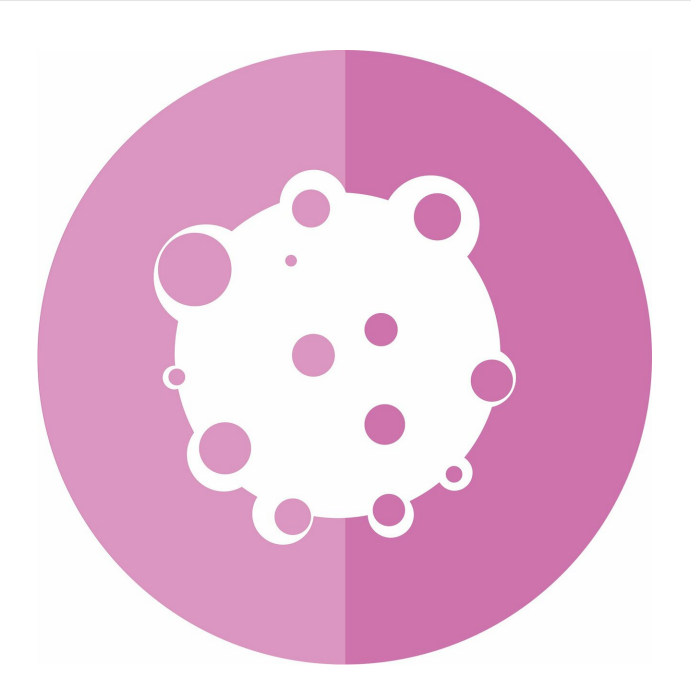


Pre-operative immunotherapy triggers encouraging response in oral cancers

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A new clinical trial suggests that immunotherapy given before other treatments for oral cavity cancers can elicit an immune response that shrinks tumors, which could provide long-term benefit for patients. Findings will be presented at the 2020 Multidisciplinary Head and Neck Cancers Symposium, taking place February 27-29 in Scottsdale, Arizona.

In the randomized phase II trial, two neoadjuvant doses of nivolumab given with or without ipilimumab led to complete or partial tumor shrinkage in most cases and did not delay any patients from continuing on to standard treatment. These promising responses could translate into improved outcomes for patients with an especially difficult and painful type of <u>cancer</u>.

"With roughly three weeks of treatment, we were able to trigger significant tumor regression. In a couple cases, there were complete pathological responses, and in other cases, there was very little tumor left. Both the single drug and the two-drug combination led to visible tumor shrinkage, and, albeit with relatively early follow-up, the majority of these patients have no evidence of disease recurrence," said lead author Jonathan D. Schoenfeld, MD, MPH, senior physician at the Dana-Farber/Brigham and Women's Cancer Center and associate professor of radiation oncology at Harvard Medical School. "Our hope is that even a couple doses of immunotherapy can stimulate an immune response that continues to prevent the cancer from coming back after patients have surgery and other therapy."

The trial enrolled 30 adults newly diagnosed with tumors in their tongue, gums or other part of the mouth; all tumors were stage T2 or higher, and



over half of the patients' cancers had spread to their lymph nodes. After receiving two doses of the PD-1 blocker nivolumab either alone or in combination with a single dose of the CTLA-4 blocker ipilimumab over the course of three weeks, no patients were delayed from surgery, the first component of standard treatment for this disease.

Most patients (52%) experienced clinical reduction of their primary tumor after immunotherapy, and four patients (one in the single-drug group, three in the combination group) had more than 90% pathologic response. While these exploratory results are promising, Dr. Schoenfeld noted that direct comparisons with the current standard of care are needed to determine whether the single-agent or combination therapy can lead to durable responses and improve patient survival. The researchers also want to understand why immunotherapy worked better with some patients than others and identify additional immune targets that could further enhance the treatment.

Twenty-one of the 30 patients experienced side effects possibly related to treatment, and grade 3-4 toxicities for five patients (33%) in the combination group and two patients (14%) in the single-drug group. Dr. Schoenfeld said his team was encouraged by these results; by comparison, for example, more than half of patients experienced serious adverse events with the same combination in trials for high-risk resectable melanoma.

While immunotherapy drugs generally are used after other treatments have failed and a patient's cancer has spread, this study adds to a growing body of research on immunotherapy given prior to surgery for patients with newly diagnosed, curable disease. "The preoperative setting is interesting because patients' immune systems haven't been affected by prior treatment. The tumor is actually in place to serve as a focal point for an immune response, so it may be easier for the body's immune system to recognize and target the <u>tumor</u>," explained Dr. Schoenfeld.



"Oral cavity cancer is a notoriously difficult cancer with high rates of disease recurrence and death, and the side effects from <u>standard</u> <u>treatment</u> tend to be particularly challenging because the treated area is essential for speaking, swallowing and breathing," he continued. "We're excited about moving immunotherapy earlier to treat more of these curative <u>patients</u> and, in the future, possibly reduce how aggressive their other treatments need to be."

Dr. Schoenfeld will present "Neoadjuvant Nivolumab +/- Ipilimumab in Patients with Oral Cavity Cancer" (Abstract 1) today during the symposium's Plenary Session. The study was funded by Bristol-Myers Squibb.

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