Combined 2-deoxy glucose and metformin improves therapeutic efficacy of sodium-iodide symporter-mediated targeted radioiodine therapy in breast cancer cells

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Abstract: Radiosensitization using either metformin or 2-deoxy-d-glucose (2-DG) in various cancer cells has been reported. The present study reveals novel information on combining these drugs to enhance radiosensitization effect in breast cancer (BC) cells. Responses to low-dose Cobalt60 radiation, as well as a newly emerged radioiodine therapy target for BC, that is, sodium-iodide symporter (NIS or SLC5A5) protein, are tested. As therapeutic potential of NIS in BC is often limited due to low uptake and fast efflux rate of iodine, the scope of these two radiosensitizers to further improve NIS-mediated 131I therapeutic efficacy is explored. Two BC cell lines, MCF-7, and MDA MB231 are tested to optimize minimal drug doses required for radiosensitization. A combination of 2 mM metformin and 20 mM 2-DG with 2 grey (Gy) Cobalt60 radiation shows significant radiosensitization effect (P=0.0002). In cells treated with the combination therapy, increased γH2A.X foci formation was noted. Further, MCF-7 BC cells overexpressing NIS (MCF-7 NIS) was established, and using the optimized drug concentrations, significant radiosensitization (P=0.0019) by 50 µCi 131I usage was found to be the case as well. Apoptosis data corroborates with the result of clonogenic assay showing significant increase in apoptotic population upon dual drug-mediated radiosensitization. In case of metformin treatment, lowered adenosine triphosphate (ATP) content of the cell has been observed. The encouraging radiosensitization effect observed using combined 2-DG and metformin may aid in reducing Cobalt60 radiation exposure or for targeted radioiodine therapy in BC cells with NIS expression. This study indicates high potential of this drug combination in sensitizing BC cells for NIS-mediated-targeted radioiodine therapy, which otherwise may have lacked efficacy. Keywords: sodium-iodide symporter, 2-DG, 131I, radiosensitizers

Introduction:
Breast cancer (BC) is the most common form of cancer in female population all over the world. Major treatment strategy for BC includes surgery, chemotherapy, hormonal therapy, and radiation therapy. Radiation therapy usually extends patients’ survival; however, it has several side effects. Thus, reducing the dose of radiation to effectively kill the cancer cells with the aid of radiosensitizers has long been of great research interest. In addition, efficacious targeted radiotherapy option for various subtypes of BC is yet to develop. One such emerging strategy under active research investigation is the endogenous aberrant expression of sodium-iodide symporter (NIS or SLC5A5) protein, which can serve the purpose of the targeted radioiodine therapy. Human NIS is an integral membrane protein that is involved in active uptake of iodide ions inside the...
thyroid cells, which has a crucial function in delivering iodine content required for the synthesis of thyroid hormones T3 and T4. NIS overexpression also serves as the basis for the targeted radioiodine therapy of patients with thyroid cancer since past 60 years.1,2 Study of NIS in BC was pioneered by Tazebay et al, showing NIS-positive expression in 80% patients with BC, whereas normal breast tissue samples examined from the patients who underwent reductive mammoplasty showed negative expression.3 Following this study, many reports across the world confirmed high NIS expression in BC by various molecular analysis methods.4-9 In our own study, NIS expression was observed in 70% BC cases, where its expression was predominantly intracellular except in a few cases where NIS staining at the cell surface was also observed.9 Translational-medicine-based studies have also been carried out for testing the efficacy of NIS-based therapy and diagnosis of malignant breast tissue. Moon et al reported significant 99mTechnetium (TcO4-) uptake in four out of 25 patients with BC by scintigraphic scanning method.10 In a very important study, Wapnir et al detected radioiodide uptake in metastatic BC by scintigraphic analysis using radioiodides (123I and 131I) or 99mTechnetium (TcO4-).7 In this study, iodine uptake was noted in 25% of NIS-positive metastatic breast tissues (two out of eight). In addition, this study also showed significant suppression of thyroid radioiodine uptake by administration of agents like thyroxine and methimazole in trial patients. In another study, NIS expression was observed in 15 out of 23 patients with triple-negative BC, and membrane expression was observed in four patients with BC.4 Further, radioiodide uptake was also demonstrated in tumor tissue by 123I scintigraphy in a patient with large primary BC; however, it was reported that had the isotope been 131I, the cumulative concentration of iodine in the tumor may not deliver sufficient therapeutic benefits.4 Overall, these reports together suggest that NIS expression is high in BC as compared with normal breast tissue, but its expression is strong in a very small subset (<25%) of positive cases. This discrepancy widens the scope of multiple studies that have focused on modulating the expression level of NIS with agents such as retinoic acids and histone deacetylases before 131I therapy.11-13 Another strategy that is still unexplored is to enhance the 131I treatment efficacy itself by using radiosensitizer drugs. Use of radiosensitizers may enhance cytotoxicity of 131I therapy in case of a very low iodine uptake when NIS expression is low. At the same time, with the use of radiosensitizers, a lower concentration of 131I can be used when NIS expression is high, which may benefit by minimizing the side effects of radiation in other NIS-expressing tissues, where entry of radioiodine cannot be avoided. Further, without the presence of a defined iodide organification mechanism in BC cells, radiosensitizers may enhance radio- ablative effect even during the short presence of 131I inside these cells. Since radiation therapy is known to show cytotoxicity via free radical-mediated oxidative damage, it has been proposed that combining radiation therapy with agents that cause oxidative stress-induced injury may sensitize cells to the cytotoxic effect of radiation.14,15 Thus, 2-deoxy-D-glucose (2-DG) shows good radiosensitization effect by disrupting thiol metabolism and increasing oxidative stress.16-19 Another drug used in this study is metformin (a bi-guanide drug and activator of AMP-activated protein kinase [AMPK]), which has been reported as a radiosensitizer for BC cells and showed eradication of radioresistant cancer stem cells by activating AMPK and suppressing mTOR.20 To date, this is the first study to show the combination of 2-DG and metformin as radiosensitizers for the enhancement of Cobalt60 or 131I radio- ablative effect on the cancer cells. We focus on combining the two radiosensitizer drugs against the different radiation exposure methods and measuring the benefit using BC cells. We show that a combination of metformin and 2-DG along with Cobalt60 radiation or NIS-mediated 131I radiation shows significantly enhanced cytotoxicity. To the best of our knowledge, this is also the first report where radiosensitization effect alone has been shown to enhance NIS-mediated 131I therapy efficacy in BC cell model.

Materials and methods

Materials

MCF-7 and MDA MB231 BC cell lines were from ATCC (Manassas, VA, USA). RPMI1640 (Cat # 31800-022) and L15 (41300021) media, fetal bovine serum (10082-147), Trypsin-ethylenediaminetetraacetic acid (EDTA) (25300-062), lipofectamine 2000 (11668-027), and Hank’s Balanced Salt Solution (HBSS, 24020-117) were obtained from Thermo Fisher Scientific (Waltham, MA, USA). pattB-CAG plasmid vector was obtained from Addgene (Cambridge, MA, USA). 2-DG (D8375), metformin (PHR 1084), propidium iodide (PI, P4170), 4’,6-diamidino-2-phenylindole (DAPI, D8417), and MTT (M5655) were obtained from Sigma-Aldrich Co (St Louis, MO, USA). Monoclonal human NIS antibody (ab17795) and annexin V FITC (fluorescein isothiocyanate) apoptosis detection kit (ab14085) were obtained from Abcam (Cambridge, UK). H2A.X monoclonal antibody (MA5-15130) and anti-mouse (35512) and anti-rabbit Dylite633 (35562) antibody were obtained from Thermo Fisher Scientific. Enliten ATP assay kit (FF2000) was from Promega
Corporation (Fitchburg, WI, USA). The authors advise no ethical approval was required for this study, as they did not use any human tissue samples, and the complete study was performed using well-established breast cancer cell lines.

**MTT assay**

A total of 5,000 cells/well were seeded in 96-well plate. Cells were allowed to adhere overnight before they were treated with mentioned drug doses of metformin or 2-DG. After 48 hours of treatment, cell viability was measured by MTT assay, and data are represented as percentage cell viability as compared with the untreated cells.

**Preparation of NIS overexpression plasmid**

NIS gene was inserted immediately downstream of chicken β-actin promoter in pcDNA3.1+ attB-CAG-FTT (Fluc2-tdtomato-thymidine kinase) vector system. Briefly, NIS gene was amplified by polymerase chain reaction using primers containing Nhe-I and Not-I restriction sites. The pcDNA3.1+ attB vector was digested with Nhe-I and Not-I, which released the FTT reporter from vector backbone. Further, NIS gene and attB vector backbone were ligated. Positive colonies were screened for NIS gene insertion by enzyme digestions and sequencing.

**Preparation of NIS expressing BC cell line**

MCF-7 BC cells were co-transfected by lipofectamine 2000 with pcDNA 3.1+ attB-CAG-NIS vector and integrase vector. We used integrase-based transfection method, where the plasmid vector contained bacterial attachment site (attB) upstream to chicken β-actin promoter. As the plasmid contained neomycin gene, positive clones were selected using 500 μg G418. The isolated cell colonies were expanded and checked for NIS expression by immunofluorescence assay and functionally validated for iodine uptake. Cells obtained showed different levels of NIS expression and thus different levels of iodine uptake. The single cell colony (MCF-7 NIS) showing high expression of NIS protein, as well as good iodine uptake, was used for radiosensitization experiments.

**Iodine uptake assay**

A total of 50,000 MCF-7 NIS cells were seeded in 24-well plate in duplicates and incubated with 125I for 1 hour. At the end of 1 hour, 125I-containing medium was removed and replaced with plain medium without 125I. Cells were collected at the mentioned time points and remaining 125I activity was measured by beta counter.

**Iodine efflux assay**

A total of 50,000 cells were seeded in 24-well plate in duplicates and incubated with 125I for 1 hour. At the end of 1 hour, 125I-containing medium was removed and replaced with plain medium without 125I. Cells were collected at the mentioned time points and remaining 125I activity was measured by beta counter.

**Clonogenic assay**

In this assay, 2×10⁶ cells were seeded in 25 mm flask and incubated with 50 μCi 125I/mL in HBSS for 5–6 hours. At the end of the time point, 125I-containing medium was removed, and cells were washed with cold HBSS. Cells were trypsinized, and 1,000 cells were seeded in six-well plates in triplicates. Medium was changed at regular intervals, and after 14 days, cells were fixed with 10% formaldehyde and stained with 4% crystal violet in methanol. Stained colonies were counted under stereomicroscope, and survival fraction was calculated according to the protocol mentioned in a previous publication. All the drug treatments were given 12 hours before the 125I treatment/radiation therapy and were replaced by plain medium after 12 hours of 125I treatment, and assay was performed after 24 hours of treatment.

**Immunofluorescence for NIS and γH2A.X foci assay**

Cells were seeded on cover slip in 35 mm plate, and after drug/radiation/125I treatment, they were washed with 1× phosphate buffered saline (PBS) for three times and fixed with 4% para formaldehyde in PBS at 37°C. An additional permeabilization step with 0.2% TritonX 100 in 4% para formaldehyde was carried out for foci staining. Non-specific sites were blocked using 2% bovine serum albumin in PBS. Cells were then incubated with γH2A.X or NIS antibody in 1:400/1:50 dilutions, respectively, for overnight at 4°C in a humid chamber. After incubation with primary antibody, cells were washed with 1× PBS and incubated with either anti-rabbit Dylight633 or anti-mouse Dylight633 secondary antibody, respectively (1:200) for 1 hour at room temperature in dark humid chamber. Cells were washed with 1× PBS, and nucleus was stained with DAPI. Cells were mounted...
with Vecta Shield, and images were captured under confocal microscope (LSM510, Zeiss). Foci assay was performed at the immediate end of radiation/131I treatment. For quantification of the foci, intensity of staining in red channel was measured by ImageJ software. For each treatment, 50 cells were analyzed, and average fluorescence intensity was plotted.

**Cell cycle analysis and apoptosis assay**

At the mentioned time points, 1×10⁶ cells were trypsinized, collected, and washed with 1× PBS. For cell cycle analysis, cells were fixed with 70% ethanol and incubated overnight at −20°C. Next day, cells were pelleted and washed with 1× PBS. Washed cells were treated with 1 µg/mL RNase A for 10 minutes at 37°C. Then, 300 µL of 50 µg/mL propidium iodide was added in each tube, incubated for 30 minutes at 37°C, and acquired in FACSCalibur™.

For apoptotic cell count, annexin V apoptosis detection kit was used, and the protocol mentioned in the kit was followed.

**ATP measurement**

Total cellular ATP was determined using manufactured recommended protocol of luciferase-based ATP detection kit. Total ATP was measured after 24 hours of radiation or 131I therapy.

**Statistics**

Data are presented as mean ± standard error. Student’s t-test was used to judge the significance of difference. All tests were two-sided, and P<0.05 was considered as significant. Graph pad Prizm software was used for all the statistical analysis.

**Results**

**A combination of 2-DG and metformin enhances tumor cell ablation upon Cobalt60 radiation treatment**

To determine the cytotoxic effect of 2-DG and metformin, BC cell lines were first exposed to various concentrations of these two drugs ranging between 0.1 mM and 100 mM, and viable cell count was measured by MTT assay (Figure S1). Further, based on the MTT data, 10–50 mM dose of 2-DG and 2–10 mM dose of metformin were combined with 1 or 2 grey (Gy) Cobalt60 radiation to determine the minimal optimum drug dose required for...
maximum radiosensitization, and cell survival analysis was performed by clonogenic assay in MCF-7 cells. We observed that a combination of 20 mM 2-DG with 1 Gy reduced twofold cell survival as compared with 1 Gy radiation alone, while its combination with 2 Gy radiation showed 1.6-fold decrease in cell survival than radiation alone (Figure 1A). Further increment in 2-DG dose did not show any significant enhancement in radiosensitization effect. Similarly, treatment of cells with metformin showed dose-dependent radiosensitization effect (Figure 1B). Combination of 2 mM metformin with 1 Gy radiation dose showed 1.5-fold decrease in cell survival as compared with 1 Gy radiation alone, while the combination of 2 mM metformin with 2 Gy radiation resulted in 1.9-fold decrease in cell survival as compared with 2 Gy radiation alone.

Upon deciding the effective minimal drug doses, we also combined both the drugs and tested in combination with 2 Gy Cobalt60 radiation. Cell survival showed significant drop in cells treated with dual drug radiosensitization to that of cells treated with radiation alone ($P=0.0094$, CI =95%) (Figure 1C). In addition to MCF-7 cells, we also evaluated the effect of radiosensitizers on Cobalt60 radiotherapy in MDA MB231 cells (Figure S2). In MDA MB231 cells, 4 mM metformin and 5 mM 2-DG dose showed radiosensitization with 1 Gy, as well as 2 Gy, radiation dose. Further, combination of both the drugs with 2 Gy radiation showed significant decrease in cell survival as compared with 2 Gy radiation alone ($P=0.0020$, CI =95%). Further, $\gamma$H2A.X foci assay was performed to ensure increased cell death is associated to increased DNA double-strand break (Figure 1D), and quantification of the immunofluorescence images indicated increase in foci intensity in cells treated with drug for 12 hours followed by the measured radiation exposure as compared with cells treated with similar radiation dose alone (Figure S3). In case of untreated cells or cells treated with drugs alone, foci formation is recorded as negligible. Since metformin and 2-DG therapy are known to reduce ATP content of the cell, we also analyzed ATP content with or without drug treatment and observed a significant drop in cells treated with metformin ($P=0.0058$, CI =95%), whereas this drop was not significant when cells were treated with 2-DG alone (Figure 1E). Further decrease in ATP content was noted when both the drugs were combined.

Cell cycle analysis was also performed at different time points after the combination therapy, which revealed S-phase arrest after 12 hours of treatment in combination as well as in case of only radiation therapy (Figure 2). It was also observed that after 24 hours and 48 hours of combination therapy, cells remained arrested in S phase, whereas in case of only radiation therapy, this arrest was relieved at these
time points, indicating prolonged cytotoxic effect of combination therapy.

Established BC cell line overexpressing cell membrane NIS protein demonstrates good iodine influx for short duration

For effective iodide accumulation inside cells, the presence of NIS transporter protein on cell membrane is crucial. Keeping this fact in mind, we engineered a mammalian expression vector for expressing NIS (Figure 3A) and isolated MCF-7 BC clonal cell variants with abundant NIS protein localized either on the cell membrane or in the cytoplasm. NIS transcript and protein expression were confirmed by real-time PCR (data not shown), western blot analysis (Figure 3B), and immunofluorescence (Figure 3C). As shown in Figure 3D, when NIS protein was expressed on the membrane, significantly high iodine accumulation took place as compared with the cells where NIS was localized predominantly in the cytoplasm ($P=0.0028$, CI=95%) or the baseline MCF-7. Treatment of NIS-positive cells with KClO$_4$, a competitive iodide inhibitor, showed complete inhibition of iodine uptake in cells, indicating NIS-specific iodine uptake. Further, to estimate the duration of iodine retention in membrane expressing clones (MCF-7 NIS), efflux assay was performed, and the results showed that 90% iodine was released within 15 minutes of incubation (Figure 3E).

We have also evaluated the therapeutic efficacy of $^{131}$I treatment on MCF-7 NIS membrane expressing cell line by measuring the survival fraction by clonogenic assay. As detailed in the “Materials and methods” section, a brief exposure of these cells to 50 $\mu$Ci $^{131}$I was found to reduce cell survival significantly ($P<0.0001$, CI=95%) than the untreated cells (Figure 4A). Radiation injury causing cellular apoptosis was assessed by annexin V staining.
protocol, which showed 46.5% live cells in case of $^{131}$I treatment as compared with 86% in control (Figure 4B). To confirm that such brief and low-dose exposure of MCF-7 NIS cell still cause DNA double-strand break, $\gamma$H2A.X foci formation assay was performed at different time intervals after $^{131}$I treatment. Formation of a significantly large number of foci in treated cells was noted as compared with the untreated cells at 12 hours post-treatment time point (Figure 4C). However, $\gamma$H2AX time kinetics did show that at 48 hours, the number and size of foci significantly diminished (data not shown). Thus, together our results confirm that although major amount of iodine...
was effluxed out by these cells within a short duration, still sufficient radio-ablative effects on MCF-7 NIS cell were evident.

**Treatment efficacy of radioiodine (\(^{131}\)I) is significantly enhanced by 2-DG and metformin combination**

The effective 2-DG and metformin dose of 20 mM and 2 mM, respectively, was tested prior to Cobalt60 radiation treatment. The similar treatment was applied in combination with \(^{131}\)I radioisotope in NIS expressing cells. Treatment of cells with 100 µCi of \(^{131}\)I alone resulted in high number of cell death (data not shown), making radiosensitization effects obscure, a lowered concentration (50 µCi) of \(^{131}\)I was used for all the experiments. Combination of dual drugs with 50 µCi \(^{131}\)I showed 9.5-fold decrease in survival fraction when compared with the cells treated with \(^{131}\)I alone (\(P=0.0019\)) and fourfold decrease when compared with cells treated with the two drugs alone (\(P=0.0036\)) (Figure 5A). The change in cell survival was further confirmed by annexin V staining to measure apoptosis in parallel (Figure 5C). Cells were analyzed for apoptosis after 48 hours of treatment, and it was observed that live cell population was 37.37% in case of \(^{131}\)I therapy, whereas it was only 11% for the combination therapy. When cells were treated with both the drugs together in absence of \(^{131}\)I, 55.54% cells were alive. Thus, the combination of both the drugs with \(^{131}\)I therapy in NIS-expressing cells results in a significant drop in live cell population. Further γH2A.X foci immunofluorescence staining showed increase in number and intensity variation of foci in case of combined treatment, while in case of \(^{131}\)I treatment alone, the number and size of foci were much lower (Figure 5B and Figure S4). We also performed cell cycle analysis after 48 hours of the combination treatment and a very slight S-phase elongation was observed, which was not significant (data not shown). Finally, we compared cytotoxic efficacy of 2 Gy Cobalt60 radiation with that of the 50 µCi \(^{131}\)I, which showed a comparable cell survival fraction (Figure 5D). However, in the presence of the radiosensitizers, a significantly lower survival fraction in \(^{131}\)I than in Cobalt60 radiation was noted (\(P=0.0201, CI=95\%\)). This result confirms that the combination of 2-DG and metformin can effectively radiosensitize BC cell and thus can be combined with either Cobalt60 radiation or NIS-targeted radioiodine therapy.

Since in majority of the patients with BC, NIS expression is localized in the cytoplasm, it is important to evaluate the effect of radiosensitizers combined with \(^{131}\)I in plain MCF-7 cells. Our data suggest that treatment with 50 µCi \(^{131}\)I results in 30% cell death (Figure 6A) as opposed to 60% death in case of MCF-7 NIS overexpressing cells. However, addition of radiosensitizers to the \(^{131}\)I therapy results in significant increase in cell death (70%) as compared with \(^{131}\)I alone (\(P=0.0110, CI=95\%\)). Thus, combination therapy of \(^{131}\)I with radio sensitizers in MCF-7 cells is as effective as \(^{131}\)I therapy alone in MCF-7 NIS cells, where NIS is localized at membrane. We also performed γH2A.X assay to evaluate DNA damage response in plain MCF-7 cells in presence or absence of radiosensitizers.
by the presence of cells at different stages of cell cycle. Previous reports suggest the action of metformin and 2-DG is due to lowered ATP content of the cell. Metformin alters mitochondrial oxidative phosphorylation, and 2-DG inhibits ATP synthesis through glycolysis. Therefore, we have also evidenced lowered ATP content of the cell in case of metformin or dual-drug treatment, but not the 2-DG treatment. Further, we looked into cell cycle status under different drug treatment conditions and observed that combined treatment resulted in elongation of S phase. This S-phase elongation was also observed in case of radiation but was relieved after 48 hours of treatment, whereas it remained nearly constant in case of combination therapy even at 48 hours.

Since 20 mM 2-DG and 2 mM metformin showed effective radiosensitization with Cobalt60 radiation, we maintained the similar dose and time points in experiments combining 2-DG and metformin with 131I treatment in MCF-7 NIS and plain MCF-7 cells. The use of 100 µCi 131I treatment alone resulted in complete cell death (data not shown). Therefore, 50% lower dose of 131I (ie, 50 µCi) was attempted, which showed 1.5-fold decrease in survival fraction in MCF-7 NIS cells as compared with untreated cells. Thus, for all the radioiodine therapy experiments, only 50 µCi of 131I was used. Even at this low radiation dose, the dual-drug use for radiosensitization showed significant decrease in cell survival, which is near-equivalent to the twofold higher dose (ie, 100 µCi alone) of 131I. Apoptosis data corroborate with the result of clonogenic assay, showing significant increase in apoptotic population with the addition of dual-drug radiosensitizers. We have also performed cell cycle analysis after 48 hours of treatment but did not observe any significant differences among samples with different treatments (data not shown). Since localization of NIS is crucial for 131I therapy and various reports suggest cytoplasmic localization of NIS in breast tumor tissues, it is important to evaluate the potential of radio sensitizers with 131I therapy in plain MCF-7 cells. Our results indicate that although 131I therapy alone shows minimal effect on cell survival, combination of radioiodine with radiosensitizers results in significant loss of live cell population in plain MCF-7cells. Thus, the present study demonstrates the potential of metformin and 2-DG combination treatment as radiosensitizer in combination with Cobalt60 or 131I radiation in NIS-expressing cells. The combination can deliver better therapeutic efficacy than using either of these drugs as a single agent. In addition to their drug toxicity, such radiosensitization effect may further boost mass reduction in cancer cases. To note, at present, there are ongoing clinical trials with both these drugs either in combination with chemotherapy or with radiation.29,30 Thus,
combination of 2-DG and metformin with Cobalt60 radiation might play an even more important role in sensitizing radioresistant population as was observed previously with combination of metformin and radiation.30 Previous reports suggest that combination of metformin and 2-DG induces p53-dependent apoptosis in prostate cancer cells through AMPK pathway and metformin inhibits 2-DG induced autophagy in these cells.31 In future, it will be interesting to understand how these mechanistic pathways are regulated when Cobalt60 radiation or 131I is combined with these two drug combination. Further, it would also be important to understand the effect of these drugs in vivo in combination with 131I in NIS-expressing breast tumor in preclinical and clinical settings.

**Conclusion**

In the present study, we have shown the use of a combination of two radiosensitizers, 2-DG and metformin, to enhance the radio-ablative effect in combination with either Cobalt60 radiation or targeted NIS-mediated radioiodine therapy. The combination of the radiosensitizers improves therapeutic efficacy effectively in BC cell lines than using any one of the radiosensitizers. Thus, in addition to the drug cytotoxicity, the radiosensitization effect may further boost effective mass reduction in pre-surgical cancer cases.

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**Disclosure**

The authors report no conflicts of interest in this work.

**References**


Supplementary materials

Figure S1 Cytotoxic effect of 2-DG and metformin in (A) MCF-7 and (B) MDA MB 231 breast cancer cell lines.
Abbreviations: 2-DG, 2-deoxy-D-glucose; Met F, metformin.

Figure S2 Effect of 2-DG, metformin, and their combination with Cobalt60 radiation on MDA MB 231 cells.
Notes: (A and B) MDA MB231 cells were treated with different dose of 2-DG or metformin, with or without radiation exposure for 24 hours, and clonogenic assay was performed. (C) Combination of 5 mM 2-DG and 4 mM metformin with 2 Gy radiation dose showing further decrease in survival fraction. **P<0.01.
Abbreviations: 2-DG, 2-deoxy-D-glucose; Ctrl, control; Gy, grey; Met F, metformin; ns, not significant.
Figure S3 Effect of Cobalt60 radiation and its combination with 2-Dg and Met F on foci formation in MCF-7 cells.

Notes: (A) Foci assay images from cells treated with Cobalt60 radiation and combination with radio sensitizers showing foci staining with dylight 633 secondary antibody in red channel, nucleus stained with DAPI in blue channel, and merged image showing colocalization of foci with nucleus. Scale bars represent 10 μm. (B) Quantification of foci from the IF images. In each sample, 50 cells were quantified for intensity of staining from various fields of the cover slip.

Abbreviations: 2-DG, 2-Deoxy-D-glucose; DAPI, 4′,6-diamidino-2-phenylindole; Gy, grey; IF, immunofluorescence; Met F, metformin; MFI, mean fluorescence intensity.
Figure S4 (A) Foci assay images from Figure 4 (MCF-7 NIS cells treated with $^{131}$I and its combination with radio sensitizers) showing foci staining with dylight 633 secondary antibody in red channel, nucleus stained with DAPI in blue channel, and merged image showing colocalization of foci with nucleus. Scale bars represent 10 μm.

(B) Quantification of foci from the IF images. In each sample, 50 cells were quantified for intensity of staining from various fields of the cover slip.

Abbreviations: 2-DG, 2-Deoxy-D-glucose; DAPI, 4,6-diamidino-2-phenylindole; IF, immunofluorescence; MET, metformin; NIS, sodium-iodide symporter; MFI, mean fluorescence intensity.
Figure S5 (A) Foci assay images from Figure 5 (MCF-7 plain cells treated with $^{131}$I and its combination with radio sensitizers) showing foci staining with dylight 633 secondary antibody in red channel, nucleus stained with DAPI in blue channel, and merged image showing colocalization of foci with nucleus. Scale bars represent 10 μm.

(B) Quantification of foci from the IF images. In each sample, 50 cells were quantified for intensity of staining from various fields of the cover slip.

Abbreviations: 2-Dg, 2-deoxy-D-glucose; DAPI, 4',6-diamidino-2-phenylindole; IF, immunofluorescence; MET, metformin; MFI, mean fluorescence intensity.