Update Breast Cancer 2018 (Part 2) – Advanced Breast Cancer, Quality of Life and Prevention

Update Mammakarzinom 2018 (Teil 2) – Fortgeschrittenes Mammakarzinom, Lebensqualität und Prävention

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ABSTRACT
The treatment of metastatic breast cancer has become more complicated due to increasing numbers of new therapies which need to be tested. Therapies are now being developed to treat special clinical or molecular subgroups. Even though intrinsic molecular subtypes play a major role, more and more new therapies for subgroups and histological subtypes are being developed, such as the use of PARP inhibitors to treat patients with BRCA mutations (breast and ovarian cancer). Supportive therapies are also evolving, allowing problems such as alopecia or nausea and vomiting to be treated more effectively. Treatment-related side effects have a direct impact on the prognosis of patients with metastatic breast cancer, and supportive therapy can improve compliance. Digital tools could be useful to establish better patient management systems. This overview provides an insight into recent trials and how the findings could affect routine treatment. Current aspects of breast cancer prevention are also presented.

ZUSAMMENFASSUNG

Introduction
Significant progress has been made in recent years in the treatment of metastatic breast cancer. The establishment of new targets and the introduction of new substance classes such as antibody-drug conjugates have significantly improved progression-free survival rates and sometimes even the overall survival of some subgroups. Interest continues to focus on understanding how side effects occur and how they should be treated as well as on maintaining patients' quality of life. As it is becoming possible to describe personal risks more precisely, prevention is also becoming more individualized. The basic approaches in metastatic breast cancer, supportive therapies and prevention presented as part of new, recently published trials and at recent conferences (including the 2017 San Antonio Breast Cancer Symposium) are discussed in more detail below.

Treatment of Metastatic HER2-positive and Triple-negative Breast Cancer (TNBC)

Data is consolidating on PARP inhibitors
New targeted therapies for metastatic TNBC (mTNBC) are urgently needed to improve the prognosis of this patient population which has shown only a limited response to other lines of therapy. Several therapeutic approaches have recently been presented at conferences and in published articles.

Last year, it was reported that PARP inhibitors yielded promising results in the treatment of TNBC. In the OlympiAD trial, the PARP inhibitor olaparib showed a benefit with regard to progression-free survival in metastatic patients with proven germline mutations in the BRCA gene compared to selected chemotherapies (capecitabine, eribulin, vinorelbine) [1]. These results led to the drug being approved for use in the USA [2]. Patients with mTNBC especially benefitted. The EMBRACA trial presented data on the PARP inhibitor talazoparib [3], which was used in an almost identical setting as olaparib in the OlympiAD trial. Here too, progression-free survival (PFS) was significantly extended (8.6 vs. 5.6 months; HR 0.54 [0.41–0.71]; p < 0.0001). The objective rate of response was 63% and therefore more than double the rate for chemotherapies (27%). Another study [4] investigated the effect of higher concentrations of talazoparib [5]. But higher systemic concentrations only resulted in more side effects but did not improve efficacy. It appears that the use of PARP inhibitors for TNBC is headed for success. It still unclear, however, whether a BRCA mutation is a precondition for this therapy.

Other antibody-drug conjugates to treat mTNBC
At the latest after the introduction of T-DM1, antibody-drug conjugates became a hot topic of discussion. Sacituzumab govitacan is an anti-Trop-2-SN-38 antibody-drug conjugate, which was used after second-line treatment in 110 patients with mTNBC until tumor progression or limiting toxicity [6]. The antibody is directed against the epithelial antigen Trop-2 and is conjugated with the active metabolite of irinotecan. The objective response rate in this heavily treated patient cohort was 34% with a median duration of
response of 7.6 months (95% CI: 4.8–11.3). The most important side effects were hematotoxicity (neutropenia grade 3/4: 41%; febrile neutropenia: 8%), fatigue and gastrointestinal symptoms. Sacituzumab govitecan appears to be an interesting drug which merits further research even if the response rates and side effects appear to be comparable to those of other monotherapies. A phase-III trial is already underway in the USA and another phase-III trial is expected to kick-off in Europe in 2018.

Immune checkpoint inhibitors headed for approval

There is a growing body of data on immune checkpoint inhibitors. The pembrolizumab (anti-PD-1) and atezolizumab (anti-PD-L1) antibodies have already been tested in numerous studies on metastatic breast cancer, and the reported response rate was around 20% in patients pretreated for mTNBC [7]. Adams and colleagues recently presented results from the KEYNOTE-086 trial, where the first-line treatment consisted of monotherapy with pembrolizumab (n = 52). The response rate was 23% and the duration of response was 10.4 months. Patients who had partial or complete response (PR, CR) showed a significantly longer duration of response and a longer overall survival [8].

Numerous studies have investigated combinations of immune checkpoint inhibitors and chemotherapy or radiotherapy, and it is hoped that the number of patients who respond to this approach will increase. A phase-I/II study was recently presented which investigated a combination of eribulin mesylate and pembrolizumab in patients with mTNBC [9]. Irrespective of the PD-L1 status, a response rate of 26.4% was reported for the 107 patients treated in the study. If treatment was successful, the duration of response was 8.3 months, with more than half of patients responding for longer than 6 months.

Another study investigated a combination of pembrolizumab and trastuzumab in patients with trastuzumab-resistant advanced HER2-positive breast cancer [10]. Some level of response (15%) was reported for PD-L1-expressing tumors but not for PD-L1-negative tumors. It appears therefore that HER2-pretreated tumors are not very immunogenic.

In summary, a clear trend has been observed with regard to the use of immune checkpoint inhibitors in breast cancer therapy: individual patients appear to respond very well and for an appreciable length of time to immunomodulatory therapy. Cytotoxic therapies might even achieve further improvements in the rate of response. But for the majority of patients, this therapy does not appear to result in any improvement. As neither positive nor negative predictive factors are known which could predict whether patients will respond, it is at present not possible to pre-select patients. Further studies, particularly in the neoadjuvant setting, will determine whether the number and composition of tumor-infiltrating lymphocytes might play a role.

Treatment of Hormone Receptor-positive, HER2-negative Advanced Breast Cancer

Data on CDK4/6 inhibitors in premenopausal patients

In recent years, therapies for patients with metastatic, hormone receptor-positive and HER2-negative breast cancer have significantly improved, although a large percentage of patients still receive chemotherapy [11]. This could be changed by the introduction of therapies which are more effective than anti-hormone monotherapy. After the introduction of everolimus, interest has focused on CDK4/6 inhibitors. More and more studies in other patient cohorts are being published. Results from the MONALEESA-7 trial which investigated the CDK4/6 inhibitor ribociclib in a population of premenopausal patients were recently presented [12]. The trial compared an anti-endocrine therapy of choice (tamoxifen, letrozole or anastrozole) plus ovarian function suppression (OFS) with the same therapy plus the addition of ribociclib. The median progression-free survival was 23.8 months in the ribociclib arm compared with 13 months in the placebo arm (Fig. 1). The overall response rate was significantly higher in patients with a measurable lesion at the start of therapy in the ribociclib arm compared with the placebo arm (51% vs. 36%). Grade 3/4 neutropenia was reported in 61% of patients in the ribociclib arm, although this side effect was not symptomatic in most patients. Patients in the ribociclib arm additionally benefited from a prolongation of the time to deterioration of their quality of life. This combination hormone therapy should therefore be recommended to treat premenopausal high-risk patients.

Studies with CDK4/6 inhibitors are combined to answer further questions

The American regulatory authority (FDA) recently presented an analysis in which all studies which filed for approval of a CDK4/6 inhibitor were pooled for analysis [13]. This included studies on palbociclib, ribociclib and abemaciclib in combination with an aromatase inhibitor as the initial therapy for metastatic or operable breast cancer. The intention-to-treat population included 1992 patients. Progression-free survival was compared for patients aged ≥70 years and patients aged <70 years. The progression-free survival benefit from CDK4/6 inhibitors was similar for both groups of patients. The side effects were only slightly higher in older patients. The conclusion is that higher age is no reason to refuse therapy with CDK4/6 inhibitors to older patients.

Another pooled analysis combined the datasets of the MONARCH-2 and MONARCH-3 trials to look for predictors which could be used to identify patients who would respond particularly well to therapy with abemaciclib and aromatase inhibitors [14]. Out of all the univariate analyses analyzing progression-free survival, only the interval between primary treatment and the start of therapy was found to be a predictor for the efficacy of abemaciclib in the setting of advanced cancer. The number of previous endocrine therapies, previous endocrine therapy for advanced disease, the time from diagnosis to metastasis, and primary metastatic status were not predictors for the efficacy of abemaciclib. Howev-
er, no confidence intervals were presented in this analysis, which limits the interpretation of the data.

**Establishment of new mTOR inhibitors unsuccessful**

The current therapy recommendations and guidelines [15, 16] recommend using three different endocrine therapies in the metastatic setting. One of these therapies includes the mTOR inhibitor everolimus. Vistusertib is a new dual mTOR inhibitor which has been tested in a single study. Vistusertib plus fulvestrant was compared with a therapy consisting of everolimus and fulvestrant [17]. Progression-free survival was longest (12.3 months) following treatment with everolimus and fulvestrant. Both therapy arms with vistusertib were inferior.

**Androgen receptor inhibition attempts to prove its worth**

The androgen receptor inhibitor enzalutamide has already shown some benefits in the treatment of TNBC [18–20]. Data on the efficacy of this therapy to treat hormone receptor-positive breast cancer have now also been presented [21]. Therapy with enzalutamide and exemestane was compared with therapy using exemestane alone in patients with advanced breast cancer with and without previous endocrine therapy. No difference between the therapy arms was reported for the overall patient population. However, when analysis was limited to a subgroup in which gene expression testing had detected an activation of the androgen receptor pathway, a significant benefit was detected for patients who had not had prior treatment (hazard ratio [HR]: 0.44; PFS 16.5 vs. 4.3 months). Enzalutamide was even found to benefit patients with prior treatment (HR: 0.55; PFS: 6 vs. 4.3 months). This therapy merits investigation in further studies.

**Supportive Therapy**

Supportive therapies are playing an increasingly important role in oncology and are being investigated in numerous scientific studies. The often significant side effects of modern multimodal therapies can only be controlled by optimal supportive treatment, and supportive therapy is needed to improve the quality of life and long-term compliance of patients.

**Benefits of acupuncture and physical exercise in therapy-induced arthralgia**

The problem of compliance is still unresolved, with high numbers of patients terminating their adjuvant endocrine treatment because of side effects such as arthralgia [22–25]. Acupuncture is an alternative, non-drug therapy for arthralgia. However, the data is inconsistent. A study was carried out to compare the use of acupuncture with feigned acupuncture (short needles at non-acupuncture sites) and a control group over a period of 12 weeks (first 6 weeks: 2 × per week, subsequently 1 × per week) [26]. A total of 226 patients with breast cancer who reported ≥ 3 points on the pain scale (Brief Pain Inventory, BPI) at the start of treatment with an aromatase inhibitor (AI) or who reported increasing

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**Fig. 1** Progression-free survival (response determined by the examiner) in the MONALEESA trials. Goserelin was administered in both arms. NR: not achieved. CI: confidence interval (based on [12]).
pain during AI therapy were included in the study. Acupuncture was carried out in 11 centers by licensed therapists who had received extensive training. Multiple measurements showed significant improvements in the strongest level of pain after 6 weeks (primary endpoint) compared to the feigned acupuncture group \((p = 0.01)\) and the control group who received no form of acupuncture intervention \((p = 0.01)\). Significant improvements were also reported with regard to average levels of pain and joint stiffness. The benefit persisted for 12 weeks after the acupuncture intervention.

Another study investigated the effect of physical exercise on arthralgia due to AI therapy [27]. A significant improvement in pain scores was reported after 12 months of exercise \((p = 0.067)\) if patients followed a program of 120–150 minutes of walking or running per week and complied with more than 70% of the planned program. This also positively affected patients’ compliance with AI therapy \((p = 0.03)\).

**Diarrhea control depends on the treating physician**

Neratinib, a pan-Her tyrosine kinase inhibitor, is another oral adjuvant therapy for patients with HER-2-positive breast cancer who have already had adjuvant therapy with trastuzumab. However, neratinib requires optimal supportive therapy. The rate of grade III diarrhea reported in the approval study was 39.8%, which led to termination of the therapy by 16.8% of patients. The CONTROL study compared an intensified loperamide regimen \((12 \text{ mg/day, d1–14, followed by 6–8 \text{ mg/day, d15–56}})\) with combinations of loperamide and the steroid budesonide or the cholesterol-lowering drug colestipol [28]. Grade III diarrhea occurred in 30.7, 25.0 and 7.7% of patients, and therapy termination rates were 20.4, 9.4 and 0%, respectively. The addition of budesonide or colestipol appears to reduce the duration and number of diarrhea episodes and could therefore improve compliance with neratinib therapy. The final analysis of the CONTROL study will be carried out when all patients have completed the 12-month neratinib therapy.

**Alopecia from chemotherapy partially avoidable**

Alopecia is an extremely onerous side effect of chemotherapy. Four studies investigated the efficacy of scalp cooling during therapy. The success rate during treatment with anthracyclines was 50–62.5% [29,30]. Menopausal status, therapy setting (neoadjuvant versus adjuvant) and dose density had no impact on alopecia rates. The problem is the high rate of around 30% of patients who discontinue scalp cooling because of the pain, with about half of these patients subsequently losing all of their hair [29]. It should be noted, however, that grade I loss of hair after scalp cooling was also rated as a success in the studies. Grade I is a loss of hair of up to 50%, which can also represent a significant burden for patients.

**Nausea and vomiting rates continue to fall**

Although there are excellent antiemetic drugs and standards, there is still potential for improvement. The use of aprepitant is standard during treatment with anthracyclines, but its use in moderately emetogenic chemotherapy has not been investigated much. A randomized, double-blinded phase-III trial which compared fosaprepitant combined with ondansetron and dexamethasone, d1, with ondansetron, d1–3, and dexamethasone, d1, in 231 patients with breast cancer showed a complete response (no vomiting or need of rescue medication during the trial) in 76.4 vs. 68.6% of patients [31]. Another multicenter study investigated a combination of 5-HT3 and NK1-receptor antagonists in a combined capsule (NEPA) to simplify therapy. The interim analysis presented the results for 2384 patients receiving highly or moderately emetogenic chemotherapy (HEC/MEC) [32]. As regards the primary endpoint, over 90% of the study participants who received HEC or MEC reported that their daily life and quality of life was not affected by vomiting. The rate of complete response (no nausea, no rescue medication) was 74%. Efficacy was rated by both the physicians and the study participants as good to very good in more than 90% of cases.

**Neuropathy continues to be a big problem**

One of the biggest current challenges is chemotherapy-related neuropathy, particularly over the longer term. A study of 238 patients in the adjuvant setting and 442 patients with metastasis showed that 30% of all studied patients were significantly or strongly affected by neuropathies in their daily life [33]. Neuropathies were significantly associated with sleep disorders, pain, impairments of physical function, restricted social life, fatigue, anxiety and depression. 32% had not received any coping strategies from their treating physician. The lack of counseling with regard to supportive therapies was associated with a significantly higher negative impact.

The above-mentioned studies on supportive therapies clearly illustrate the importance of a holistic approach for patients and the importance of targeting potential side effects both prophylactically and therapeutically.

**Bone Oncology**

The preventive effect of bisphosphonates on bone metabolism and bone density and their contribution to reducing the risk of fractures has been known since many years. An Oxford meta-analysis once again showed that the adjuvant use of bisphosphonates to prevent osteoporosis after menopause additionally resulted in a 34% improvement in breast cancer-specific disease-free survival (DFS) and even a 17% improvement in overall survival [34]. This has led to a positive recommendation for adjuvant bisphosphonate therapy in parallel to treatment AI in the AGO and the current DGGG S-3 guidelines [15,16].

**Duration of adjuvant bisphosphonate therapy**

But the question remains how long the respective bisphosphonate therapy should be administered. In this context, the SUCCESS study investigated the impact of 2 years’ \((4 \text{ mg zoledronic acid every 3 months})\) vs. 5 years’ \((4 \text{ mg zoledronic acid every 3 months for 2 years, then every 6 months})\) bisphosphonate therapy on DFS and overall survival (OS) [35]. The results for 2802 patients out of the original 3754 patients were included in the analysis. The two study groups showed no significant differences with regard to their basic characteristics or tumor characteristics. The maximum observation period was 4 years (the median observation period was 2.95 years for DFS and 3 years for OS). DFS and OS were defined based on the STEEP criteria. The results of the
analysis showed no significant differences in terms of DFS (p > 0.827) or OS (p > 0.713) between the 2-year and the 5-year bisphosphonate therapy [35]. Multiple Cox regression analysis adjusted for a number of risk factors also showed no significant differences between groups with regard to DFS, bone recurrence or OS in either premenopausal or postmenopausal women with breast cancer. No differences were found between the two groups with regard to circulating tumor cells (CTC). The rate of side effects was significantly higher after 5 years of treatment compared to 2 years of bisphosphonate therapy. Osteonecrosis of the jaw (ONJ) occurred in 11 and 5 cases, respectively, during 5-year or 2-year bisphosphonate therapy. The limitations of the study listed by the authors included the relatively short observation period, the unbalanced rate of “loss to follow up” and the relatively rare events. Unfortunately this study also lacked a control group which did not receive bisphosphonate therapy [35].

Aromatase inhibitors must be monitored continuously for bone protection

As part of the ABCSG-16 trial, the impact of extending adjuvant endocrine therapy (EAT) with anastrozole by a further 2 vs. 5 years after 4–6 years of primary endocrine therapy was investigated. As regards the primary endpoint of disease-free survival, the report [36] found that after 10 years, DFS for the group which had two additional years aromatase inhibitor therapy was 71.1%, while DFS for the group with an additional 5 years was 70.3% and therefore not significantly different (HR: 0.997; 95% CI: 0.86–1.15; p = 0.982). As regards the side effect “fractures” the group with 5 years of EAT had a higher rate of fractures compared to the group which received 2 years of anastrozole therapy (Fig. 3) [36]. The figures were not adjusted for potential bone protective therapies.

Molecular pathways of breast cancer risk, bone metabolism and breast development are linked

It has been known for some time that the signaling pathways which regulate bone metabolism (RANKL/RANK/OPG) also regulate progesterone-based proliferation in the breast and can have an impact on the pathogenesis and progression of breast cancer [37–39]. It was recently reported that this signaling pathway is particularly important in patients with a BRCA 1 mutation. RANKL appears to be the main mediator in the pathogenesis of breast cancer in this patient population [40–42], meaning that a blockade of this signaling pathway might be a useful strategy for breast cancer prevention. A study with denosumab is planned to investigate this point further [42]. Other substances such as the selective progesterone receptor modulator telapristone may also be of interest, as preclinical models have shown that these substances inhibit the paracrine expression of RANKL [43].

Quality of Life and Digitization of Medicine

Quality of life, therapies with few side effects and survival are among the most important therapy goals for patients [44,45], and patient well-being, therapy, compliance and overall well-being are mutually dependent. Side effects, e.g. due to therapy with aromatase inhibitors, can result in the patient terminating their therapy [22–25], which in turn is likely to result in a poorer prognosis [46]. This means that taking the patient’s quality of life...
into consideration is also important in the context of improving her prognosis.

**Digital medicine to improve communication, quality of life and prognosis**

Digitization in medicine is constantly increasing [47]. There are still huge opportunities to optimize the healthcare for patients by improving therapy selection, patient care and the management of related areas. Much has been written in this context about “big and smart data”. Given the rapid advances in machine learning and the interoperability of databases (which has done much to simplify their use), it is expected that central patient care processes will be automated [47] and electronic tools will increasingly be used to communicate with patients [11, 48]. But it will also be necessary to ensure that interests of the patients (data confidentiality, self-determination and data sovereignty) are maintained [49].

The field has huge potential and presents many opportunities. To take just one example, it has been shown that the use of electronic tools to provide a constant line of communication between the treating medical staff and patients as a means of monitoring side effects can result in better overall survival [50]. The integration of such platforms in various studies across the German-speaking world is increasing. Electronic patient-reported outcome systems have already been tested since several years in studies such as PRAEGNANT [51, 52], Seraphina [53] and Precycle [54].

**Knowledge processing and knowledge exchange for physicians**

As the breadth of knowledge continues to grow by leaps and bounds, it cannot be ruled out that in the long run knowledge will be permanently managed using digital means. The App “Mammakarzinom Transparent” (https://mammakarzinom.onkowissen.de/[55]) is just one attempt to make therapy algorithms available and include additional information on efficacy and side effects. Examples of clickable icons are shown in ▶Figs. 4 to 6; clicking on the icon will call up further information about the respective therapy.

**Quality of life in the metastatic setting**

In Germany, quality of life is an accepted endpoint in addition to overall survival for the assessment of therapies after they have been approved. In recent evaluations, progression-free survival was not accepted for assessments about the additional benefit of a specific therapy. However, it is well known that for the patient disease progression has a negative impact on their quality of life and a delay in disease progression will therefore improve patients’ quality of life [56, 57]. Particularly in the metastatic therapy setting where there are frequent changes of therapy, data about the extent to which therapy sequences influence the further course of disease and quality of life have not been thoroughly investigated. New studies which take a comprehensive look at therapy lines [51, 58] could help to collect new data and improve the care of these patients.
Prevention

Importance of panel genes

In the last few years, the medical understanding of genetic and non-genetic risk factors has significantly improved [59, 60]. In addition to the BRCA 1 and BRCA 2 genes, the clinical importance of so-called panel genes (e.g., the following genes which are associated with a high or moderate risk of breast cancer: CHEK2, PALB2, ATM, RAD51D, BARD1, MSH6 and others) is being investigated [61 – 67]. One study reported that PALB2 (OR: 7.5) might need to be classified as belonging to the group of high-risk genes while ATM, BARD1 and RAD51D could probably be classified as moderate risk genes [62]. No increased risk was reported for the genes BRIP1, RAD51C, MRE11A, RAD50, NBN, MLH1 and PMS2 [62]. It is not yet clear to what extent this information will have an impact on prevention.

▶ Fig. 4  Current therapy algorithms for metastatic breast cancer (schematic) (based on [55]).
After primary adjuvant anthracycline/taxane-based therapy

1. Carboplatin + gemcitabine
   - Doxorubicin (non-pegylated liposomal) + cyclophosphamide
   - Mitoxantrone + cyclophosphamide
   - Paclitaxel (q1w) + bevacizumab
   - Paclitaxel + capecitabine
   - Re-induction anthracycline: epirubicin or pegylated liposomal doxorubicin
   - Re-induction taxane monotherapy: paclitaxel q1w or docetaxel q3w or nab-paclitaxel
   - Vinorelbine
   - Paclitaxel + epirubicin

2. Capecitabine
   - Eribulin
   - Capecitabine
   - Vinorelbine
   - Taxane re-challenge
   - Capecitabine + bevacizumab
   - Mitoxantrone + cyclophosphamide
   - Paclitaxel + bevacizumab

3. Capecitabine + bevacizumab
   - Eribulin
   - Capecitabine
   - Vinorelbine
   - Taxane re-challenge
   - Eribulin
   - Mitoxantrone + cyclophosphamide
   - Paclitaxel + bevacizumab

After primary adjuvant platinum/taxane-based therapy

1. Carboplatin + gemcitabine
2. Capecitabine
3. Capecitabine + bevacizumab

TNBC

Treatment naïve/primary metastasis

After primary adjuvant platinum/taxane-based therapy

1. Carboplatin + gemcitabine
   - Eribulin
   - Capecitabine
   - Vinorelbine
   - Paclitaxel + bevacizumab
   - Paclitaxel + capecitabine
   - Mitoxantrone + cyclophosphamide
   - Re-induction anthracycline: epirubicin or pegylated liposomal doxorubicin
   - Re-induction taxane monotherapy: paclitaxel q1w or docetaxel q3w or nab-paclitaxel
   - Vinorelbine
   - Paclitaxel + epirubicin

Fig. 5 Current therapy algorithms for patients with metastatic triple-negative breast cancer (schematic) (based on [55]).
Fig. 6 Current therapy algorithms for patients with hormone receptor-positive, HER2-negative breast cancer (schematic) (based on [55]).
Which population is suitable for mutation testing?

The approval of the first PARP inhibitor to treat metastatic breast cancer in the USA [2] raised the question whether genotyping should be carried out in all patients who could be considered for this therapy. A recent study presented the mutation frequencies for BRCA 1 and BRCA 2 and other panel genes irrespective of the usual test criteria used in predictive genetic diagnostics [68] (see chapter “Prognostic and Predictive Factors”). The question about the extent to which predictive genetic testing could and should be expanded also includes the debate about whether patients with low mutation rates needed to be informed about any variants of unclear significance (VUS) detected during testing. In the case of genetic changes where not much is known about their clinical relevance, the uncertainty could lead to anxiety, worry and even unjustified measures. At present, more than 3000 of such VUS in the BRCA 1 and BRCA 2 genes are known. Significant progress has also been achieved recently in this area. Studies have shown that functional in vitro analysis of 139 VUS in BRCA 2 was able to identify 13 more pathogenic mutations, while 12 could be classified as harmless variants [69]. More analyses of this type could significantly decrease the diagnostic uncertainty.

Importance of low-penetrance risk genes

In addition to high and moderate risk genes, the importance of common genetic changes with significantly lower risk modifications is becoming increasingly clear. A further 75 variants have just been recently validated [70, 71]. Other variants and their clinical importance are also gradually being characterized [72 – 76]. The data obtained could be summarized in scores and tested in clinical practice [77]. Combinations with clinical risk factors such as mammographic density, size or BMI could also be of clinical use [78 – 82]. However, such studies still need to be carried out.

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Conflict of Interest

A. D. H. received honoraria from AstraZeneca, Genomic Health, Roche, Novartis, Celgene and Pfizer. N. N. received consultancy honoraria from Janssen-Cilag and travel support from Novartis. F. O. received speaker and consultancy honoraria from Amgen, Celgene, TEVA, AstraZeneca, Novartis, Roche, and MSD. F. A. T. received honoraria from Astra Zeneca, Genomic Health and Novartis. H. -C. K. received honoraria from Carl Zeiss meditec, TEVA, Theraclion, Novartis, Amgen, Astra Zeneca, Pfizer, Janssen-Cilag, GSK, LUV Pharma, Roche and Genomic Health. P. H. received honoraria, unrestricted educational grants and research funding from Amgen, AstraZeneca, Eli Lilly, MSD, Novartis, Pfizer and Roche. P. A. F. received honoraria from Roche, Pfizer, Novartis and Celgene. His institution conducts research for Novartis. H. T. received honoraria from Novartis, Roche, Celgene, TEVA, Pfizer and travel support from Roche, Celgene and Pfizer. J. E. received honoraria from Roche, Celgene, Novartis, Pfizer, Pierre Fabre, TEVA and travel support from Celgene, Pfizer, TEVA and Pierre Fabre. M. P. L. has participated on advisory boards for AstraZeneca, MSD, Novartis, Pfizer, Genomic Health and Roche and has received honoraria for lectures from Lilly, Roche, Novartis, Pfizer, Genomic Health, AstraZeneca, medac and Eisai. M. W. received speaker honoraria from Astra Zeneca, Celgene and Novartis. V. M. received speaker honoraria from Amgen, Astra Zeneca, Celgene, Daiichi-Sankyo, Eisai, Pfizer, Pierre-Fabre, Novartis, Roche, Teva, Janssen-Cilag and consultancy honoraria from Genomic Health, Roche, Pierre Fabre, Amgen, Daiichi-Sankyo and Eisai. E. B. received honoraria from Novartis, Riemser and Hexal for consulting and clinical research management activities. A. S. received honoraria from Roche, Celgene, AstraZeneca, Novartis, Pfizer, Zuckerschwerdt Verlag GmbH, Georg Thieme Verlag, Aurinkamed GmbH, MCI Deutschland GmbH, bsh medical communications GmbH and promedics GmbH. W. J. received honoraria and research grants from Novartis, Roche, Pfizer, Lilly, AstraZeneca, Chugai, Sanofi, Daichi, Tesaro. F. S. participated on advisory boards for Novartis, Amgen and Roche and received honoraria for lectures from Roche, Novartis and Pfizer.

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