 2A, 2D, 2F, 3B, 4A, 4C, S3, S4, S5, S7, S11, S12, S14, S21, S22A, S24, S26, S27)
  - n: number of individual TCGA samples within each class. n = 716 radiation-sensitive samples, n = 199 radiation-resistant samples.
  - Boxplots: box = 25th, 50th, and 75th percentiles, whiskers = 1.5 times the interquartile range.
  - Statistical test: Analysis of covariance (ANCOVA).

- Comparison of experimental glutathione half-cell potentials between radiation-sensitive and -resistant cell lines (Fig. 2C)
  - n: number of biological replicates performed for each cell line. n = 12 for all cell lines except rSCC-61, where n = 10 (erroneously-negative GSSG values were obtained for 2/12 samples due to being below the limit of detection, and these were removed from the analysis).
  - Boxplots: box = 25th, 50th, and 75th percentiles, whiskers = 1.5 times the interquartile range.
  - Statistical test: 2-sided 2-sample Welch’s t-test without assumption of equal population variance.

- Comparison of gene knockout effects between radiation-sensitive and -resistant FBA tumor models (Fig. 3C, 3D, 4D, S22B)
  - n: number of individual FBA tumor models in each class. n = 716 radiation-sensitive tumor models, n = 199 radiation-resistant tumor models.
  - Statistical test: Analysis of covariance (ANCOVA). p-values for each of the 3,268 simulated gene knockouts performed were subsequently adjusted for multiple hypothesis testing using the Benjamini-Hochberg procedure for independent tests.

- Comparison of model-predicted and experimentally-validated effect of gene knockdown on GSH production (Fig. 3F)
  - n1: number of biological replicates performed for each cell line, for each gene knockdown and negative control. n1 = 3 for all cases
  - n2: number of cell line pairs for comparing to model predictions. n2 = 3
  - Statistical test: 2-sided 1-sample t-test with null hypothesis population mean = 0

- Comparison of experimental glucose oxidase response between radiation-sensitive and -resistant cell lines (Fig. 4B)
n: number of biological replicates performed for each cell line, for both 0 mU/mL and 10 mU/mL glucose oxidase treatment. Outliers were automatically detected and removed as determined by Grubbs’ 2-tailed test with $\alpha=0.05$. n = 5 for GBM and HNSC cell lines, except for rSCC-61 10 mU/mL (n = 4) due to outlier removal. n = 10 for BRCA cell lines, except for NQO1(−) 0 mU/mL (n = 9) due to outlier removal.

Bar chart: mean ± 1 standard error.

Statistical test: 2-sided 2-sample t-test from mean, standard deviation, and number of observations from both samples

Comparison of model-predicted and experimentally-validated effect of gene knockdown on H$_2$O$_2$ response (Fig. 4F)

$n_1$: number of biological replicates performed for each cell line, for both glucose oxidase treatments, for each gene knockdown and negative control. Outliers were automatically detected and removed as determined by Grubbs’ 2-tailed test with $\alpha=0.05$. $n_1 = 5$ for GBM and HNSC cell lines, except for 1) M059K 10 mU/mL TKT KD (n=4) and 2) rSCC-61 10 mU/mL PGAM2 KD (n=4) due to outlier removal. $n_1 = 10$ for BRCA cell lines, except for 1) NQO1(−) 0 mU/mL PGD KD (n=9) and 2) NQO1(−) 0 mU/mL TKT PD (n=9) due to outlier removal.

$n_2$: number of cell line pairs for comparing to model predictions. $n_2 = 3$

Statistical test: 2-sided 1-sample t-test with null hypothesis population mean = 0

Correlation between patient clinical factors and FVA-predicted metabolic fluxes in radiation-resistant tumor models (Fig. 5A, S16, S17, S18)

$n$: number of radiation-resistant tumor models. $n = 199$

Statistical test:

♦ Numerical factors: univariate regression

♦ Categorical factors: 1-way ANOVA

Regression of metabolic reaction flux versus gene expression value for NADPH-generating reactions/genes (Fig. S20)

$n$: number of NADPH-generating reactions/genes. $n = 12$

Statistical test: univariate regression

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Highlights

• Multi-omics genome-scale models provide a pan-cancer analysis of redox metabolism
• Radiation-resistant tumors produce elevated levels of reduced redox cofactors
• Rerouted metabolic fluxes increase ROS clearance in radiation-resistant tumors
• Models illustrate redox metabolic heterogeneity across similar clinical phenotypes
Context and Significance

Although many cancer patients receive radiation therapy, certain patient tumors continue to progress following radiation. Tumor production of antioxidant molecules may help explain this lack of response. In this study, the authors create computational models to show that tumors which respond poorly to radiation are capable of increasing antioxidant production to protect against damaging effects of radiation treatment. Specific drivers of antioxidant production in radiation-resistant tumors were identified, and these computational predictions were validated experimentally by utilizing a panel of cancer cells with varying radiation sensitivities. Finally, this approach identified the most important antioxidant pathways in individual patient tumors, which could lead to personalized treatment strategies for improving cancer patient response to radiation therapy.
Figure 1. An automated bioinformatics pipeline integrates multi-omics data into personalized FBA models of TCGA patient tumors.

(A) Implementation of flux balance analysis (FBA), including utilization of a stoichiometric representation of the Recon3D human metabolic network reconstruction, application of reaction constraints to obtain a solution space of flux values, and maximization of an objective function within this valid solution space. (B) Flux balance analysis (FBA) calculates the objective value, i.e., the maximum value of the objective function. Flux variability analysis (FVA) calculates the minimum and maximum possible fluxes through each metabolic reaction while maintaining the objective function at its maximum value. (C) Pipeline for integrating multi-omics data from The Cancer Genome Atlas (TCGA) and publically-available repositories into personalized FBA models of TCGA patient tumors. (D) Classification of TCGA patient tumors into radiation-sensitive and -resistant classes based on observed decrease/increase in tumor size following radiation therapy. See also Figures S1–S2.
Figure 2. Radiation-resistant tumors display compartmental differences in redox metabolic fluxes compared to radiation-sensitive tumors.

(A) Comparison of FBA-predicted production of reduced NADPH, NADH, and GSH between radiation-sensitive and -resistant TCGA tumors. (B) Matched pairs of radiation-sensitive and -resistant cell lines across multiple cancer types for experimental validation. (C) Experimentally-measured glutathione half-cell potential in cancer cell lines. (D) Comparison of FBA-predicted cytosolic and mitochondrial production of reduced NADPH. (E) Major NADPH-producing reactions with their associated cellular compartments and metabolic pathways. (F) FVA-predicted fluxes through major NADPH-producing reactions. IDH1 fluxes were separated between tumors with IDH1 R132 mutations (left) and wild-type IDH1 tumors (right). Reaction names are colored based on cellular compartment. (G)
Hierarchical clustering of FVA-predicted fluxes based on TCGA patient tumor (rows) and NADPH-producing reaction (columns). Values are the Z-score of reaction fluxes across all tumors for each individual reaction. ns: not significant, *: \( p \leq 0.05 \), **: \( p \leq 0.01 \), ***: \( p \leq 0.001 \), ****: \( p \leq 0.0001 \). Boxplots: box = 25th, 50th, and 75th percentiles, whiskers = 1.5 times the interquartile range. See also Figures S3–S8, Table S1.
Figure 3. Simulated genome-wide knockout screen identifies targets of redox metabolism in radiation-resistant cancers.

(A) Schematic comparing maximum total NADPH production between WT (left) and G6PD-knockout (right) models for individual patient tumors. Note that for reactions with more than one associated gene (e.g. GLUD1/2 reaction), only one gene is knocked out at a time. (B) Effect of simulated knockout of each individual gene in Recon3D on total NADPH production in TCGA tumors. Values are the ratio of total NADPH production after versus before knockout. Genes are rank ordered based on increasing mean KO/WT ratio (decreasing gene knockout effect) across all tumor models. Outset: KO/WT ratios are averaged across all tumor models. Inset: For the top 15 genes, KO/WT ratios from individual patient tumor models are shown, along with the comparison between radiation-sensitive and -resistant cohorts. (C) Volcano plot comparing the effect of each simulated gene knockout (individual dot) on total NADPH production between radiation-sensitive and -resistant
tumors. X-axis: $\log_2(\text{Resistant}/\text{Sensitive})$, where “Resistant” equals the mean $(\text{WT-KO})/\text{WT}$ ratio in radiation-resistant tumors, and “Sensitive” equals the mean $(\text{WT-KO})/\text{WT}$ ratio in radiation-sensitive tumors; values $< 0$ (green dot on the left of the dotted line) signify knockouts with greater effects on total NADPH production in radiation-sensitive tumors, whereas values $> 0$ (red dot on the right of the dotted line) signify knockouts with greater effects on total NADPH production in radiation-resistant tumors. Y-axis: statistical significance (false discovery rate-adjusted p-values based on the Benjamini-Hochberg procedure) comparing knockout effects between radiation-sensitive and -resistant tumors; values above the dotted line (FDR-adjusted p-value $\leq 0.05$) are statistically-significant. The size of each dot is proportional to the overall effect size (mean $(\text{WT-KO})/\text{WT}$ ratio across all tumor models regardless of radiation sensitivity). (D) Volcano plot comparing the effect of each simulated gene knockout on total reduced GSH production between radiation-sensitive and -resistant tumors. Genes tested by experimental siRNA knockdown studies are bolded. (E) Schematic demonstrating the measurement of $\Delta E_{hc}$ GSH/GSSG (difference in glutathione half-cell potential between siRNA knockdown and negative control) in radiation-sensitive and -resistant cancer cell lines. (F) Comparison of model-predicted and experimentally-measured effects of gene knockdown on reduced GSH production. Top 3 rows: $\Delta E_{hc,\text{Res}} - \Delta E_{hc,\text{Sens}}$ in siRNA knockdowns across all three cell line pairs. $\Delta E_{hc,\text{Res}} - \Delta E_{hc,\text{Sens}} > 0$ for gene knockdowns causing greater oxidation in the radiation-resistant cell line, corresponding to a model-predicted $\log_2(\text{Resistant}/\text{Sensitive}) > 0$. Middle row: t-statistic from 1-sample t-test comparing the three experimentally-measured values of $\Delta E_{hc,\text{Res}} - \Delta E_{hc,\text{Sens}}$ to the null hypothesis population mean of zero (equal effect in radiation-sensitive and -resistant cell lines). Bottom row: model-predicted log fold change in gene knockout effect on reduced GSH production between radiation-resistant and -sensitive TCGA tumor models. ns: not significant, *: $p \leq 0.05$, **: $p \leq 0.01$, ***: $p \leq 0.001$, ****: $p \leq 0.0001$. Boxplots: box = 25th, 50th, and 75th percentiles, whiskers = 1.5 times the interquartile range. See also Figures S9–S10, S11A, S12–S13, Tables S2–S3.
Disparities in redox metabolism and $\text{H}_2\text{O}_2$-scavenging systems between radiation-sensitive and -resistant tumors determine differential ROS response. (A) Comparison of FBA-predicted total clearance of $\text{H}_2\text{O}_2$ between radiation-sensitive and -resistant TCGA tumors. (B) Experimentally-measured response of radiation-sensitive and -resistant cancer cell lines to 2 hr treatment of 10 mU/mL glucose oxidase, calculated as the relative cell viability compared to 0 mU/mL glucose oxidase. Data are represented as mean ± 1 standard error. (C) FVA-predicted fluxes through major $\text{H}_2\text{O}_2$-clearing reactions. Inset: Major $\text{H}_2\text{O}_2$-clearing reactions with their associated cellular compartments and compartment-specific isoforms. (D) Volcano plot comparing the effect of each gene knockout on total $\text{H}_2\text{O}_2$ clearance between radiation-sensitive and -resistant tumors. Genes tested by experimental siRNA knockdown studies are bolded. (E) Schematic showing the measurement of Bliss independence scores in radiation-sensitive and -resistant cancer cell lines. (F) Comparison of model-predicted and experimentally-measured effects of gene knockdown on $\text{H}_2\text{O}_2$ clearance. Top 3 rows: $\text{Bliss}_{\text{Sens}} - \text{Bliss}_{\text{Res}}$ in siRNA knockdowns across all three cell line pairs. $\text{Bliss}_{\text{Sens}} - \text{Bliss}_{\text{Res}} > 0$ for gene knockdowns causing a greater decrease in cell viability with glucose oxidase treatment in the radiation-resistant cell line, corresponding to a model-predicted $\log_2(\text{Resistant}/\text{Sensitive}) > 0$. Middle row: t-statistic from 1-sample t-test comparing the three experimentally-measured values of $\text{Bliss}_{\text{Sens}} - \text{Bliss}_{\text{Res}}$. 

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BlissRes to the null hypothesis population mean of zero (equal effect in radiation-sensitive and -resistant cell lines). Bottom row: model-predicted log fold change in gene knockout effect on H$_2$O$_2$ clearance between radiation-resistant and -sensitive TCGA tumor models. ns: not significant, *: $p \leq 0.05$, **: $p \leq 0.01$, ***: $p \leq 0.001$, ****: $p \leq 0.0001$. Boxplots: box = 25th, 50th, and 75th percentiles, whiskers = 1.5 times the interquartile range. Bar chart: mean ± 1 standard error. See also Figures S11B, S14–S15, Table S4.
Figure 5. Personalized metabolic flux profiles highlight heterogeneity in redox metabolism between radiation-resistant tumors.  

(A) Correlation between patient clinical factors and predicted fluxes through major NADPH-producing reactions among radiation-resistant patients. Values are represented as the p-value of either the univariate regression (for numerical factors) or 1-way ANOVA (for categorical factors) between reaction fluxes and clinical factor values. Statistically significant (p ≤ 0.05) associations are represented with black borders. (B) Personalized NADPH-generating flux profiles of 3 radiation-resistant HNSCC patients with their associated clinical factors. Radar chart values are the percentiles of reaction fluxes across all radiation-resistant tumors for each individual reaction. Patient profiles (red, filled) are shown overlaid on top of the profiles of all other radiation-resistant HNSCC tumors (black, not filled). See also Figures S16–S19.
# KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Antibodies          |        |            |
| Mouse monoclonal anti-alpha Tubulin (DM1A) | Thermo Fisher Scientific | Cat#62204; RRID: AB_1965960 |
| Rabbit monoclonal anti-GAPDH (14C10) | Cell Signaling Technology | Cat#2118 |
| Chemicals, Peptides, and Recombinant Proteins | | |
| 2-Vinylpyridine     | Sigma-Aldrich | Cat#132292; CAS: 100–69-6 |
| 5-Sulfosalicylic acid dihydrate | Sigma-Aldrich | Cat#S2130; CAS: 5965–83-3 |
| 5,5'-Dithiobis(2-nitrobenzoic acid) | Sigma-Aldrich | Cat#D8130; CAS: 69–78-3 |
| CellTracker Red CMTPX Dye | Thermo Fisher Scientific | Cat#C34552 |
| Ethylenediaminetetraacetic acid disodium salt dihydrate | Sigma-Aldrich | Cat#E5134; CAS: 6381–92-6 |
| Glucose oxidase from *Aspergillus niger* | Sigma-Aldrich | Cat#G7141; CAS: 9001–37-0 |
| Glutathione Reductase from baker’s yeast (*S. cerevisiae*) | Sigma-Aldrich | Cat#G3664; CAS: 9001–48-3 |
| L-Glutathione oxidized | Sigma-Aldrich | Cat#G6654; CAS: 27025–41-8 |
| L-Glutathione reduced | Sigma-Aldrich | Cat#S4251; CAS: 70–18-8 |
| N-TER Nanoparticle siRNA Transfection System | Sigma-Aldrich | Cat#N2913 |
| Potassium phosphate dibasic | Sigma-Aldrich | Cat#P8281; CAS: 7758–11-4 |
| Potassium phosphate monobasic | Sigma-Aldrich | Cat#P5655; CAS: 7778–77-0 |
| Triethanolamine | Sigma-Aldrich | Cat#T58300; CAS: 102–71-6 |
| Triton X-100 | Sigma-Aldrich | Cat#X100; CAS: 9002–93-1 |
| β-Nicotinamide adenine dinucleotide 2'-phosphate reduced tetrasodium salt hydrate | Sigma-Aldrich | Cat#N7505; CAS: 2646–71-1 (anhydrous) |
| Critical Commercial Assays | | |
| CellTiter-Glo 2.0 Cell Viability Assay | Promega | Cat#G9241 |
| Deposited Data | | |
| Enzyme turnover numbers | BRENDA database | https://www.brenda-enzymes.org |
| FBA models of TCGA tumors | This paper | https://github.com/kemplab/FBA-pipeline |
| Mutational effect predictions | Envision database | https://envision.gs.washington.edu/shiny/envision_new/ |
| TCGA clinical data | The Cancer Genome Atlas; Synapse TCGA_Pancancer project | https://www.synapse.org/#!Synapse:syn300013/wiki/27406 |
| TCGA genomic data | The Cancer Genome Atlas; GDC data portal | https://portal.gdc.cancer.gov |
| TCGA transcriptomic data | The Cancer Genome Atlas; (Rahman et al., 2015) | GEO: GSE62944 |
| Thermodynamic parameters | Virtual Metabolic Human database | https://www.vmh.life |
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Human: MDA-MB-231 NQO1(-) | Dr. David Boothman, Indiana University (Huang et al., 2016) | Derived from CVCL_0062 |
| Human: MDA-MB-231 NQO1(+) | Dr. David Boothman, Indiana University (Huang et al., 2016) | Derived from CVCL_0062 |
| Human: M059J | ATCC | CRL-2366 |
| Human: M059K | ATCC | CRL-2365 |
| Human: rSICC-61 | Dr. Cristina Furdui, Wake Forest University (Bansal et al., 2014) | N/A |
| Human: SCC-61 | Dr. Jeffrey N. Myers, UT MD Anderson Cancer Center | CVCL_7118 |
| Oligonucleotides | | |
| siRNA panel – See Table S7 | Sigma-Aldrich | N/A |
| siRNA: MISSION siRNA Universal Negative Control #1 | Sigma-Aldrich | Cat#SIC001 |
| Software and Algorithms | | |
| Source data and original code | This paper | https://github.com/kemplab/FBA-pipeline |
| Optimization software | Gurobi 8.0 | https://www.gurobi.com |