When It’s Really Good, We Know It!

A few weeks ago, an extraordinary announcement came from the FDA. Pembrolizumab, a PD-L1 inhibitor already approved for selected disease indications, received accelerated approval for second-line therapy for any unresectable or metastatic tumor demonstrating microsatellite instability–high (MSI-H) or mismatch DNA repair. This is the first time on record that a drug has been approved for a specific tissue phenotype rather than a specific disease.

The classic condition that shows this phenotype is Lynch syndrome. Years ago, Dr. Henry Lynch of Creighton University observed an unusual clustering of colon cancer, as well as an excess of other cancers, in certain families. This was decades before the molecular underpinnings of the disease were understood. And until now, the primary importance of knowing who carried pathologic germline mutations was for screening and early detection, not disease management.

The success of checkpoint inhibitors (antibodies to CTLA-4, PD-1, or PD-L1) has been transformative in some diseases, generally those with a high mutation burden. Some cancers, however, such as colon and pancreatic cancers, are considered “cold tumors,” unresponsive to checkpoint blockade. When I think of how this story unfolded in MSI-H tumors, I almost break into a sweat. Were it not for one unusual patient and some very inquisitive scientists, we might never have gone down this road.

As the story goes, investigators at Johns Hopkins had completed treatment in a cohort of patients with colorectal cancer, with only one response. Not enough, usually, to be a signal. But this patient was special, and it was an exquisite response. The physicians discussing the case were puzzled. A pathologist asked if the patient might have Lynch syndrome. The treating physician noted that he did, and the pathologist remarked that these patients have an unusually dense infiltration of T lymphocytes, a disease hallmark. Therefore, such an individual might be expected to show response. And the rest is history.

This story has several lessons. The first is about team science; the participating pathologist brought information that the average oncologist might not know. That one contribution paved the way for 2 subsequent trials in MSI-H colorectal and noncolorectal tumors. The second is about clinical observation; when you see something extraordinary, ask why. Throughout history, we have learned so much from astute clinicians who took the time to dig deeper and understand better. The third lesson is that the FDA can move quickly. The approval was based on a single-arm trial involving 90 patients with colorectal cancer and 59 patients with 1 of 14 other cancer types. Patients with end-stage cancer experienced responses, even complete responses, and the responses were durable.

Looking back, it was serendipity that allowed that one patient with Lynch syndrome in the early trial to show us the way. Thank goodness for that! Going forward, the FDA will require more confirmative studies. What that confirmation might look like is a bit mind-boggling, so I’ll leave it to the experts. In the meantime, we’ve got a new option for an uncommon syndrome. And that’s a very good thing.

Reference


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