

## Radiation redux

*Could traditional radiation treatments work in concert with immunotherapy to mount a more effective assault on cancer?*

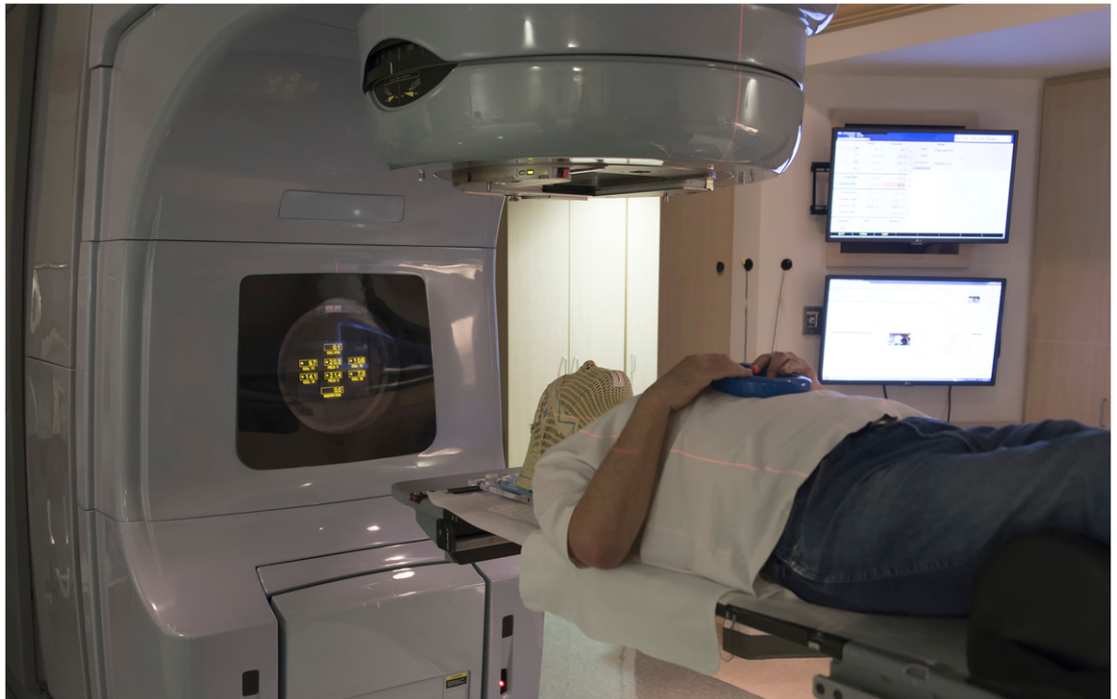
Elie Dolgin, *Science Writer*

When Michael Postow started his fellowship training in medical oncology at the Memorial Sloan Kettering Cancer Center (MSKCC) in 2010, he wasn't familiar with the term "abscopal." Coined decades before he was born and then largely forgotten by physicians, it describes a phenomenon in which radiation delivered at one site in the body has an effect on distant tissues that are, literally, "outside" the "scope" of the radiation blast. "It wasn't something I had ever heard about," Postow says. That changed when a patient at MSKCC had an unexpected response to radiation therapy.

The young woman had been receiving a then-experimental immunotherapy drug, called ipilimumab, to treat her advanced case of melanoma. The drug, now marketed by Bristol-Myers Squibb as Yervoy, blocks a so-called checkpoint protein that normally blunts immune activation. Ipilimumab, in effect, takes the brakes off the immune system, freeing cancer-killing T cells to mount an attack against the tumor.

But this therapeutic strategy only works when there are tumor-targeted T cells in the immune system's tank, otherwise there's nothing to rev into action. That seemed to be the case for Postow's patient. Tumors continued to grow throughout her spleen, in a lymph node in her chest, and near her spine. The pain from the tumor in her back became excruciating.

Postow and his colleagues, led by physician-scientist Jedd Wolchok, chief of the melanoma and immunotherapeutics service at MSKCC, arranged for the woman to receive radiation. They expected to shrink the tumor near her spine and offer some relief from the back pain, nothing more. But scans came back showing tumor regression throughout her body. After more than a year of worsening disease, the patient was finally in remission (1). "That's when all the bells started going off for everyone," Postow says. "Maybe there was something we did with the radiation."



**Fig. 1.** Researchers are combining radiation therapy, here targeting brain cancer, with immunotherapy in hopes of finding a robust strategy for stymying various cancers. Image courtesy of Shutterstock/John Panella.

The MSKCC team's 2012 case report was the first demonstration in a patient that radiation can synergize with a checkpoint inhibitor like ipilimumab. The report showed, says Sandra Demaria, a cancer immunologist at Weill Cornell Medicine, that eliciting a strong enough immune response will lead to the abscopal effect. "From there," Demaria adds, "everyone became interested, and things have progressed fast."

Just 5 years after that initial report, around 100 clinical trials are ongoing to test the mix of radiation and checkpoint inhibitors in patients with cancers of every stripe. The word abscopal is finally on the tongues of oncologists and drug executives everywhere.

"It's energized the field of radiation oncology and broadened the mindset of pharma as well," says William McBride, a tumor immunologist at the University of California, Los Angeles. "It's very early days to know where this is going to go, but it's certainly exciting."

### Joining Forces

Around 60% of all cancer patients receive radiation at some point in their course of treatment, mainly to shrink tumors so drugs can finish them off or just to reduce pain and buy time. However, the therapy itself can be completely curative if given in the earliest stages of the disease, when tumors remain contained to one site in the body. That's why Jonathan Schoenfeld from the Dana-Farber Cancer Institute describes radiation as "one of the most effective forms of cancer treatment we have." But it's mainly local treatment. Adding immunotherapy to the mix, Schoenfeld says, should help give radiation system-wide powers.

Renewed interest in the abscopal effect comes as pharmaceutical companies all jockey to expand the markets for their competing—and lucrative—immunotherapy drugs. One way they hope to bolster both effectiveness and sales is by boosting response rates to these agents.

When taken alone, checkpoint inhibitors generally only work for around 20–30% of patients. To get those numbers up, drug firms are actively testing immunotherapeutic cocktails, either combining checkpoint inhibitors with one another or with other types of treatments, such as cancer-killing viruses, engineered T cells, and radiation.

The radiation-augmented strategy rests on the idea that high-energy blasts of X-rays or other particles act, at least in part, like a natural cancer vaccine. With any vaccine, a bolus of antigen prompts the immune system to recognize the same antigen later and to be on alert to destroy it. The radiation does the same by setting in motion the process of cancer cell death, during which the floundering tumor cells release bits of cancer debris—the antigens—that prime T cells not only for local clean-up but also for an onslaught against any tumor cells elsewhere in the body. Those immune cells just need to be given the chance. Add in a checkpoint inhibitor to unleash them, and they can surge into action.

"The radiation is changing what the immune cells are doing," says Christopher Barker, a radiation oncologist at MSKCC, who collaborated with Postow and Wolchok on the 2012 report (1). "And it's modulating

them in a way that makes them more likely to have anticancer effects."

### Slow Acceptance

This idea is now pretty well accepted, but it wasn't always so. Pathologist Robin Mole of the Medical Research Council's Radiobiology Unit at Harwell, United Kingdom, was clearly proud when he introduced the term "abscopal" in a 1952 lecture. It "conveys the exact meaning required," he noted (2).

Yet, it was a word ahead of its time. Clinicians simply had no use for it. Only rarely did they ever administer localized radiation to cancer patients and then see untargeted tumors disappear as well. Such responses were so uncommon, in fact, that a recent systematic review of the scientific literature by researchers at the Moffitt Cancer Center could find only 46 clinical case reports on the abscopal effect between 1969 and 2014 (3).

So, perhaps it's no surprise that Demaria was also unaware of the abscopal effect when in the early 2000s, shortly after completing her medical training in anatomic pathology at the New York University (NYU) School of Medicine, she started studying the immune

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**—Silvia Formenti**

effects of chemotherapy given to women with breast cancer ahead of surgery. Demaria's research showed that immune cells often infiltrated the tumors of women who responded to the therapy (4).

Silvia Formenti, a radiation oncologist who had joined NYU around that time, had her own investigation testing the benefits of adding radiation to chemotherapy for this exact kind of patient, many of whom had tumors that had spread locally to nearby lymph nodes in the underarm area (5). The therapy was remarkably effective, and when Formenti heard about Demaria's immunological findings, she wondered if radiation might be having a similar immune-stimulating effect.

"It seemed like whatever we were doing locally in the tumor was converting the tumor into an immunogenic hub," recalls Formenti. She didn't dare make the analogy to a vaccine among fellow oncologists, Formenti adds, but that's what she was thinking.

The two women decided to team up to test the idea. Demaria and Formenti implanted mice with breast tumors in both flanks, one to represent the primary cancer, the other a metastatic site. They then irradiated the primary tumor and saw that the distant tumor shrank when the mice also received a growth factor that increases numbers of the antigen-presenting immune cells that activate T cells. But the distant tumor grew unabated in mice that got the drug without radiation (6). The abscopal effect was indeed immune-mediated, contrary to other leading explanations for the distant antitumor effects.

Few in the radiation oncology community initially accepted the conclusions. "They treated us like we





