Chemically enhanced radiotherapy: visions for the future

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Abstract: Radiotherapy (RT) is an important part of cancer management, with more than a third of all cancer cures being attributable to RT. Despite the advances in RT over the past century, the overall outcomes in a majority of malignancies are still unsatisfactory. There has been a constant endeavor to enhance the outcome of RT, and this has been in the form of altered fractionation, oxymimetic radiosensitizers, the use of concurrent chemotherapy, anti-angiogenic therapy and anti-growth factor receptor targeted therapies. This article presents a vision for the future, with emphasis upon emerging prospects which could enhance RT outcomes. Positive speculations regarding the use of immunological aspects, the use of nanoscale technology and the adoption of metronomic concurrent chemotherapy have been presented. Also, the potential with the use of low dose hyperradiosensitivity in enhancing chemotherapy outcomes too has been discussed. In this era of evidence based clinical practise, there exists a strong obsession towards the ‘present’ with ‘contempt towards the future’. Accepting the shortcomings of the existing modalities, there must be a strong zeal towards discovering better methodologies to enhance radiotherapeutic outcomes for the sake of a better future.

Keywords: Novel therapeutics; radiosensitizers; gold nanoparticles; hyperradiosensitivity; metronomic chemotherapy; immunoradiotherapy

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Introduction

Radiotherapy (RT) is an important aspect of cancer care, with estimates suggesting that it is currently responsible for 40% of all cancer cures (1). Ever since RT has been used for treating cancer, there has been a constant quest to improve effectiveness while reducing toxicity. The earliest methods included experimentation with radiation time-dose-fractionation based on radiobiological modelling. Then, understandings about predictors of radiation response, such as anaemia and tumour hypoxia generated a quest for combating hypoxia by addition of experimental oxymimetic radiosensitizers (such as misonidazole & nimorazole) (2). That era also ushered the combination of chemotherapy with RT, earlier with agents such as hydroxyurea and methotrexate, and later with 5-FU analogs and platinum agents. The discovery of monoclonal antibodies and small molecule tyrosine kinase inhibitors against various growth factor receptors and angiogenic targets had encouraged the combination of RT with targeted therapy. The combination of hormonal therapy prior to RT in prostate cancer, the use of neoadjuvant chemotherapy prior to RT in head and neck squamous cancers have been other recent attempts intended at improving RT outcomes (3-5).

Despite the modern advances in RT such as the adoption of conformal and intensity modulated treatments, cancer patients in significant numbers continue to fall prey to the formidable malady. Thus it would be realistic to admit at present, that the current accomplishments are far from
adequate, if not trivial. But there is hope for a better future. Newer understandings on cancer biology and newer perspectives in cancer therapies have opened up scope for dramatic improvements in outcomes for the future.

Recent understandings about the interactions between RT, chemotherapy and the host immune system has encouraged the development of strategies to utilize these to unlock new opportunities to improve RT outcomes. The advent of nanoscale medicine has allowed better tumour targeting of chemotherapeutic agents. The fact that RT and nanoscale medicines augment the efficacy of each other makes their concurrent use an excellent proposition worth exploration. The use of nanoscale particles with high atomic number elements as radiosensitizers is another exciting prospect worthy of investigation, as this approach can be expected to provide effects similar to heavy ion therapy, all while using a conventional photon beam. Newer perspectives in delivery of chemotherapy have explored the rather unconventional form of chemotherapy such as metronomic chemotherapy. These used with, or after RT may help enhance outcomes while sparing the patient from the toxicities of conventional chemotherapy. Also, the discovery of hyperradiosensitivity at low doses of radiation has encouraged the use of low dose RT as a chemopotentiator.

This editorial review touches upon the most promising prospects which may revolutionize radiotherapeutic treatment of cancer patients. Less emphasis has been placed on on-going trials involving newer conventional chemotherapeutics and existing molecular targeted therapies, and instead, focus has been placed upon novel ideas and technologies that are likely to be available in the foreseeable future.

**Immunology, immunological interactions and prospects**

While some components of the immune system, such as the cytotoxic T lymphocytes and NK cells strive hard to eliminate tumours, cancer cells enjoy the protective effect of certain immune suppressive cells and suppressive cytokines. Recently it has been understood that a variety of tumour protective immune factors exist, such as T-reg (CD4⁺CD25⁺ regulatory T-cells), MDSC (myeloid derived suppressive cells), certain cytokines (IL-10, TGF-beta) and regulatory tumor associated macrophages (TAMs) (6). It has been demonstrated that chemotherapy and RT alter tumours’ immune tolerance and that this could indeed be an important, albeit less recognized mechanism of action of these modalities.

Of late, it has been established that while maximally tolerated dose (MTD) chemotherapy depletes all immune cells, low dose chemotherapy selectively depletes T-reg cells and hence enhance antitumor immune response. For example, low dose cyclophosphamide reduces suppressive cells selectively without depleting cytotoxic T cells; but at higher doses, cyclophosphamide loses this specificity (7). Other agents at lower doses such as paclitaxel and 5-FU are known to cause MDSC apoptosis. The same is discussed later under the heading dedicated to metronomic chemotherapy.

Recent research has focussed upon the function of CTLA-4 and PD-1 receptors. These receptors present on cytotoxic T cells act in a suppressive manner so as to prevent autoimmunity. However, this inhibition also allows cancer cells to survive killing by these cytotoxic T-cells. Monoclonal antibodies against CTLA-4 (ipilimumab) and PD-1 (nivolumab) have demonstrated excellent results in melanoma and lung carcinoma. RT and anti-CTLA-4 and PD-1 inhibitors are likely to be synergistic. An interesting case report described a patient who had progressive disease despite initial immunotherapy with ipilimumab and a sudden dramatic systemic response after localized RT (8).

**Prospects with nanomedicine**

Nanoscale technology aims to enhance drug targeting, improve biodistribution, overcome resistance mechanisms and reduce toxicity of therapeutic molecules. Various nanomedicines utilizing forms of liposomes, polymers, micelles, dendrimers and others are observed to passively accumulate in tumours owing to their vasculature which is leakier due to wider fenestrations. This enhanced accumulation at tumour sites is called ‘enhanced permeability and retention’ (EPR) effect (9). Though obstacles to the EPR effect do exist, most significantly in the form of reticuloendothelial system (RES) capture, this can be reduced by the use of PEGylation which by producing a hydrated barrier causes hindrance to the attachment of phagocytes. The EPR effect along with PEGylation can increase tumour drug concentration by 10-100 times in comparison to that with the use of free drugs (10). There is immense potential for the use of nanoscale technology in cancer treatment (Table 1). First of all, the availability of agents such as liposomal doxorubicin and nano-albumin bound (nab-) paclitaxel has already enhanced
efficacy and safety in comparison to the more traditional forms of doxorubicin and paclitaxel. Thus, the use of nanoparticle bound chemotherapy in place of conventional free forms of chemotherapy will render the delivery of concurrent chemoradiotherapy much more efficacious and tolerable (11).

In a potential ‘eureka moment’ for oncology, it was observed that the use of poly-L glutamic acid bound paclitaxel as a radiosensitizer could reduce the TCD50 dose in a preclinical model from 53.9 Gray (Gy) to just 7.9 Gy (12). Another interesting observation has been that the tumour targeting of liposomal doxorubicin could be enhanced by the use of a peptide (HVGGSSV) which would bind selectively to irradiated tumours. Thus, this selective binding suggests that irradiation can be utilized to guide drug delivery to tumours (13).

The use of high atomic number (Z) nanoparticles as radiosensitizers is another extremely attractive approach on the horizon. The high Z atoms interact in a very different manner to ionizing radiation in comparison to low Z atoms. Since biological tissues are mostly made up of low Z atoms, the introduction of high Z nanoparticles into tumours can dramatically intensify response to RT. High Z atoms during photoelectric effect undergo inner shell ionization wherein one of the deeply bound electrons is ejected. This results in a highly unstable atomic system which stabilizes by the emission of low energy photons (fluorescence) and auger electrons. Several auger electron emissions can occur from single inner shell ionization and these auger electrons have a range of 10-100 nm and hence deposit energy very locally. Thus, a very high energy deposition which can be comparable to heavy ion therapy can be achieved with the use of a high Z nanoparticle radiosensitizer (14,15).

The most commonly studied high Z nanoparticle radiosensitizers in preclinical models include gold, iron oxide and hafnium oxide nanoparticles. An in-silico simulation showed that the use of gold nanoparticles as radiosensitizer could lead to higher dose enhancement with kilovoltage range ionizing radiation. For example, while the dose-enhancement with 2 megavolt photons was 53%, the dose enhancement with kilovoltage range photons was as high as 560% (16). While nanoparticles are known to enhance RT, the favour is returned by RT in that it enhances nanoparticle accumulation into tumours. In a mouse breast tumour model, it was observed that iron oxide nanoparticle accumulation was doubled after a single 15 Gy dose (17).

**Metronomic chemotherapy to enhance RT**

Metronomic chemotherapy is the chronic administration of chemotherapy at low doses which are minimally toxic, in a schedule of administration without prolonged drug-free breaks. Commonly used agents include low dose versions of conventional chemotherapy agents such as cyclophosphamide, methotrexate, capecitabine, etoposide, etc. In addition, a few non-cancer drugs such as celecoxib, metformin, valproate and such are being used (termed ‘drug-repositioning’, wherein drugs approved for non-oncological indications are re-positioned for oncological use (18).

In contrast to conventional MTD chemotherapy, the new approach of metronomic chemotherapy is very less toxic inherently by its design. Many subtle properties of chemotherapeutic agents could possibly have been masked by the MTD approach, but are now being unravelled in the metronomic approach (19). While MTD chemotherapy solely aims towards killing of malignant cells, the metronomic approach owes its efficacy to numerous other effects. Metronomic chemotherapy has anti-angiogenic effects and this has been demonstrated experimentally in that it was able to reduce angiogenic factors such as thrombospondin-1 (20). Another study in patients undergoing thoracic irradiation for lung cancer revealed

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**Table 1 Nanoscale technologies and prospects for enhancing outcomes**

<table>
<thead>
<tr>
<th>Aspects of nanoscale medicine worthy of investigation for use with radiotherapy</th>
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<tbody>
<tr>
<td>Nanoparticle bound chemotherapy such as paclitaxel and doxorubicin may greatly reduce concurrent radiotherapy doses required to achieve tumour control</td>
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<tr>
<td>The HVGGSSV peptide bound liposomal doxorubicin binds specifically to irradiated tissues; hence, nanoparticle bound chemotherapy can be ‘guided’ to tumor targets painted with radiation doses</td>
</tr>
<tr>
<td>High-atomic number element nanoparticles as radiosensitizers can cause a drastic dose enhancement effect for kilovoltage range photon beams, thus enabling effectiveness comparable to that with the use of heavy ion therapy</td>
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that the addition of metronomic chemotherapy caused a marked reduction of VEGF (21). In addition to its anti-angiogenic effects, metronomic chemotherapy also has immunoregulatory functions, wherein the elimination of immunosuppressive cells (such as T-reg & MDSC) and the increased MHC-1 molecule expression, increased dendritic cell maturation are all known to enhance anti-tumour immune response (22).

Metronomic chemotherapy has already proven beneficial in often difficult situations involving patients who were heavily treated with conventional regimens (23,24). At present there is a paucity of clinical trials combining concurrent metronomic chemotherapy with RT. If not in every patient, this approach could help improve outcomes at least in patients who are unlikely to tolerate use of conventional concurrent chemotherapy (Table 2). Lastly, it is very much feasible that the use of metronomic chemotherapy adjuvant after a course of standard treatment may help reduce distant recurrences. A study utilizing metronomic tegafur-uracil observed a very significant reduction in distant failures among nasopharyngeal carcinoma patients who had persisting plasma EBV DNA levels after completion of definitive chemoradiotherapy (25).

**Prospects with antiangiogenic therapy**

The use of RT can unintentionally enhance the process of angiogenesis by up-regulating factors such as VEGF in tumor cells, VEGFR in endothelial cells and αVβ3 integrin in tumor endothelial cells. Thus, the quest for combining antiangiogenic use with RT is very much rational (26). Recently, the results of the RTOG 0615 study regarding the addition of bevacizumab (monoclonal antibody targeting VEGF) with chemoradiation for nasopharyngeal carcinoma was made available. The phase II study demonstrated feasibility for adding bevacizumab to standard chemoradiation in treating nasopharyngeal carcinoma, and suggested that there may be a slight benefit in the form of delaying progression of subclinical distant disease (27). However the overall results with bevacizumab so far have been far from satisfactory, and this is despite the great zeal that existed for integrating bevacizumab into anti-cancer regimens a decade ago (28).

The toxicity and questionable efficacy of bevacizumab has now led to the quest for newer anti-angiogenic agents. The antiangiogenic effects of small molecule tyrosine kinase inhibitors such as sunitinib, sorafenib and pazopanib has roused interest, but their efficacy is yet to be established, especially in conjunction with RT.

The discovery of RGD (arginine-glycine-glutamine) as an αVβ3 integrin antagonist has opened another window of opportunity. Since RT in itself up-regulates αVβ3 integrin in tumor endothelial cells, despite tumour cell kill RT can unintentionally promote angiogenesis mediated progression/metastases in tumours such as glioma. The development of integrin antagonists such as the RGD peptidomimetic agent S247 and the cyclic RGD pentapeptide cilengitide may help enhance RT outcomes by antagonizing RT induced αVβ3 integrin upregulation (29). However, as with all anti-angiogenic agents, optimism must be guarded until when efficacy can be proven beyond doubt.

**Prospects with newer conventional chemotherapy and targeted therapy agents**

The expected increments in outcomes with the combinations of RT with conventionally delivered ‘newer’ chemotherapy agents (e.g., docetaxel, pemetrexed, gemcitabine) and molecular targeted therapies (e.g., cetuximab, nimotuzumab, erlotinib) are small, even if statistically significant. This is in contrast to the prospects to dramatic improvements potentially feasible with
novel agents and newer methodology available at the horizon, for example with the use of gold nanoparticles as radiosensitizers, or with the use of immunological methods of enhancing RT. This editorial review does not consider these currently existing agents as ‘novel’ agents, and thus, little emphasis is placed upon these.

Utilizing low dose radiation as a chemopotentiator

Conventionally, RT is delivered with dose-fraction sizes of 1.8–3 Gy per fraction. Lower doses per fraction are generally avoided given the undue prolongation of overall treatment time, as well as the prediction of lesser cell kill at lower fraction sizes. It was, however, observed that cell killing at doses <1 Gy is greater than that predicted by the linear-quadratic model, hence the nomenclature ‘low dose hyperradiosensitivity’. It is postulated that this phenomenon is present below the threshold dose that would be required to initiate cellular radiation response mechanisms. In particular, a radiation inducible ATM gene dependent G2 phase checkpoint was found to have a threshold activation dose <0.4 Gy. Thus at such low doses, hyperradiosensitivity occurs due to failure of cell cycle arrest of these radiation-damaged G2-phase cells (30-32).

Despite the enhanced cell killing with low-dose fractionated radiotherapy (LDFRT), it would not be practical to utilize LDFRT alone as a treatment regimen, since the entire course of RT would be likely to be long enough to cause accelerated tumour cell repopulation to negate any cell killing. Hence, an innovative approach has been experimented, that involving the use of LDFRT to enhance chemotherapeutic outcomes.

Various studies have utilized LDFRT to potentiate various regimens of chemotherapy in difficult clinical situations. For example, LDFRT was used to potentiate chemotherapy with pemetrexed in recurrent lung carcinoma, and a dramatic enhancement in response rates was observed (33). LDFRT when used with gemcitabine in pancreatic carcinoma was associated with promising response rates (34). LDFRT was also observed to enhance response rates in head and neck carcinoma when used with neoadjuvant chemotherapy with paclitaxel and carboplatin (35). Among patients with breast carcinoma, when LDFRT was used with neoadjuvant chemotherapy, there was very good tolerability, and a good pathological complete response (pCR) rate. Among those patients who did not attain pCR, a fibrotic reaction was found to encase the residual tumour, which could potentially be inhibitory for residual cells to proliferate or metastasize (36).

Despite very good outcomes without any noticeable additional toxicity with the use of LDFRT as a chemopotentiator, it is rather unfortunate that very few clinical trials have focused upon its prospects. It is hoped that future research will help us refine the logic and the technicalities associated with the use of LDFRT as a chemopotentiator.

Summary and conclusions

While the prospects of promising modalities and methods to improve radiotherapeutic outcomes have been touched upon in this review, it must be said that there could be, and there will be many more innovations which could brighten up prospects of better outcomes for cancer patients (Table 3). The development very precise technologies of RT delivery, combined with the discovery of newer radiopharmaceuticals for positron emission tomography for functional imaging may enable extremely intelligent and biologically adaptive treatment delivery. The discovery of newer radioprotectors is largely welcome too, given that the existing radioprotector namely amifostine holds limited efficacy in limited sites. Also, advances in epigenetics will inevitably be attempted to augment RT outcomes.

Going by existing trends, it is unfortunate that very little interest and funding is dedicated to studies using chemotherapy to enhance RT (37). It must also be remarked that despite extremely good prospects with technologies such as high-Z radiosensitizers, and despite very promising results with initial trials with low dose hyperradiosensitivity for chemopotentiation, there has been very few trials to continue the progress (this is in sharp contrast to the amount of funding and emphasis received by futile trials involving antiangiogenic therapies). It can only be hoped that in the future, trials dedicated to chemically enhance RT will receive their due share of funding.

While clinical research follows a strict rational approach for conceptual development, occasional ideas may be found through ‘out of the box, lateral thinking’, too. Too much of adherence to current standards may be harmful, as potential discoveries are lost due to our staunch obsession with the present, and neglect for the future. All said and done, progress, big or small can only be expected to be achieved through efforts, and with an open mind. While the world remains obsessed with the practise of ‘evidence based medicine’, it can only be hoped that new innovations do get their legitimate share of optimism and emphasis.
Table 3 ‘Chemical enhancement’ of radiotherapy: summary of future prospects

<table>
<thead>
<tr>
<th>Modality</th>
<th>Remarks</th>
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<tbody>
<tr>
<td>Nanoparticle bound chemotherapy as radiosensitizers</td>
<td>Very high prospects for revolutionizing radiation based therapy</td>
</tr>
<tr>
<td>Nanoparticle bound high-Z radiosensitizers</td>
<td>Groundbreaking potential can be unlocked</td>
</tr>
<tr>
<td>Radiotherapy to guide nanoparticle bound chemotherapy to tumors</td>
<td>A novel way to use radiotherapy to paint tumor so as to guide nanoparticle bound chemotherapy to tumor</td>
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<tr>
<td>Metronomic chemotherapy concurrently with radiotherapy</td>
<td>Very useful, especially in patients unable to tolerate conventional concurrent chemotherapy. May hold additional benefits due to antiangiogenic and immunomodulatory properties</td>
</tr>
<tr>
<td>Concurrent chemotherapy with newer agents such as taxanes</td>
<td>Already in clinical use. Small outcome improvements. Toxicity concerns</td>
</tr>
<tr>
<td>Concurrent therapy with molecular targeted monoclonal antibodies and tyrosine kinase inhibitors against growth factor receptors</td>
<td>Already in use. Newer agents actively being investigated</td>
</tr>
<tr>
<td>Anti-CTLA4 and anti-PD-1 monoclonal antibodies with radiotherapy</td>
<td>Synergy may exist. Abscopal responses can be expected</td>
</tr>
<tr>
<td>Antiangiogenic therapy with radiotherapy</td>
<td>Existing agents such as bevacizumab hold questionable efficacy with considerable toxicity. Newer agents awaited</td>
</tr>
<tr>
<td>Use of low dose radiation as a chemopotentiator</td>
<td>The phenomenon of hyperradiosensitivity at low doses to enhance chemotherapy outcomes has been experimented with promising results</td>
</tr>
<tr>
<td>Radioprotectors to widen therapeutic window</td>
<td>Currently, amifostine is the only approved radioprotector, however, has limited activity, confined to limited tissues. Newer radioprotectors yet to be discovered. Concerns of tumor protection must be addressed beyond doubt</td>
</tr>
<tr>
<td>Integration of newer radiopharmaceuticals for better radiotherapy targeting and delivery</td>
<td>Myriad new radiopharmaceuticals utilizing various targets have already revolutionized cancer diagnosis and treatment. Further refinements expected</td>
</tr>
<tr>
<td>Adoptive immunotherapy, viral vectors, cyclin dependent kinase inhibitors, and other novel therapies to enhance radiotherapy</td>
<td>Ongoing in various preclinical/early-clinical trials</td>
</tr>
<tr>
<td>Epigenetic manipulation to enhance radiotherapy outcomes</td>
<td>Yet to be explored</td>
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Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

References


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