Intimate Relationships Affected by Breast Cancer: Interventions for Couples

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Introduction

Breast cancer is the most common cancer in women worldwide. Globally, it is also the principal cause of death from cancer among women. 1 in 8 women will develop breast cancer in their lifetime [1]. While only a few decades ago the majority of women did not survive breast cancer, advances in medicine and diagnostics and the higher aging of the population have led to an increased number of persons diagnosed with breast cancer as well as to increased survival rates. In Western countries, 89% of women diagnosed with breast cancer are still alive 5 years after their diagnosis. Therefore, breast cancer survivors are the largest group of cancer survivors [2]. Cancer is increasingly viewed and treated as a chronic disease, due to advances in treatment efficacy and improvements in survival rates. Still, despite the growing number of people treated or followed up for cancer or having survived the disease, a cancer diagnosis remains a life-threatening disease that can affect psychosocial functioning even years after the medical treatment has ended. Cancer patients can experience psychosocial, emotional, and physical distress [3].

There is a growing awareness that a cancer diagnosis and its treatment not only influence the patients but also their families, friends and caregivers [4]. Usually, the primary caregiver of the woman is her spouse. In addition, cancer patients identify their partners as their most important source of support [5]. Thus, due to more women surviving longer and being in a committed relationship at the time of diagnosis, an ongoing challenge is how to cope with the disease, its treatment, and its long-term sequel within the context of their relationships. Breast cancer introduces individual and relationship challenges for both the patient and her partner [6]. Research has shown that some of these issues can serve to bring a couple closer together (cohesion) while others can disrupt relationship functioning in different ways (e.g., by communication problems, less intimacy, and decreased relationship satisfac-
tion) at different stages of the disease-recovery timeline. Therefore, the disease can affect both members of the dyad and can induce serious distress and negatively influence successful coping throughout the cancer journey [6].

**Impact on Individual and Dyadic Functioning**

Both members of the dyad can experience fear, uncertainty, depression, and anxiety. In women with early-stage breast cancer, 7–46% and 32–45% show clinically significant levels of depression and anxiety, respectively [7, 8]. The prevalence rates of depression and anxiety among spouses of cancer patients are in the range of 10–53% and 16–56%, respectively [9]. The partner has to adjust to the woman’s disease, has his own worries, and is confronted with the potential loss of his wife; he is burdened by taking on the partner’s tasks and by providing support and caring. Partners often experience a ‘double role’: On the one side, they may suffer from their own psychological distress, and on the other side, they are the main source of support for the distressed women [10]. The diagnosis and medical treatment of breast cancer also lead to additional problems with regard to body image, sexual functioning, and intimate relationships.

**Body Image and Sexuality**

Women with breast cancer have to cope with significant changes in body appearance (e.g., scarring, hair loss, disfigurement) and functioning (e.g., numbness, pain). Breast cancer and its physical side effects can profoundly affect a woman’s body image and can contribute to lower self-esteem, less desire for sexual activity, lower libido, and depression [11]. A negative body image has strong implications for psychosocial adjustment and social functioning. In women with breast cancer, the iatrogenic menopause can cause significant detriment to sexuality by, e.g., a lowered libido, decreased vaginal lubrication and dyspareunia, as well as numbness in previously sensitive breasts [12]. In addition, psychosocial aspects such as depression or anxiety can contribute to sexual disturbances. Sexual problems are frequent in cancer survivors [2] and can negatively affect the relationship satisfaction [13]. Otherwise, for women with breast cancer, the relationship quality has emerged as a strong predictor of sexual functioning [14]. Therefore, sexual functioning and the enhancement or preservation of relationship satisfaction should be addressed in interventions for couples coping with cancer.

**Dyadic Stress and Coping**

Emotional and instrumental social support from partners appear to be important contributors to the women’s adjustment to breast cancer and thus to their quality of life after the diagnosis. Despite this need for support, a cancer diagnosis nevertheless frequently leads to maladaptive patterns of interaction between the patient and her partner that negatively influence the couple’s relationship functioning. They must cope with distress induced by the physical side effects and a potentially increased functional disability associated with cancer and its treatment, and also with role changes in their relationship. Furthermore, the patient and her partner can experience important restrictions in their social life in addition to interruptions of their daily life. Economic or financial consequences, household and child care responsibilities, and the insecure future are further stresses and strains [15]. A challenge for couples could also be to cope with changes in established communication patterns, responsibilities, or roles [16]. Thus, for some couples, the cancer experience can lead to significant adjustment and relationship difficulties, which may result in feelings of greater conflict and less intimacy. Despite the negative consequences of a cancer diagnosis, a growing amount of literature also reports positive aspects of the cancer experience, such as greater appreciation of life, changed priorities, closer relationships, and enhanced spirituality [17–19].

For a long time, cancer was considered as an individual stress experience demanding individual coping strategies. However, both members of the dyad and the couple’s functioning can be profoundly affected by the cancer experience, with interrelations between the psychological distress states of patients and partners [20]. For most of the women with breast cancer, the men are the main source of instrumental and emotional support during a time when the men themselves could be under extreme stress. Both partners of the dyad interact with and mutually influence each other. There is some evidence emerging that dyadic processes, specifically communication and dyadic coping, may be involved in the adjustment process for both partners [21]. Consequently, it is obvious to consider the individual and dyadic coping of the patient in relation to the individual psychological distress, and vice versa. Therefore, cancer can be a stressor concerning both partners simultaneously, and coping with cancer should be viewed as a dyadic affair [20, 22].

There is growing evidence that dyadic processes as dyadic coping and communication may influence the adjustment to cancer for both partners. Couples may tend to be kind to each other under the stressful influence of the cancer diagnosis, which may result, e.g., in increased positivity in the relationship. However, this may occur within the context of avoidance (e.g., not talking about cancer and disease-related stressors, mutual withdrawal, hiding concerns, denying worries, and not upsetting the other or hostility due to the stressors associated with the disease). Thus, despite what might appear to be an increase in explicit positive interaction between partners, underlying negative behaviors can be deleterious to couple functioning and may jeopardize the long-term adjustment to the cancer. This phenomenon is known as ‘protective buffering’ and shows strong associations with higher levels of patient and partner distress [23]. Therefore, dyadic coping as a stress management strategy supports couples in being mutually involved in the stress coping process by providing and receiving support from each other and engaging in joint problem-solving as well as shared emotion regulation, to act as a team rather than as individuals [21]
Psychosocial interventions should include both partners of the couple and should be aimed at supporting and strengthening the coping abilities of both partners.

Interventions for Couples

Given that coping with cancer should be characterized as a dyadic affair, a growing amount of literature involving couple-based interventions came up over the past 2 decades. Nevertheless, the majority of these studies were published only in the last decade. The most promising results in enhancing the well-being of both partners are shown by interventions with multiple targets (e.g., improvements in communication and sexuality, reduction of emotional distress, strategies to enable partners to express their distress [25], and stress communication (e.g., expressing worries, needs, and feelings regarding cancer- or relationship-related topics), supportive behavior, and effective dyadic coping [15]). Intimacy emerged as a key component of relationship quality and could serve as a buffer for psychological distress [26].

Women with cancer who are in committed relationships express a great need for support from their partners, even more than from others [27]. Although partners show high motivation to offer this support, they may have difficulties in providing social support because of their own distress or because they do not know how to be the type of support person the woman needs. In addition, the needs of the patient change across time, making support even more complicated. Hence, many women feel disappointed by their interactions with their partners when addressing cancer. This dissatisfaction can occur even within the context of an overall high relationship satisfaction. The challenges are likely to persist beyond the completion of treatment, but are most salient during the treatment phase, due to the acute burden of the diagnosis and treatment these couples have to confront.

Components of Couple Interventions

The content of interventions for couples coping with cancer is heterogeneous. A combination of skill training and psycho-educational intervention is recommended [28]. The components can be discussed according to the development-contextual model of couples coping with chronic illness [29] in 3 dimensions: dyadic appraisal, dyadic coping, and dyadic adjustment. Dyadic appraisal was conceptualized at the individual and dyadic levels and contains the appraisal of disease, self-efficacy, and communication [28]. How do both partners perceive and understand their coping abilities and their emotional state? How do they appraise the disease as a unit? In this process, the quality of communication influences the couple’s appraisal of their disease and efficacy [28]. Dyadic coping supports couples in coping with the disease as a team [15]. Enhancing communication skills, self-efficacy, and dyadic coping should influence the dyadic adjustment measured in quality of life and mental, physical, and relationship satisfaction [30]. Dyadic coping may reduce stress but, even more importantly, may increase intimacy, cohesion, and mutual confidence [24].

Thus, given the challenges that breast cancer poses for the couple’s relationship, in addition to the central importance of the couple’s relationship to the patient’s and partner’s adjustment, it is critical to determine the most efficacious way to assist couples in facing a breast cancer diagnosis. There is increasing recognition that the quality of marital interactions, rather than global social support, the mere presence of a partner, or even overall marital satisfaction, is essential to achieve positive patient outcomes. Patient-partner interaction patterns that have been associated with positive patient adaptation include candid communication about cancer-related issues, the ability to express emotions to a partner who is able to listen supportively, and effective problem-solving skills. For instance, in patients with breast cancer, high levels of empathy from spouses were a stronger predictor of patient psychological adjustment than overall marital satisfaction [31]. In addition, the ability to express emotions and to communicate openly with partners about cancer has been associated with fewer emotional and physical complaints and higher levels of self-esteem and perceived control [32], in addition to higher relationship satisfaction [33, 34]. Conversely, negative or unhelpful interaction patterns, such as partner avoidance and criticism, hiding concerns from each other, mutual denying of worries, and avoidance of shared discussion, are associated with poorer patient adjustment, including increased distress, maladaptive coping strategies, and intrusive thoughts about the illness [15] along with lower relationship functioning [21]. A recent review illustrates the importance of stress communication, supportive behavior and effective dyadic coping for relationship satisfaction in couples coping with cancer [15]. In addition, the scope of communication should be enlarged from cancer-related issues to the couple’s life in general [34]. Helping couples to maintain relationship normalcy and quality during the cancer trajectory and to cope as a unit may lead to better communication and enhanced relationship intimacy [26].

An often neglected subject of couple-based interventions is the caregivers’ self-care [28]. Partners often put their own needs on hold, resulting in less time spent on maintaining their own mental, social and physical health [35]. This behavior could affect the partners’ own health.

Effectiveness of Interventions for Couples

The benefit of psychosocial interventions for couples coping with cancer still remains unclear. Some reviews and meta-analyses report that these interventions are effective in improving individual and dyadic functioning and have beneficial effects on relationship quality and psychosocial adjustment, albeit with small to medium effect sizes [19, 36, 37]; others show heterogeneous evidence [38]. An often expressed critique is that the studies have only small sample sizes and are therefore underpowered to examine changes in the multiple outcomes measured, which results in only small to medium effect sizes [39]. For example, in a recent meta-analysis [19] with 20 randomized controlled trials, 9 studies included in the meta-analysis had 35 or less couples per group. One reason for the small sample size could be the challenging recruitment process. Badr and Krebs [19] reported refusal rates ranging from 3 to 82%.
Indeed, only few studies report a detailed description of the reasons for refusal. Distance from the study center, fear of randomization, and the perceived burden of study participation are documented barriers. Timing or scheduling issues and age (i.e., younger couples are more likely to participate than older ones) emerged as key factors [19, 37]. More research is required to identify barriers and obstacles for couples to participate in couple-based interventions (e.g., by approaching couples at routine clinic visits, scheduling study-related appointments with medical appointments, decreasing the number of sessions, expanding the delivery methods by conducting trials by phone, the Internet or in the couples’ homes, enhancing the cooperation with physicians or clinic staff, and increasing the perception that psychosocial interventions are an important part of overall medical care) [19, 40] and to identify the couples’ preferences for interventions considering factors such as disease stage, age, feasibility, and cost effectiveness [19].

Another reason for the heterogeneity of evidence could be that only a subgroup of patients with cancer and partners suffers from increased distress [20]. Given that most interventions were offered to all patients regardless of their distress level, floor effects occur [38]. In addition, there is a strong association between baseline and outcome distress [41]. Badr and Krebs [19] suggest the identification of profiles of couples at risk who may benefit from couple-based interventions. Moreover, screening for relationship and/or psychological distress can be useful. In a recent meta-analysis, only 19 of 198 retrieved studies preselected patients according to their psychological distress, but these interventions reported large effects [38].

Most studies include couples with newly diagnosed patients with early-stage cancer (patients with poor prognosis are underrepresented) [20] and only short-term follow-up. Therefore, the influence of the length or timing of the intervention on the outcome remains unclear, besides the need of booster sessions to maintain the positive impact of the intervention.

Researchers criticize the variation or absence of theoretical models or frameworks, the variation in the intervention approaches, and the diversity of reported outcomes [19, 28]. The majority of interventions were modified from marital therapy interventions developed for healthy people or couples with relationship problems, or from existing cognitive behavioral therapy (CBT) interventions developed for individuals. Thus, it remains unclear whether this approach is appropriate or whether couple-based intervention may be best applied in different ways, depending on the difficulties and issues encountered by the couple [42]. For example, for a dissatisfied couple, the focus of the intervention could be on the relationship and techniques from couple therapy, whereas in couples who are satisfied with their relationship it may be more

<table>
<thead>
<tr>
<th>Component</th>
<th>Description</th>
<th>Goal</th>
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<tr>
<td>Intervention techniques</td>
<td>Communication and problem-solving skills</td>
<td>open communication and expression of worries (e.g., fear of disease progression or recurrence), emotions, and needs regarding cancer-related and relationship-related topics; open communication with children; supportive listening, self-disclosure and empathy; “relationship talk”</td>
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<td></td>
<td>Dyadic coping</td>
<td>both partners are mutually involved in the stress coping process; receiving and providing support from each other with joint problem-solving and shared emotion regulation</td>
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<td></td>
<td>Supportive behavior</td>
<td>thoughtful, respectful and appreciative attitude toward the patient/partner; approving the other’s self-efficacy, resources, and strengths</td>
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<td></td>
<td>Psycho-education</td>
<td>providing information on the diagnosis, medical treatment and its side effects as well as on how to manage the challenges together as a couple; symptom management</td>
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<tr>
<td>Intervention topics that should be addressed</td>
<td>Caregivers’ self-care</td>
<td>consideration of the caregiver’s burden and needs</td>
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<td></td>
<td>Body image/sexual functioning/ intimacy</td>
<td>addressing topics related to changes in body appearance and functioning and its impact on sexual functioning; sensate focus and communication about sexual problems; incorporating the cancer into the relationship to sustain normalcy</td>
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<tr>
<td></td>
<td>Positive aspects</td>
<td>addressing positive aspects of the disease such as greater appreciation of life, changed priorities, closer relationship, and enhanced spirituality</td>
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helpful to focus on the medical and psychosocial challenges of the cancer by providing, e.g., information on the medical treatment and side effects and on how to manage the challenges together.

Other aspects that could influence the efficacy of interventions for couples could be the timing, dosage, and delivery of the intervention and program flexibility. Couples often experience the time immediately after the diagnosis as the most emotionally taxing [43], whereas others emphasize the time when the medical treatment has ended and the transition back to ‘normal’ life is expected, including reprioritizing life goals and management of healthy lifestyle changes [31, 44]. Thus, the content of an intervention should vary depending on the couple’s needs at different time points during the cancer journey [37]. Li and Loke [28] and Regan et al. [37] recommended an appropriate dosage of sessions (e.g., 6 sessions), delivery by trained professionals (higher effect sizes for professionals with mental health background [19]), and greater variation in the contents and mode of delivery of couple-based interventions. Furthermore, more studies are needed to identify gender differences to tailored interventions to the specific wants and needs of women with breast cancer and their partners. Table 1 gives an overview of useful components for couple-based interventions emerging from existing research, which provide flexible modularity.

Nevertheless, the effect sizes for couple-based interventions are comparable to other psycho-oncologic interventions [38] and to those reported in meta-analyses of patient-only and partner-only interventions [37]. In sum, couple-based interventions have promising effects on multiple aspects of psychosocial distress for both the patients and their partners. However, more research is necessary to strengthen future trials methodologically as well as conceptually and to determine ways of integrating couple-based interventions into routine clinical cancer care.

**Disclosure Statement**

The author has no conflicts of interest to declare.

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**References**


