The Landscape of One Breast: Empowering Breast Cancer Survivors through Developing a Transdisciplinary Intervention Framework in a Jiangmen Breast Cancer Hospital in China

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Abstract
Women with Breast Cancer (WBC) in China needs to pay much effort into resisting stigma, they suffer from bodily disfigurement and it will be unbeneﬁcial for their mental and sexual health. However, related studies in this area are rare in China. The research objectives are to understand what kinds of support should be contained in a transdisciplinary intervention framework (TIF) for Chinese WBC. The feminist participatory action research (FPAR) was adopted in this study. Oral history, Peer support group’s materials, and reflection meeting notes as important data. This study revealed that WBC faces diﬃculties returning to the job market and discrimination, and gender stereotypes are commonly found in the whole treatment process. WBC suffered from structural stigma, public stigma, and self-stigma. The findings revealed that forming a critical timeline for intervention is essential, risk factors for coping are side effects, body image, fear of being stigmatized, and lack of personal care during hospitalization. Protective factors for coping with breast cancer are the support of health professionals, spouses, and peers with the same experience and reduction of symptom distress. Benefit finding is crucial for WBC to rebuild their self-respect and identity. Collaboration is essential between 1) Health and medical care, 2) Medical social work, 3) Peer counselor network, and 4) self-help organization to form the TIF for quality care. The research ﬁndings are crucial for China Health Bureau to develop medical social services through a lens that is sensitive to gender, societal, cultural, and practical experiences of WBC survivors and their families.

Keywords: Gender Sensitive, Women With Breast Cancer, Transdisciplinary Intervention Framework, Empowerment, Feminist Participatory Action Research
1. Background to the Study

Like other developing countries, for cancer patients in Mainland China “cancer” is not just a discourse regarded as a medical problem, they will be suffering from financial, social, psychological, and emotional problems as well. WBCs are suffering from psychological stress, self-stigma, and social exclusion. Wang (2014) reported that WBC received less social support and that at least one-third of WBCs in China suffer from higher psychological distress. Social support has also been connected with the quality of life of patients with breast cancer. Higher levels of social support have been associated with better adjustment for WBC (Simpson, Carlson, Beck, & Patten, 2002). The impact of bodily disfigurement on breast cancer survivors’ experiences of change in personal and sexual relationships. Compared to Israeli husbands, Chinese husbands were also more dissatisfied with their sexual relationships and marital adjustment after their wives’ breast cancer experience (Woloski-Wruble & Kadmon, 2002). Another literature review undertaken by O’Cathain et al. (2007) found that one of the barriers for breast cancer screening in Asian women was their “inability to act without husband permission” (Yusuf et al., 2013, p.3695). Therefore we concern about whether a Chinese WBCs had the right or not to participate in making their medical decision. A survey by the Southern Metropolis Daily in 2011 found that China Government has begun to purchase social services to provide medical social work services to patients. This survey published on Nandu.com (2014) shows that 68% of respondents (patients) are unaware of the existence of medical social workers in their area. Thus, social support from medical social services for WBC is insufficient and limited. According to a Consultant Medical Officer of breast cancer ward in China, lack of a gender-centered perspective is still evident at every treatment stage in Mainland China, there is an urge to educate the breast cancer doctors on how to better release the confirmation of bad news for the WBC and their families with gender sensitivity. Worse still, breast cancer doctors tend not to maximize the rights of breast cancer patients in choosing whether to have breast reconstruction or a mastectomy. "Safe enough space” will be given space to share decision-making and develop self-disclosure among the WBC so as not to silenced feminist standpoints. Although, there is abundant research evidence in Western countries. Indeed, breast cancer survivors have reported that if they are well-informed about the disease, related treatment and prognosis are beneficial for them to develop adequate coping strategies at different stages of the disease, decrements to quality of life are highly associated with their cancer experience, level of anxiety, depression, feelings of social isolation and fear of cancer recurrence (Stanton et al., 2002; Stanton & Snider, 1993). However, there is no formal helping profession, such as medical social workers and clinical psychologists for breast cancer patients or well-organized patient self-help organizations in Mainland China. Therefore, there is an urge for more researchers in this area to advocate more appropriate and prompt medical care and social services for breast cancer patients and their families.

2. Research Objectives

Based on the above discussion, through this research for policymakers in a Breast Cancer Hospital in China to modify their health care support services for WBC, therefore the objectives of this research are: 1. What are WBC’s treatment experiences, resistance to problems, meanings of their symptoms? 2. What kinds of services or support should be contained in a transdisciplinary intervention framework for Chinese WBC through a lens that is sensitive to gender, societal, cultural, and practical experiences of WBC for policymakers in a Breast Cancer Hospital in China to modify their health care support service for WBC?
3. Research design and methods

Health care researchers of Mainland China are still dominated by logical positivism (Yu & Liu, 2008) and its successors of medical hegemonism. This study was highly influenced by the concept of “an ethics of Care Epistemology” (Kim, 2006). The knowledge construction should include the voice of the patients and the minority group. How does the medical system disregard the treatment process, construct the powerless, oppress or cause inferiority of the woman through socialization or distribution of power and privilege? In view of this, the feminist perspective is considered the most appropriate lens to understand the oppressed voices of WBC in medical settings in Mainland China. Therefore, this study was adopted the Feminist Participatory Action Research. This FPAR generally includes a spiral process of (a) planning for change, (b) acting and observing, (c) reflecting on the processes and consequences, and then (d) beginning the process again through re-planning (Mertler, 2017). As in many action research projects, these stages are not always linear and often overlap.

4. Sampling and Data Collection

The scope of this study is located in Jiangmen, which is a prefecture-level city in Guangdong Province in southern China. It is a third-tier City now part of the Guangzhou-Shenzhen conurbation. The research project will cooperate with the Jiangmen Maternity and Child Health Care Hospital (JMCHCH). Six core action team members were recruited, 14 Participants and a carer (eligible women aged 18 or above) were outpatients with distress (8 participants who would like to join self-help groups or peer counselor networks) receiving chemotherapy, radiation or medication at JMCHCH. They were identified and invited through purposive sampling and referred by the hospital. As PAR is committed to anti-oppression, empowerment and takes the role of change. The results of action research are intended to directly benefit the situation of ordinary people and the collaborator, research findings are supposed to encourage interaction and self-reflection. Thus, methods requiring a high level of training or high cost of resources are not often employed. According to Tuhiwai Smitch (2001), the results of our studies (anti-oppressive research) should be “disseminated back to the people in culturally appropriate ways and in a language, that can be understood” (Brown & Strega, 2016, p.141). Thus, this is one of the reasons to adopt qualitative research methods in this project because it is more user-friendly, so as not to create oppression and feelings of powerlessness both in the reflective and investigative processes. For collecting the data, in action Stage I using oral history and participatory observation, and group materials such as expressive art products, drawings, themes songs, letter, poetry, handicraft as data in Stage II, storytelling and public speech content will be the data in action Stages III and IV. Reflection meetings notes in all action stages as a very important data.

5. Finding and Discussion

5.1. Experiences of WBC at Every Treatment Stage

We began our journey of oral history in early 2018, following discussion with the action team members, it was classified into the treatment stages could be divided into four stages. For stage 1: Stage of suspected breast cancer (SS), in the interview, the WBCs felt very stressed and helpless in this stage but both formal and informal network are rare. It is vital if he health professionals’ positive attitudes having a great influence on the willingness of the WBC to receive cancer treatment. WBCs and their families were well-informed about their illness, its treatment and prognosis, which could reduce refusal of cancer treatment. they had to live with
their difficulties and tried to comfort themselves. For stage 2: Stage of diagnosis (SD), the biggest impact for breast cancer patients and their families is harmful to their lives, followed by the impact on livelihoods and their body image. Although compared with the other two factors, we found a negative impact on body image to be comparatively low. Nevertheless, body image is one of the biggest barriers to receive the breast cancer removal surgery of WBCs. At stage 3: Stage of treatment and prognosis (ST), the Risk factors for coping of the cancer is suffering from distressing and traumatizing treatment side effects, fear of being stigmatized/discriminated about the state of their health and the changes to their bodily appearance as well as lacking family personal care during hospitalization. For the Protective factors for coping, the caring responsibility of elderly parents and children would increase the resilience and optimistic thinking of WBC. Social support of spouse and peer support from those with the same cancer experience was greatly helpful. Support of health professionals, we found attitudes and communication skills crucial for supporting WBC to continue with cancer treatment. Due to WBC facing such critical moments, in this study, it revealed that they lacked coping strategies to deal with breast cancer treatment, so that the protective factors for coping are enhancing coping strategies as well as releasing symptom distress. Stage 4: Stage of rehabilitation and integration (SRI), In this research, among the interviewees, were facing difficulties returning to the job market and social inclusion, they concerned about their livelihoods in the future and fears of finding employment, they worried about not finding a job and worry about their livelihoods was more important than the illness itself.

5.2. Relationships

On Changing in their Relationships with Others and community after Surgery, the current study supports previous literature in that these women experience changes in the whole of their life (Yusuf et al., 2013); from caregiver to care receiver; changes in their social role; changes in body and personal image, changes in personal relationships; either positive or negative. In this research revealed that breast cancer in Mainland China is perceived as a shameful disease. The dominant pathologizing and cultural discourses in Mainland China constructs them as “Bringing bad luck and disgrace to the family and village’ and thus they are treated with negative stigmatizing attitudes. For WBC who choose not to speak to their children, they worried about them and being a burden to the family. Some parents choose to deceive their children until their death, only revealing the fact that their parents died of cancer at this time; making it difficult for children to accept that their parents died of cancer. Many children were not allowed or informed to see the “last face” of their parents, the right to grief should be advocated. In this current study we found that Chinese families especially husbands, brothers, sons and fathers play a crucial role in medical decision making.

5.3. Body Image

On Changing in body image, according to Schilder (1950), body image includes psychological, physical and social dimensions (Yang, 2003). We will examine it by using scars (body), self-concept (psychology), intimate sexual (social) contacts with WBC and consider how society understands how breast-conserving surgery may lead to barriers in social interaction and by understanding the changes in the psychological and physical status of women with cancer and their experience of gender issues.
5.3.1 Physical Level - Mastectomy Scar. From the interviews, we found that seeing wounds is a very difficult moment for WBC. Some women are even more afraid of family members and even their husbands seeing or touching their wounds:
“I dared not look at the wound even when I was in the bathroom alone. It was quite disgusting indeed as part of the breast was removed....there was a bunch of things.” (Ah Xiu)

5.3.2 Psychological Level - Self-Identity. WBC commonly avoid looking in the mirror and experience common sentiments including decreased self-worth and attractiveness and feeling deformed. WBC who had a mastectomy commonly has doubts about whether they are “real women”:
“Without the breasts, a woman seemed as unlike a woman. ... When going to swim or hot springs in the future, it would be so ugly/odd if there is no breast on one side.” (Ah Xiu)

5.3.3 Social Level - Intimate Relationships. Decreased or lack of sexual desire after cancer treatment was a common problem for WBC couples:
“I did not think about having sex. Of course, he would have the thought (having sex) after abstinence for quite a while.” (Ah Xiu)
From the stories, it reflected that WBCs’ changes to their sexual identity was noticeable in the Chinese society. Doctor did not aware the importunateness to explain to breast cancer patients their rights to breast-conservation and non-breast conservation, as well as the related physically consequences of this operation, this is obviously further exploitation of the rights of women with breast cancer.

5.4 Beneficial Findings of Breast Cancer Survivors through the Peer Support Network

Tedeschi and Calhoun’s (1995) study illustrated that benefit finding may relate to improved quality of life and that can help them overcome personal and environmental challenges. As mentioned in the literature, WBC also benefitted from their fight with cancer and its empowering process, which increased their “individual sense of control, as well as enhancing positive coping and well-being” (Stang & Mittelmark, 2009). However, the benefit finding of WBC is seldom explored in Chinese research. In current study, through the empowering peer-support group which is an effective way to facilitate WBC to be transformed as peer counselors, WBC could gain their power through finding positive changes in the whole treatment process. Ah Xia’s changes have summarized in Table 1:

<table>
<thead>
<tr>
<th>Named the picture: Slacken herself</th>
<th>1. Improvements in social relations:</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Improvements in social relations:</td>
<td>-Accompanied by a few good friends (with same cancer experience) to go out for a walk, maybe on the grass and sit down to take rest and chat.</td>
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<tr>
<td>2. A greater appreciation for life:</td>
<td>-I think that I'm different, I'm more optimistic.</td>
</tr>
<tr>
<td>3. A changed sense of priorities:</td>
<td>-I feel less burdened to view things and work now. It’s not that serious to see things and work. I have been gradually slowing down my pace in these two years.</td>
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<tr>
<td>4. Self-love:</td>
<td>-I feel it’s a new me, I’ll treat myself good and to love myself more</td>
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<td>5. Beneficial spiritual changes:</td>
<td></td>
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Table 1: Beneficial Finding of the Peer Counselors
In this FPAR, we found that all stages of treatment, such as pre-treatment, diagnosis and treatment (surgery, chemotherapy) lack gender-centeredness in every treatment stage in hospitals in China. Most women with breast cancer reported that problems are not only caused by physiological and psychological factors but also suffered from structural stigma (Chinese Culture), public stigma (Relationships) and self-stigma (body image). Structural stigma is “societal-level conditions, cultural norms, and institutional practices” (Hatzenbuehler & Link, 2014, p. 2). In current study, WBC were full of problems in their life narratives. Breast cancer in Mainland China is perceived as a shameful disease. The dominant pathologizing and cultural discourses in Mainland China give them a mark of “Bringing bad luck and disgrace’, which leads to bad treatment through negative attitudes and stigmatization. Public stigma refers to a set of negative attitudes and beliefs that motivate individuals to fear, reject, avoid, and discriminate against people (Parcesepe & Cabassa, 2013), in current study revealed that WBC face difficulties in returning to the job market, this illustrated that discrimination from their communities and social stigma are barriers to making it possible for them to better integrate into their communities. Self-stigma is sourced from the body image of WBC, WBCs are affected by the dominant discourse, that is, how beauty of the female body is affected by the male vision. The research team reflected on the meaning of body image. When a woman lost a breast, it also affected the interaction between herself as a breast cancer patient and society, or it became a way of domesticating/assimilating the body through discriminatory eyes or gossip of others; merely perceived as the one that lost a breast.

The action team has decided to draft-up a preliminary intervention framework to advocate empowering medical social services for WBS and their families. The action team has proposed to conduct study tour in Hong Kong to learn supportive services for breast cancer patients in Hong Kong and study the intervention framework in the Breast Cancer Ward. We hope to find out the questions are: If we want to set up a Medical Social Service Unit (MSSU) in the hospital, what kinds of support would be helpful for WBC to better reintegrate them into the community, ease their embarrassment and self-stigmatize identity? What kinds of training/services/resources should be offered for helping professionals and peer counselors? What resources could fund this sector? How to obtain managerial support and proceed with reform of the whole treatment process.

5.5 Construction of the Transdispliary Intervention Framework(TIF) Using the Participatory Approach

Under collaboration with peer counselors, breast cancer doctor, psychologist, front-line social worker and social work teacher, researcher and the consultant by using the participatory approach, reflection in action. Through this participatory process, every member has raised their ideas on service design, service content and service-workflow, they also highlighted the urgency and feasibility to form the peer support system in Jiangmen. We had to prioritize the core interventions at every treatment stage. The TIF workflow was subsequently constructed and the crucial findings of the Role & functions of different parties are summarized in Table 2.
Table 2: Role & Functions of Different Parties

<table>
<thead>
<tr>
<th>Role &amp; functions of the Health and medical care domains (doctors and nurses)</th>
<th>SS</th>
<th>Gatekeeper</th>
<th>1. Preliminary medical consultation, 2. Make referral to the Medical Social Service Unit (MSSU) based on their willingness.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SD</td>
<td>Educator</td>
<td>Well-informed and explain the treatment plan, side effects and opportunities for cure for WBCs and family members.</td>
</tr>
<tr>
<td></td>
<td>ST</td>
<td>Therapist</td>
<td>1. Standardized caring service, 2. On-going explanation of illness and its treatment plan, 3. Refer WBCs in need to MSSU.</td>
</tr>
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<thead>
<tr>
<th>Roles &amp; functions of the Medical social work domains - social worker and clinical/counseling psychologist</th>
<th>SS</th>
<th>Broker</th>
<th>1. Emotional supported, 2. Refer to welfare unit and/or Family and Patient Resource Centers (FPRC).</th>
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<tbody>
<tr>
<td></td>
<td>SD</td>
<td>Accessor</td>
<td>Assess the resources of the WBC</td>
</tr>
<tr>
<td></td>
<td>ST</td>
<td>Case manager, Broker and Therapist</td>
<td>1. Provide case management (Encourage patients to continue treatment, provide the nonhospital-based home-help service for the WBCs in need), 2. Provide support to peer counselors, 3. Sustain the WBC’s social support (e.g., home visit) for WBCs, 4. Carer support, 5. Refer the rehabilitation resources, 6. Therapeutic intervention (e.g. Sexual and hospice care, etc.)</td>
</tr>
<tr>
<td></td>
<td>SRI</td>
<td>Enabler, Advocator, Advisor, organizer and trainer</td>
<td>1. Nurture the peer support network and the FPRC, 2. Provide carer support, 3. Facilitate the community integration of the WBCs, 4. Regularly organize community education to public, e.g. Anti-discrimination education, Employment opportunities for WBCs, Maximize the rights of WBCs, Provide illness information, Side effect’s management, Symptoms management and its prevention and recurrence.</td>
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<tr>
<th>Roles &amp; functions of the Peer counselor network - peer counselor</th>
<th>SS</th>
<th>Broker</th>
<th>1. Encourage the WBCs to receive treatment, 2. Provide medical resources</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>SD</td>
<td>Accompanying person</td>
<td>1. Pair-up peer counselor support network, 2. Share treatment experience in fighting cancer, 3. Provide emotional support and induce hope to the WBCs.</td>
</tr>
<tr>
<td></td>
<td>ST</td>
<td>Educator and supporter</td>
<td>1. Share health care or healthy diet’s information, 2. Share the road of fighting breast cancer, 3. Encourage the WBCs to maintain social support.</td>
</tr>
<tr>
<td></td>
<td>SRI</td>
<td>Accompanying person</td>
<td>Support and encourage breast cancer survivors to become peer counselors to help new breast cancer patients in fighting cancer.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Roles &amp; functions of the self-help organization</th>
<th>SS</th>
<th>Broker</th>
<th>Linking the care and health resources</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SD</td>
<td>Accompanying person</td>
<td>1. On-going emotional support, 2. Report the situation of the WBCs to MSSU if any crisis</td>
</tr>
<tr>
<td></td>
<td>ST</td>
<td>Educator</td>
<td>Provide stress management activies to family members, carer</td>
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During the research process, our team members had different views and understandings on what transdisciplinary, multidisciplinary as well as interdisciplinary approaches were. This is why we adopted the TIF because it is neither a multidisciplinary nor interdisciplinary nor solely an intervention framework. We had several considerations here, we found that in the Hong Kong model of breast cancer support services all professionals share the roles and functions, through the Hong Kong study tour, the multidisciplinary team able to understand the knowledge and skills of another helping professionals. The second consideration was the needs of WBC in every treatment stage, which involved different professionals, family systems and health care systems, such as the family support system, peer support system, the Clinical Psychology Service system, Medical and Nursing Care system, the Medical Social Service system. All systems had a great influence on breast cancer patients and their families to overcome the side effects of treatment, psychological hardship and/or social problems. These systems are crucial for WBC to better re-integrated into their communities during the rehabilitation phase. These considerations directed our action team to reflect that the “Transdisciplinary” approach should be promoted in medical social service settings in Mainland China.

6. Conclusion

In the current research, we found that all stages of treatment, such as pre-treatment, diagnosis, and treatment (surgery, chemotherapy) as well as rehabilitation lack gender-centeredness in every treatment stage in hospitals in China. Throughout the whole journey of the FPAR, I had a question, am I to “rethink” what is “gender bias” in the Chinese context? Gender insensitivity? Over-generalization? Double standards and androcentrism? (Jahn et al. 2017). What is the meaning of these professional jargon for WBCs? Caring is essential to the experience of women. The concepts of care and responsibility are primary to Chinese women’s construction of their moral domain (Gilligan, 1982) as demonstrated by the notion that women are the primary caregivers in the home, caring for children and other family members, such as the elderly (Walker & Thompson, 1983). It was inspiring that this current study revealed caring duties as one of the protective factors for WBC to continue treatment and fight cancer. When “caregivers” were very sick, who can replace their role in the family? These stories of WBCs revealed that in Chinese culture they do not allow their children to get involved in their treatment process and caring duties even when WBC faces death in the terminal stage of cancer. Do the children of WBC and their relatives have the right to grief (bereavement counseling for WBC’s children) and how can they get the children of WBC involved; this could be one arguable issue for providing counseling for a person who is dying because it is important to be able to say goodbye to their beloved which is crucial to health grief. Grief is a normal and natural response to loss and we often expect to grieve the death of a family member. These WBS’s stories revealed that the dominant cultural conception of breast cancer in Mainland China is “shameful”, such discrimination is unspoken, the suffering is wounded not just in the body, the breast scar, but in the voice. Why they need to
treat disrespectful, why they need to bear the discrimination, for a WBC if, without structural changes, they may suffer in a triple way, the illness itself, the discrimination caused by the cancer stigma, and no chance to be given to say goodbye to the beloved that is the triple suffering and as triple oppressions.

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