Treatment of Patients with Early Breast Cancer: Evidence, Controversies, Consensus

German Expert Opinions on the 17th International St. Gallen Consensus Conference

Behandlung von Patientinnen mit frühem Mammakarzinom: Evidenz, Kontroversen, Konsens

Meinungsbild deutscher Expert*Innen zur 17. Internationalen St.-Gallen-Konsensuskonferenz

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Introduction

The motto of this year’s 17th St. Gallen (SG) Conference on “Primary Treatment of Early Breast Cancer” (SG-BCC) was “Customizing local and systemic therapies for women with early breast cancer.” Targeting the treatment of early breast cancer more and more to the specific disease situation of each patient is a clinical challenge. The 60 breast cancer experts came from 25 countries, including five panel members from Germany (see Table 1). The SG-BCC recommendations are based on a majority vote of the panelists aiming to establish an international consensus for everyday clinical practice. The panelists come from a number of different countries with different health systems and resources. It is not surprising that this would also be reflected in the consensus. For some years now, a German working group has been commenting on the voting results of the SG-BCC panel and their agreement with the treatment recommendations of the Breast Commission of the "Arbeitsgemeinschaft Gynäkologische Onkologie e.V." (AGO Mamma) [1], which updates its recommendations every year.

Genetic Testing for High-Risk Mutations

General considerations

Genetic testing for mutations in high-risk genes (e.g. BRCA1/2) requires appropriate patient information and counselling. The German experts agree with the majority vote (78%) of the SG-BCC panelists that patients with a calculated risk of a pathogenic germline mutation > 10% should be offered genetic testing (Level of Evidence [LoE] 2bB AGO++). For further details, the German experts refer to the current recommendations of the AGO Mamma [1].

Genetic testing of which genes?

In addition to mutations in the known risk genes BRCA1/2, the SG-BCC panelists (67%) recommend mutation analysis of other
The clinical benefit of genetic testing is highest for the high-risk BRCA1/2 mutations with high and moderate risk of disease. The moderately penetrant genes mentioned (ATM, BARD1, BRIP1, CDH1, CHEK2, NBN, PALB2, PTEN, STK11, RAD51C/RAD51D, and TP53) are among the core genes of common panels and will therefore usually be analysed as well. Clinical consequences should preferably be studied in the context of prospective trials or clinical registries (LoE 3aB AGO+).

The German experts agree [1]. The AGO Mamma differentiates between mutations with high and moderate risk of disease. The clinical benefit of genetic testing is highest for the high-risk BRCA1/2 genes (LoE 1bA AGO++), as it results in effective preventive measures. The AGO has upgraded PARP (poly ADP-ribose polymerase) inhibition as an effective treatment option in metastatic breast cancer with germline BRCA1/2 mutation (gBRCA1/2). In early breast cancer, the OlympiA trial [3] (NCT02032823) with the PARP inhibitor olaparib has reached its primary endpoint according to a press release. The scientific data will be presented at the 2021 annual meeting of the American Society of Clinical Oncology (ASCO). The SG-BCC panelists (56%) voted in favour of future genetic testing of all patients eligible for adjuvant treatment with olaparib. The German experts agree in principle with the majority vote.

### Testing for adjuvant olaparib?

PARP (poly ADP-ribose polymerase) inhibition is an effective treatment option in metastatic breast cancer with germline BRCA1/2 mutation (gBRCA1/2). In early breast cancer, the OlympiA trial [3] (NCT02032823) with the PARP inhibitor olaparib has reached its primary endpoint according to a press release. The scientific data will be presented at the 2021 annual meeting of the American Society of Clinical Oncology (ASCO). The SG-BCC panelists (56%) voted in favour of future genetic testing of all patients eligible for adjuvant treatment with olaparib. The German experts agree in principle with the majority vote.

### Procedure in confirmed disease-causing mutation

Drawing on fictitious cases with different clinical scenarios, the SG-BCC panelists discussed how to counsel patients at genetically increased risk of breast cancer.

- If a disease-causing mutation in BRCA1, BRCA2 or PALB2 – high penetrance (odds ratio [OR] > 3) were detected, almost 85% of SG-BCC panelists would advise a 40-year old woman to undergo risk-reducing bilateral mastectomy. For 60-year old women, barely half the SG-BCC panelists (46%) voted in favour of this measure.

The German experts recommend that the decision-making process include a thorough counselling. Risk-reducing mastectomy is an effective procedure. Regular radiological monitoring, including MRI, may be an option. In some older women, prophylactic tamoxifen may also be an option [1].
• If a disease-causing mutation of intermediate penetrance (OR 2–3) was detected in the genes BARD1, CHEK2, CDH1, TP53, two-thirds of the SG-BCC panellists would recommend intensified screening, including MRI, for the 40-year-old woman, compared with only 42% of the SG-BCC panellists for the 60-year-old woman. A good third (35%) consider routine monitoring to be adequate in older women.

• If disease-causing mutations were detected in the low-risk genes ATM, BRRP1, NF1, RAD51C, RAD51D, FRANCC, STK11 (low penetrance; OR 1–2), 50% of the SG-BCC panellists voted for intensified screening, including MRI, for the 40-year-old woman, while 40% considered routine monitoring to be adequate. In the 60-year-old woman, the SG-BCC panellists voted by majority (62%) for routine breast screening. Only 30% recommended intensified monitoring, including MRI. The German experts agree with the SG-BCC majority votes on the genes with medium or low penetrance. The lower the penetrance and the higher the age of the patient, the less aggressive the screening should be.

Pathology

Relevance of the proliferation index (Ki-67)

For years, the significance and validity of Ki-67 testing for treatment decisions in early oestrogen receptor-positive (ER+) and HER2-negative (HER2−) breast cancer have been discussed. Recently, an international working group [4] recommended Ki-67 testing, stating that patients with early ER+/HER2− breast cancer (T1−2 N0−1) and Ki-67 ≤ 5% do not need adjuvant chemotherapy, whereas Ki-67 > 30% would warrant chemotherapy. Almost two thirds (62%) of SG-BCC panellists agreed with this statement.

The German experts basically agree with the SG-BCC vote, but points out that the question involves extreme “cut-off” values with high “inter-observer” concordance [4, 5–7].

For patients with ER+/PR+/HER2− breast cancer without lymph node involvement (N0), according to the SG-BCC vote (42%), a Ki-67 level of 30% and above indicates a high risk and thus the need for chemotherapy. The German experts do not agree with this majority vote, but agree with those panellists (36%) who state that no definitive Ki-67 cut-off in this situation exists (N0) indicating chemotherapy per se. For intermediate Ki-67 levels between 10 and 25%, it is also necessary from the German point of view to include other criteria for risk assessment.

Ki-67 measurement before and during neoadjuvant endocrine therapy

Almost two thirds of the SG-BCC panellists (61%) and the German expert group agree that Ki-67 testing should be performed in routine clinical practice for all patients with early ER+/HER2− breast cancer. Likewise, two thirds (68%) of SG-BCC panellists recommend Ki-67 testing during or after neoadjuvant endocrine therapy (NET) to assess treatment response. They also voted by a majority (70%) that the prognosis of patients with ER+/HER2− ductal breast cancer can be assessed by the change in Ki-67 levels after a 2–4 week course of endocrine therapy (NET). The German experts agree that a 2–4 week NET to assess endocrine sensitivity, as used for example in the German ADAPT trial [8] and in the POETIC trial [9], is an appropriate measure [1].

Focus on multigene signatures

In certain circumstances, multigene signatures can support the treatment decision for/against chemotherapy in early ER+/HER2− breast cancer. Based on various clinical scenarios, the SG-BCC panellists voted on when multigene signatures would be helpful. The starting point in each case is a patient with early ER+/HER2− breast cancer (tumour size 1–3 cm) eligible for chemotherapy. The clinical situation of this patient varied in terms of gender (male/female), menopausal status and age (pre/postmenopausal), axillary lymph node (LN) involvement (pN0, 1–3 LN, ≥ 4 LN), and tumour grading (G1, G2, G3).

The majority of SG-BCC panellists recommended gene expression analysis in selected patients (Table 2). A majority of the panellists (79%) rejected multigene signature in patients with ER+/HER2− primary breast cancer (1–3 cm) with four or more affected lymph nodes.

The German experts basically agree and emphasise that because of the available prospective data gene expression analyses are only indicated in patients with a maximum of three affected lymph nodes. Moreover, gene expression analyses should only be performed if the decision for/against chemotherapy cannot be based on the usual clinical and pathological factors. The voting of the SG-BCC panellists is in line with the AGO recommendations [1]. In addition, the German experts point out the poor data in men, which is why the argument for them can only be made by

>Table 2 In early ER+/PR+/HER2− breast cancer (T 1–3 cm), the multigene signature is recommended by the SG-BCC panel in selected cases. Consent of the German experts.

<table>
<thead>
<tr>
<th>Patient profile</th>
<th>Gene expression analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender</td>
<td>Never</td>
</tr>
<tr>
<td>Male</td>
<td>x (56%)*</td>
</tr>
<tr>
<td>Female</td>
<td>x (72%)</td>
</tr>
<tr>
<td>Menopause status</td>
<td></td>
</tr>
<tr>
<td>Premenopausal</td>
<td>x (67%)</td>
</tr>
<tr>
<td>Postmenopausal</td>
<td>x (64%)</td>
</tr>
<tr>
<td>Nodal status</td>
<td></td>
</tr>
<tr>
<td>Negative (N0)</td>
<td>x (63%)</td>
</tr>
<tr>
<td>1–3 involved lymph nodes</td>
<td>x (83%)</td>
</tr>
<tr>
<td>≥ 4 involved lymph nodes</td>
<td>x (79%)*</td>
</tr>
</tbody>
</table>

Tumour grading

| G1 | x (60%) |
| G2 | x (72%) |
| G3 | x (61%) |

* majority vote in each case
analogy. In the context of the SG-BCC recommendation, it should be noted that in Germany the multigene testing recommended by the AGO is reimbursed for routine care in NO patients, but in patients with LN involvement only under special contract arrangements (e.g., outpatient specialist management) [1].

Early TNBC: no PD1/PD-L1 testing

The SG-BCC panellists and the German experts agree that neither PD1/PD-L1 testing (majority vote: 93%) nor tumour-infiltrating lymphocytes (TILs; majority vote: 61%) are routinely indicated in patients with early triple-negative breast cancer (TNBC: ER−/PR−/HER2−) undergoing systemic treatment [1].

Neoadjuvant systemic therapy

General considerations

The concept of neoadjuvant therapy is recognised and favoured in Germany as a standard approach in early breast cancer as soon as adjuvant chemotherapy is indicated under the same treatment regimen [1]. In contrast, the SG-BCC panellists (60%) do not favour the concept of neoadjuvant therapy (60%). Pathological complete response (pCR) following neoadjuvant systemic therapy (NAST) may be used as a surrogate endpoint for drug approval in early breast cancer. With a clear majority (83%), the SG-BCC panellists put this approach into perspective: Achieving pCR is promising but inadequate when defining standard treatment. This can only be defined based on survival data. The German experts agree.

NACT or NET?

The SG-BCC panellists voted almost unanimously (98.21%) that in postmenopausal patients with ER+/HER2− breast cancer and low risk, based on clinical pathological criteria or gene expression analysis, for whom neoadjuvant therapy is planned, endocrine therapy should be preferred over chemotherapy. The German experts agree that chemotherapy is not indicated here. In Germany, neoadjuvant endocrine therapy (NET) is not standard [1].

In early ER+/HER2− breast cancer, 74% of SG-BCC panellists supported gene expression analysis of core biopsies to decide whether to administer NET or neoadjuvant chemotherapy (NACT). From the German perspective, the question – NET or NACT – is not significant clinically, as NET is not a standard regimen in Germany. Multigene signature is only reasonable in those patients where the indication for chemotherapy is questionable.

Neoadjuvant systemic therapy in HER2+ breast cancer

With a clear majority (85%), the SG-BCC panellists saw no need for anthracyclines in addition to neoadjuvant taxane/anti-HER2-based regimens in stage II breast cancer patients without LN involvement (cN0). In contrast, in LN involvement (c/pN+), 54% of SG-BCC panellists advocated neoadjuvant anthracycline/taxane-based chemotherapy plus anti-HER2 therapy.

In the modified case of a stage I/IICcN0 patient with neoadjuvant taxane/trastuzumab therapy, 35% of the panellists voted to administer an anthracycline plus pertuzumab in addition to the neoadjuvant taxane/trastuzumab regimen. Just under 30% (27%) chose pertuzumab/platinum and 24% opted to add pertuzumab only.

The AGO Mamma [1] rates anthracycline-free and anthracycline-containing standard regimens in combination with trastuzumab plus pertuzumab as equally effective. However, various long-term sequelae have been described, and this needs to be discussed with the patient. The German experts therefore agree with the SG-BCC vote that cN0 patients can receive a standard anthracycline-free regimen. This is also true for patients with LN involvement regardless of the stage (for example: six cycles of TcH [docetaxel, carboplatin, trastuzumab] or six cycles of TcHP [TcH + pertuzumab]). Some German experts, however, prefer treatment with anthracyclines in cases of higher risk, e.g., with lymph node involvement.

Neoadjuvant therapy in TNBC

In early TNBC, carboplatin is an effective treatment option alongside anthracyclines and taxanes. About 60% of SG-BCC panellists rejected neoadjuvant carboplatin in addition to anthracycline/cyclophosphamide/taxane-based treatment. The German experts see a difference to the German recommendations. According to the AGO Mamma [1], neoadjuvant platinum-containing chemotherapy can be used in early TNBC depending on patient risk profile and possible side effects (LoE 1a A AGO+). In Germany, carboplatin is usually combined with a taxane.

No checkpoint inhibition in early TNBC

The German experts agree with the SG-BCC panellists (90%) that at present checkpoint inhibitors should not be given in early TNBC outside of clinical trials [1]. Trial participation (GeparDouze, Alexandra, Neo Mono) is recommended [10–12].

Local Treatment Following NAST

Residual axillary tumour

The SG-BCC panellists (73%) and the German experts agreed that axillary lymph node dissection (ALND) is indicated if a macrometastasis (> 2 mm) was confirmed in the sentinel LN (SLN) or target LN (= biopsied and labelled LN; TLN) following NAST. The majority of SG-BCC panellists recommend ALND if 1/3 of the SLN is affected. The German experts agree in principle, but refers to the differentiated recommendations of the AGO Mamma [1] (Fig. 1). If a SLN or the TLN is “positive” following NAST, ALND is indicated irrespective of the number of LN examined and the size of the detected metastasis. There is agreement that axillary dissection is not justified when only isolated tumour cells are detected (ypN0[i+]).

The German experts criticise that the issues voted on do not contain any information on the axillary status before NAST. They add that ALND, just like SLND or TAD (“targeted axillary dissection”, i.e., TLN plus SLN excision [SLNE]), also serve diagnostic objectives. At present, however, there is no evidence for superiority of a regional therapy option (ALND vs. radiotherapy) in patients...
with ycN0 or ypN1 status. Basically, according to the German experts, the evidence is limited and partly based on empirical data. The AXSANA/EUBREAST 3 trial [13] undertaken by the AGO Breast study group and the AWOgyn (Arbeitsgemeinschaft für ästhetische, plastische und wiederherstellende Verfahren in der Gynäkologie e.V.) will close these gaps in knowledge.

Is it possible to omit ALND?

There was no consensus SG‑BCC vote on the question of whether ALND can be avoided in patients with positive nodal status (cN1) before treatment and marked, histologically positive TLN who convert to ycN0 status and whose lymphatic drainage areas (LDA) must be irradiated. 41% of SG‑BCC panellists agreed to omit ALND when 3/3 SLNs were tumour-free and 37% of panellists agreed when 1/1 SLN was tumour-free.

From the German perspective, the question cannot be answered because it is unclear whether the TLN had been removed. The evidence on this is sparse. Although the German experts emphasise that the AGO recommends both: TAD and ALND in the situation of ycN0. If the TLN corresponds to the SLN, a negative SLN suffices [14, 15]. More than 80% of SG‑BCC panellists (82%) recommend ALND when a patient with cN1 status (biopsy confirmed) has not responded or has only responded marginally to NAST (ypN1). For this situation (ypN+), the AGO Mamma recommends ALND (▶ Fig. 1) [1].

Patient with unsuspicious nodes at presentation (cN0) with positive SLN following NAST

The SG‑BCC panellists voted on whether radiotherapy of the axilla can replace ALND in selected patients with initial cN0 but positive SLN (ypN1).

- The majority of SG‑BCC panellists (62%) recommend ALND instead of radiotherapy in patients with 2/3 “positive” SLN, if there was at least one macrometastasis (> 2 mm).
If only 1/3 SLN demonstrates macrometastasis (> 2 mm), 48% of SG-BCC panellists voted for axillary radiotherapy; 52% voted for ALND.

When ypN1mic (> 0.2–2 mm) or ypN0(i+) (≤ 0.2 mm) was detected in 1/3 SLN, the majority of SG-BCC panellists (72% and 88%, respectively) favoured axillary radiotherapy over ALND.

The more residual tumour following NAST, the more SG-BCC panellists felt that level I/II radiotherapy alone did not have an adequate therapeutic effect, and therefore ALND should be performed. The German experts agree in principle, but refer to the markedly more differentiated AGO recommendations [1]. Due to the limited evidence, the AGO Mamma still recommends ALND. There is an increased risk of additional lymph node involvement following NAST and that – unlike in primary surgery [16, 17] – those are treatment-resistant cells with questionable response to radiotherapy. Due to the poor data in general, new recommendations must be postponed until the ongoing trials [13, 16, 18] have been completed. Overall, the SG-BCC questions on LN staging following NAST do not reflect the complexity of the situation.

TAD following NAST?
The surgical approach in the axilla following NAST does not depend on tumour biology. The German experts agree with the respective majority vote of the SG-BCC panellists on the TAD indication – assuming optimised and standardised technique [13].

- TAD is an adequate alternative to standard ALND (60%).
- TAD is an option in c/pN1 patients with conversion to ycN0 (90%).
- TAD is an option regardless of breast cancer subtype (85%).

Beside that “no surgery” is not an option in cases of presumed pathological complete remission, no new statement on breast surgery following NAST was presented at the SG-BCC this year. From the German perspective, however, it should be noted that surgery in the new, shrunken tumour volume remains standard according to the AGO Mamma, thus facilitating a very high breast conserving surgery rate [1].

**Surgery, Radiotherapy and Breast Reconstruction**

Management following mastectomy
On the question of how and when patients should undergo post-mastectomy radiotherapy (PMRT), 32% of SG-BCC panellists voted for an expander during PMRT (before planned reconstruction), while 20% would irradiate first and reconstruct later. The remainder favoured immediate reconstruction with autologous tissue (25%) or an implant (single- or two-stage procedure; 23%).

From the German perspective, all other options are also possible and should be discussed by the tumour board and with the pa-

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**Algorithm of breast reconstruction**

- **Patient wishes to undergo breast reconstruction**
  - N.B.: Habitus, breast volume, wishes, previous surgery

- **No postmastectomy radiotherapy**
  - SSM/NSM and implantation or MRM + tissue expander → implant or not suitable for alloplastic reconstruction or wish of patient → autologous reconstruction

- **Postmastectomy radiotherapy indicated**
  - Mastectomy
    - Radiotherapy
    - Delayed autologous reconstruction
  - Not suitable for autologous reconstruction
    - e.g., too little subcutaneous fat, wishes of patient
  - Direct prosthesis reconstruction or two-staged implant-based reconstruction: MRM → tissue expander → implant + radiotherapy
    - N.B.: Increased complication rate, particularly capsular fibrosis

**To be discussed in individual cases:**
- Immediate autologous reconstruction
  - N.B.: Increased fibrosis rate
- Delayed prosthesis reconstruction
  - N.B.: Increased complication rate

**Fig. 2 AGO Mamma algorithm for breast reconstruction following mastectomy, from: [1]. Source: Courtesy of AGO Mamma.**
tient (AGO ++) [1]. Delayed autologous reconstruction is preferred, with temporary expander or implant if needed. ▶ Fig. 2 illustrates the breast reconstruction algorithm recommended by the AGO Mamma [1]. The decision for breast reconstruction, especially delayed reconstruction, must be discussed with each patient individually. The German experts note an increased risk of complications (e.g., risk of capsular fibrosis) if radiotherapy is performed after implant reconstruction.

Hypofractionated radiotherapy following reconstruction?

If a patient with immediate breast reconstruction requires PMRT, the SG-BCC panelists (64%) without restrictions consider moderate hypofractionated radiotherapy a suitable option. The German experts note the clearly limited data available. The AGO Mamma has not commented on hypofractionated PMRT [1].

Radiotherapy Following Breast-Conserving Surgery

Hypofractionated radiotherapy as standard?

In stage I/II ER+/HER2– breast cancer following breast-conserving surgery (BCS) with negative resection margins, the majority of SG-BCC panelists (72%) voted for moderately hypofractionated whole-breast radiotherapy (WBRT; 15–16 fractions) as the preferred fractionation regimen, regardless of patient age. This is consistent with the S3 guideline and the recommendation of the AGO Mamma [1, 2]. 9% of the SG-BCC panelists favoured ultra-short course WBRT, in line with the FAST and FAST-Foward trials [19, 20]. The AGO Mamma defines ultra-short course WBRT as an alternative in selected cases (LoE 1bB AGO+). Trial enrollment is recommended (LoE 5D AGO ++). 8% of the SG-BCC panelists (70%) recommended continuing post-neoadjuvant treatment with trastuzumab and pertuzumab. If the patient was cN0 at initial diagnosis, the SG-BCC panelists (70%) recommend adding pertuzumab in high risk patients (cN+: LoE 1bB AGO+). Trial enrollment is recommended (LoE 5D AGO ++). If a patient does not achieve pCR after standard NAST, further post-neoadjuvant treatment with trastuzumab emtansine (T-DM1) is the therapy of choice for almost all SG-BCC panelists (89%). This corresponds to the AGO recommendation (LoE 1bB AGO+). According to the SG-BCC majority vote (77%) and the German experts, post-neoadjuvant T-DM1 is also indicated in small tumour residuals (<5 mm) [28].

Focus on partial breast irradiation

For partial breast irradiation (PBI) in stage I/II ER+/HER2– breast cancer without LN involvement, the majority of SG-BCC panelists saw no PBI indication in patients with lobular breast cancer (80%), or lymphovascular invasion (87%), or germline mutation (85%), and/or in patients <40 years (92%). The German experts agree in principle, but notes the limited data available, especially on germline mutations, which is why there is no guideline recommendation [1, 22]. However, the increased risk of ipsilateral secondary breast cancer due to genetic predisposition and the protective effect of WBRT support the recommendation [23].

Importance of multigene signatures in radiotherapy

According to the SG-BCC majority vote, commercially available multigene signatures do not provide a basis for deciding whether regional nodal irradiation (RNI: 92%) or chest wall irradiation (89%) is indicated. This also applies to the decision to forego radiotherapy in invasive breast cancer following breast-conserving surgery (84%). The German experts agree in each case. The AGO Mamma [1] and DEGRO [24] advise against the use of multigene signatures in these situations.

Post-neoadjuvant Systemic Therapy

General consideration

The prognostic significance of pathological complete response (pCR: ypT0/is pN0) following neoadjuvant chemotherapy (NACT) is undisputed. The question of whether the tumour stage at initial diagnosis or the intrinsic tumour subtype also affect the future outcome of a patient with pCR was supported by two-thirds of the SG-BCC panelists (65%). The German experts agree and add that the prognosis following NACT can be estimated with different models [25–27].

Post-neoadjuvant therapy in HER2 positive breast cancer

For patients with HER2+ breast cancer and clinically suspect lymph nodes at initial diagnosis (cN+) who achieve pCR with neoadjuvant chemotherapy plus trastuzumab and pertuzumab, the SG-BCC panelists (56%) recommend continuing post-neoadjuvant treatment with trastuzumab and pertuzumab. If the patient was cN0 at initial diagnosis, the SG-BCC panelists (70%) recommend additive administration of pertuzumab is not needed in the post-neoadjuvant setting. The German experts agree with both majority votes. The AGO Mamma recommends post-neoadjuvant trastuzumab (LoE 2aC AGO+++) in low risk of recurrence patients and additional pertuzumab in high risk patients (cN+: LoE 2bc AGO++) [1].

If a patient does not achieve pCR after standard NAST, further post-neoadjuvant treatment with trastuzumab emtansine (T-DM1) is the therapy of choice for almost all SG-BCC panelists (89%). This corresponds to the AGO recommendation (LoE 1bB AGO+). According to the SG-BCC majority vote (77%) and the German experts, post-neoadjuvant T-DM1 is also indicated in small tumour residuals (<5 mm) [28].

Post-neoadjuvant therapy in TNBC

In patients with early TNBC and pCR following NACT plus immunotherapy, the SG-BCC panelists (85%) see no indication for post-neoadjuvant checkpoint inhibitors. The German experts agree. No survival data from clinical trials are available yet.

If patients who do not achieve pCR, continued treatment with capecitabine is a post-neoadjuvant option. The German experts agree with the majority vote of the SG-BCC panelists (88%). The AGO-Mamma recommends up to eight cycles of capecitabine (LoE 1bB AGO+). Trial enrollment is recommended (LoE 5D AGO+) [1], for example, in the SASCIA trial [29].

Post-neoadjuvant therapy in ER+/HER2– breast cancer

The SG-BCC panelists stated unanimous (100%) that patients with early, hormone-sensitive (ER+/HER2–) breast cancer who have not achieved pCR in the breast but a good response in the axilla (pN0) after NET should receive post-neoadjuvant chemotherapy. The German experts agree with the statement, with the comment that conventional (4–6 months) NET is rarely used in
Germany and is primarily reserved for older patients or those with significant comorbidities.

No pCR following NET

Further SG-BCC questions on post-neoadjuvant chemotherapy in patients (ER+/HER2−) without pCR following NET rarely arise in Germany, as NET is not a standard treatment regimen in Germany. Moreover, the probability of pCR following NET is very low (5%).

Ductal Carcinoma in Situ (DCIS)

Postoperative radiotherapy in ER+ DCIS?

According to the SG-BCC panelists (majority vote: 58%), omitting postoperative radiotherapy following BCS of ER+ DCIS with adequate resection margin is justified in all patients over 70 years of age, and in principle (no age limit in the question) in patients at low biological (“luminal-like”) or genomic (multigene testing) risk (70%) as well as in low grade tumours (G1, 74%). Two-thirds of the SG-BCC panelists (67%) recommend omitting postoperative radiotherapy only in older patients (> 70 years) with at least one of the low-risk factors noted above. A simple majority (53%), on the other hand, saw an indication for postoperative radiotherapy in unifocal DCIS (≤ 2 cm) without necrosis.

The German experts point out that postoperative radiotherapy following BCS of ER+ DCIS with adequate free resection margin reduces the recurrence rate in the affected breast, but has no effect on overall survival [30]. This should be discussed with each patient individually. In the view of the Germans group, the voting results reflect that the overall risk profile should be taken into account when deciding for or against radiotherapy following BCS in DCIS.

Endocrine therapy following BCS and R0 situation

Various options are available if a patient with ER+ DCIS and postoperative radiotherapy also desires endocrine therapy as recurrence prophylaxis [1]. The vast majority of the panel favoured endocrine therapy (83%) with standard-dose (20 mg/day) or lower-dose (5 mg/day) tamoxifen or an aromatase inhibitor (AI).

The AGO Mamma recommends endocrine therapy in addition to adjuvant radiotherapy as an option in individual cases (AGO+/−). All the choices listed (tamoxifen 20 mg, tamoxifen 5 mg, AI [only in postmenopausal patients]) are an option [1]. The German experts emphasise that the indication for endocrine therapy depends on possible risk factors, potential side effects and the patient’s wishes. The patient should be informed that endocrine therapy is not associated with an overall survival benefit, but may in particular reduce the risk of secondary cancer in the other breast (LoE 1a) [1].

Radiotherapy

Importance of the boost

The SG-BCC panel heterogeneously discussed the issue of routine boost irradiation following BCS and WBRRT in patients with early invasive breast cancer. Nearly half (47%) would boost the tumour bed only in younger patients (18% in < 40 years, 29% in < 50 years), 31% would base their indication on tumour biology alone (G3, extensive intraductal component, HER2-positive, TNBC), while 20% regard the boost as indicated in every patient. From the German perspective, the heterogeneous vote is also reflected in the recommendations of the AGO Mamma (Fig. 3) [1]. The AGO Mamma clearly recommends the boost in premenopausal patients (LoE 1b B AGO++). In postmenopausal patients, boost is only indicated in patients at increased risk (LoE 2b B AGO+) [1].
Radiotherapy following NACT

<table>
<thead>
<tr>
<th>Pretreatment</th>
<th>Posttreatment</th>
<th>RT-BCS</th>
<th>PMRT</th>
<th>RT-RN*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Locally advanced</td>
<td>pCR/no pCR</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>cT1/2 cN1+</td>
<td>ypT1+ or ypN1+ (no pCR)</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>cT1/2 cN1+</td>
<td>ypT0/is ypN0</td>
<td>Yes</td>
<td>Increased risk of relapse</td>
<td></td>
</tr>
<tr>
<td>cT1/2 cN0 (ultrasound mandatory)</td>
<td>ypT0/is ypN0</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>

Local advanced: T3–4 or cN2–N3

1 Criteria for increased risk of relapse:
- pN0 premenopausal high risk: central or medial tumor localization, and (G2–3 and ER/PR negative)
- Pretreatment pN1a/cN+ high risk: central or medial tumor localization and (G2–3 or ER/PR negative) or premenopausal, lateral tumor localization and (G2–3 or ER/PR negative)

* Regarding coverage of axilla level I/II please also see slides “Additional RT of the axilla after primary surgery”

** confirmed by core biopsy

Boost in DCIS?
The SG-BCC panelists reject routine radiation boost both generally in DCIS (89%) and also in low-risk DCIS patients (96%). A slight majority (55%) rejects routine radiation boost in DCIS patients < 50 years of age. In contrast, two-thirds of SG-BCC panelists (65%) recommend routine radiation boost in DCIS patients at increased risk, for example, due to necrosis, close resection margins (< 2 mm), and large lesions.

The AGO Mamma recommends radiation boost in DCIS patients only in special cases at increased risk (LoE 1bB AGO+/−). According to the AGO, this includes patients < 50 years of age or those ≥ 50 years if additional risk factors are present (e.g., symptoms, G2/3, central necrosis, close resection margins, multifocal tumour, etc.) [1]. The German experts point out that this is an individual decision that should be discussed in the multispecialty tumour board and with the patient. Data from a randomised trial [31] on this question was presented for the first time at the San Antonio Breast Cancer Symposium 2020 showing no difference between hypofractionated radiotherapy and conventional radiotherapy and a moderate advantage of adding the boost.

Moderate hypofractionation in invasive cancer

The SG-BCC panelists regard hypofractionation as adequate radiotherapy modality following mastectomy (90%) and in regional nodal irradiation (RNI) (76%). – The AGO Mamma has not issued a statement regarding fractionation in PMRT without RNI. For combined irradiation of the chest wall with RNI, the German experts recommend conventional fractionation due to the limited data available on hypofractionation.

The AGO Mamma regards hypofractionated RNI as an option in selected cases (LoE 2bB AGO+). The publication by Wang et al. [32] is based solely on patients with locally advanced breast cancer with a short follow-up period. In Germany, the standard is conventional fractionation over a period of five weeks (LoE 1aA AGO++). Current ongoing trials will clarify this question [33–36].

The majority of SG-BCC panelists (59%) voted in favour of hypofractionated radiotherapy as the standard option for the chest, thoracic wall and regional lymph nodes – only rare circumstances such as repeat (second) radiotherapy were excluded. Due to the limited data on hypofractionation, the AGO Mamma recommends conventional fractionated radiotherapy in this indication, but considers hypofractionated irradiation an option (LoE 2bB AGO+/−). Only 21% of SG-BCC panelists considered hypofractionated radiotherapy a standard option only following BCS, regardless of the patient’s age. This vote corresponds to the recommendations of the AGO Mamma [1]. Regardless of this, it was pointed out at the SG-BCC that obstacles to the implementation of hypofractionation, for example, billing models based on the number of radiation fractions, should be reduced [37, 38].

Regional node irradiation (RNI) following NAST

Another focus of the SG-BCC vote was the importance of RNI following NAST in TNBC or HER2+ breast cancer stage II and above. In clinically suspicious LN prior to NACT (cN0), a clear majority of SG-BCC panelists saw no indication for RNI in pCR of the primary tumour (TNBC: 86%; HER2+: 90%). In contrast, in patients with stage II/III, pCR and clinically suspicious LN (cN1) prior to NACT, 70% (TNBC) and 65% (HER2+) routinely recommended RNI despite pCR. Only 26% (TNBC) and 30% (HER2+) restricted this to stage III patients.

With reference to the recommendation of the AGO Mamma, the German expert group emphasises that the indication for RNI should be risk-adapted (Fig. 4) [1]. They therefore fully agree with the SG-BCC vote in the cN0 patient. In contrast, RNI is indicated in cN1 patients with initial stage III despite pCR. In stage II, the indication for RNI should be discussed with the patient depending on other risk factors.

Fig. 4 Recommendations of the AGO Mamma on radiotherapy following neoadjuvant systemic chemotherapy, from: [1]. Source: Courtesy of AGO Mamma.
Elderly patients with life expectancy > 10 years

According to the AGO Mamma, adjuvant radiotherapy can be omitted in patients with small invasive ER+/HER2− breast cancer (pT1pN0) who have undergone breast-conserving surgery and have a life expectancy of less than 10 years, after individual consultation and accepting an increased risk of intramammary recurrence. This requires that the cancer has been resected completely (R0) and that the patient receives adjuvant endocrine therapy (LoE 1aB AGO++) [1].

The SG-BCC also discussed the question of adjuvant radiotherapy in older patients (> 70 years) with breast-conserving surgery in ER+/HER2− breast cancer and a life expectancy of more than 10 years. The majority of SG-BCC panelists did not recommend this option in general (90%). The SG-BCC panelists (88%) consider to omit additional radiotherapy, especially in patients with small ER+/HER2− breast cancers (< 2.5 cm) and low clinical or genomic risk. According to the SG-BCC majority vote, radiotherapy should not be omitted in larger tumours (> 2.5 cm/n0; majority vote: 80%), in the case of a positive SLN (90%) and in the case of unfavourable clinical/biological factors or high genomic risk (92%).

From the German perspective, the decision for or against adjuvant radiotherapy in older women with low-risk breast cancer requires an individual risk-benefit analysis. Trial data are so far only available with a follow-up period of up to about ten years [39]. Important options in older patients, which may affect the risk-benefit analysis in the future, are partial-breast radiotherapy and, potentially ultra-hypofractionated irradiation [1].

Intraoperative radiotherapy

Overall, 61% of SG-BCC panelists agreed that there are patients eligible for intraoperative radiotherapy (IORT) as sole modality. The German experts agree. The AGO Mamma considers intraoperative radiation alone to be a therapeutic option in patients > 70 years of age with a low risk of recurrence (LoE 1bA AGO+) and in specific cases in patients > 50 years of age (LoE 1bA AGO+/−) [1]. Due to the methodical limitations of the TARGIT-A trial [40,41], no general recommendation favouring IORT as sole modality can be given [22].

Adjuvant Systemic Therapy in ER+ Breast Cancer

General considerations

Half of the SG-BCC panelists defined the threshold of ER-positive cells detected by immunohistochemistry at ≥ 1% and ≥ 10% respectively. From the German perspective, the cut-off ≥ 1% is adequate for adjuvant endocrine therapy, although knowing that 1–10% ER-positive cells are considered as “questionably endocrine sensitive”. Adjuvant endocrine therapy may be offered to these patients (LoE 3bD AGO+). With ER-positive cells > 10%, there is a clear indication for treatment (LoE 1A AGO++) [1].

Patients with “questionably endocrine sensitive” ER+/HER2− breast cancer must be informed accordingly. The German experts add that nowadays more sensitive antibodies are used in the detection of ER-positive cells and that it can be assumed that patients with ER-low expression (“low expressers”) may have more biologically aggressive “basal-like” breast cancer.

The German experts and the SG-BCC panelists agree that patients with luminal A- or luminal B-like breast cancer without lymph node involvement (pN0) may benefit from adjuvant endocrine therapy regardless of tumour size – even in the case of microinvasion in the sentinel node (SG majority vote: 59 and 58%, respectively).

In case of ER+/HER2+ breast cancer (pN0), the majority of SG-BCC panelists (51%) voted in favour of adjuvant anti-HER2-based therapy if the tumour size was 5 mm or larger. A minority of 15% would administer adjuvant anti-HER2-based therapy regardless of tumour size. – The AGO Mamma recommends the use of trastuzumab (Lo2bB AGO+) in pN0 patients and a tumour size > 5 mm and for tumour sizes > 10 mm (LoE 1aA AGO++). In tumours ≤ 5 mm in diameter, adjuvant trastuzumab is possible on an individual risk-adapted decision (LoE 2bB AGO+/−) [1].

Duration of endocrine therapy in the premenopausal patient

If a premenopausal patient with ER+/HER2− breast cancer and high risk of recurrence received adjuvant tamoxifen plus OFS (ovarian function suppression) for five years, almost 90% of SG-BCC panelists recommend continuing endocrine therapy for another five years. A slight majority (45%) favour monotherapy with tamoxifen, 41% want to switch to an AI (plus OFS, if the patient remained premenopausal). Only four percent would continue tamoxifen/OFS.

The German experts emphasize that there is no valid data for extended adjuvant endocrine therapy (EAT) after five years of tamoxifen/OFS. From the German perspective, continued treatment with tamoxifen is an option. The AGO Mamma recommends continued treatment with tamoxifen (years 6–10) (LoE 1aA AGO++) after initial five years of tamoxifen and regards continued treatment with tamoxifen alone as a “possible” option (LoE 5D AGO+) after initial five years of endocrine therapy plus OFS [1].

Duration of endocrine therapy in lymph node involvement

According to the recommendation of the AGO Mamma, the standard is adjuvant endocrine therapy for five years (AGO++) [1]. For patients with lymph node involvement at initial diagnosis, the majority of SG-BCC panelists vote for endocrine therapy beyond five years: 34% recommend 7–8 years and 53% would treat for a total of ten years. The AGO Mamma recommends treatment duration beyond five years after individual risk-benefit analysis (AGO++). Duration and choice of treatment and, if necessary, the sequence (AI or tamoxifen) depend, among other things, on patient menopausal status, treatment tolerance, risk of recurrence, and the patient’s wishes [1].

Adjuvant therapy with CDK4/6 inhibitors?

CDK4/6 inhibitors have not yet been approved in Germany for adjuvant therapy in ER+/HER2− breast cancer. The only positive trial data currently available – albeit with a still short follow-up period (<20 months) – is for adjuvant abemaciclib from the monarchE trial [42]. In the SG-BCC vote, 54% of panelists favour abemaciclib
in addition to adjuvant endocrine therapy in patients with at least four involved lymph nodes. In patients with 1–3 involved LN and additional risk factors (e.g. G3 and/or T3 or high Ki-67), an equally narrow majority of SG-BCC panelists (54%) reject adjuvant abemaciclib. About 60% of the SG-BCC panelists do not see additional Ki-67 testing (in addition to other prognostic markers) as an option to allow patients the adjuvant treatment with a CDK4/6 inhibitor.

The AGO Mamma considers adjuvant abemaciclib over two years in addition to standard endocrine therapy to be an option in patients with an increased risk of recurrence and if the inclusion criteria of the monarchE study are met (LoE 2bC AGO+−) [1].

**Use of gene expression signatures**

The majority of SG-BCC panelists (79%) reject the indication for chemotherapy in postmenopausal patients if their genomic risk according to the clinical criteria of the MINDACT [43], TAILORx [44] or RxPonder [45] or similar trials is low and/or their recurrence score (RS) is ≤ 25.

The German experts agree in principle. Gene expression analysis should only be used if the traditional clinical-pathological factors (tumour size, nodal involvement, grading, Ki-67, ER/PR as well as HER2) do not allow decision-making for or against chemotherapy followed by endocrine therapy (vs. endocrine therapy alone). If gene expression analysis is indicated, the recommendation resulting from the analysis should be followed. In addition, the German experts refer to the current data of the ADAPT trial [8] as well as to the ADAPTtate [46] and ADAPTcycle [47] trials on this issue, which are currently recruiting patients with intermediate and higher clinical risk.

**Focus on OFS**

Standard adjuvant endocrine therapy in premenopausal patients (ER+/HER2−) is tamoxifen for 5 years (AGO 1aA++) if the risk of recurrence is low, plus OFS (LoE 2bC AGO++) in higher risk of recurrence. Tamoxifen/OFS treatment should only be given as long as it is tolerated by the patient and she is clearly premenopausal. According to the AGO recommendation, tamoxifen/OFS or AI/OFS is an option following chemotherapy once ovarian function returns within 24 months [1].

The German experts do not agree with the majority vote of the SG-BCC panelists recommending in principle an OFS in premenopausal patients with clinical stage II (71%). In patients < 40 years of age, as many as 94% would expand treatment with OFS. From a German perspective, the issue is not differentiated enough. The indication for OFS is based on the risk of recurrence.

If a patient with stage II ER+/HER2 breast cancer is premenopausal after initial chemotherapy, the question of further endocrine therapy was addressed. 43% of SG-BCC panelists see the indication for OFS (plus tamoxifen) in all patients, while 52% rely on supplementary OFS only in “high risk” cases (age < 40 years, lymph node involvement [N+], high Ki-67 and/or luminal B carcinoma, or intermediate or high risk according to gene expression analysis). In a separate vote, 94% of the SG-BCC panelists favour OFS in principle as part of endocrine therapy in patients with a risk of recurrence that justifies the indication for chemotherapy, as long as the patient remains premenopausal.

From the perspective of the German experts, the voting results reflect that, regardless of the stage, it must be differentiated between patients with low and high risk of recurrence and that an OFS is only indicated with increased risk [1]. The previous chemotherapy is a surrogate marker for high risk.

**Role of multigene signatures in endocrine therapy**

In premenopausal patients with ER+/HER2− breast cancer without lymph node involvement and with low/intermediate genomic risk, e.g. RS 16–25, the majority of SG-BCC panelists (53%) agree on OFS in addition to tamoxifen or an AI. Almost a quarter recommended only tamoxifen (22%) or endocrine therapy plus chemotherapy (24%).

The German experts cannot completely agree with the majority vote. According to the AGO Mamma [1], chemotherapy plus endocrine therapy can be useful in this group with individually increased risk of recurrence [44]. This must be discussed with the informed patient and decided individually.

In patients with 1–3 positive LN and low genomic risk (e.g., RS ≤ 25), 30% of SG-BCC panelists recommend chemotherapy followed by oral endocrine therapy, while 17% refuse oral endocrine therapy plus OFS. About one quarter consider both treatment options to be adequate, and 26% would prefer chemotherapy or endocrine monotherapy.

With regard to the indication for OFS, the German experts add that this should be independent of the genomic risk. So far, there is no data clearly proving a correlation between OFS indication and multigene test result.

**Oestradiol level during OFS?**

For patients under OFS, 53% of SG-BCC panelists recommend routine measurement of oestradiol levels, while 47% reject this step. From the German perspective, there is no reason for routine testing. This corresponds to the 50:50 vote of the panelists. Testing should be done after hysterectomy to determine menopausal status and may also be useful during endocrine therapy with an AI plus GnRH analogue to verify endocrine suppression.

**Chemotherapy effect in premenopausal patients**

Chemotherapy efficacy in premenopausal patients is based not only on the cytotoxic effect but also on the ovarian suppression induced by chemotherapy – especially in patients with favourable biological factors (positive ER/PR status, well-differentiated cancer, low Ki-67 score, low genomic risk). The extent of an endocrine effect of the chemotherapy itself is under discussion. The voting result of the SG-BCC panelists was quite heterogeneous. From the German perspective, it is impossible to differentiate between cytotoxic and chemotherapy-induced endocrine effects.

**Effective chemotherapy regimens**

In patients with stage I/II ER+/HER2 breast cancer without LN involvement and chemotherapy indication, the majority of SG-BCC panelists (34%) recommend an anthracycline/cyclophosphamide/taxane-based regimen, plus 6% who chose a dose-dense anthracycline-containing regimen. An anthracycline-free regimen with either four (32%) or six cycles (12%) of taxane/cyclophosphamide (TC) was favoured by 44% of panelists. The broad vote is in
line with the recommendations of the AGO Mamma [1]. The German experts refer to the standard treatments recommended by the AGO Mamma [1]. The chemotherapy regimen to be used must be decided individually with the patient taking into account potential adverse events. In patients with low volume LN involvement, anthracycline-free standard chemotherapy regimens are generally considered to be equivalent to anthracycline-containing standard regimens. Equal efficacy with the standard anthracycline-taxane sequences has been demonstrated for the TC regimen only if six cycles (6× docetaxel/cyclophosphamide) are administered.

Focus on the postmenopausal patient

Triage as such ADAPT [8], MINDACT [43], TAILORx [44] and RxPONDER [45] have studied the impact of endocrine therapy ± chemotherapy in ER+/HER2− breast cancer. Based on the trial outcomes, the SG-BCC panelists see the indication for chemotherapy in addition to endocrine treatment (vs. endocrine treatment alone) in the majority of postmenopausal stage III patients, regardless of biomarkers (68%) and in the case of large tumour volume, for example, N3 (> 10 affected lymph nodes) or T3N2 (96%). The German experts agree in each case here, since in the high-risk clinical situation there is the basic indication for chemotherapy.

When asked whether chemotherapy is indicated in the same patient – postmenopausal, stage III – with G1/2 cancer and lobular histology, 48% SG-BCC panelists agree, while 52% reject this recommendation. In case of low-risk G1 cancer with Ki-67 < 10%, 63% do not recommend chemotherapy. In terms of the RS, 61% of SG-BCC panelists reject chemotherapy for RS < 11, while 58% see an indication for chemotherapy for RS > 25. The German perspective it is impossible to comment on the indication for chemotherapy due to the limited information in the question. This is also reflected in the ambivalent outcome of the voting. The German experts add that the decision on chemotherapy does not depend on histology (NST or lobular), but on the known clinical-pathological factors and, if needed, on gene expression analysis.

Chemotherapy for high tumour burden

If chemotherapy is indicated in ER+/HER2 negative breast cancer with locally advanced stage or with a high tumour burden, the German experts refer to the standard chemotherapies as recommended by the AGO Mamma [1].

Adjuvant Systemic Therapy in Estrogen Receptor-negative (ER−) Breast Cancer

TNBC and ER−/HER2+ breast cancer

In pN0 patients with ER-negative (ER−) and HER2+ breast cancer and tumour size of 5–6 mm and larger, a majority of SG-BCC panelists (52%) recommend adjuvant anti-HER2-based systemic therapy. Almost as many (46%) would also start adjuvant anti-HER2 therapy in smaller lesions (including 12% even in microinvasion). The German experts add that the prognostic data of this patient group reveals a significant risk of recurrence regardless of tumour size [48 – 50]. Recent retrospective data [51] suggests an effective effect of adjuvant anti-HER2 therapy even in very small HER2+ breast cancers (pT1a).

In case of TNBC without LN involvement (pN0), the majority of SG-BCC panelists (46%) favour adjuvant systemic therapy if the tumour size is 5 mm or larger. The German expert group agrees with reference to the AGO Mamma [1].

Adjuvant Systemic Therapy in HER2-positive Breast Cancer

Trastuzumab ± Pertuzumab

Patients with HER2+ breast cancer usually also require adjuvant anti-HER2 targeted treatment if chemotherapy is indicated. The AGO Mamma recommends trastuzumab-based adjuvant therapy in patients without LN involvement and tumour size larger than 5 mm (6–10 mm: LoE 2bB AGO+; > 10 mm: LoE 1aA AGO++). This decision must be re-evaluated on a case-by-case basis in HER2+ breast cancer ≤ 5 mm (LoE 2bB AGO+/-). Adjuvant trastuzumab plus pertuzumab is recommended in patients with lymph node involvement (pN+) (LoE 1bB AGO+) and is an option only in some patients without lymph node involvement (LoE 1bB AGO+/-) but at increased risk [1]. The German experts therefore agree with the majority vote of the SG-BCC panelists (94%) that patients with HER2+ breast cancer without LN involvement should not receive adjuvant pertuzumab in addition to trastuzumab.

Adjuvant use of neratinib?

The adjuvant use of neratinib in patients with prior (neo)adjuvant trastuzumab/pertuzumab and/or trastuzumab emtansine (T-DM1) regimen is supported by 63% of the SG-BCC panelists in the positive ER (ER+) and high risk of recurrence (for example, ≥4 involved LN). The AGO Mamma recommends that patients with ER+/HER2+ breast cancer who have already received one year of trastuzumab should continue treatment with neratinib for one year in combination with standard endocrine therapy as an option (LoE 1bB AGO+), and in the post-neoadjuvant setting on an individual basis in non-pCR patients (LoE 2bB AGO+/-) [1]. Due to a lack of data, there is no recommendation on the use of neratinib following trastuzumab/pertuzumab and T-DM1 treatment.

The German experts agree with the majority vote of the SG-BCC panelists, as the AGO Mamma sees a possible indication in ER-positive cases at increased risk. The German experts note the potential adverse events of neratinib, which must be discussed with the patient.

Anthracyclines and anti-HER2 therapy

A clear majority (76%) of SG-BCC panelists agree that there are patients with HER2+ breast cancer who can receive an anthracycline sequentially in addition to anti-HER2 therapy combined with anthracycline-free chemotherapy. The German experts agree with this vote. Sequential anthracycline administration is an option depending on individual risk and individual (in particular cardiac) comorbidities.
T-DM1 instead of trastuzumab/paclitaxel?

Two-thirds of SG-BCC panellists (69%) see no indication for T-DM1 in the adjuvant setting in patients with HER2+ stage I breast cancer, while 31% would consider adjuvant T-DM1 in special circumstances. The German experts agree and add that T-DM1 is not approved for this situation because there are no study data for such an approach.

Adjuvant Systemic Therapy in TNBC

Adjuvant PD1-/PD-L1-targeted therapy in TNBC?

About 90% of SG-BCC panellists see no indication for PD/PD L1-targeted immune checkpoint inhibitors in addition to adjuvant chemotherapy in patients with stage II/III TNBC. The German experts agree with the SG-BCC majority vote and note the insufficient data and lack of approval of immune checkpoint inhibitors in early TNBC. In terms of adjuvant treatment, reference is made to the currently ongoing Alexandra trial [10]. They add that patients with early TNBC and chemotherapy indication should primarily receive neoadjuvant treatment [1].

Adjuvant use of PARP inhibition?

One hopeful therapeutic perspective in early BRCA1/2-associated breast cancer is the use of PARP inhibitors. In terms of the future outcomes of the OlympiA trial [3], 48% of SG-BCC panellists would support adjuvant olaparib in BRCA1/2-associated breast cancer if after three years of follow-up the OlympiA trial will show an absolute benefit in invasive disease-free survival (iDFS) of > 5% in the olaparib arm versus the control arm. From the German perspective, we should wait for the presentation of the outcome data at the ASCO meeting in June 2021.

Surgical Issues

BCS plus radiotherapy in locally recurrent breast cancer

If a patient is diagnosed with intramammary recurrence and/or ipsilateral second breast cancer more than five years after surgery for the primary tumour followed by radiotherapy, 63% of the SG-BCC panellists see BCS plus radiotherapy as a suitable alternative to mastectomy. The German experts agree in principle, but still recommend as primary option mastectomy (LoE 3b AGO++). According to the AGO Mamma, BCS is an option if subsequent (partial) irradiation of breast is possible (LoE 2bB AGO+) [1].

BCS for patients with recurrence?

The majority of SG-BCC panellists see BCS as an option in patients with intramammary recurrence if it is a low-risk situation (small cancer of the luminal A type) (majority vote: 81%) or if the initial diagnosis occurred at least five years previously (majority vote: 64%). The German experts agree that BCS is justifiable in small cancers and/or luminal A type. Moreover, a time limit cannot be defined. The longer since the initial diagnosis, the less likely is a local recurrence, but rather a second cancer. In everyday clinical practice, however, the distinction is not always possible.

If repeat radiotherapy is not possible in a patient with intramammary recurrence, the question arises as to whether BCS is appropriate. The SG-BCC vote was 50:50, reflecting the complexity of decision-making. If radiotherapy following BCS is not an option, the German experts recommend the discussion with the patient should primarily focus on mastectomy with possible reconstruction (LoE 3bB AGO++) [1].

No axillary intervention?

The majority (83%) of the SG-BCC panellists feel that no axillary intervention is needed in patients over the age of 70 if there are no clinically suspect lymph nodes (cN0). The German experts point out that the decision does not depend solely on the age of the patient, but that comorbidities, the risk of recurrence and possible therapeutic consequences must be considered and discussed with the patient. The decision for or against axillary surgery should only be taken after detailed information. This is also reflected in the recommendations of the AGO Mamma [1]. The German experts also refer to the INSEMA trial [52].

No surgery following NAST?

With a clear majority (86.00%), the SG-BCC panellists reject omitting that surgery in patients with early breast cancer and clinical as well as radiological complete response with NAST. The German experts agree with this vote (majority vote: 84%).

Surgical approach in the axilla

The SG-BCC panellists did not agree on the question of whether to remove more than ten LN in high axillary tumour burden (>5 LN involved). The German experts point out that ALND is defined in terms of its anatomical boundaries and is not based on the number of removed lymph nodes. The goal is to clear the axilla of all tumour manifestation.

The SG-BCC panellists also failed to reach a consensus on the question of surgical approach in the axilla in patients who, after BCS and in an N0 situation (“sentinel node mapping”), currently present with ipsilateral recurrence without LN involvement on imaging. About one third of the SG-BCC panellists recommend SLNE with or without frozen section and 12% favour ALND, while 20% are against axillary surgery. The German experts do not agree with the interventions in the axilla. According to AGO Mamma recommendations, SLNE in cN0 patients after primary SLNE (LoE 2aB AGO−) is not indicated [1].

Oligometastasis

Curative intent in isolated metastasis

In patients with clinical stage T2N1 breast cancer and isolated bone metastasis, the SG-BCC panellists (85%) recommend a curative therapeutic approach with optimal systemic therapy and radiotherapy of the isolated metastasis. The German experts agree with the maximum curative approach whenever there is a chance of cure or long-term survival.
Intensive treatment even in case of 3–5 metastases
If a patient (cT2N1) is diagnosed with multiple (> 3) bone metastases confirmed by fine-needle biopsy and the tumour is no longer clinically detectable in the breast and axilla after 6 months of systemic therapy with excellent clinical response in the bone, the SG-BCC panelists (69%) recommend a palliative concept with regard to local and loco-regional control. The German experts do not agree with the SG-BCC majority vote for a palliative loco-regional concept, but recommend at least considering the continuation of a multimodal approach with curative intent.

Follow-up Care and Quality of Life in Breast Cancer Patients

Intravaginal oestrogens for mucosal dryness
If a patient on adjuvant AI therapy experiences mucosal or vaginal dryness that cannot be adequately relieved with moisturisers or lubricants, SG-BCC panelists (73%) recommend intravaginal oestrogens for symptomatic relief. The German experts agree in principle, but caution that only topical vaginal oestriol containing oestrogens may be used. These are also considered safe for patients with ER+ breast cancer and do not negatively affect treatment success [53]. The issue should be addressed proactively. It is important to maintain compliance with the systemic therapy.

Scalp cooling to prevent alopecia
SG-BCC panelists (69%) recommend that patients receiving chemotherapy associated with a risk of alopecia be routinely offered scalp cooling. The German experts agree in principle [54]. This prevents higher-grade alopecia in 40–50% of patients [55,56]. The patients must be informed that side effects, especially headaches, may occur as a result of the strong cooling of the scalp. However, coverage by health insurance is still limited in Germany at present.

Aiming for physical activity
The SG-BCC panelists particularly recommend physical activity and exercise (44%) as well as acupuncture (20%) and normalisation of body weight (20%) to alleviate disease- and treatment-related secondary symptoms of breast cancer. From the German perspective, all the above proposals are important options and should be pursued according to individual needs and preferences. The best data is on 3–5 hours of physical activity per week (LoE 1bA AGO++). Other measures can be found in the recommendations of the AGO Mamma [1].

Reducing alcohol consumption
Reduced alcohol consumption can help reduce the risk of recurrence in breast cancer patients. The majority of SG-BCC panelists (57%) recommend a maximum of one alcoholic drink per day. The AGO Mamma recommends limiting daily alcohol consumption to a maximum of 6 g/day (LoE 2bB AGO+) [1]. Regardless of this recommendation, it is important to set realistic goals. The general rule is that the less alcohol the better.

Comments
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References


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[34] Alliance for Clinical Trials in Oncology. Hypofractionated Radiation Therapy After Mastectomy in Preventing Recurrence in Patients With Stage IIa-IIIa Breast Cancer. Accessed April 18, 2021 at: https://clinicaltrials.gov/ct2/show/NCT03414970


[45] Kalinsky K, Barlow WE, Meric-Bernstam F et al. First results from a phase III randomized clinical trial of standard adjuvant endocrine therapy (ET) +/- chemotherapy (CT) in patients (pts) with 1–3 positive nodes, hormone receptor-positive (HR+) and HER2-negative (HER2-) breast cancer (BC) with recurrence score (RS) <25: SWOG S1007 (RxPonder). San Antonio Breast Cancer Symposium 2020; Abstract No. GS3-00


