Recent Advances in Combined Modality Therapy

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Key Words. Radiation • Chemotherapy • Targeted therapy • Radiosensitization

Disclosures: Michelle L. Mierzwa: None; Mukesh K. Nyati: None; Meredith A. Morgan: None; Theodore S. Lawrence: None.

The content of this article has been reviewed by independent peer reviewers to ensure that it is balanced, objective, and free from commercial bias. No financial relationships relevant to the content of this article have been disclosed by the authors or independent peer reviewers.

ABSTRACT

Combined modality therapy emerged from preclinical data showing that carefully chosen drugs could enhance the sensitivity of tumor cells to radiation while having nonoverlapping toxicities. Recent advances in molecular biology involving the identification of cellular receptors, enzymes, and pathways involved in tumor growth and immortality have resulted in the development of biologically targeted drugs. This review highlights the recent clinical data in support of newer generation cytotoxic chemotherapies and systemic targeted agents in combination with radiation therapy. The Oncologist 2010;15:372–381

INTRODUCTION

Radiation therapy (RT) is used to control locally confined tumors with organ preservation. The ability of radiation alone to control soft tissue tumors is often limited by the tumor volume or the surrounding normal tissue tolerance to radiation. Cytotoxic chemotherapy drugs and biologic agents have been given before, during, or after RT in order to improve tumor responses. Neoadjuvant or induction chemotherapy can be employed to decrease micrometastases and decrease tumor size prior to RT with the hopes of improving tumor control with RT and/or decreasing the amount of normal tissue irradiated. Adjuvant chemotherapy is given after RT primarily to decrease systemic micrometastases. Concurrent chemoradiotherapy (chemo-RT) emerged from preclinical data showing that carefully chosen drugs could enhance the sensitivity of tumor cells to RT while having nonoverlapping toxicities. This spatial cooperation was the initial rationale for combining chemotherapy and RT, in which each agent had an independent mechanism of action at a different anatomic target. Since that time, concurrent chemoradiation has been used to enhance radioresponse locally within the primary tumor. Here, we review the recent clinical data in support of newer generation cytotoxic chemotherapies and systemic targeted agents in combination with RT.

Combined modality therapy enhances RT or systemic therapies alone at both the tissue and cellular levels. At a tissue level, radiation can increase vascular permeability and allow greater drug delivery to the tumor as well as increase drug concentration by promoting drug retention within the tumor. At a cellular level, systemic therapies can enhance radiation sensitivity by inhibiting DNA repair mechanisms, enhancing oxygen radical formation to promote DNA double strand breaks, inhibiting progression...
through the cell cycle to lock cells in a radiosensitive phase, inducing apoptosis, and inhibiting cellular signaling cascades. Recent advances in molecular biology involving the identification of cellular receptors, enzymes, and pathways involved in tumor growth and immortality have resulted in the development of biologically targeted drugs. The use of targeted therapies in conjunction with RT promises to enhance the therapeutic ratio by increasing the efficacy of RT without significantly increasing treatment-related side effects. As individual tumor mutations and molecular markers are better understood, it may also be possible to improve patient selection by determining which patients will benefit from a given agent. (See Table 1.)

It is appropriate to have an article on combined modality therapy in this issue, because Dr. Glatstein (or Eli, as we shall call him; he has never stood for the formality of being called Dr. Glatstein during the standard work day) felt that an important part of the reason he had been brought to the National Cancer Institute (NCI) was to “marry NCI chemotherapy with Stanford radiation therapy.” When the senior author started his residency in radiation oncology with Eli in 1984, “combined modality” for lymphomas, a major emphasis at the NCI, meant chemotherapy first, followed by RT. We had a particularly intensive study for advanced-stage follicular lymphoma that involved full-dose chemotherapy with a nine-drug myelosuppressive regimen followed by total nodal radiation. After all that chemotherapy, the patients’ counts would drop after the dose delivered by simulation and would vaporize after the first week of treatment. It would take us months to get patients through this. In contrast, local toxicities (esophagitis and pneumonitis) dominated the treatment of limited-stage small cell lung cancer with concurrent aggressive chemotherapy and RT. Eli was among the early users of “adaptive therapy.” Treatment with chemoradiation would start on Monday, and we would resimulate patients every Friday in order to shrink the fields. (The total course was a spicy 40 Gy in 15 fractions with concurrent doxorubicin-containing chemotherapy!) Combined modality therapy was superior, but there was significant morbidity and even mortality from treatment.

**Cytotoxic Chemotherapy**

**The Platinos**

Cisplatin is one of the most commonly used chemotherapy agents for radiosensitization. It forms both inter- and intrastrand DNA adducts that produce single-strand breaks when removed by DNA mismatch repair processes. These single-strand breaks can be converted to lethal double-strand breaks by radiation. Thus, mismatch repair defective cells are not radiosensitized by cisplatin and carboplatin [1]. Phase III trials have proven longer survival times for patients treated with concurrent cisplatin-based chemo-RT than for those treated with RT alone in cervical cancer, head and neck cancer, non-small cell lung cancer (NSCLC), and esophageal cancer.

Oxaliplatin is a third-generation platinum derivative that has been found to be a potent radiosensitizer [2]. It is a diaminocyclohexane that causes DNA adduct formation and prevents DNA synthesis [3]. Although oxaliplatin appears to cause DNA adduct formation like other platinum derivatives, oxaliplatin adducts are not repaired by the same DNA mismatch repair systems as other platinum adducts. Oxaliplatin, therefore, has been found to have activity in cisplatin- and carboplatin-resistant cells [4, 5]. Oxaliplatin results in longer survival times in metastatic colorectal cancer patients in combination with 5-fluorouracil (5-FU) than with 5-FU alone [6]. Oxaliplatin is under investigation in phase I/II studies in combination with RT and 5-FU or capecitabine for the neoadjuvant treatment of rectal cancer patients, with an 81% rate of R0 resection. The pathologic complete response rate has been in the range of 8%–26%, with rates of grade 3 or 4 diarrhea of 16%–38% [7–9]. Phase III studies need to be conducted prior to endorsing the routine use of oxaliplatin with concurrent RT.

| Therapy/Agent | Disease Sites                                      |
|---------------|----------------------------------------------------|
| **Cytotoxic chemotherapies** |                                      |
| Oxaliplatin   | Rectal cancer                                      |
| 5-FU/capecitabine | Rectal cancer, pancreatic cancer                   |
| Gemcitabine   | Pancreatic cancer, NSCLC                           |
| Paclitaxel    | NSCLC, SCC of head and neck                        |
| Temozolomide  | High grade glioma, brain metastases                |
| **Targeted therapies** |                                      |
| Cetuximab     | NSCLC, SCC of head and neck                        |
| Gefitinib     | NSCLC, SCC of head and neck                        |
| Erlotinib     | Rectal cancer, GBM, NSCLC                          |
| Bevacizumab   | Rectal cancer, GBM, NSCLC, pancreatic cancer       |

Abbreviations: GBM, glioblastoma multiforme; NSCLC, non-small cell lung cancer; SCC, squamous cell carcinoma.

Table 1. Disease sites for which newer cytotoxic chemotherapy or targeted agents are currently under investigation concurrently with radiation
5-FU and Capecitabine

5-FU is a pyrimidine analog of uracil and functions as an antimetabolite. Its radiosensitizing effects are a result of its ability to prevent DNA synthesis through inhibition of thymidylate synthase [10], rather than its RNA-dependent effects. The combination of 5-FU and RT was superior to RT alone in randomized trials for patients with head and neck and gastrointestinal (GI) cancers, including esophageal, gastric, pancreatic, and rectal cancers.

Capecitabine is an oral 5-FU prodrug; it is converted to 5-FU by the enzyme thymidine phosphorylase (TP). TP catalyzes the mutual transformation of the pyrimidine nucleosides thymidine and thymine in nucleic acid metabolism and also converts the FU-based drugs, including capecitabine, into active 5-FU. It is, therefore, a limiting factor of the mechanism of action for capecitabine. A proposed mechanism of selectivity is the preferential conversion in the liver, which is relatively resistant, and at the tumor site, because these tissues often have higher TP activity than normal tissue.

RT was reported to lead to better efficacy with capecitabine through tumor-associated induction of TP, especially within the radiation portal. However, this was not found to be consistently true in other preclinical studies of colon cancer cells [11], in tumor xenografts [12], or in a phase I dose-escalation study from the University of Alabama [13]. Although these results support the concurrent use of capecitabine and RT in pancreatic cancer, there appear to be additional genes (other than TP) associated with response to capecitabine alone and with RT.

Because of its ease of administration compared with continuous infusion 5-FU, attempts have been made to substitute capecitabine for infused 5-FU in combination with RT in pancreatic cancer and rectal cancer patients [14, 15]. We are unaware of any head-to-head comparisons of infused 5-FU versus capecitabine, but our overall impression is that these are probably approximately equivalent.

Gemcitabine

2′,2′-Difluoro-2′-deoxycytidine, or gemcitabine, requires intracellular phosphorylation in a rate-limiting step by deoxycytidine kinase to form its active metabolites difluorodeoxycytidine diphosphate (dFdCDP) and difluorodeoxycytidine triphosphate (dFdCTP). These metabolites of gemcitabine produce two distinct mechanisms of action contributing to gemcitabine’s radiosensitizing and cytotoxic properties, respectively. DFdCDP is a direct inhibitor of ribonucleotide reductase and thus inhibits deoxynucleotide triphosphate synthesis whereas dFdCTP is incorporated into DNA, both leading to the inhibition of DNA synthesis. Early preclinical studies demonstrated that gemcitabine was an effective radiosensitizer in a variety of cell types [16, 17]. Subsequent studies investigating the mechanism(s) of radiosensitization suggested that dATP pool depletion (via ribonucleotide reductase inhibition) as well as redistribution of cells into the early S-phase of the cell cycle underlie gemcitabine-mediated radiosensitization [18]. Based on the observed preclinical radiosensitization by gemcitabine as well as the clinical efficacy of gemcitabine as a single agent, gemcitabine has been combined with RT in the clinic to treat many solid tumor types.

Gemcitabine has become the standard of care for metastatic pancreatic cancer [19]. We feel it has become a standard component for locally advanced disease. In this latter group, a recent trial phase III trial of 74 patients with resectable pancreatic cancer randomized patients to either gemcitabine or gemcitabine and concurrent RT, showing a statistical survival advantage for patients in the chemo-RT arm (11 months versus 9.2 months; \( p < .04 \)), but higher GI and hematologic toxicity rates in the chemo-RT arm [20]. A French phase III trial of induction chemo-RT followed by gemcitabine was compared with gemcitabine alone, showing the chemo-RT arm to be more toxic and less effective than gemcitabine alone. The 1-year overall survival rates were 32% versus 53% favoring gemcitabine alone [21]. In that trial, however, patients in the chemo-RT arm had significantly shorter survival than historical controls. This could be because concurrent, high-dose gemcitabine and RT is toxic when large radiation fields are used [22]. However, the use of conformal radiation to the primary tumor site without elective nodal radiation permits the safe administration of full systemic doses of gemcitabine or gemcitabine plus oxaliplatin [23, 24]. Although Eli has been noted to say, “Local control does not guarantee immortality,” local failure does occur in these patients [25] and can be the cause of death. This has motivated our recent attempts to dose escalate radiation using intensity-modulated RT with full-dose gemcitabine. Results are promising, with a median survival duration of 23 months [26].

In a phase III trial of adjuvant therapy for locally advanced resected pancreatic cancer, the addition of adjuvant gemcitabine after 5-FU and concurrent RT showed longer survival than with 5-FU and RT alone for patients with pancreatic head tumors [27]. Although a phase I/II study showed that concurrent full-dose gemcitabine and RT can be given safely as adjuvant treatment [28], it is not known whether this combination is superior to the more commonly used 5-FU or capecitabine plus RT combination.

Phase I/II studies have also evaluated the role of gemcitabine (alone or in combination with a platinum or paclitaxel) and concurrent RT in locally advanced NSCLC
patients. Preliminary data have shown high rates of pulmonary toxicity, although gemcitabine, at a dose of 150 mg/m², was tolerated in the setting of three-dimensional (3D) RT planning [29–33]. Caution is advised when gemcitabine is given with concurrent RT because of gemcitabine’s enhanced radiosensitization effects in the lung and esophagus, and appropriate chemotherapy doses have not been well established.

**Paclitaxel**

The taxanes are mitotic spindle inhibitors that bind to the N-terminal amino acid in β-tubulin and stabilize tubulin polymers, thereby promoting microtubule assembly and inhibiting disaggregation [34]. This action causes cellular arrest in G2/M, the most radiosensitive phase of the cell cycle. Paclitaxel is the most commonly used taxane in combination with RT and appears to act by causing mitotic arrest and inducing apoptosis, and also may also induce reoxygenation within the tumor [35, 36].

Paclitaxel has been used in combination with carboplatin currently with RT in the setting of locally advanced NSCLC because it was thought to have a favorable toxicity profile. Initially, concurrent chemo-RT with cisplatin and etoposide was found, in trials, to be superior to sequential therapy in unresectable NSCLC patients [37]. Phase II data then showed that RT with low-dose weekly concurrent paclitaxel plus carboplatin and induction or paclitaxel plus carboplatin was feasible [38, 39]. Specifically, the Cancer and Leukemia Group B (CALGB) initially evaluated the combination of induction carboplatin and paclitaxel for two cycles followed by low-dose weekly concurrent chemotherapy with RT. They found a median survival time of 15.1 months, and the trial demonstrated the feasibility of this regimen [39]. A phase III trial, CALGB 39801, was then completed, in which all patients received low-dose weekly carboplatin and paclitaxel with concurrent RT to 66 Gy and were randomized to two cycles of induction chemotherapy. Both arms of that trial showed disappointing results, with a median survival time of 11–13 months, demonstrating that this was not an efficacious regimen [40]. The CALGB 30105 trial evaluated induction chemotherapy followed by concurrent carboplatin, paclitaxel, and RT or concurrent carboplatin, gemcitabine, and RT in stage IIIA/IIIB NSCLC patients. Both arms required 3D RT planning, and all patients received 74 Gy. They found that patients in the paclitaxel arm had a median overall survival time of 24 months whereas the gemcitabine arm was closed early secondary to a high rate of grade 4 or 5 pulmonary toxicity. The hypothesis generated by that trial is that the longer median survival time seen in the carboplatin–paclitaxel arm was a result of the higher RT dose [36].

To date, carboplatin and paclitaxel have not been compared with cisplatin-based chemo-RT, which continues to be considered the standard as definitive or neoadjuvant treatment in NSCLC patients.

In locally advanced and unresectable head and neck cancer, paclitaxel has been used concurrently with RT. The Eastern Cooperative Oncology Group (ECOG) E2399 study was a phase II trial of induction carboplatin plus paclitaxel followed by concurrent carboplatin, paclitaxel, and RT for patients with resectable stage III/IV squamous cell carcinoma of the larynx and oropharynx. That study showed an 81% organ preservation rate overall at 2 years, with a higher organ preservation rate in oropharynx patients, and the regimen was well tolerated [41]. Another phase II trial in unresectable head and neck cancer patients, including all sites, showed that RT with concurrent weekly carboplatin plus paclitaxel was well tolerated, but 42% of patients (n = 50) had some component of locoregional failure at 5 years, and the overall survival rate was 35% [42]. Chougule et al. [43] conducted a trial of concurrent carboplatin plus paclitaxel and RT, demonstrating a 91% response rate with a 26% locoregional failure rate in locally advanced head and neck cancer patients. This regimen, however, required that 31% of patients be hospitalized because of toxicity. Salama et al. [44] conducted a phase I/II trial at the University of Chicago using induction carboplatin and paclitaxel followed by concurrent carboplatin, paclitaxel, fluorouracil, and hydroxyurea and twice daily RT in stage III/IV head and neck cancer patients, showing a high locoregional control rate and overall survival rate of 91% and 62%, respectively, at 5 years. Ninety-one percent of patients experienced grade 3 or 4 acute toxicity during chemo-RT. As in locally advanced NSCLC patients, concurrent carboplatin and paclitaxel regimens with RT have not been prospectively compared with cisplatin-based chemo-RT in locally advanced head and neck cancer patients.

**Temozolomide**

Temozolomide is an orally administered alkylating agent that induces apoptosis by methylation of DNA at nucleophilic sites, causing the formation of O6-methylguanine. This DNA adduct induces futile cycling of the mismatch repair pathway, but the adduct can also be repaired by the enzyme O6-methylguanine-DNA methyltransferase (MGMT). Patients with high levels of MGMT methylation, which indicates that the gene has been silenced, have longer progression-free and overall survival times after treatment with alkylating agents [45].

In preclinical studies, temozolomide was found to have schedule-dependent activity against recurrent high-grade gliomas and advanced melanoma, to have oral bioavailabili-
ity, and to penetrate the central nervous system [46, 47]. Its use in the concurrent setting with RT is largely based on early clinical data supporting a small advantage over current treatment at that time. A large phase II trial of temozolomide versus procarbazine in recurrent glioblastoma multiforme (GBM) patients demonstrated a significant 6-month progression-free and overall survival advantage for patients treated with temozolomide as a single agent, with similar toxicity [48]. Following this, a pilot phase II trial of concurrent temozolomide and RT followed by adjuvant temozolomide was then conducted, demonstrating a promising overall survival rate at 2 years of 31% [49]. Concurrent temozolomide and RT followed by temozolomide alone in GBM was then studied in a large phase III trial and found to result in longer survival versus RT alone [50], thereby establishing the standard of care.

Temozolomide with concurrent RT is currently being investigated for patients with high-risk, low-grade gliomas (Radiation Therapy Oncology Group [RTOG] 0424 trial). Additionally, temozolomide is being examined for use with concurrent RT in the setting of brain metastasis. A phase II single-institution trial of whole-brain RT and concurrent as well as adjuvant temozolomide in 27 patients with at least one brain metastasis from breast cancer or NSCLC revealed complete and partial response rates of 7% and 41%, respectively. The regimen was well tolerated [51]. RTOG 0320 is a phase III trial that is under way investigating whole-brain RT and stereotactic radiosurgery alone versus the same radiation with concurrent temozolomide or erlotinib.

**Molecular Targeted Therapies**

**Epidermal Growth Factor Receptor–Directed Therapies**

The human epidermal growth factor receptor (EGFR) family includes EGFR, human epidermal growth factor receptor 2 (HER-2), HER-3, and HER-4. (See Figure 1.) EGFR family members have been identified to play a critical role in development, progression, and resistance to chemo-RT in NSCLC, colorectal cancer, pancreatic cancer, squamous cell carcinoma of the head and neck, brain cancer, and breast cancer. Additionally, EGFR is either overexpressed or hyperphosphorylated in the majority of cancers of epithelial origin. Several clinical studies have suggested that EGFR expression is a powerful prognostic factor for response to both RT and chemo-RT. In a retrospective review, Kumar et al. [52] found that a high EGFR expression level predicts poor response to therapy and poor overall survival in head and neck cancer patients. Similarly, Pivot et al. [53] found that lower EGFR levels predicted better disease-free survival in patients with laryngeal and hypopharyngeal cancers treated with induction chemotherapy followed by RT. The idea that blocking these receptors produces selective response in tumors driven by these molecules has led to the development of several monoclonal antibodies (mAbs) and small molecules that inhibit receptor kinase activity [54].

**Cetuximab**

Cetuximab binds to the EGFR with approximately tenfold greater affinity than the mouse mAb C225 from which it is derived. Cetuximab not only blocks EGF-induced autophosphorylation of the EGFR but also induces its internalization and degradation, which may contribute to the outcome of treatment. It has also been reported that cetuximab inhibits nuclear translocation of EGFR, which may be particularly effective in combination with RT, because nuclear EGFR can induce DNA protein kinase activity, leading to efficient DNA repair. Therefore, inhibition of EGFR is considered an excellent target for radiosensitization. Still, many aspects of this therapy have yet to be elucidated. For example, only modest radiosensitization in cultured cells can be achieved via EGFR inhibitors, and even this requires prolonged inhibition of EGFR phosphorylation. However, xenograft data show much more promising results; the cause of this difference is not yet clearly understood. It has been postulated that, in addition to blocking direct effects on EGFR, cetuximab may have effects on cell cycle redistribution, angiogenesis, and immune reaction. This argument is supported by clinical findings suggesting that: (a) cetuximab can produce a response in tumors with negative EGFR staining by immunohistochemistry and (b) the strength of the EGFR staining does not predict response to EGFR inhibitors combined with chemotherapy. Although we cannot yet predict patient response based on the presence or absence of the target itself, recent findings suggest that patients carrying a *KRAS* mutation do not benefit from treatment with anti-EGFR agents.

A randomized trial in locally advanced head and neck cancer patients demonstrated that patients receiving the combination of RT and cetuximab had a statistically significant better locoregional disease response without significantly worse acute toxicities [55]. This is the first study to show that the combination of a molecularly targeted therapy with RT can result in better local control and survival than with RT alone. Based on this success, the addition of cetuximab to concurrent chemo-RT as definitive treatment in locally advanced head and neck cancer patients is currently being investigated in the RTOG 0522 study. In the postoperative setting, RTOG 0234 is a phase II trial now closed to accrual that evaluated cetuximab with concurrent
chemo-RT for patients with head and neck cancer and high-risk pathologic features.

In NSCLC patients, preliminary phase II studies have shown that cetuximab can be safely administered with RT and that this does not appear to lead to higher rates of acute treatment toxicities. The SCRATCH trial, a phase II trial in stage III NSCLC patients treated with weekly cetuximab and thoracic RT to 64 Gy demonstrated acceptable toxicity for this regimen [56]. The Non-small cell lung cancer, Erbitux And Radiotherapy (NEAR) trial is an ongoing, phase II, feasibility trial of concurrent locoregional RT and cetuximab in stage III NSCLC patients [57].

Cetuximab in combination with gemcitabine and RT has been assessed in pancreatic cancer. A randomized phase II trial was conducted in inoperable patients receiving gemcitabine, cetuximab, and RT, who then were randomized to adjuvant gemcitabine or adjuvant gemcitabine plus cetuximab. No statistical difference was seen between arms; the 2-year overall survival rate was 20% [58]. This result is not surprising; the presence of mutated RAS in >90% of pancreatic cancer cases suggests that EGFR inhibition is unlikely to be successful [59].

**Gefitinib and Erlotinib**

Gefitinib and erlotinib are small molecule tyrosine kinase inhibitors used clinically. They inhibit unregulated EGFR signaling.
signaling by binding to the intracellular ATP binding site, which prevents tyrosine kinase activity.

Recent data have demonstrated that schedule plays a critical role in the interaction between gefitinib and erlotinib with chemotherapy and with RT. When given prior to gemcitabine and cisplatin, gefitinib and erlotinib antagonize chemotherapy toxicity, whereas the small molecule tyrosine kinase inhibitors produce substantial synergy when given after cytotoxic chemotherapy [37, 60]. The molecular mechanism for this schedule dependence is now understood. Both gemcitabine and cisplatin cause initial EGFR phosphorylation, followed by EGFR degradation and cell death. Therefore, giving an EGFR inhibitor prior to chemotherapy blocks this initial phosphorylation, inhibits degradation, and increases survival. Indeed, this is likely the explanation for the negative results of the Iressa NSCLC Trial Assessing Combination Treatment (INTACT)-1 and INTACT-2 studies [61, 62], and a recent clinical trial that prospectively used chemotherapy followed by small molecule tyrosine kinase inhibition has produced promising results [63]. In contrast, administration of an EGFR inhibitor after chemotherapy-induced EGFR phosphorylation potentiates degradation and the subsequent cytotoxicity. A corollary of the hypothesis that EGFR degradation causes cytotoxicity is that a proteosome inhibitor such as bortezomib might antagonize chemo-RT by inhibiting EGFR degradation. In addition, we have recently shown that EGFR-driven log-phase and plateau-phase cells have a dramatically different response to an EGFR inhibitor combined with radiation. A brief (<2 hours) pretreatment with erlotinib can protect plateau-phase EGFR-expressing cells from radiation by inhibiting EGFR activation, thus preventing cells from being driven into a fatal S phase after radiation [64]. In contrast, long-term exposure (24 hours) of log-phase cells to EGFR inhibition ultimately arrests cells in late G1, which is a relatively sensitive phase of the cell cycle. These findings demonstrate the importance of schedule and cell cycle in the response to combination treatment with RT, chemotherapy, and EGFR inhibitors.

Gefitinib has been used in phase I/II trials with RT in NSCLC and head and neck cancer patients. In stage III NSCLC patients, a phase II trial of induction carboplatin, paclitaxel, and irinotecan followed by chemo-RT to 74 Gy with concurrent carboplatin, paclitaxel, and gefitinib (250 mg daily) showed that the regimen was well tolerated, but disappointing results of a 24% partial response rate and median OS time of 16 months were reported [65]. In stage III and IV head and neck cancer patients, a phase II trial using two cycles of neoadjuvant carboplatin and paclitaxel followed by concurrent RT, 5-FU, hydroxyurea, and gefitinib showed an estimated 2 year overall survival rate of 83% with a low rate of grade 4 toxicities [66].

Erlotinib has been combined with RT in the phase I/II setting in rectal cancer, glioma, and NSCLC patients, for which EGFR expression is commonly seen. Erlotinib was tried in a dose-escalation study of patients with locally advanced or recurrent rectal cancer with concurrent RT, demonstrating excellent local control but only a 5% pathologic complete response rate at a maximum-tolerated dose of 100 mg daily for 45 days [67]. In GBM patients, the North Central Cancer Treatment Group reported a phase I/II trial in which patients received concurrent RT, temozolomide, and erlotinib. The regimen was well tolerated, but the median survival time of 15.3 months did not show an advantage over historic controls in the European Organization for Research and Treatment of Cancer 26981 study [52].

It is noteworthy that despite the high frequency of mutant RAS in pancreatic cancer and lack of difference in response with cetuximab, erlotinib is the only molecularly targeted agent shown thus far to result in statistically longer survival in patients with advanced disease treated with gemcitabine [69].

**Bevacizumab**

Expression of angiogenic factors is required for tumor growth, and the most commonly studied factor is vascular endothelial growth factor (VEGF). Bevacizumab is a monoclonal antibody that binds to the VEGF receptor on the extracellular surface and is thought to normalize tumor vasculature, thereby potentially allowing enhanced oxygen and chemotherapy drug delivery locally. In rectal cancer, bevacizumab has been found to have antivascular effects, decreasing perfusion and intrastitial fluid pressure within the tumor [70]. In locally advanced rectal cancer, phase I and II trials have been conducted to evaluate the addition of bevacizumab to neoadjuvant therapy. When bevacizumab was added to neoadjuvant capecitabine and RT for locally advanced rectal cancer, the pathologic complete response rate was 32%, with a 72% sphincter preservation rate and no grade 3 GI or hematologic toxicities [71].

Bevacizumab is also under investigation with concurrent RT in the treatment of GBM patients. Preliminary phase II data with concurrent temozolomide and RT have shown that the regimen is well tolerated without a higher rate of acute toxicity [72].

Bevacizumab has also been evaluated with gemcitabine and RT in locally advanced pancreatic cancer patients. This regimen is now approached with caution, however, because of high toxicity manifest as surgical complications and ulceration or bleeding in the radiation field [73–75].

Despite some encouraging results for bevacizumab plus
RT in small trials, there is no current recognized role for this therapy off protocol.

**Future Directions**

How can we best build on our knowledge of chemo-RT to design more effective combinations (with and without targeted therapies)? First, we feel that preclinical data may guide us in optimizing therapy and patient selection. For instance, preclinical studies show that it is possible for targeted therapies that block cell cycle progression (such as cetuximab) to antagonize some chemotherapies that depend on cell cycle progression (such as gemcitabine) [76]; it is possible that such effects may explain some of the negative results seen in adding targeted therapies to lung cancer treatment [77]. In another example, targeted therapies have typically been tested in unselected populations or, even worse, in populations in which preclinical data suggest that the patient population will be resistant to the therapy (erlotinib in pancreatic cancer, which commonly displays RAS mutations). It is possible that the addition of RT could alter signaling pathways so that these issues don’t matter, but this is speculation.

A second possible focus for improvement would be to devote more attention to determining if the target is actually being hit. The senior author learned this directly from Eli during his residency, during which special efforts were made to obtain tumor and bone marrow biopsies to determine incorporation of the radiation sensitizers iododeoxy and bromodeoxyuridine. These studies are difficult, to be sure, but an understanding of how, for instance, cetuximab affects downstream signaling in tumors from even four to six patients might provide us with more understanding about how to optimize this agent in combination with RT or chemo-RT than a randomized phase II trial of 100 unselected patients without pharmacodynamic studies.

Third, we feel that these studies suggest a role for highly conformal RT to enable the use of chemotherapy and targeted therapies. We can’t discuss this without bringing Eli into the picture. Eli has been unfairly characterized as “anti-technology”; in fact, he (and we) are opposed to the mindless application of technology that, sadly, often characterizes our field. Highly conformal therapy, which excludes clinically uninvolved regions, can permit full doses of chemotherapy and targeted therapies to be administered safely. Eli has been known to complain that radiation oncologists and medical oncologists are sometimes engaged in “a battle for the last megakaryocyte.” The increasing use of conformal RT and targeted therapies will, we hope, replace this battle with a collaboration over how to cure patients while treating acne and mild diarrhea.

Finally, we feel that these developments give the opportunity to expand the scope of practice of the radiation oncologist, and, here too, we celebrate Eli’s lead. Eli always told his trainees that we are not “modality salesmen” but we need to be oncologists and physicians. Indeed, he would say that radiation oncologists are the only physicians who are consistently called on to treat both early-stage disease (like the surgeon) and metastatic disease (like the medical oncologist). During his medical oncology training, the senior author was struck by how much of his training was focused on ameliorating the toxicity of chemotherapy. The introduction of far less toxic targeted therapy permits the radiation oncologist to lead efforts to develop new radiation sensitizer strategies, as part of multidisciplinary care, without subjecting patients to severe toxicity.

**Author Contributions**

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