Clinical Data on Immunotherapy in Breast Cancer

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Abstract
Background: Breast cancer has traditionally been considered to have a low immunogenic potential compared to other tumor entities. Summary: The most extensively studied immunotherapeutic agents for breast cancer to date are immune checkpoint inhibitors, with the results of the IMpassion130 trial leading to the approval of atezolizumab plus nab-paclitaxel for first-line treatment of programmed cell death ligand 1-positive, metastatic, triple-negative breast cancer, and studies in earlier stages have yielded promising results. Other immunotherapeutic options being assessed in phases 2 and 3 trials include vaccine-based therapies and treatment with anti-human epidermal growth factor receptor 2 (H-directed immune-linked antibodies) and substances evaluated in early clinical trials as cellular therapies (adoptive cell therapy and chimeric antigen receptor T cells). Key Messages: Immunotherapy is an emerging modality for the treatment of breast cancer, as evidenced by the plethora of preclinical and clinical concepts and ongoing trials. Early studies established the role of immunotherapeutic agents in the metastatic setting. Ongoing studies will expand our knowledge about the timing of administration, best partners for combination therapy, and predictive biomarkers to guide immunotherapy for breast cancer. © 2020 The Author(s) Published by S. Karger AG, Basel

Introduction
The immune system plays a crucial role in cancer development and inherits tumor-stimulating and tumor-impeding capacities. Immunologically, tumor progression can be suppressed by the prevention or inhibition of tumor growth or promoted by the facilitation of cellular transformation or influencing of the tumor microenvironment to create favorable conditions [1, 2]. These unique, host-protecting and cancer-promoting features of the immune system are encompassed by the concept of “immunoediting.” This dynamic process has 3 phases (elimination, equilibrium, and escape), which describe how tumor cells escape from immune surveillance or are eradicated through the activation of innate and adaptive immune mechanisms [3]. Breast cancer has traditionally been considered to be a disease with a low immunogenic potential (a so-called “cold” tumor) compared to other tumor entities, such as melanoma and lung cancer, given its lower mutational burden, the smaller number of tumor-infiltrating lymphocytes (TIL), and the lesser programmed cell death protein/ligand (PD-1/L1) expression [4, 5]. However, with a better understanding of breast cancer biology and tumor and immunological profiling, this view began to change, leading to the development of new concepts of potential mechanisms of immune evasion and aspects of the tumor microenvironment. Although the breast cancer immune landscape is dynamic and heterogeneous, with considerable variation among
tumor stages and between subtypes (triple-negative vs. other subtypes) and disease settings (early vs. metastatic), the clinical evidence of responsiveness to various immunotherapies is accumulating [6]. An emerging body of preclinical and clinical data and ongoing studies addresses immunotherapy in neoadjuvant, adjuvant, and metastatic disease settings and for different subtypes of breast cancer and immunotherapy combinations. This review summarizes key clinical trials, recent findings, and emerging therapeutic concepts in the field of immunotherapy for breast cancer.

**Immunotherapy in Breast Cancer**

**Immune Checkpoint Inhibition**

Important targets in the cancer-immunity cycle are immune checkpoints; interactions with cell surface proteins can be blocked by immune checkpoint inhibitors (ICI). The most promising and most studied ICI in breast cancer to date are anti–PD-1/L-1 antibodies [7]. The PD-1/PD-L1 pathway represents an adaptive immune resistance mechanism of tumor cells. Binding between PD-1, which is expressed on T cells, and PD-L1, which is expressed on tumor cells, inhibits the antitumoral T-cell-adapted immune response. Blockage of this pathway via checkpoint inhibitors “releases the brakes” of the immune system in the battle against cancer cells and can reactivate immune surveillance and antitumoral immune activity [1, 8, 9]. Immunotherapy in general and ICI specifically have shown the most promising results to date for triple-negative breast cancer, which is characterized by a greater PD-L1 expression relative to other breast cancer subtypes [5, 10].

**Metastatic Breast Cancer**

The use of several ICI as monotherapy for metastatic triple-negative breast cancer has been studied, with moderate but durable response rates and tolerable safety profiles observed [11–13]. These effects seem to be restricted mostly to first-line treatment of metastatic disease and to PD-L1-positive tumors. For example, the phase 2 KEYNOTE-086 study assessed pembrolizumab monotherapy (200 mg i.v. every 3 weeks) with regard to PD-L1 status and line of therapy in 170 patients with metastatic triple-negative breast cancer receiving pretreatment (cohort A, 62% PD-L1 positive) and 84 such patients receiving first-line treatment (cohort B, 100% PD-L1 positive) [14]. Objective response rates were 5.3% in cohort A (5.7% in PD-L1-positive patients) and 21.4% in cohort B, with a median response duration of 10.4 months in cohort B (not achieved in cohort A). The median progression-free survival (PFS) was 2 months (95% CI 1.9–2.0 months) in cohort A and 2.1 months (95% CI 2.0–2.2 months) in cohort B, and the median overall survival (OS) was 9.0 (95% CI 7.6–11.2) and 18.0 (95% CI 9.0–23.0) months, respectively [14]. Similar results were obtained for atezolizumab monotherapy in a phase 1 study conducted in patients with metastatic or locally advanced triple-negative breast cancer; objective response rates were 24% for first-line treatment and 6% for later lines, and 12% for PD-L1-positive tumors and 0% for PD-L1-negative tumors [11]. Studies comparing ICI monotherapy with chemotherapy, such as the KEYNOTE-012 study, in which second-/third-line pembrolizumab treatment was compared with single-agent chemotherapy (capecitabine, eribulin, gemcitabine, or vinorelbine) of the physicians’ choice for patients with metastatic triple-negative breast cancer, have revealed no significant difference in OS between groups [15]. These studies have shown that the effects of PD-L1 inhibitors for the treatment of metastatic triple-negative breast cancer are strongest for early treatment lines and PD-L1-overexpressing cancers. Given the immunomodulatory effects of chemotherapy, which are based partly on the induction of tumor-associated antigen (TAA) release, the next step was to examine the use of ICI combined with chemotherapy [16]. IMpassion130, a phase 3 study, evaluated the use of nab-paclitaxel (100 mg/m² i.v. on days 1, 8, and 15) plus atezolizumab (840 mg i.v. on days 1 and 15 of every 28-day cycle) or placebo for the treatment of first-line metastatic or locally advanced triple-negative breast cancer [17]. The enrolled patients had tumor relapse ≥12 months after completion of (neo)adjuvant chemotherapy. PFS was significantly longer in the atezolizumab-plus-chemotherapy arm of the intention-to-treat (ITT) population (7.2 vs. 5.5 months; p = 0.0025) and in the subgroup with PD-L1-positive tumors (7.5 vs. 5.0 months). OS did not differ significantly in the ITT population (21.3 vs. 17.6 months), but it reflected a “clinically meaningful” benefit in patients with PD-L1-positive tumors treated with combination therapy (25.0 vs. 18.0 months; HR = 0.71; 95% CI 0.54–0.94) [17]. However, formal statistical testing of these data could not be performed because OS was not predefined as a primary outcome in case of a nonsignificant difference in the ITT population. Grade 3 and 4 adverse events occurred in 48.7% of patients receiving atezolizumab and in 42.2% of those receiving placebo. The most frequent adverse events were neutropenia, peripheral neuropathy, decreased neutrophil count, and fatigue. Two treatment-related deaths (caused by autoimmune hepatitis and septic shock, respectively) occurred in the experimental arm and 1 death (caused by hepatic failure) occurred in the placebo arm. Adverse events leading to the discontinuation of therapy occurred in 15.9% of patients receiving atezolizumab and in 8.2% of those receiving a placebo [17]. The IMpassion130 study led to US Food and Drug Administration (FDA) approval of atezolizumab for patients with unre-
Table 1. Immune checkpoint blockade trials for metastatic breast cancer in the recruitment phase

| Agent | Trial / Phase | Setting |
|-------|---------------|---------|
| Pembrolizumab | Focused Ultrasound and Pembrolizumab in Metastatic Breast Cancer / Phase 1 | Metastatic |
| | Study of Pembrolizumab Plus Fulvestrant in Hormone Receptor Positive, HER-2-Negative Advanced/Metastatic Breast Cancer Patients / Phase 2 | Metastatic |
| | PVX-410 Vaccine plus Pembrolizumab in HLA-A2+ Metastatic Triple-Negative Breast Cancer / Phase 2 | Metastatic |
| | Establishing the Recommended Biological Dose for AE37 Peptide Vaccine in Combination with Pembrolizumab that Will Enhance the Tumor-Specific Immune Response and Demonstrate Efficacy in Patients with Advanced Triple-Negative Breast Cancer / Phase 2 | Metastatic |
| | Phase 1b Study of Pegylated Liposomal Doxorubicin and Pembrolizumab in Endocrine-Resistant Breast Cancer / Phase 1 | Metastatic |
| | A Study to Evaluate Concurrent VRP-HER2 Vaccination and Pembrolizumab for Patients with Breast Cancer / Phase 2 | Metastatic |
| | Pembrolizumab and Exemestane/Leuprolide in Premenopausal HR+/HER2- Locally Advanced or Metastatic Breast Cancer / Phases 1 and 2 | Metastatic |
| | Her2-BATS and Pembrolizumab in Metastatic Breast Cancer/ Phases 1 and 2 | Metastatic |
| | Phase II Study of Pembrolizumab and Nab-Paclitaxel in HER-2-Negative Metastatic Breast Cancer / Phase 2 | Metastatic |
| | A Study Of Pembrolizumab in Combination with Trastuzumab-DM1 / Phase 1 | Metastatic |
| | Safety and Efficacy of SGN-LIV1A plus Pembrolizumab for Patients with Locally Advanced or Metastatic Triple-Negative Breast Cancer / Phases 1 and 2 | Metastatic |
| | Pilot Study of Paclitaxel plus Pembrolizumab in Metastatic HER2-Negative Breast Cancer / Phase 2 | Metastatic |
| | Study of Olaparib plus Pembrolizumab versus Chemotherapy plus Pembrolizumab after Induction with First-Line Chemotherapy plus Pembrolizumab in Triple-Negative Breast Cancer (TNBC) (MK-7339-009/KEYLYNK-009) / Phases 2 and 3 | Metastatic |
| | Intratumoral Tavo and Pembro in Patients with Inoperable Locally Advanced or Metastatic TNBC (KEYNOTE-890) / Phase 2 | Metastatic |
| | A Study of Pembrolizumab with Carboplatin and Gemcitabine in Patients with Metastatic Triple-Negative Breast Cancer / Phase 2 | Metastatic |
| | Pembrolizumab, Letrozole, and Palbociclib in Treating Postmenopausal Patients with Newly Diagnosed Metastatic Stage IV Estrogen Receptor-Positive Breast Cancer / Phase 2 | Metastatic |
| | Pembrolizumab and Ruxolitinib Phosphate in Treating Patients with Metastatic Stage IV Triple-Negative Breast Cancer / Phase 1 | Metastatic |
| | Chemokine Modulation Therapy and Pembrolizumab in Treating Participants With Metastatic Triple-Negative Breast Cancer / Phase 1 | Metastatic |
| | SBRT and Oncolytic Virus Therapy before Pembrolizumab for Metastatic TNBC and NSCLC / Phase 2 | Metastatic |
| | Pembrolizumab and Carboplatin in Treating Patients with Circulating Tumor Cells Positive Metastatic Breast Cancer / Phase 2 | Metastatic |
| | A Study of EDP1503 in Patients with Colorectal Cancer, Breast Cancer, and Checkpoint Inhibitor Relapsed Tumors / Phases 1 and 2 | Metastatic |
| | Vaccination with Flt3L, Radiation, and Poly-ICLC / Phases 1 and 2 | Metastatic |
| | DS8201a and Pembrolizumab in Participants with Locally Advanced/Metastatic Breast or Non-Small Cell Lung Cancer / Phase 1 | Metastatic |
| | Tesetaxel plus 3 Different PD-(L)1 Inhibitors in Patients with Metastatic TNBC and Tesetaxel Monotherapy in Patients with HER2 Negative MBC / Phase 2 | Metastatic |
| | Phase 1a/1b Study of TPST-1495 Alone and with Pembrolizumab in Subjects with Solid Tumors / Phase 1 | Metastatic |
| | CPI-006 Alone and in Combination with Ciforadenant and with Pembrolizumab for Patients with Advanced Cancers / Phase 1 | Metastatic |
| | Pembrolizumab in Treating Participants with Metastatic, Recurrent or Locally Advanced Cancer and Genomic Instability / Phase 2 | Metastatic |
| | Immunotherapy Using Tumor Infiltrating Lymphocytes for Patients with Metastatic Cancer / Phase 2 | Metastatic |
| Agent | Trial / Phase                                                                 | Setting  |
|-------|------------------------------------------------------------------------------|----------|
|       | A Study of XmAb®22841 Monotherapy and in Combination w/ Pembrolizumab in Subjects w/ Selected Advanced Solid Tumors / Phase 1 | Metastatic |
|       | Study of SO-C101 and SO-C101in Combination with Pembro in Adult Patients with Advanced/Metastatic Solid Tumors / Phase 1 | Metastatic |
|       | A Study of ZN-c3 in Participants with Solid Tumors / Phases 1 and 2          | Metastatic |
|       | A Personalized Medicine Study for Patients with Advanced Cancer of the Breast, Prostate, Pancreas or Those with Refractory Acute Myelogenous Leukemia / Phase 1 | Metastatic |
|       | A First-in-Human Study Using BDC-1001 in Advanced and HER2-Expressing Solid Tumors / Phase 1 | Metastatic |
|       | Pembrolizumab with Intratumoral Injection of Clostridium Novyi-NT / Phase 1   | Metastatic |
|       | Malignant Pleural Disease Treated With Autologous T Cells Genetically Engineered to Target the Cancer-Cell Surface Antigen Mesothelin / Phases 1 and 2 | Metastatic |
|       | GB1275 Monotherapy and in Combination with an Anti-PD1 Antibody in Patients with Specified Advanced Solid Tumors or in Combination with Standard of Care in Patients with Metastatic Pancreatic Adenocarcinoma / Phases 1 and 2 | Metastatic |
|       | Study of ONCR-177 Alone and in Combination with PD-1 Blockade in Adult Subjects with Advanced and/or Refractory Cutaneous, Subcutaneous or Metastatic Nodal Solid Tumors / Phase 1 | Metastatic |
|       | Trans-Artery/Intra-Tumor Infusion of Checkpoint Inhibitors for Immunotherapy of Advanced Solid Tumors / Phases 2 and 3 | Metastatic |
|       | Administration of Autologous T-Cells Genetically Engineered to Express T-Cell Receptors Reactive against Mutated Neoantigens in People with Metastatic Cancer / Phase 2 | Metastatic |
|       | A Study of CDX-1140 as Monotherapy or in Combination in Patients with Advanced Malignancies / Phase 1 | Metastatic |
|       | FT500 as Monotherapy and in Combination with Immune Checkpoint Inhibitors in Subjects with Advanced Solid Tumors / Phase 1 | Metastatic |
|       | Dose Escalation and Expansion Study of FLX475 Monotherapy and in Combination with Pembrolizumab / Phases 1 and 2 | Metastatic |
|       | A Phase 1/2 Safety Study of Intratumorally Dosed INT230-6 / Phases 1 and 2    | Metastatic |
|       | Study of DF1001 in Patients with Advanced Solid Tumors / Phases 1 and 2      | Metastatic |
|       | Open-Label, Dose-Escalation Study of Pemigatinib in Subjects with Advanced Malignancies – (FIGHT-101) / Phases 1 and 2 | Metastatic |
|       | Dose Escalation Study of GSK3326595 in Subjects with Solid Tumors and Non-Hodgkin’s Lymphoma / Phase 1 | Metastatic |
|       | Focused Ultrasound Ablation and PD-1 Antibody Blockade in Advanced Solid Tumors / Phase 1 | Metastatic |
|       | Nivolumab /— Nivolumab in Metastatic Triple-Negative Breast Cancer / Phase 2  | Metastatic |
|       | NIMBUS: Nivolumab plus Ipilimumab in Metastatic Hypermutated HER2-Negative Breast Cancer / Phase 2 | Metastatic |
|       | Nivolumab and Eribulin in HER2-Negative Metastatic Breast Cancer / Phases 1 and 2 | Metastatic |
|       | Pre-Operative Trial for Breast Cancer with Nivolumab in Combination with Novel IO / Phase 2 | Metastatic |
|       | Stereotoxic Radiation and Nivolumab in the Management of Metastatic Breast Cancer Brain Metastases / Phase 1 | Metastatic |
|       | Phase IIb Study Evaluating Immunogenic Chemotherapy Combined with Ipilimumab and Nivolumab in Breast Cancer / Phase 2 | Metastatic |
|       | Nivolumab, Ipilimumab, and Bicalutamide in Human Epidermal Growth Factor (HER) 2-Negative Breast Cancer Patients / Phase 2 | Metastatic |
|       | Immune Induction Strategies to Improve Response to Immune Checkpoint Blockade in Triple-Negative Breast Cancer (TNBC) Patients / Phase 2 | Metastatic |
|       | Trastuzumab Deruxtecan with Nivolumab in Advanced Breast and Urothelial Cancer / Phase I | Metastatic |
|       | Sequential Immuno Apheresis Plasma Volume Escalation Cohort Study of Removal of Soluble Tumor Necrosis Factor Receptors 1 and 2 (sTNFR1/2) with or without Nivolumab in Patients with Inoperable or Metastatic Solid Tumors | Metastatic |
|       | Tesetaxel plus 3 Different PD-(L)1 Inhibitors in Patients with Metastatic TNBC and Tesetaxel Monotherapy in Patients with HER2-Negative MBC / Phase 2 | Metastatic |
| Agent | Trial / Phase | Setting |
|-------|--------------|---------|
| Study BT5528-100 in Patients with Advanced Solid Tumors Associated with EphA2 Expression / Phases 1 and 2 | Metastatic |
| A Study of NKTR-262 in Combination with NKTR-214 and with NKTR-214 plus Nivolumab in Patients with Locally Advanced or Metastatic Solid Tumor Malignancies / Phases 1 and 2 | Metastatic |
| A Dose-Escalation Study to Evaluate the Safety, Tolerability, Pharmacokinetics, and Pharmacodynamics of IPI-549 / Phase 1 | Metastatic |
| A Dose Escalation and Cohort Expansion Study of NKTR-214 in Combination with Nivolumab and Other Anti-Cancer Therapies in Patients with Select Advanced Solid Tumors (PIVOT-02) / Phases 1 and 2 | Metastatic |
| A Personalized Medicine Study for Patients with Advanced Cancer of the Breast, Prostate, Pancreas or Those with Refractory Acute Myelogenous Leukemia / Phase 1 | Metastatic |
| A Study of Gene Edited Autologous Neoantigen Targeted TCR T Cells with or without Anti-PD-1 in Patients with Solid Tumors / Phase 1 | Metastatic |
| A Study of ASP1948, Targeting an Immune Modulatory Receptor, in Subjects with Advanced Solid Tumors / Phase 1 | Metastatic |
| Phase 1/2 Study Exploring the Safety, Tolerability, and Efficacy of INCAGN01876 Combined with Immune Therapies in Advanced or Metastatic Malignancies / Phases 1 and 2 | Metastatic |
| Atezolizumab + Stereotactic Radiation in Triple-Negative Breast Cancer and Brain Metastasis / Phase 2 | Metastatic |
| 89Zr-Atezolizumab PET Scan and Lobular Breast Cancer | Metastatic |
| Assessing Efficacy of Carboplatin and Atezolizumab in Metastatic Lobular Breast Cancer / Phase 2 | Metastatic |
| A Study of Atezolizumab and Paclitaxel Versus Placebo and Paclitaxel in Participants with Previously Untreated Locally Advanced or Metastatic Triple-Negative Breast Cancer (TNBC) / Phase 3 | Metastatic |
| A Study of Atezolizumab plus Nab-Paclitaxel or Paclitaxel in the Treatment of Unresectable Locally Advanced or Metastatic Triple-Negative Breast Cancer / Phase 3 | Metastatic |
| Stereotactic Radiation and Immunotherapy in Patients with Advanced Triple Negative Breast Cancer / Phase 2 | Metastatic |
| Triple-B Study; Carboplatin-Cyclophosphamide versus Paclitaxel with or without Atezolizumab as First-Line Treatment in Advanced Triple-Negative Breast Cancer / Phase 2 | Metastatic |
| Clinical Trial of Atezolizumab with Paclitaxel, Trastuzumab, and Pertuzumab in Patients with Metastatic HER-2-Positive Breast Cancer / Phase 2 | Metastatic |
| A Study Of Ipatasertib in Combination with Atezolizumab and Paclitaxel as a Treatment for Participants with Locally Advanced or Metastatic Triple-Negative Breast Cancer / Phase 3 | Metastatic |
| Atezolizumab Combined with Immunogenic Chemotherapy in Patients with Metastatic Triple-Negative Breast Cancer / Phase 2 | Metastatic |
| A Study to Evaluate the Safety and Efficacy of Ipatasertib in Combination with Atezolizumab and Paclitaxel or Nab-Paclitaxel in Participants with Locally Advanced or Metastatic Triple-Negative Breast Cancer / Phase 1 | Metastatic |
| A Study of the Efficacy and Safety of Atezolizumab plus Chemotherapy for Patients with Early Relapsing Recurrent Triple-Negative Breast Cancer / Phase 3 | Metastatic |
| Carboplatin with or without Atezolizumab in Treating Patients with Stage IV Triple-Negative Breast Cancer / Phase 2 | Metastatic |
| Ipatasertib in Combination with Carboplatin, Carboplatin/Paclitaxel, or Capecitabine/Atezolizumab in Treating Patients with Metastatic Triple-Negative Breast Cancer / Phases 1 and 2 | Metastatic |
| Atezolizumab + Pertuzumab + Trastuzumab in CNS Mets in BC / Phase 2 | Metastatic |
| Cryoablation, Atezolizumab/Nab-Paclitaxel for Locally Advanced or Metastatic Triple-Negative Breast Cancer / Phase 1 | Metastatic |
| Olaparib with or without Atezolizumab in Treating Patients with Locally Advanced Unresectable or Metastatic Non-HER2-Positive Breast Cancer / Phase 2 | Metastatic |
| Atezolizumab, Cobimetinib, and Eribulin in Treating Patients with Chemotherapy-Resistant Metastatic Inflammatory Breast Cancer / Phase 2 | Metastatic |
| Testing the Drug Atezolizumab or Placebo with Usual Therapy in First-Line HER2-Positive Metastatic Breast Cancer / Phase 3 | Metastatic |
| A Study of Multiple Immunotherapy-Based Treatment Combinations in Hormone Receptor (HR)-Positive Human Epidermal Growth Factor Receptor 2 (HER2)-Negative Breast Cancer / Phases 1 and 2 | Metastatic |
 sectable, locally advanced, or metastatic triple-negative breast cancer, with PD-L1-stained tumor-infiltrating immune cells of any intensity covering ≥1% of the tumor area. Two ongoing studies are evaluating the use of ICI plus chemotherapy in patients with early (<12 months after [neo]adjuvant chemotherapy) relapse who were ineligible for the Impassion130 study [18, 19]. KEY-NOTE-355 is a phase 3 study assessing the use of pembrolizumab, paclitaxel, and gemcitabine/carboplatin for first-line treatment of locally recurrent inoperable or metastatic triple-negative breast cancer with relapse <6 months after the initial diagnosis, according to PD-L1 expression (no expression, combined positive score ≥1, and combined positive score ≥10) [18]. That study is currently recruiting patients, but an interim analysis revealed a significant and clinically meaningful improvement in PFS in patients receiving pembrolizumab relative to those receiving chemotherapy, with a safety profile consistent with previously published data [20, 21]. Impassion132 is enrolling a similar group of patients with relapse ≤12 months after...
curative-intent chemotherapy, with the comparison of atezolizumab or placebo with the investigators’ choice of chemotherapy (gemcitabine plus carboplatin or capecitabine) [19]. In addition to the question of which chemotherapeutic backbone will be the optimal partner for ICI combination therapy, the timing of chemotherapy is being examined. Given the rationale that chemotherapy has immunostimulatory effects, but also leads to lymphodepletion, the combined administration of ICI with low-dose induction chemotherapy is an approach that has shown promising results in early-phase clinical trials [22]. Similarly, the SAFIR-02 trial demonstrated that ICI maintenance monotherapy following a response to induction chemotherapy can significantly improve survival relative to the continuation of chemotherapy in patients with metastatic triple-negative PD-L1-positive breast cancer [23]. Regarding biomarkers for treatment response, line of treatment and PD-1/L1 expression seem to be the most suitable to date in the metastatic setting, with patients receiving first-line ICI and those with PD-1/L1-positive tumors benefitting the most from ICI therapy. Several ongoing clinical trials are assessing combination therapies with different ICI or ICI and other immunogenic agents (Table 1).

Early Breast Cancer

The observation that PD-1/L1 expression and the number of TIL are greater in early-stage versus metastatic breast cancer, and promising results from phases 1 and 2 studies, have provided the rationale for evaluation of ICI in the neoadjuvant setting [24–26]. The recently presented phase 3 KEYNOTE-522 study assessed the effects of pembrolizumab (200 mg, 4 cycles, i.v.) and placebo every 3 weeks plus neoadjuvant chemotherapy with 12× paclitaxel (80 mg/m²) and carboplatin (area under the curve [AUC] = 5, every 3 weeks or 1.5 mg/mL/min once weekly), followed by 4 cycles of pembrolizumab or placebo plus doxorubicin (60 mg/m²) or epirubicin (90 mg/m²) plus cyclophosphamide (600 mg/m²) every 3 weeks in 1,174 patients with newly diagnosed triple-negative breast cancer [27]. Pathologic complete response (pCR) rates were 64.8% (95% CI 59.9–69.5) in the pembrolizumab arm and 51.2% (95% CI 44.1–58.3) in the placebo arm (p < 0.001). This effect was strongest in patients with node-negative disease and more advanced tumor stages, and it was independent of PD-L1 expression. Grade 3+ adverse events occurred in 76.8% of patients in the pembrolizumab arm and in 72.2% of those in the placebo arm, and discontinuation of any trial drug due to treatment-related adverse events occurred in 23.3 and 12.3% of cases, respectively [27].

A similar study (NeoTRIP) assessed the effects of 8 cycles of atezolizumab (1,200 mg i.v. every 3 weeks) or placebo plus neoadjuvant chemotherapy with 8 cycles of carboplatin (AUC = 2, i.v., on days 1 and 8 every 3 weeks) and nab-paclitaxel (125 mg/m² i.v. on days 1 and 8 every 3 weeks), followed by definitive breast surgery and 4 cycles of doxorubicin (60 mg/m²) or epirubicin (90 mg/m²) plus cyclophosphamide (600 mg/m²) every 3 weeks in 280 women with triple-negative breast cancer [28]. The pCR rate did not differ significantly between the atezolizumab (43.5%) and placebo (40.8%) arms in the ITT analysis. Among PD-L1-positive patients, the pCR rate was higher in the atezolizumab group (51.9%) compared to the placebo group (48%), but this difference was not significant. However, PD-L1-positive status was the only variable associated with the pCR rate in the multivariate analysis (p < 0.01) [28]. The results of these 2 neoadjuvant chemotherapy studies seem to conflict, but correct interpretation of the outcomes requires consideration of differences in study design, primary endpoints (pCR in KEYNOTE-522 vs. event-free survival in NeoTRIP), and chemotherapy timing (before surgery in KEYNOTE-522 vs. after surgery in NeoTRIP). Further analyses of the long-term NeoTRIP outcomes will show whether atezolizumab plus the chemotherapy combination has a role in the neoadjuvant setting. The importance of immunotherapy timing was demonstrated in the phase 2 GeparNuevo trial, which assessed the effects of neoadjuvant durvalumab (1.5 g i.v. every 4 weeks) or placebo plus nab-paclitaxel (125 mg/m² weekly for 12 weeks), followed by 4 cycles of epirubicin/cyclophosphamide (90/600 mg/m² every 2 weeks) [29]. The trial also included a “window cohort” that received durvalumab or placebo monotherapy 2 weeks before the initiation of chemotherapy. In the ITT population; the pCR rate did not differ significantly between patients receiving durvalumab (54.3%; 95% CI 42.5–61.4) and those receiving a placebo (44.2%; 95% CI 33.5–55.3); in the window cohort, however, the pCR rate was significantly higher in patients treated with durvalumab (61%) compared to the placebo arm (41.4%; p = 0.035). Several ongoing neoadjuvant, adjuvant, and post-neoadjuvant chemotherapy trials are further evaluating the role of ICI in breast cancer treatment (Table 2).

Other ICI

Proteins other than PD-1/PD-1 that might be targetable with checkpoint inhibitors have been identified; they include lymphocyte-activation gene 3, B7-H4, and T-cell immunoreceptor with immunoglobulin (Ig) and immunoreceptor tyrosine-based inhibition motif domains (TIGIT) [4]. The use of these ICI in combination with PD-L1 ICI is currently being studied in phase 1 trials for solid tumors, and studies for breast cancer are planned [30]. Another possible ICI target is CTL-associated antigen 4 (CTLA-4), which can limit T-cell activation by interaction with its ligand, CD80, or CD86. Increased T-cell activation due to CTLA-4 inhibition has been shown to
Table 2. Immune checkpoint inhibitor trials for breast cancer in the recruitment phase

| Agent | Trial / Phase | Setting |
|-------|--------------|---------|
| Pembrolizumab | Breast Cancer Study of Preoperative Pembrolizumab + Radiation / Phase 1 | Neoadjuvant |
| | A Study of Changes in PD-L1 Expression during Preoperative Treatment with Nab-Paclitaxel and Pembrolizumab in Hormone Receptor-Positive Breast Cancer / Phase 1 | Neoadjuvant |
| | Effects of MK-3475 (Pembrolizumab) on the Breast Tumor Microenvironment in Triple-Negative Breast Cancer / Phase 1 | Neoadjuvant |
| | Study of Immunotherapy in Combination with Chemotherapy in HER2-Negative Inflammatory Breast Cancer / Phase 2 | Neoadjuvant |
| | Neoadjuvant Phase 2 Study of Pembrolizumab and Carboplatin plus Docetaxel in Triple-Negative Breast Cancer / Phase 2 | Neoadjuvant |
| | Neoadjuvant Her2-Targeted Therapy and Immunotherapy with Pembrolizumab / Phase 2 | Neoadjuvant |
| | Neoadjuvant Pembrolizumab + Decitabine Followed by Std Neoadj Chemo for Locally Advanced HER2- Breast Ca / Phase 2 | Neoadjuvant |
| | I-SPY 2 TRIAL: Neoadjuvant and Personalized Adaptive Novel Agents to Treat Breast Cancer / Phase 2 | Neoadjuvant |
| | Pembrolizumab in High-Risk Ductal Carcinoma in situ (DCIS) / Phase 1 | Neoadjuvant |
| | Study of Pembrolizumab (MK-3475) versus Placebo in Combination with Neoadjuvant Chemotherapy and Adjuvant Endocrine Therapy in the Treatment of Early-Stage Estrogen Receptor-Positive, Human Epidermal Growth Factor Receptor 2-Negative (ER+/HER2-) Breast Cancer (MK-3475-756/KEYNOTE-756) / Phase 3 | Neoadjuvant/ adjuvant |
| | Pembrolizumab in Treating Patients with Hormone Receptor-Positive, Localized Inflammatory Breast Cancer Who Are Receiving Hormone Therapy and Did Not Achieve a Pathological Complete Response to Chemotherapy / Phase 2 | Adjuvant |
| | Testing MK-3475 (Pembrolizumab) as Adjuvant Therapy for Triple Receptor-Negative Breast Cancer / Phase 3 | Adjuvant |
| | Pembrolizumab with Carboplatin Compared to Carboplatin Alone in Breast Cancer Patients with Chest Wall Disease / Phase 2 | Adjuvant/ metastatic |
| | Study of GX-I7 in Combination with Pembrolizumab in Refractory or Relapsed (R/R) TNBC Subjects(GX-I7-CA-006/KEYNOTE-899) / Phases 1 and 2 | Adjuvant/ metastatic |
| | Pembrolizumab in Treating Patients with Stage IV Metastatic or Recurrent Inflammatory Breast Cancer or Triple-Negative Breast Cancer Who Have Achieved Clinical Response or Stable Disease to Prior Chemotherapy / Phase 2 | Adjuvant/ metastatic |
| | Galipepimut-S in Combination with Pembrolizumab in Patients with Selected Advanced Cancers / Phases 1 and 2 | Adjuvant/ metastatic |
| Nivolumab | Peri-Operative Ipilimumab+Nivolumab and Cryoablation versus Standard Care in Women with Triple-Negative Breast Cancer / Phase 2 | Neoadjuvant |
| | Trial of Nivolumab with Chemotherapy as Neoadjuvant Treatment in Inflammatory Breast Cancer (IBC) / Phase 2 | Neoadjuvant |
| | A Study of Neoadjuvant Nivolumab + Palbociclib + Anastrozole in Post-Menopausal Women and Men with Primary Breast Cancer / Phase 2 | Neoadjuvant |
| | Ipilimumab, Nivolumab, and Talimogene Laherparepvec before Surgery in Treating Participants with Localized, Triple-Negative or Estrogen Receptor-Positive, HER2-Negative Breast Cancer-Deleted / Phase 1 | Neoadjuvant |
| | Study of Nivolumab versus Placebo in Participants with High-Risk Breast Cancer / Phase 3 | Neoadjuvant/ adjuvant |
| | OXEL: Immune Checkpoint or Capecitabine or Combination Therapy as Adjuvant Therapy for TNBC with Residual Disease / Phase 2 | Adjuvant |
| | TP5T-1120 as Monotherapy and in Combination with (Nivolumab, Docetaxel or Cetuximab) in Subjects with Advanced Cancers / Phase 1 | Adjuvant/ metastatic |
| | COM701 in Subjects with Advanced Solid Tumors / Phase 1 | Adjuvant/ metastatic |
| | Nivolumab and Ipilimumab in Treating Patients with Rare Tumors / Phase 2 | Adjuvant/ metastatic |
| | FT500 as Monotherapy and in Combination with Immune Checkpoint Inhibitors in Subjects with Advanced Solid Tumors / Phase 1 | Adjuvant/ metastatic |
have antitumoral effects [31, 32]. Two monoclonal antibodies, i.e., tremelimumab and ipilimumab, have been assessed clinically. A phase I study evaluated tremelimumab (3–10 mg/kg i.v. every 4 weeks for 3 months) plus exemestane (25 mg orally daily) in 26 patients with advanced hormone receptor-positive breast cancer [33]. The best overall response was stable disease at ≥12 weeks in 11 (42%) patients, with a tolerable safety profile [33]. Ipilimumab has been approved by the FDA for the treatment of metastatic melanoma, and its use for breast cancer has
been assessed in early clinical phase studies. A pilot study evaluated preoperative treatment with tumor cryoablation and/or single-dose ipilimumab (10 mg/kg i.v.) in 19 patients undergoing mastectomy [34]. The researchers examined safety, tolerability, and immune activation and found both individual treatments and their combination to be safe, with intratumoral and systemic immunological effects [34]. Other trials assessing the use of chemotherapy plus multiple ICI combination therapies for metastatic breast cancer are currently underway (Table 2).

ICI Combination Therapies

ICI Plus Chemotherapy. Most clinical data on combination therapies are for ICI combined with chemotherapy, including those generated in the above mentioned large clinical studies IMPASSION130 and KEYNOTE-355, which showed durable response rates and increased survival outcomes [17, 18]. The molecular mechanisms underlying this positive synergistic effect are not understood precisely, but they are based on complex drug- and dose-dependent interactions between chemotherapeutic agents and the immune system. These interactions include chemotherapy-induced cytotoxic immune responses, induction of proinflammatory cytokine secretion, inhibition of myeloid-derived suppressor cells, upregulation of receptors enhancing tumor cell susceptibility to lysis by natural killer cells, and recruitment and activation of immune cells such as dendritic, natural killer, and T cells. Several ongoing trials are seeking to determine the optimal chemotherapeutic agents to combine with ICI and the optimal sequence of application (Tables 1, 2) [35–39].

ICI Plus Anti-HER2 Targeted Therapy. Several other attempts to add immunotherapy to established treatment regimens have been made. Targeted therapeutic agents, such as the anti-HER2 antibody trastuzumab, have been found to mediate antitumoral activity, partly through the modulation of immune activity, and thus might be beneficial when used in combination with immunotherapeutic agents [40, 41]. The single-arm phase IB/2 PANACEA study evaluated the effects of trastuzumab (6 mg/kg i.v. every 3 weeks) combined with pembrolizumab (200 mg i.v. every 3 weeks) in patients with advanced HER2-positive breast cancer who had progressed during trastuzumab-based therapy, according to PD-L1 status [42]. Six out of 40 (15%) patients in the PD-L1-positive cohort and none of 12 patients in the PD-L1-negative cohort showed an objective response. Pembrolizumab was tolerated well and the antitumoral activity was durable, with a median treatment response duration of 11.2 months (range: 6.2 to not reached) [42]. In a similar phase 1 trial assessing the effects of the ICI durvalumab (1,125 mg i.v. every 3 weeks) combined with trastuzumab (6 mg/kg i.v. every 3 weeks) in extensively pretreated patients with metastatic HER2-positive breast cancer, the researchers observed no objective response to therapy and all 15 patients had <1% PD-L1 expression [43]. The KATE2 trial also demonstrated the dependency of antitumoral activity induced by ICI combination therapy on the PD-L1 expression status [44]. This phase 2 study was the first to compare the effects of the ICI atezolizumab (1,200 mg i.v. every 2 weeks) plus the antibody drug conjugate trastuzumab emtansine (3.6 mg/kg i.v. every 3 weeks) with those of trastuzumab emtansine (3.6 mg/kg every 3 weeks) alone in 202 patients with advanced or metastatic HER2-positive breast cancer who progressed after trastuzumab-based treatment, stratified by PD-L1 immune cell status (<1 vs. ≥1%). In the ITT population, PFS and OS did not differ significantly between groups. In the subgroup of PD-L1-positive patients, however, PFS and 1-year OS were numerically greater in the combined therapy group (PFS = 8.5 months, range: 5.7 to not evaluated, HR = 0.60; 95% CI 0.32–1.11; 1-year OS = 94.3%, HR = 0.55; 95% CI 0.22–1.38) compared to the trastuzumab emtansine group [PFS = 4.1 months, range: 2.7–11.1; 1-year OS = 87.9%] [44]. These trials show that ICI plus anti-HER2 targeted therapy has a potential role in the treatment of PD-L1-positive tumors, but their samples are small and this research is in an early phase. Larger, ongoing clinical trials assessing this therapeutic approach will aid in the assessment of its effectiveness (Table 1).

ICI Plus Poly(ADP-Ribose) Polymerase Inhibitors. Olaparib and talazoparib are the 2 poly(ADP-ribose) polymerase (PARP) inhibitors that have been approved for the treatment of patients with metastatic breast cancer with germline BRCA1 or 2 mutations. As BRCA loss of function causes altered homologous recombination, these tumors with DNA repair deficiencies are highly susceptible to genomic instability, leading to the activation of intracellular cytotoxic interferon (IFN) genes and the accumulation of immunogenic neoantigens [45, 46]. In vivo experiments conducted with breast cancer cell lines and animal models have shown that treatment with PARP inhibitors leads to upregulation of PD-L1 expression [45, 47, 48]. These findings provided the rationale for the first clinical studies assessing ICI plus PARP inhibitor therapeutic combinations. The MEDIOLA trial, a single-arm phase 1/2 basket study, evaluated the effects of treatment with olaparib (300 mg orally, twice a day, for 4 weeks) followed by olaparib plus durvalumab (15 mg i.v. every 4 weeks) in patients with metastatic HER2-negative, germ-line BRCA-mutated solid tumors [49]. Disease control was achieved at 28 weeks in 15 out of 30 patients (50%), with a tolerable safety profile [49]. Similarly, the TOPACIO/KEYNOTE-162 study, an open-label phase 2 trial, evaluated the combination of niraparib (200 mg orally) and pembrolizumab (200 mg i.v. every 3 weeks) in patients with advanced or metastatic triple-negative breast cancer.
cancer. The overall response rate (ORR) in the whole cohort was 21%, and response rates were greater among patients with BRCA mutations (47%) and those with PD-L1-positive tumors (32%) than among those without BRCA mutation (11%) and PD-L1-negative patients (8%), respectively. These findings suggest the existence of synthetic lethal interaction with PARP inhibitor-ICI combination therapy, which might be most effective in PD-L1-positive patients, but it would be premature to draw definite conclusions at this stage of research. Additional studies are underway to further explore this possibility (Table 1) [50].

ICI plus Cyclin-Dependent Kinases 4/6 Inhibitors. Cyclin-dependent kinase 4/6 (CDK4/6) inhibitors have shown significant clinical activity in patients with metastatic hormone receptor-positive, HER2-negative breast cancer [51–53]. Although their antitumoral effect is achieved primarily by the suppression of retinoblastoma phosphorylation, which leads to cell cycle arrest and inhibition of cell proliferation, they also enhance tumor immunogenicity [54–56]. Thus, a possible synergistic effect of combination therapy with ICI has been proposed. An initial phase 1/2 study assessed the effect of the CDK4/6 inhibitor abemaciclib (150 mg orally twice a day) plus pembrolizumab (200 mg i.v. every 3 weeks) in 28 patients with hormone receptor-positive, HER2-negative metastatic breast cancer [57]. At 24 weeks, the researchers observed an overall response rate of 14% (4 patients) and a manageable safety profile [57]. These results seem promising when compared with the 11% early response rate achieved with abemaciclib monotherapy in the MONARCH 1 trial [58], a phase 2 study of abemaciclib as a single agent used to treat refractory hormone receptor-positive, HER2-negative metastatic breast cancer.

ICI plus Other Immunotherapies. As the immune system is complex, intertwined, and multidimensional, the combination of ICI with other immunotherapies seems to be a promising approach to account for potential escape mechanisms and to increase the treatment effectiveness relative to monotherapy. Such combinations are applied with the goals of enhancing immune cell infiltration, increasing immune cell activity, and augmenting cancer antigen presentation [46]. The combined use of anti-CTLA-4 and anti-PD-1 agents has been proven to be clinically effective and lead to durable response rates in patients with melanoma [59, 60]. In these clinical trials, the combination of these 2 agents increased the toxicity, but also the response rate, relative to monotherapy. A single-arm study assessed the effects of the anti-PD-L1 agent durvalumab (15 mg/kg i.v. every 4 weeks) combined with the anti-CTLA-4 agent tremelimumab (1 mg/kg i.v. every 4 weeks) in patients with metastatic hormone receptor-positive or triple-negative breast cancer [61]. Of the 18 enrolled patients (7 with triple-negative and 11 with hormone receptor-positive breast cancer), 3 (17%) patients with triple-negative breast cancer showed an overall response (43% response rate for this subgroup). These results permit speculation about the possible role of this combination for patients with triple-negative breast cancer, but further trials are warranted to clarify these findings.

Vaccine-Based Therapies

Examination of the use of the immunoregulatory capacities of antigen-presenting cells for vaccination and induction of an immune response in patients with cancer was initiated decades ago [62]. In subsequent years, further developments such as next-generation sequencing and nanotechnology led to the development of new therapeutic and diagnostic options [63, 64]. With these technologies, therapies based on sequencing of cancer cell DNA and encoding of tumor-specific mutations became possible [63]. Cancer treatment vaccines are based on the concept of strengthening the patient’s immune system to fight cancer cells. The key to this approach is to enable immune cells to recognize the tumor and be activated through substances in cancer cells [65]. Such vaccines can be derived using different approaches. Tumor-specific antigens (neoantigens) that arise from cancer-specific mutations and are specific to tumor cells can be used, but most of these neoantigens are unique to individual patients, necessitating a personalized approach to vaccination therapy [66, 67]. Another approach involves TAA, which are expressed mainly on cancer cells of specific cancer entities but may also be expressed at lower levels on normal cells. As T cells with a strong TAA affinity are normally removed by central and peripheral immune tolerance mechanisms, TAA-based vaccines must be sufficiently immunogenic to activate the remaining low-affinity TAA-reactive T cells [68, 69]. In vitro and in vivo research has produced promising results regarding the application of this vaccination therapy to breast cancer [70]. Chablaní et al. [71] reported a significant increase in tumor suppression capability after vaccination in murine breast cancer models. The first human studies on the use of personalized RNA mutanome vaccines in combination with PD-1 blockade therapy have been successful for melanoma [72]. Administration of an INVAC-1 vaccine to 26 patients with solid tumors (5 with breast cancer) led to stable disease in 58% of the patients, and 42% of the patients received multiple vaccinations (up to 12 cycles) because of an initial tumor regression [73].

A promising approach for breast cancer immunotherapy is represented by vaccines that use HER2-derived peptides as TAA. Mittendorf et al. [74] assessed the ability of vaccination with E75, a human leukocyte antigen (an A2/A3-restricted HER2 peptide), and granuloce-
macrophage colony-stimulating factor (GM-CSF) versus placebo to prevent disease recurrence in the adjuvant setting in patients with HER2-expressing breast cancer (immunohistochemistry [IHC] score 1–3). As they had determined that E75 vaccination safely and effectively elicited HER2-specific immunity in previous studies, they further examined dose escalation and schedule optimization according to lymph node status and the risk of disease recurrence in that phase 1/2 clinical trial [75]. The disease-free survival (DFS) rate was 94.3% in the vaccinated group (106 patients) and 86.8% in the control group (76 patients; \( p = 0.08 \)). Subgroup analysis showed that the patients who benefited most from vaccination were those with positive lymph nodes (DFS = 90.2% [vaccinated] vs. 79.1% [control]; \( p = 0.13 \)), a low HER2 expression (IHC score 1+ or 2+; DFS = 94% [vaccinated] vs. 79.4% [control]; \( p = 0.04 \)), or grade 1 or 2 tumor (DFS = 98.4% [vaccinated] vs. 86.0% [control]; \( p = 0.01 \)). The observation that the HER2 expression level affects the response to E75 vaccination, with patients with a low HER2 expression (IHC score 1+ or 2+ with fluorescence in situ hybridization [FISH] negativity) showing better immunological responses than patients with HER2-overexpressing tumors, has been reported previously and might guide further definition of the subgroups that benefit most from this vaccination therapy [76]. Given these promising results, Mittendorf et al. [77] conducted the PRESENT trial, a phase 3 study assessing the effects of vaccination with 1,000 μg nelipepimut-S (NP-S/E75; monthly for 6 months, then every 6 months through 36 months) plus GM-CSF (250 μg subcutaneously) relative to a placebo in patients with T1-T3 low HER2-expressing (IHC score 1+ or 2+ with FISH negativity) node-positive breast cancer in the adjuvant setting, after completion of neoadjuvant or adjuvant chemotherapy. An interim analysis revealed no significant difference in DFS between the NP-S group (24 patients; 6.3%) and the control group (37 patients; 0.8%) [77]. Approximately half of the recurrences observed in this trial were diagnosed by protocol-specific annual imaging of asymptomatic patients, without biopsy confirmation, and 74% of these recurrences were seen in the NP-S group, which led to termination of the trial after a median follow-up period of 16.8 months. As recurrence occurred 3 times more frequently in the NP-S arm than in the control arm, the researchers suspected that it was related to vaccination, but whether it represents a form of pseudoprogression or a true vaccine induced-recurrence remains unknown.

In addition to “preventive vaccination,” several attempts have been made to evaluate vaccination therapy in the metastatic setting. Miles et al. [78] conducted the largest phase 3 study to date assessing vaccination with sialyl-TN (STn) conjugated to the carrier protein keyhole limpet hemocyanin (KLH) versus placebo in 1,028 women with metastatic breast cancer who had previously received chemotherapy and had shown a response or disease stability. STn is a carbohydrate epitope found on a variety of glycoproteins, including mucin 1 (MUC-1) [78]. It is expressed by various tumor cells and seems to have functional significance in tumor growth and metastasis. Increased STn expression has been associated with the progression and poor prognosis of breast cancer, and STn is believed to be an important TAA [79–83]. In the phase 3 trial, patients received 1 intravenous dose of cyclophosphamide (300 mg/m²) 3 days before subcutaneous injection of 100 μg STn-KLH or placebo at weeks 0, 2, 5, and 9 [84]. Subsequently, the vaccine or placebo was administered without cyclophosphamide monthly for 4 months and then quarterly until disease progression. The vaccine was tolerated well and induced a considerable immune response, as reflected by serum titers and IgM and IgG antibodies. However, no significant difference was observed between the vaccine and placebo groups in the time to progression (3.4 vs. 3 months) or OS (23.1 vs. 22.3 months) [84].

Given the observation in preclinical studies that the MUC-1 protein might act as an oncoprotein for the activation of estrogen receptor-a function [85], Ibrahim et al. [86] conducted a retrospective subgroup analysis of data from the study of Miles et al. [84] to determine whether patients who received concurrent STn-KLH and endocrine therapy benefitted from vaccination. In the subset of patients receiving endocrine therapy (\( n = 350 \)), the median OS was significantly longer in the vaccination group than in the placebo group (36.6 vs. 30.7 months; \( p = 0.036 \)). The response to vaccination therapy depended on the immune response; the median OS of patients with greater antibody responses (anti-oncostatin M IgG titers >1:320) was significantly longer than that of patients with lesser responses (41.3 vs. 25.4 months; \( p = 0.0147 \)). Several other ongoing trials are further assessing the application of various promising vaccination therapies in early and metastatic disease (Table 3).

### Cellular Therapy

Adoptive cell therapy is based on isolation of immune cells (T cells in most cases) from a patient. These cells are then enriched ex vivo for tumor-specific cloning, expanded, activated, and autologously readministered to the patient [87, 88]. Zacharakis et al. [89] reported a case in which a patient with repeatedly treated metastatic hormone receptor-positive breast cancer showed durable disease regression for >22 months after the adoptive transfer of mutant protein-specific TIL in conjunction with interleukin-2 and checkpoint blockade. The concept of adoptive cell therapy using mutation-specific TIL for
Table 3. Tumor vaccination trials for breast cancer in the recruitment phase

| Agents                                                                 | Trial / Phase                                                                 | Setting   |
|------------------------------------------------------------------------|------------------------------------------------------------------------------|-----------|
| Dendritic cell vaccine (DC1)                                           | HER2-Directed Dendritic Cell Vaccine during Neoadjuvant Therapy of HER2+Breast Cancer / Phase 1 | Neoadjuvant |
| Mammaglobin-A DNA vaccine                                              | Safety and Immune Response to a Mammaglobin-A DNA Vaccine in Breast Cancer Patients Undergoing Neoadjuvant Endocrine Therapy / Phase 1 | Neoadjuvant |
| PVSRIPO                                                                | Open Label Immunotherapy Trial for Breast Cancer / Phase 2                     | Neoadjuvant |
| V3-MOMMO                                                               | Open Label Immunotherapy Trial for Breast Cancer / Phase 2                     | Neoadjuvant |
| Granulocyte-macrophage colony-stimulating factor, multi-epitope HER2 peptide vaccine H2NVC | A Vaccine (H2NVC) before Surgery for the Treatment of HER2-Expressing Ductal Carcinoma in situ / Phase 1 | Neoadjuvant |
| Poly ICLC                                                              | Safety and Immunogenicity of a Personalized Synthetic Long Peptide Breast Cancer Vaccine Strategy in Patients with Persistent Triple-Negative Breast Cancer following Neoadjuvant Chemotherapy / Phase 2 | Adjuvant |
| Neo-antigen pulsed dendritic cell                                      | Breast Cancer Neoantigen Vaccination with Autologous Dendritic Cells / Phase 1 | Adjuvant |
| Neoantigen DNA vaccine dnrvalumab                                      | Neoantigen DNA Vaccine Alone vs. Neoantigen DNA Vaccine plus Durvalumab in Triple-Negative Breast Cancer Patients following Standard of Care Therapy / Phase 1 | Adjuvant |
| P10s-PADRE with MONTANIDE™ ISA 51 VG                                   | Vaccination of Triple-Negative Breast Cancer Patients / Phase 2               | Adjuvant |
| DC1 vaccine WOKVAC vaccine                                             | Vaccine to Prevent Recurrence in Patients with HER-2 Positive Breast Cancer / Phase 2 | Adjuvant |
| Cyclophosphamide Multi-epitope folate receptor-a peptide vaccine       | Multi-Epitope Folate Receptor-a Peptide Vaccine, GM-CSF, and Cyclophosphamide in Treating Patients with Triple-Negative Breast Cancer / Phase 2 | Adjuvant |
| Typhoid vaccine                                                        | Typhoid Vaccine in Testing Response to Immune Stress in Patients with Stage I-III A Breast Cancer | Adjuvant |
| Pertuzumab Sargramostim                                                | TPIV100 and Sargramostim for the Treatment of HER2-Positive, Stage II-III Breast Cancer in Patients with Residual Disease after Chemotherapy and Surgery / Phase 2 | Adjuvant |
| Activated CIK and CD3-MUC1 bispecific antibody cyrotherapy            | Study of Activated Cytokine-Induced Killer Armed with Bispecific Antibody for Advanced Breast Cancer / Phase 2 | Adjuvant |
| YE-NEO-001                                                             | QUILT-2.025 NANT Neoepitope Yeast Vaccine (YE-NEO-001): Adjuvant Immunotherapy Using a Personalized Neoepitope Yeast-Based Vaccine to Induce T-Cell Responses in Subjects w/ Previously Treated Cancers / Phase 1 | Adjuvant |
| CD105/Yb-1/SOX2/CDH3/MDM2-polyepitope plasmid DNA vaccine              | Vaccine Therapy in Treating Patients with HER2-Negative Stage III-IV Breast Cancer / Phase 1 | Adjuvant/metastatic |
| HER2 DC1 vaccine                                                       | Immune Response and Potential Booster for Patients Who Have Received HER2-Pulsed DC1 / Phase 2 | Adjuvant/metastatic |
| HER-2 vaccine                                                          | Vaccine Therapy in Treating Patients with Metastatic Solid Tumors / Phase 1   | Adjuvant/metastatic |
| VRP-HER2 Pembrolizumab                                                | A Study to Evaluate Concurrent VRP-HER2 Vaccination and Pembrolizumab for Patients with Breast Cancer / Phase 2 | Metastatic |
| AE37 peptide vaccine Pembrolizumab                                     | Establishing the Recommended Biological Dose for AE37 Peptide Vaccine in Combination with Pembrolizumab that Will Enhance the Tumor-Specific Immune Response and Demonstrate Efficacy in Patients with Advanced Triple-Negative Breast Cancer / Phase 2 | Metastatic |
| PVX-410 pembrolizumab                                                 | PVX-410 Vaccine plus Pembrolizumab in HLA-A2+ Metastatic Triple-Negative Breast Cancer / Phase 1 | Metastatic |
| PF-06936308                                                           | A Study to Evaluate Escalating Doses of a Vaccine-Based Immunotherapy Regimen for NSCLC and TNBC / Phase 1 | Metastatic |
| SV-BR-1-GM INCMDGA000012 Cyclophosphamide                             | Combination Study of SV-BR-1-GM in Combination with INCMGA000012 and Epacadostat / Phases 1 and 2 | Metastatic |
the treatment of metastatic triple-negative breast cancer is currently being evaluated [90]. Another adoptive cell therapy involves the use of chimeric antigen receptor (CAR) T cells, for which T cells are genetically designed to express receptors against specific targets [91]. CAR T cells are being used currently for the treatment of hematopoietic malignancies, such as lymphoma [92]. Their application in solid tumors is being examined in experimental and clinical research [93]. Szőőr et al. [93] assessed the effect of HER2-specific CAR T-cell therapy in vitro and in xenografts with trastuzumab-resistant breast cancer. They observed that HER2-specific CAR T cells induced tumor regression and proved that antibody resistance can be overcome by targeting the same epitope with CAR T cells. Promising therapeutic approaches of this type are also available for triple-negative tumors. Song et al. [94] showed that T cells expressing folate receptor-α CAR inhibited the outgrowth of triple-negative breast cancer in vitro and in xenografts. In an early-phase clinical study, CAR T-cells specific for mesenchymal-epithelial transition factor (cMets) were injected into accessible cutaneous or lymph-node metastases in patients with metastatic breast cancer [95]. The researchers observed extensive necrosis and immune cell invasion in the excised lesions and concluded that the intratumoral injection of cMet-specific CAR T cells generated an in-
flammatory response within the tumors. Table 4 provides an overview of studies examining CAR T-cell therapy for breast cancer that are currently in the recruitment phase.

**Anti-HER2 Targeted Antibody Therapy**

Monoclonal antibodies are identical Ig produced by clonal immune cells [96]. Although their antitumoral effect is caused by their binding ability and the inhibition of their corresponding receptor, an additional immunological pathway exists for anti-HER2-targeted monoclonal antibodies by which the adaptive and innate immune systems are activated mainly through the antibody-dependent cell-mediated cytotoxicity of natural killer cells and monocytes against tumor cells [97–100]. Margetuximab, a novel chimeric anti-HER2 IgG1 antibody [101, 102], binds to the same receptor and has anti-proliferative effects similar to those of trastuzumab, but it is intended to have a stronger effect on the immune system due to modification of the Fc region. This effect is mediated by the increased binding capacity of both alleles of the activating CD16A receptor and the decreased binding capacity of the negative regulator CD32B receptor [102].

The low-affinity CD16A-158F allele has been associated with a decreased clinical response to trastuzumab [99, 103]. CD16A is an Fc receptor expressed on the surfaces of natural killer cells and macrophages that plays roles in the signal transduction antibody-dependent immune response and phagocytosis [104, 105]. In a phase 1 study, Bang et al. [105] first investigated the toxicity profile, optimal dosing schedule, pharmacokinetics, and antitumoral activity of single-agent therapy with margetuximab (0.1–6.0 mg/kg i.v. every 4 weeks or 10–18 mg/kg i.v. every 3 weeks) in patients with HER2-positive metastatic solid carcinomas. A tumor response was seen in 18 out of 23 (76%) patients with breast cancer, with a tolerable safety profile [105]. In the SOPHIA study, a phase 3 clinical trial, 536 patients with metastatic breast cancer who had received therapy with pertuzumab and 1–3 lines of chemotherapy for metastatic disease were randomized to receiving combinations of margetuximab (15 mg/kg i.v. every 21 days) or trastuzumab (6 mg/kg i.v. every 21 days) with chemotherapy (capecitabine, eribulin, gemcitabine, or vinorelbine) [106]. PFS was longer in the margetuximab arm than in the trastuzumab arm (5.8 vs. 4.9 months; HR = 0.76; 95% CI 0.59–0.98; p = 0.033). The safety profiles in the 2 arms were comparable, with grade 3 and 4 adverse events occurring in 52.3% of the patients receiving margetuximab and 48.3% of those receiving trastuzumab. In the ITT population, the OS duration was 18.9 months in the margetuximab arm and 17.2 months in the trastuzumab arm (HR = 0.95; 95% CI 0.69–1.31; p = 0.95). In a planned exploratory analysis conducted in patients who carried the CD16A-158F allele (which has a lower ligand-binding ability), the benefit of margetuximab for PFS was more pronounced (6.9 vs. 5.1 months; HR = 0.68; 95% CI 0.52–0.90; p = 0.005). Several ongoing clinical trials are assessing the use of margetuximab for metastatic breast cancer (Table 5).

Ertumaxomab, a trifunctional bispecific antibody currently under clinical testing, targets HER2, CD3, and the Fcy receptors I, IIa, and III and forms a tri-cell complex of tumor, T, and accessory cells [107, 108]. In a phase 1 study, Kiewe et al. [109] found that ertumaxomab elicited a clinical response in 5 out of 15 patients with metastatic breast cancer, as well as a strong T cell-associated immune response, with an acceptable number of drug-related adverse events.

**Cytokine-Activated Mediation Therapy**

Another treatment strategy is based on the assumption that monoclonal antibody therapy activates a signal that stimulates the release of type I IFN and INF-γ-producing CD8+ T cells [97, 110]. Continuing this thought, concepts are being developed and early studies are being conducted to examine the combination of IFN-γ and anti-HER2 antibodies [111]. Zhang et al. [111] produced an anti-HER2 single-chain variable fragment-IFN-γ fusion protein that showed activity superior to that of anti-HER2 antibodies in xenografts and was even effective on tumors with anti-HER2 resistance. In a phase 1 study, Han et al. [112] assessed the effect of IFN-γ administered weekly in combination with paclitaxel, trastuzumab, and pertuzumab in patients with HER2-positive breast cancer. The 9 enrolled patients tolerated the therapy well, but oncological outcomes from that study remain to be published [112].

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Table 5. Anti-HER2 targeted antibody trials for breast cancer in the recruitment phase

| Agent         | Trial / Phase                                                                 | Setting       |
|---------------|-------------------------------------------------------------------------------|---------------|
| Margetuximab  | A Study to Evaluate the Efficacy and Safety of Margetuximab plus Chemotherapy | Metastatic    |
|               | in the Treatment of Chinese Patients with HER2+ MBC / Phase 2                  |               |
|               | A Study of MGD013 in Patients with Unresectable or Metastatic Neoplasms / Phase |               |
|               | 1                                                                              |               |
Oncolytic Virus Immunotherapy

Oncolytic viruses (OV) are being used in a novel approach to immunotherapy, as they can induce cancer cell death, and to enhance the immune response to cancer [113]. They replicate in cancer cells and induce cell death without harming physiologic cells. Infection of the cancer cell can be achieved via membrane fusion or binding to surface receptors [114, 115]. Various mechanisms underlie the antitumoral activity of different OV. The induction of lysis of cancer cells and cytotoxic proteins against them, insertion of therapeutic genes into the viral genome, sensitization of tumor cells to other treatments, enhancement of tumor apoptotic activity, and amplification of the antitumoral immune response are possible mechanisms [114, 116]. The FDA approved talimogene laherparepvec, the first herpes simplex OV, for the treatment of metastatic melanoma in 2015. OV for breast cancer are being tested in preclinical models and very early-stage clinical trials. Because of their immune-stimulating effects, OV are being assessed as monotherapy and in combination with other immunotherapeutic agents [117]. A phase 1 dose-escalation study assessed the clinical efficacy and safety of single and recurrent intratumoral doses of the herpes simplex OV HF10 in 6 patients with recurrent breast cancer; the researchers reported that the agent was safe and well tolerated, with therapeutic potential [118, 119]. In a similar phase 1 clinical trial, the antitumoral activity of combination therapy with docetaxel (75 mg/m² i.v. every 3 weeks) and escalating doses of reovirus type 3 Dearing (i.v., median tissue culture infectious dose up to $3 \times 10^{10}$ on days 1–5 every 3 weeks) was assessed in 25 oncological patients; 1 patient had metastatic breast cancer and showed a complete response to treatment [120, 121]. In vivo studies have investigated the effects of OV as monotherapy, in combination with chemotherapy, and as sensitizers for ICI therapy in the neoadjuvant setting on triple-negative breast cancer, and they have shown durable response rates [122, 123]. Thus, data on OV to date have been generated mainly in preclinical models; a few clinical trials have been conducted, albeit with small samples of patients with various types of cancer. Additional studies are currently underway and will better characterize the clinical effectiveness of this approach (Table 6).

Conclusion

Immunotherapy is a rapidly emerging field in breast cancer, as evidenced by the plethora of preclinical and clinical concepts and ongoing trials. Initial studies established the role of immunotherapeutic agents as checkpoint inhibitors in the metastatic setting. Questions that remain to be answered include which chemotherapeutic backbone the optimal partner for combination therapy is, which combinations with other immune-oncologic substances are effective, and which timing of application is optimal. As immunogenic factors, such as PD-L1 and TIL

### Table 6. Oncolytic virus therapy for breast cancer in the recruitment phase

| Agents | Trial / Phase | Setting |
|--------|--------------|---------|
| Pelareorep, letrozole, atezolizumab, trastuzumab | A Window-of-Opportunity Study of Pelareorep in Early Breast Cancer / Phase 1 | Neoadjuvant |
| Ipilimumab, nivolumab, talimogene laherparepvec | Ipilimumab, Nivolumab, and Talimogene Laherparepvec before Surgery in Treating Participants with Localized, Triple-Negative or Estrogen Receptor-Positive, HER2-Negative Breast Cancer-Deleted / Phase 1 | Neoadjuvant |
| Paclitaxel, pelareorep, avelumab | A Study to Assess Overall Response Rate by Inducing an Inflammatory Phenotype in Metastatic Breast Cancer with the Oncolytic Reovirus PelareorEp in CombinaTion with Anti-PD-L1 Avelumab and Paclitaxel – BRACELET-1 Study / Phase 2 | Metastatic |
| ADV/HSV-tk, valacyclovir, radiation: SBRT, pembrolizumab | SBRT and Oncolytic Virus Therapy before Pembrolizumab for Metastatic TNBC and NSCLC / Phase 2 | Metastatic |
| PVSRIPO | Examining Bioactivity of PVSRIPO in Invasive Breast Cancer / Phase 1 | Metastatic |
| Pelareorep, retifanlimab | INCMGA00012 and Pelareorep for the Treatment of Metastatic Triple-Negative Breast Cancer, IRENE Study / Phase 2 | Metastatic |
| Cyclophosphamide and JX-594 dose escalation, cyclophosphamide and JX-594, cyclophosphamide | A Study of Metronomic CP and JX-594 in Patients with Advanced Breast Cancer and Advanced Soft-Tissue Sarcoma (METROmaX) / Phases 1 and 2 | Metastatic |
| TBio-6517, pembrolizumab | Study of TBio-6517, Given Intratumorally, Alone or in Combination with Pembrolizumab, in Solid Tumors / Phases 1 and 2 | Metastatic |
| ONCR-177, pembrolizumab | Study of ONCR-177 Alone and in Combination with PD-1 Blockade in Adult Subjects with Advanced and/or Refractory Cutaneous, Subcutaneous or Metastatic Nodal Solid Tumors / Phase 1 | Metastatic |
expression, decrease over the course of disease, an important aspect of immunotherapy will be its effectiveness for early-stage breast cancer. Another important aspect will be the identification of suitable biomarkers to identify patients who will benefit from certain treatment approaches. Understanding of the tumor microenvironment, the roles of the innate and adaptive immune systems in the development and progression of breast cancer, and factors that account for responses to immunotherapeutic agents is necessary to enable immunotherapy to come of age fully in breast cancer treatment.

**Conflict of Interest Statement**

The authors declare the following conflicts of interests. JCR has received travel grants from Medac GmbH (Wedel, Germany), Gedeon Richter (Budapest, Hungary), Celgene (Summit, NJ, USA), Daiichi Sankyo (Tokyo, Japan), and Pfizer (New York City, NY, USA) and has been an honorary speaker for Pfizer. L.S. has received travel grants from Medac GmbH (Wedel, Germany) and Celgene (Summit, USA) outside the scope of this work. E.-F.S. is receiving: grants from the University of Saarland, Storz, and Erbe; personal fees and other compensation from Roche (Basel, Switzerland), Pfizer (New York City, NY, USA), Celgene (Summit USA), Aigen (Thousand Oaks, CA, USA), and Astra Zeneca (Cambridge, UK); and other fees from Esai (Tokyo, Japan), Ethicon (Somerville, NJ, USA), Johnson & Johnson (New Brunswick, NJ, USA), Novartis (Basel, Switzerland), Tesaro (Waltham, MA, USA), Teva (Petch Tikwa, Israel), Medac GmbH (Wedel, Germany), MSD (Kenilworth, NJ, USA), Vifor (Sankt Gallen, Switzerland), Gedeon Richter (Budapest, Hungary), Takeda (Tokyo, Japan), and AGE (Buchholz, Germany).

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**Author Contributions**

J.C.R. and M.P.R. were responsible for the conception and design of this study, interpretation of data, and drafting of this paper. L.S., C.M., and A.C.K. contributed substantially to the drafting and editing of this work. E.F.-S. was involved in the conception of this study and critical revision of this paper. All of the authors reviewed this work and contributed to the final version submitted.

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