Cryoablation and Immunotherapy for Breast Cancer:
Overview and Rationale for Combined Therapy

Helaina C. Regen-Tuero, BS • Robert C. Ward, MD • William M. Sikov, MD • Peter J. Littrup, MD

From the Department of Diagnostic Imaging, Warren Alpert Medical School of Brown University, Rhode Island Hospital, 593 Eddy St, Providence, RI 02903 (H.C.R.T., R.C.W.); Department of Diagnostic Imaging, Women and Infants Hospital of Rhode Island, Providence, RI (R.C.W.); Program in Women’s Oncology, Warren Alpert Medical School of Brown University, Women and Infants Hospital of Rhode Island, Providence, RI (W.M.S.); and Department of Diagnostic Radiology, Wayne State University, Ascension Providence Rochester Hospital, Rochester Hills, Mich (P.J.L.). Received September 25, 2020; revision requested November 9; revision received November 17; accepted December 10. Address correspondence to H.C.R.T. (e-mail: helaina_regen-tuero@brown.edu).

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Cryoablation is a well-tolerated outpatient procedure that has been used to treat metastatic sites as well as small breast cancers in patients who are considered poor candidates for surgery. Recent studies suggest that cell disruption caused by cryoablation may increase the expression and immunogenicity of tumor neoantigens, which could enhance the ability of the immune system to recognize and attack cancer cells at both local and distant sites. Such an approach might broaden the role of immunotherapy for the treatment of breast cancer, which has previously demonstrated limited response to these agents, likely owing to the modest immunogenicity of most breast cancer subtypes. If cryoablation can induce a systemic tumor-specific response, it could enhance tumor susceptibility to immunotherapy agents. This review briefly summarizes the necessary components for generating an immune response against tumor cells, reviews the tumor microenvironment of breast cancer, describes the rationale for and limitations of immune checkpoint inhibition, highlights the potential for cryoablation to induce a systemic tumor-specific immune response, and describes the rationale for combining cryoablation and immune checkpoint inhibitors for the treatment of breast cancer.

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Cryoablation, the use of extreme cold to kill cancer cells, is a well-tolerated outpatient procedure that has been used to eradicate metastatic disease and treat small breast cancers in patients who are considered poor candidates for surgery (1–3). Recent studies demonstrate that, in addition to causing direct damage to neoplastic tissue, cryoablation induces a systemic tumor-specific immune response (4,5). These findings are of particular significance as agents targeting the immune system have rapidly become a mainstay in cancer treatment and are now the standard of care in the treatment of several malignancies, including melanoma, lung, and bladder cancer. The goal of using immunotherapy agents is to flag cancer cells as foreign so they are recognized by the immune system, while also modulating immune-regulating signals toward an inflammatory state directed against the tumor. Investigated strategies have included priming the immune system against tumor-associated antigens with cancer vaccines, injecting oncolytic viruses into tumors, delivering chimeric antigen receptor T-cell therapies, and administering antibodies against various immune system targets to exploit existing immunity (6). When antibodies are used to inhibit immune checkpoint molecules, this removes the physiologic brake on the activated immune system to allow a sustained antigen-specific immune response.

Studies examining the efficacy of immune checkpoint inhibition in breast cancer have primarily focused on antibodies targeting cytotoxic T-lymphocyte antigen-4 (CTLA-4), programmed cell death 1 protein (PD-1), and programmed cell death ligand 1 (PD-L1). However, while preclinical studies have been promising, early clinical studies of immune checkpoint inhibition for breast cancer have shown relatively modest responses (7,8). This has been primarily attributed to the modest immune response elicited by most breast cancers, which typically demonstrate lower mutational rates and lower expression of tumor-associated neoantigens (9). Emerging data suggest that this relative resistance of breast cancer to immunotherapy agents may be overcome by combining multiple strategies, particularly those that enhance tumor immunogenicity such as cryoablation. In this review, we describe how cryoablation and immune checkpoint inhibitors interact with the immune system, summarize recent data on their efficacy in the treatment of breast cancer, and outline the rationale for combined therapy.

Principles of an Antigen-specific Immune Response

The efficacy of the immune system in mediating tumor regression depends on the efficient induction and maintenance of tumor antigen–specific T-cell responses. The inherent genetic instability of most tumor cells results in the expression of aberrant antigenic proteins or the overexpression of normally repressed genes that ultimately provide a target for T-cell recognition.

The activation of effector T cells depends on two signals. The process is initiated once the T-cell receptor engages its cognate antigen through interaction with the major histocompatibility complex on antigen-presenting cells (APCs). The second signal is delivered when the B7 costimulatory molecule expressed on APCs binds to the CD28 ligand on the surface of T cells. This results in a proinflammatory state and T-cell expansion. If the costimulatory signal is not received, T cells presented with antigen become anergic.

After a T cell has been activated by these two signals, a physiologic negative feedback loop is initiated to prevent...
Tumor-infiltrating Lymphocytes

Tumor-infiltrating lymphocytes (TILs) have been identified in some breast tumors, and when present, predict greater response to neoadjuvant chemotherapy and better overall survival rates (13–15). This advantage appears to be directly associated with the amount of lymphocytic infiltrate, with tumors demonstrating greater than 50% TILs (ie, lymphocyte-predominant breast cancers [LPBCs]) deriving greatest benefit. For example, a study of 2009 patients with node-positive breast cancer found that in estrogen receptor–negative/human epidermal growth factor receptor 2 (HER2)–negative breast cancers, every 10% increase in TILs was associated with a 15%–17% reduction in risk of recurrence and a 17%–27% reduction in risk of death (13). In another study of patients with locally advanced breast cancer treated with neoadjuvant chemotherapy, patients with LPBC had a 40% pathologic complete response as compared with a 7% pathologic complete response in patients with tumors that had no lymphocytic infiltrate (15). Unfortunately, while triple-negative and HER2-positive breast cancers have most consistently been associated with the presence of TILs and higher rates of LPBCs, lymphocyte-predominant infiltrates are relatively uncommon: across all subtypes, they are seen in less than 10% of breast cancers (16,17).

The composition of the lymphocytic infiltrate is important as well. Increased CD8+ effector T cells have been shown to predict improved clinical outcome, and higher numbers of intratumoral CD8+ effector T cells are associated with improved breast cancer–specific survival (18). Conversely, the presence of CD4+ regulatory T cells in the tumor has been associated with worse prognosis, including decreased disease-free and overall survival (19).

Checkpoint Inhibition

Immune checkpoint inhibitors represent a major disruptive breakthrough in cancer care. CTLA-4 blockade and its clinical success in melanoma therapy pioneered the field of checkpoint inhibition, which in turn led to the development of other targets, such as those of the PD-1/PD-L1 axis (20, 21). Response to these agents is likely affected by a number of both tumor- and host-related factors. For example, studies suggest that patient sex, age, gut microbiome, and human leukocyte antigen class I (HLA-I) genotype may have a role (22). In patients with melanoma, resistance to immune checkpoint inhibition has been associated with Wnt/β-catenin pathway activation, tumor loss of phosphatase and homolog expression, and mutations in interferon receptor–associated Janus kinase 1 (JAK1) or JAK2 (23). In breast cancer specifically, studies of immunotherapy response predictors have been largely limited to tumor PD-L1 expression, TIL density and composition, and tumor mutation load. A recent systematic review and meta-analysis demonstrated that PD-L1 positivity, higher TIL levels, and higher CD8+ T-cell levels predict better response to immunotherapy (24). Higher tumor mutational burden has also correlated with improved response, which has been discouraging given that most breast cancers demonstrate low tumor mutational burden (9,23,25).

Recent clinical trials have investigated the efficacy of monotherapy with anti–PD-1/PD-L1 agents in breast cancer (7,8,24). The proportion of patients who respond to these agents has been modest, with objective response rates ranging from 6% to 19% in patients with PD-L1–positive tumors, and rates of 0%–4.7% in patients with PD-L1–negative tumors (7, 8,27,28) (Table 1). However, in patients who respond, the treatment effects are durable. PD-L1 expression is challenging to measure, and reporting has not been standardized, which poses an additional challenge to determining which patients might benefit from anti-PD-1/ PD-L1 monotherapy (29). Regardless, the addition of agents that elicit an immune response and upregulate PD-L1 may enable those with PD-L1–negative breast cancers to clinically benefit from PD-1 inhibitors (30).
There have been far fewer clinical trials investigating CTLA-4 antagonist monotherapy in breast cancer. In one phase 1 study of 26 patients with hormone receptor–positive metastatic breast cancer, tremelimumab (anti–CTLA-4) was combined with the aromatase inhibitor exemestane (26). Although disease stability was observed in 42% of patients, no patients showed partial or complete response. However, it is important to note that this subtype has historically low levels of TILs, and so may have been less likely to respond to immune modulation than those with higher levels of TILs.

Combination therapy with CTLA-4 and PD-1/PD-L1 antagonists is also currently being investigated. An ongoing phase I/II trial is comparing nivolumab (anti-PD-1) alone to combination therapy with ipilimumab (anti–CTLA-4) in solid cancers, including triple-negative breast cancer (ClinicalTrials.gov identifier NCT01928394). Two other trials are investigating the efficacy of durvalumab (anti–PD-L1) and tremelimumab (anti–CTLA-4) in patients with metastatic HER2-negative breast cancer (ClinicalTrials.gov identifier NCT02536794) and hormone receptor–positive, HER2-negative stage II or III breast cancer (ClinicalTrials.gov identifier NCT03132467).

Cryoablation

Breast cryoablation is an office-based US-guided percutaneous procedure that uses extremely cold temperatures to freeze tissue surrounding the tip of the ablation probe needle (31). It is well-tolerated by patients and achieves cosmetically satisfactory results (32). Cryoablation has been successfully used to target metastatic sites in the liver, kidney, lung, bone, and soft tissues, and clinical trials examining its efficacy in treating small, early-stage invasive ductal carcinoma have been promising (1–3). The Figure provides a representative example of cryoablation for the purpose of intraductal carcinoma treatment. Recently, clinical interest has emerged in using cryoablation to also elicit an immune response that acts on metastatic sites distant from the treated lesion.

The rationale for this interest can be explained by review of the mechanism of cryoablation. Two short freeze-thaw cycles at a high freeze rate result in coagulative necrosis of cells closest to the probe. In the freeze phase, water freezes in the extracellular space faster than it does in the intracellular space, which sets up an osmotic gradient that drives fluid out of cells, resulting in cellular dehydration. During the thaw phase, the osmotic gradient reverses so that water rapidly enters cells, causing cell rupture. Additional injury results from the formation and growth of ice crystals, as well as from endothelial cell dysfunction, which causes vascular stasis and ischemia of tumor cells (33,34).

The resultant coagulative necrosis of tumor cells elicits a local inflammatory response, and importantly, also causes the release of intact tumor antigens, cellular stress signals, and type 1 cytokines into circulation (35,36). These signals result in the recruitment of APCs to the tumor and enhanced presentation of tumor-specific antigens to T cells, thereby inducing a tumor-specific T-cell response (37,38). This has been demonstrated in animal models of breast cancer, specifically, in which high freeze rate cryoablation resulted in an increase in tumor-specific T cells in tumor-draining lymph nodes, a decrease in the number of regulatory T cells, and improved survival (4,5). Essentially, local ablative therapies convert the tumor into an in situ vaccine that induces a systemic antitumor immune response.

Notably, when compared with heat-based modalities such as radiofrequency and microwave ablation, cryoablation induces a greater postablative immune response as demonstrated by markedly elevated levels of proinflammatory cytokines (39–42). Partly because of the analgesic effects of ice, it is also better tolerated than heat-based ablation, and so only local anesthesia is required for targets in the breast (43).
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In preclinical studies of cryoablation in breast cancer, the elicited immune response has been associated with regression of tumors distant from the treated lesion, a phenomenon known as the abscopal effect (4,5,34). However, in practice, this is a rare phenomenon when using local ablation agents alone for the treatment of breast cancer and is limited to case reports (44,45).

**Combined Cryoablation and Checkpoint Inhibition**

Since cryoablation induces a systemic tumor-specific T-cell response and checkpoint inhibitors remove the breaks on the immune system, combining the two therapies may have synergistic effects. Cryoablation may enhance immune recognition of breast cancer, facilitating tumor response to immunotherapy. In turn, checkpoint inhibition may allow the body to mount a robust immune response to the tumor-specific antigens released by cryoablation, enabling destruction of tumor cells in not only the breast, but also in regional lymph nodes and occult metastatic sites (36). Essentially, combination therapy could allow the abscopal effect (36).

Studies of combination therapy in animal models of solid tumors have demonstrated efficacy. In a mouse model of prostate cancer, cryoablation of the primary tumor followed by anti–CTLA-4 therapy slowed growth or triggered rejection of injected second tumors modeling micrometastasis (46). Clinical trials studying combination therapy for the treatment of various cancers, including breast cancer, are ongoing (47) (Table 2). While the results of these larger trials are not yet available, a few pilot studies have published preliminary data that show promise. A study combining cryoablation with either ipilimumab (anti–CTLA-4) or pembrolizumab (anti–PD-1) for the treatment of metastatic melanoma is reporting responses in both local (cryoablated) and distant lesions (48).

In breast cancer specifically, a treat-and-resect pilot study has demonstrated that preoperative cryoablation and single-dose ipilimumab are safe alone and in combination (49). Furthermore, the combination therapy was associated with a potentially favorable intratumoral and systemic immunologic response as demonstrated by sustained elevations in type 1 cytokines, activated and proliferating CD8+ T cells, and posttreatment proliferative effector T cells relative to regulatory T cells within the tumor bed (49). In addition, this combination therapy was associated with upregulation of interferon gamma, which in turn has been shown to increase expression of PD-L1, and thus could benefit patients with low TILs and low PD-1/PD-L1 expression (49–51).

Given that breast cancer is generally less immunogenic than most cancers, it may require the combination of both anti–CTLA-4 and anti–PD-L1 agents with cryoablation to induce the abscopal effect. Accordingly, investigators are currently recruiting patients for clinical trials that use both nivolumab (anti–PD-1) and ipilimumab (anti–CTLA-4) with cryoablation for the treatment of breast cancer (ClinicalTrials.gov identifiers NCT02833233 and NCT03546686). The results of these studies will help determine if the theory behind combined therapy translates to improved clinical outcomes. Should any beneficial clinical immune response be noted, thorough analyses of multiple treatment parameters and additional adjuvants (eg, toll-like receptors) will be needed to further improve upon those responses. Finally, some checkpoint inhibitors are dose dependent, and a combined percutaneous approach may allow the administration of lower doses, which may help decrease adverse effects while maintaining effective clinical outcomes (52).

**Conclusion**

Immunotherapy is a rapidly evolving field that seeks to harness the potential of the body’s immune system to recognize and eradicate neoplastic tissue. While the efficacy of these agents in breast cancer has previously been limited by the modest immune response mounted toward most tumors, cryoablation has the potential to enhance tumor immunogenicity, which may facilitate response to these agents. In turn, immune checkpoint inhibition removes the brakes on
the activated immune system and by so doing may allow the body to mount a robust immune response to tumor-specific antigens released by cryoablation. Thus, combining cryoablation with immune checkpoint inhibition is a rational strategy aimed at improving immune recognition and activation, and may result in an augmented tumor-specific immune response that acts on both local and distant disease. The results of ongoing studies will determine the trajectory with which this combination approach may be used in the future to treat breast cancer.

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