



Lymphatic pain in breast cancer survivors: An overview of the current evidence and recommendations



Jeanna Mary Qiu^a, Mei Rosemary Fu^{b,*}, Catherine S. Finlayson^c, Charles P. Tilley^d,
Rubén Martín Payo^{e,f}, Stephanie Korth^g, Howard L. Kremer^g, Cynthia L. Russell Lippincott^b

^a Harvard Medical School, Boston, MA, 02115, USA

^b School of Nursing and Health Studies, University of Missouri-Kansas City, Kansas City, MO, 64108, USA

^c Lienhard School of Nursing, College of Health Professions, Pace University, Pleasantville, NY, 10570, USA

^d School of Nursing-Camden, Rutgers University, Camden, NJ, 08102, USA

^e Faculty of Medicine & Health Sciences, University of Oviedo, Cristo Campus, 33006, Oviedo, Spain

^f Principality of Asturias Health Research Institute (ISPA), 33011, Oviedo, Spain

^g University Health-UMKC Health Sciences District, Kansas City, MO, 64108, USA

ARTICLE INFO

Keywords:

Lymphatic
Pain
Assessment
Behavioral
Breast cancer
Fluid accumulation

ABSTRACT

Among the 7.8 million women with breast cancer worldwide, at least 33%–44% of them are affected by lymphatic pain. Lymphatic pain refers to co-occurring pain (e.g., pain, aching or soreness) and swelling. Pharmacological approaches, such as the uses of NSAIDs, opioids, antiepileptics, ketamine and lidocaine, have very limited effects on lymphatic pain. Limited research in this field has made it difficult for patients and clinicians to differentiate lymphatic pain from other types of pain. Precision assessment to distinguish different types of pain is essential for finding efficacious cure for pain. Innovative behavioral interventions to promote lymph flow and reduce inflammation are promising to reduce lymphatic pain. The goal of this review is to provide a comprehensive understanding of lymphatic pain through research evidence-based knowledge and insights into precision assessment and therapeutic behavioral intervention for lymphatic pain.

1. Introduction

Lymphatic pain refers to co-occurring pain, or sensations of aching, soreness, or tenderness, and swelling.^{1,2} For breast cancer survivors, lymphatic pain occurs usually in the ipsilateral body or upper limb.^{1,2} While breast cancer mortality has been significantly decreased over years,^{3,4} lymphatic pain remains one of the most common and long-term and debilitating effects of cancer treatment.^{5,6} Currently, at least 33%–44% of the more than 7.8 million women treated for breast cancer worldwide are affected by lymphatic pain.^{1,3,5}

The concept of lymphatic pain has emerged in recent research.^{1,2,5,6} Historically, the concept of cancer-related pain has been used to study chronic pain associated with cancer or cancer treatment.^{7–9} Cancer-related pain refers to persistent pain that continues more than three months after active cancer treatment.^{7–10} Researchers have operationalized cancer-related pain in terms of occurrence and severity of

general bodily pain in any body location.^{7–10} This line of research has increased our understanding of cancer-related chronic pain, yet, it has not been able to distinguish different types of pain after cancer treatment, such as lymphatic pain due to fluid accumulation and inflammation, general bodily pain, postmastectomy pain, chemotherapy-induced peripheral neuropathy, or arthralgias related to hormonal treatments.^{7–10} Consequently, opportunities are limited to explore the underlying physiological and psychosocial mechanisms of different types of pain and develop efficacious pain treatments.

Lymphatic pain is caused by abnormal lymph fluid accumulation.^{1,2} Mainstream pharmacological approaches (e.g., NSAIDs, opioids, anti-epileptics, ketamine and lidocaine) have very limited effects on lymphatic pain.^{7,11,12} Recent research demonstrates that some behavioral interventions that promote lymph flow and reduce inflammation are efficacious for lymphatic pain.² However, limited research has made it difficult for patients and clinicians to differentiate lymphatic pain from

* Corresponding author.

E-mail addresses: jeanna.qiu@hms.harvard.edu (J.M. Qiu), mei.fu@umkc.edu (M.R. Fu), cfinlayson@pace.edu (C.S. Finlayson), ct770@camden.rutgers.edu (C.P. Tilley), martinruben@uniovi.es (R. Martín Payo), Stephanie.Korth@UHKC.org (S. Korth), howard.kremer@uhkc.org (H.L. Kremer), russellc@umkc.edu (C.L. Russell Lippincott).

<https://doi.org/10.1016/j.wcn.2024.04.001>

Received 19 December 2023; Received in revised form 13 April 2024; Accepted 22 April 2024

Available online 1 July 2024

2949-7515/© 2024 West China Second Hospital of Sichuan University and China Science Publishing & Media Ltd. Publishing services by Elsevier B.V. on behalf of KeAi Communications Co. Ltd. This is an open access article under the CC BY-NC-ND license (<http://creativecommons.org/licenses/by-nc-nd/4.0/>).

other types of pain. The goal of this review is to provide a comprehensive understanding of lymphatic pain through research evidence-based knowledge and insights into precision assessment and therapeutic behavioral interventions for lymphatic pain.

2. Impact of lymphatic pain

Lymphatic pain negatively impacts breast cancer survivors' physical function, emotional health, and overall health. Lymphatic pain significantly interferes individuals' activities of daily living (ADLs).¹ ADLs are essential daily activities for individuals to live an independent life and are important measures for daily living function.^{13–15} Impairment in ADLs occurs when individuals are not able to perform the essential daily living activities, resulting in poor quality of life (QOL) and lack of independent living. A recently study of 568 breast cancer survivors detailed that patients with lymphatic pain reported impairments in 45% of ADLs and had a significantly increased risk of having difficulty in performing all the 13 ADLs (i.e., cooking, using a knife, writing/typing, cleaning, vacuuming, laundry, carrying objects, yard work, dressing self, bathing self, driving, making bed, and taking care of children) when compared to patients with only pain but no swelling.¹ Patients with lymphatic pain were 4.74 times more likely to have impaired ADLs (OR = 4.74, 95% CI = [2.65–8.50], $P < 0.001$) compared to patients with only pain but no swelling.¹⁶ Noticeably, when patients with lymphatic pain reported co-occurring fatigue, their risk of having impaired ADLs increased 24.43 times (OR = 24.43, 95% CI = [5.44–109.67], $P < 0.001$).¹⁶ This undergirds the importance of assessing the incremental impact of co-occurring lymphatic pain and other symptoms.

Lymphatic pain also increases emotional distress among breast cancer survivors.^{1,16} Emotional distress are negative emotions evoked by an individual's experience of physical symptoms, such as pain.^{17–20} These negative emotions include frustration, sadness, guilt/self-blame, being worried, irritation, fear, anger, loneliness, helplessness, hopelessness, anxiety, and depression.^{17–20} A recent study of 354 breast cancer survivors found that the odds of having emotional distress were 12.82 times higher in patients with lymphatic pain (OR = 12.82, 95%CI = [6.72–24.46], $P < 0.001$). The risk of having emotional distress for patients with lymphatic pain increased tremendously if fatigue also co-occurred (OR = 26.52, 95%CI = [9.64–72.90], $P < 0.001$).¹⁶ While emotional distress is influenced by several factors related to cancer treatment, such as having a mastectomy and a lumpectomy, recent research demonstrates that lymphatic pain is a very important and significant predictor for emotional distress in breast cancer survivors.^{1,16} Further research is needed to detail how lymphatic pain impacts breast cancer survivors' physical functioning, emotional well-being, and overall health.

3. Etiology of lymphatic pain

Lymphatic pain and lymphedema share similar etiologies of abnormal fluid accumulation.^{1,6,21–25} Breast cancer treatment interrupts normal functions of lymphatic system, which causes an accumulation of lymph fluid and creates pain, or aching, or soreness along with swelling.^{1,6,21,22} Lymphatic pain can occur in patients either with or without a diagnosis of lymphedema.^{1,2,23,24} Lymphedema following breast cancer treatment is a chronic and incurable condition.^{11,22} The hallmark of lymphedema is swelling which is often defined and quantitatively operationalized as an increase in limb size or girth or lymph fluid level.^{11,23,24} Lymphatic pain often indicates an early stage of lymphedema for breast cancer survivors without a diagnosis of lymphedema.^{1–6,23–27} For breast cancer survivors with lymphedema, lymphatic pain is part of “living with a perpetual discomfort” and the exacerbation of lymphatic pain indicates the worsening of lymphedema.²⁸ Future research is needed to further explicate the etiology and underlying mechanism of lymphatic pain.

4. Risk factors for lymphatic pain

The major risk factors for lymphatic pain are associated with cancer treatment, such as surgical removal of lymph nodes which interrupt the normal function of lymphatic system and radiation exposure which is associated with trauma to the lymphatic system.¹ Other non-cancer treatment related risk factors include lymphedema diagnosis, financial hardship, obesity, and younger age.¹ The odds of having lymphatic pain were 9.68 times higher in breast cancer survivors with a lymphedema diagnosis (OR = 9.68, $P < 0.001$, 95% CI = [5.78–16.63]).¹ Limited research has explored the genetic or genomic influence on lymphatic pain. Research on heterogeneity of lymphedema phenotype²¹ found that the odds of having pain were 4.70 times higher in patients with the genotype VEGF-C rs3775203 heterozygous A/C compared to those with homozygous C/C genotype. Similarly, the odds of having pain were 6.29 times higher in patients with genotype IL13 rs1800952 homozygous T/T and 2.04 times higher in patients with heterozygous T/C had 2.04 compared to the homozygous C/C genotype. Notably, patients whose genotype contained both VEGF-C rs3775203 and IL13 rs1800952 variants had 12.86 times higher odds of experiencing pain. More research is needed to elucidate the genetic and genomic impact on lymphatic pain.

It is noteworthy that a recent study of 568 breast cancer survivors was the first to identify financial hardship (i.e., not having enough income to make ends meet) as one of the major risk factors for lymphatic pain.¹ Patients with financial hardship were 4.64 times more likely to have lymphatic pain (OR = 4.64, $P = 0.001$, 95% CI = [1.99–11.32]). This provides evidence that social determinants of health (e.g., financial status) have negative impact on lymphatic pain, suggesting that assessment of patient's financial status is important in identifying a patient's risk of lymphatic pain. In the United States, racially minoritized women have delays in breast cancer treatment as well as inadequate treatment.^{29,30} Later stage diagnosis and delayed treatment lead to higher risk of pain and lymphedema due to the need to have more aggressive surgical treatment, more lymph nodes removed, and radiation.^{22,29,30}

Obesity increases the risk of lymphatic pain.^{1,5,11} Patients with a body mass index (BMI) ≥ 30 kg/m² were 3.49 times more likely to have lymphatic pain (OR = 3.49, 95%CI = [1.87–6.50]; $P < 0.001$).⁵ Obesity and lymphatic pain are inflammatory conditions, and more research studies are needed to explore the role of inflammatory pathways that contribute to lymphatic pain. In the United States, obesity is more likely to occur among women living in rural and economically disadvantaged communities.^{31,32} Thus, assessing place-based disadvantages for the risk of lymphatic pain is essential.

An early study³³ found that younger age was a risk factor for lymphedema. Similarly, younger age is also a significant risk for lymphatic pain (OR = 0.97, $P = 0.011$, 95% CI = [0.96–0.99]).¹ Compared to older women, women of younger age may have jobs outside the home, share more parenting responsibilities, care for elderly parents, and do more household chores; all of which may contribute to stress and chronic inflammation that could lead to increased risk of lymphatic pain and lymphedema. More research is needed to explore stress and actual physical labor on the lymphatic system and fluid accumulation.

5. Precision assessment of lymphatic pain

As a cluster of co-occurring symptoms of pain, or sensations of aching, soreness, or tenderness and swelling, lymphatic pain is a subjectively perceived indicator that reflects abnormal biological or physiological changes that may or may not be observed objectively.³⁴ The subjective nature of lymphatic pain entails its assessment using patient-reported outcome measures (PROMs). PROMs are patient's direct reports about his/her health condition without clinician or other people's interpretation of the patient's response.^{35,36} As a component of lymphatic pain, swelling may be measured by an objective measure of limb volume or circumference differences or fluid level. However, these objective measurements of swelling are less associated with QOL than patient-reported

symptoms (e.g., pain, aching, soreness, and swelling).¹¹ Therefore, PROMs are optimal measures for lymphatic pain, a subjective phenomenon.^{35,36}

Lymphatic pain following breast cancer has been operationalized as the patient-report of co-occurring pain, or aching, or soreness, or tenderness and swelling.¹ *The Breast Cancer and Lymphedema Symptom Experience Index (BCLE-SEI) Part I* has been used to assess lymphatic pain (i.e., pain, aching, soreness, tenderness, and swelling) and additional symptoms related to lymph fluid accumulation or lymphedema.^{1,2,5} BCLE-SEI is valid and reliable patient-report instrument with a Cronbach's alpha of 0.92 for symptom occurrence.^{6,19,20,26,27} A response frame can be adjusted (e.g., "now," "past seven days", or "past three months") to indicate whether lymphatic pain occurs currently or has been persistent. Each item is rated on a 5-point Likert scale (i.e., 0 = no presence of a given symptom to 4 = greatest severity of a given symptom). Higher scores indicate more severe lymphatic pain. A recent large study of 568 breast cancer survivors used BCLE-SEI and identified four pain phenotypes: 33.1% were classified as lymphatic pain phenotype (i.e., presence of pain, aching, or soreness, and swelling), 35.9% as pain without swelling phenotype (i.e., only pain, aching, or soreness without arm/hand swelling), 5.9% as only swelling phenotype (i.e., only arm/hand swelling without pain, aching, or soreness), and 25% as phenotype of no symptom (i.e., absence of pain, aching, soreness, and arm/hand swelling). This study found that 41% of patients in the lymphatic pain group had >5% interlimb volume differences using an objective measure of limb volume by infra-red perometer (Perometry 350S). This supports the clinical observation that lymphatic pain often precedes changes in limb volume or lymph fluid. Thus, self-reported lymphatic pain can be considered an important marker for early lymphedema. Given that the major causes of lymphatic pain are inflammation and fluid accumulation; future research should explore genotypes, biomarkers, and fluid levels as potential objective markers of lymphatic pain.

6. Therapeutic behavioral intervention

Pharmacologic interventions have very limited effects on cancer-related pain and lymphatic pain, including the use of NSAIDs, opioids, antiepileptics, ketamine and lidocaine and long-term use of the medications in breast cancer patients can be problematic.¹¹ Behavioral strategies are also used for cancer-related pain.^{7–10} Lymphedema is usually treated through manual lymph drainage, physical therapy, compression garments, upper extremity exercise, focusing on swelling reduction.¹¹ Like lymphedema, lymphatic pain is associated with increased lymph fluid accumulation evidenced by increased lymph fluid level and inter-limb volume differences;^{1,2} and associated inflammatory responses.^{21,25} Effective treatment of lymphatic pain can decrease the risk of developing lymphedema and lessen lymphedema severity.^{26,27} However, there are persistent and worldwide challenges for patients (e.g., availability of interventions, cost and time for clinical visits) to receive timely and effective interventions for managing lymphatic pain.

The Optimal-Lymph-Flow (TOLF) is a web- and mobile-based program and one of the available behavioral interventions for lymphatic pain. TOLF intervention is based on physiological (fluid accumulation and inflammation) and cognitive principles (low self-efficacy for pain management) to promote lymph flow.^{2,26,27,37–40} TOLF program consists of therapeutic lymphatic exercises, healthy diet (i.e., nutrition-balanced, portion-appropriate diet, adequate hydration), and proper sleep. **Table 1** details the TOLF intervention strategies. TOLF therapeutic lymphatic exercises to promote lymph flow include muscle-tightening deep breathing, muscle-tightening pumping, and limb mobility exercises. In addition, TOLF provides patients with knowledge about lymphatic system, lymphedema, and daily self-assessment.

Guided by the self-efficacy theory (Fig. 1) and based on physiological-cognitive-behavioral principles, TOLF provides self-management strategies to activate the lymphatic system and promote lymph flow to decrease lymphatic pain and reduce the risk and severity of

Table 1
The-Optimal-Lymph-Flow (TOLF) program.^{2,26,27,37–40}

Keep a Healthy Weight		
Strategies	Rationale	Actions
Home-Based Muscle-Tightening Exercises	✓ Muscle-tightening-deep-breathing stimulates lymphatic ducts and help the lymphatic system to absorb lipids and decrease inflammation.	✓ Twice a day in the morning and at night before brushing teeth or as much as the patient wants throughout the day.
> Muscle-Tightening Deep Breathing	✓ Muscle-tightening pumping exercises create muscle pumping. This helps lymph fluid flow and decreases inflammation and helps to absorb lipids.	✓ Sedentary lifestyle: At least every 4 h
> Muscle-Tightening Pumping	✓ Muscle-tightening-deep-breathing and pumping exercises help to relax the body and decrease stress and pain.	
Diet	✓ Eat a nutrition-balanced diet: more vegetables and fruits as well as quality proteins.	✓ Each meal
> Eat nutrition-balanced Diet	✓ Maintain portion-appropriate diet: Cease eating when feeling 75% full for each meal.	
> Maintain Portion-appropriate Diet	✓ People may actually be thirsty, not hungry.	✓ Drink 6 to 8 glasses of water daily in the morning, before and during meals, and throughout the day. ✓ Avoid drinks with calories (e.g. juices). ✓ Drink green tea to boost metabolism.
Stay Hydrated		✓ 30-min 3 times a week or daily.
Large Muscle Exercises	✓ Large muscle exercises (e.g. walking, running, swimming, dancing, Yoga) help to burn more calories.	
> Walking, running, dancing, swimming, Yoga	✓ Large muscle exercises also promote lymph flow by creating muscle pumps.	
Get Enough Sleep	✓ Lack of sleep increases the production of the stress hormone cortisol, creates hunger, and leads to overeating. ✓ Getting just one more hour of sleep per night reduces belly fat accumulation.	✓ 7–8 h of sleep per night.

lymphedema.^{2,26,27,37–40} TOLF intervention focuses on training patient to implement self-management skills in their daily lives outside clinical settings without professionally administered therapy (e.g., by therapists or nurses). In other words, to achieve the therapeutic effects, patients execute home-based self-management skills. Self-efficacy for TOLF intervention refers to a breast cancer survivor's belief in her ability to execute TOLF self-management skills.^{2,26,27,38–42} Therefore, building patients' skills to manage lymphatic pain would increase their self-efficacy for TOLF interventions.

In a clinical trial, TOLF intervention was effective in preventing lymph fluid accumulation and maintaining pre-surgical limb volume among over 90% of 140 patients after breast cancer surgery.³⁸ A recently single-arm trial focusing on the immediate effects of TOLF lymphatic exercises² demonstrated a significant reduction in lymph fluid levels in bioimpedance mean L-Dex scores (M_{Δ} (Mean Changes) = -2.68 , 95% CI = $[-4.67, -0.69]$, $P = 0.010$). TOLF intervention demonstrated greater

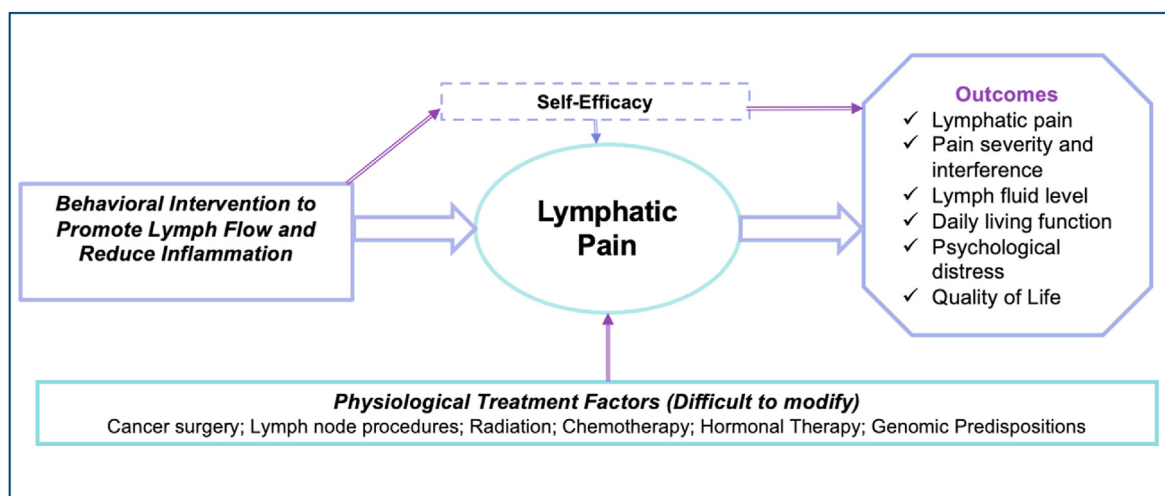


Fig. 1. Theoretical model for lymphatic pain management.

effects in patients with abnormal lymph fluid levels (i.e., L-Dex ≥ 7.1) in mean L-Dex scores ($M_{\Delta} = -5.19$, 95% CI = $[-1.75, -8.63]$, $P = 0.008$).²

A larger, 12-week, parallel Randomized Clinical Trial (RCT)³⁷ randomized 120 patients either into Arm Precaution (AP) control to improve limb mobility or TOLF intervention to promote lymph flow. At the end of the trial, significantly fewer patients in the TOLF intervention group reported chronic pain (i.e., lymphatic pain) (49% vs. 71%; OR = 0.39, 95% CI = $[0.17, 0.90]$, $P = 0.021$). TOLF intervention was effective in achieving complete pain reduction in 50% of patients (50% vs 22%; OR = 3.56, 95% CI = $[1.39, 9.76]$, $P = 0.005$) compared to the AP control group of 22%. TOLF intervention was effective in lowering significantly median severity scores (Med) of pain ($Med_{TOLF} = 0$, Interquartile Range [IQR] = 0–1 vs $Med_{AP} = 1$, IQR = 0–2; $P = 0.024$) and general bodily pain ($Med_{TOLF} = 1$, IQR = 0–1.5 vs $Med_{AP} = 1$, IQR = 1–3; $P = 0.040$).¹⁴ In addition, significantly fewer patients in TOLF group reported arm/hand swelling ($P = 0.038$). TOLF intervention achieved a 13% reduction in the proportion of patients who took pain medications compared to the AP control group which had a 5% increase. Results of a single-arm trial² supported the immediate effects of TOLF lymphatic exercises¹³ on reduction in lymphatic pain ($Med_{\Delta} = -1.00$, 95% CI = $[-1.5, -0.1]$, $P = 0.004$), and arm/hand swelling ($Med_{\Delta} = -1.00$, 95% CI = $[-1.5, -0.5]$, $P = 0.004$).

7. Recommendations for research and clinical practice

To deploy the emerging knowledge regarding lymphatic pain and different types of pain, the following recommendations are posited to translate and integrate this knowledge into research and clinical practice.

Recommendation for research. Given that the mechanism of lymphatic pain is associated with lymph fluid accumulation and inflammation, it is possible that lymphatic pain may have different underlying mechanisms compared to other pain phenotypes (e.g., chemotherapy-induced peripheral neuropathy, or arthralgias related to hormonal treatments, pain without swelling). Emerging research evidence presents new insights to differentiate lymphatic pain from other types of pain, which is essential in finding a cure. Future research should explore the unique underlying mechanisms of lymphatic pain through biomarker and genomic approaches as well as associated demographic and socioeconomic determinants (e.g., age, financial status, ethnicity).

Current research suggests that behavioral interventions (e.g., TOLF intervention) designed to promote lymph flow produce significant benefits in reducing lymphatic pain, general bodily pain, and specific lymphedema symptoms (e.g., arm/hand swelling, heaviness, limited movement in shoulder and arm).^{2,26,27,37–40} Future research should

investigate the effects of different behavioral interventions on different types of pain, including lymphatic pain. Research should also focus on developing targeted and effective therapeutic interventions for pain by investigating which therapeutic approaches, such as pharmacological, low-dose laser, or behavioral approaches, are most efficacious for different types of pain with different etiologies.

Recommendations for clinical practice. To differentiate lymphatic pain from other types of pain, clinicians may use the reliable and valid patient outcome measures (e.g., BCLE-SEI) to evaluate and track lymphatic pain in clinical practice to ensure timely interventions.^{43,44} Research-based behavioral interventions to promote lymph flow and inflammation (e.g., TOLF intervention) are able to induce immediate and long-term therapeutic effects to relieve lymphatic pain, swelling, and lymphedema symptoms.^{2,26,27,37–40} Such interventions can be prescribed to patients to reduce not only lymphatic pain, but also general bodily pain and other symptoms associated with fluid accumulation.

8. Conclusion

At least one-third of breast cancer survivors have suffered lymphatic pain that is defined as co-occurring pain or sensations of aching, soreness, or tenderness and swelling.^{1,16} Importantly, lymphatic pain results in greater impairments in ADLs, emotional distress, and overall health.^{1,5,16} Precision assessment that enables clinicians to distinguish different types of pain is imperative to find a cure for pain. Behavioral interventions to promote lymph flow are safe, efficacious, and affordable. Such interventions are beneficial for millions of women treated for breast cancer worldwide to reduce lymphatic pain, general bodily pain, and other symptoms related to fluid accumulations and chronic inflammation.

Conflicts of interest

The authors declare no conflict of interest.

Human and animal rights and informed consent

Not Applicable.

Funding

This review was part of study funded by Oncology Nursing Foundation (2022 ONF RE33) with Mei R Fu as the principal investigator and the National Institute of Health/National Science Foundation/National Cancer Institute (1R01CA214085-01) with Mei R Fu and Yao Wang as the

multiple principal investigators. Its contents are solely the responsibility of the authors and do not necessarily represent the official views of the funders. The funders had no role in the preparation of the manuscript and decision to publish.

Compliance with ethical standards

Not Applicable.

CRediT authorship contribution statement

Jeanna Mary Qiu: Writing – review & editing, Writing – original draft, Formal analysis, Data curation, Conceptualization. **Mei Rosemary Fu:** Writing – review & editing, Writing – original draft, Project administration, Funding acquisition, Formal analysis, Data curation, Conceptualization. **Catherine S. Finlayson:** Writing – review & editing, Conceptualization. **Charles P. Tilley:** Writing – review & editing, Conceptualization. **Rubén Martín Payo:** Writing – review & editing, Conceptualization. **Stephanie Korth:** Writing – review & editing, Conceptualization. **Howard L. Kremer:** Writing – review & editing, Conceptualization. **Cynthia L. Russell Lippincott:** Writing – review & editing, Conceptualization.

References

- Fitzgerald Jones K, Fu MR, McTernan ML, et al. Lymphatic pain in breast cancer survivors. *Lymphatic Res Biol.* 2022;20(5):525–532. <https://doi.org/10.1089/lrb.2021.0017>.
- Fu MR, McTernan ML, Qiu JM, et al. The effects of kinect-enhanced lymphatic exercise intervention on lymphatic pain, swelling, and lymph fluid level. *Integr Cancer Ther.* 2021;20:15347354211026757. <https://doi.org/10.1177/15347354211026757>.
- World Health Organization. *Breast Cancer*; 2023. Available on <https://www.who.int/news-room/fact-sheets/detail/breast-cancer>. Accessed February 3, 2024.
- American Cancer Society (ACS). *Breast Cancer Facts & Figures*. Atlanta, 2022–2024: American Cancer Society, Inc. Accessed February 3, 2024. URL: <https://www.cancer.org/research/cancer-facts-statistics/breast-cancer-facts-figures.html>.
- Fu MR, Axelrod D, Guth A, et al. The effects of obesity on lymphatic pain and swelling in breast cancer patients. *Biomedicine.* 2021 Jul 14;9(7):818. <https://doi.org/10.3390/biomedicine9070818>. PMID: 34356882; PMCID: PMC8301355.
- Fu MR, Axelrod D, Cleland CM, et al. Symptom report in detecting breast cancer-related lymphedema. *Breast Cancer.* 2015 Oct 15;7:345–352. <https://doi.org/10.2147/BCTT.S87854>. PMID: 26527899; PMCID: PMC4621182.
- Paice JA. Pain in cancer survivors: how to manage. *Curr Treat Options Oncol.* 2019; 20(6):48. <https://doi.org/10.1007/s11864-019-0647-0>.
- Treede RD, Rief W, Barke A, et al. Chronic pain as a symptom or a disease: the IASP classification of chronic pain for the international classification of diseases (ICD-11). *Pain.* 2019;160(1):19–27. <https://doi.org/10.1097/j.pain.0000000000001384>.
- Jiang C, Wang H, Wang Q, Luo Y, Sidlow R, Han X. Prevalence of chronic pain and high-impact chronic pain in cancer survivors in the United States. *JAMA Oncol.* 2019; 5(8):1224–1226. <https://doi.org/10.1001/jamaoncol.2019.1439>.
- Jones KF, Fu MR, Wood Magee L, et al. "It is so easy for them to dismiss": a phenomenological study of cancer survivors with chronic cancer-related pain [published online ahead of print, 2023 mar 21]. *J Palliat Med.* 2023. <https://doi.org/10.1089/jpm.2022.0538>.
- Armer JA, Ostby P, Ginex P, et al. ONS Guidelines for Cancer treatment-related lymphedema. *Oncol Nurs Forum.* 2020;47(5):518–538. <https://doi.org/10.1188/20.ONF.518-538>. PMID: 32830794.
- Dowell D, Ragan KR, Jones CM, et al. Prescribing opioids for pain—the new CDC clinical practice guideline. *N Engl J Med.* 2022;387(22):2011–2013. <https://doi.org/10.1056/NEJMp2211040>.
- Sweeney C, Schmitz KH, Lazovich D, Virnig BA, Wallace RB, Folsom AR. Functional limitations in elderly female cancer survivors. *J Natl Cancer Inst.* 2006;98:521–529.
- O'Toole JA, Ferguson CM, Swaroop MN, et al. The impact of breast cancer-related lymphedema on the ability to perform upper extremity activities of daily living. *Breast Cancer Res Treat.* 2015;150:381–388.
- Park JH, Merriman J, Brody A, et al. Limb volume changes and activities of daily living: a prospective study. *Lymphatic Res Biol.* 2020. <https://doi.org/10.1089/lrb.2020.0077>.
- Fu MR, McTernan ML, Qiu JM, et al. Co-Occurring fatigue and lymphatic pain incrementally aggravate their negative effects on activities of daily living, emotional distress, and overall health of breast cancer patients. *Integr Cancer Ther.* 2022;21: 15347354221089605. <https://doi.org/10.1177/15347354221089605>.
- Fu MR, Kang Y. Psychosocial impact of living with cancer-related lymphedema. *Semin Oncol Nurs.* 2013;29(1):50–60. <https://doi.org/10.1016/j.soncn.2012.11.007>.
- Reis JC, Antoni MH, Travado L. Emotional distress, brain functioning, and biobehavioral processes in cancer patients: a neuroimaging review and future directions. *CNS Spectr.* 2020;25(1):79–100. <https://doi.org/10.1017/S1092852918001621>.
- Shi S, Lu Q, Fu MR, et al. Psychometric properties of the breast cancer and lymphedema symptom experience index: the Chinese version. *Eur J Oncol Nurs.* 2016;20:10–16. <https://doi.org/10.1016/j.ejon.2015.05.002>.
- Cachero-Rodríguez J, Menéndez-Aller Á, Fu MR, Llaneza-Folgueras A, Fernández-Alvarez MM, Martín-Payo R. Psychometric properties of the Spanish version of breast cancer and lymphedema symptom experience index. *Psicothema.* 2022;34(2): 291–298. <https://doi.org/10.7334/psicothema2021.388>.
- Fu MR, Conley YP, Axelrod D, et al. Precision assessment of heterogeneity of lymphedema phenotype, genotypes and risk prediction. *Breast.* 2016;29:231–240. <https://doi.org/10.1016/j.breast.2016.06.023>.
- Kwan ML, Yao S, Lee VS, et al. Race/ethnicity, genetic ancestry, and breast cancer-related lymphedema in the Pathways Study. *Breast Cancer Res Treat.* 2016 Aug; 159(1):119–129. <https://doi.org/10.1007/s10549-016-3913-x>. Epub 2016 Jul 22. PMID: 27449493; PMCID: PMC5010992.
- Fu MR, Wang Y, Li C, et al. Machine learning for detection of lymphedema among breast cancer survivors. *mHealth.* 2018;4:17. <https://doi.org/10.21037/mhealth.2018.04.02>. Published 2018 May 29.
- Wei X, Lu Q, Jin S, et al. Online ahead of print). Developing and validating a prediction model for lymphedema detection in breast cancer survivors. *Eur J Oncol Nurs.* 2021. <https://doi.org/10.1016/j.ejon.2021.102023>.
- Fu MR, Aouizerat BE, Yu G, et al. Model-based patterns of lymphedema symptomatology: phenotypic and biomarker characterization. *Curr Breast Cancer Rep.* 2021;13(1):1–18. <https://doi.org/10.1007/s12609-020-00397-6>.
- Liu F, Li F, Fu MR, et al. Self-management strategies for risk reduction of subclinical and mild stage of breast cancer-related lymphedema: a longitudinal, quasi-experimental study. *Cancer Nurs.* 2021;44(6):E493–E502. <https://doi.org/10.1097/NCC.0000000000000919>.
- Du X, Li Y, Fu L, et al. Strategies in activating lymphatic system to promote lymph flow on lymphedema symptoms in breast cancer survivors: a randomized controlled trial. *Front Oncol.* 2022;12:1015387. <https://doi.org/10.3389/fonc.2022.1015387>.
- Fu MR, Rosedal M. Breast cancer survivors' experiences of lymphedema-related symptoms. *J Pain Symptom Manag.* 2009;38(6):849–859. <https://doi.org/10.1016/j.jpainsymman.2009.04.030>.
- Buki LP, Rivera-Ramos ZA, Kanagui-Muñoz M, et al. I never heard anything about it: Knowledge and psychosocial needs of Latina breast cancer survivors with lymphedema. *Women's Health (Lond).* 2021;17:17455065211002488. <https://doi.org/10.1177/17455065211002488>. PMID: 33764235; PMCID: PMC8010798.
- Eversley R, Estrin D, Dibble S, Wardlaw L, Pedrosa M, Favila-Penney W. Post-treatment symptoms among ethnic minority breast cancer survivors. *Oncol Nurs Forum.* 2005;32(2):250–256. <https://doi.org/10.1188/05.ONF.250-256>. PMID: 15759063.
- Miller JW, Smith JL, Ryerson AB, Tucker TC, Allemani C. Disparities in breast cancer survival in the United States (2001–2009): findings from the CONCORD-2 study Suppl 24. *Cancer.* 2017;123(24):5100–5118. <https://doi.org/10.1002/ncr.30988>. PMID: 29205311; PMCID: PMC5826549.
- Wielen LM, Gilchrist EC, Nowels MA, Petterson SM, Rust G, Miller BF. Not near enough: racial and ethnic disparities in access to nearby behavioral health care and primary care. *J Health Care Poor Underserved.* 2015;26(3):1032–1047. <https://doi.org/10.1353/hpu.2015.0083>. PMID: 26320931; PMCID: PMC4962556.
- Armer J, Fu MR. Age differences in post-breast cancer lymphedema signs and symptoms. *Cancer Nurs.* 2005;28(3):200–209. <https://doi.org/10.1097/00002820-200505000-00007>.
- Fu MR, LeMone P, McDaniel RW. An integrated approach to an analysis of symptom management in patients with cancer. *Oncol Nurs Forum.* 2004;31(1):65–70. <https://doi.org/10.1188/04.ONF.65-70>.
- Washington AE, Lipstein SH. The patient-centered outcomes research institute—promoting better information, decisions, and health. *N Engl J Med.* 2011;365:e31. <https://doi.org/10.1056/NEJMp1109407>.
- Helyer LK, Varnic M, Le LW, Leong W, McCready D. Obesity is a risk factor for developing postoperative lymphedema in breast cancer patients. *Breast J.* 2010;16: 48–54. <https://doi.org/10.1111/j.1524-4741.2009.00855.x>.
- Fu MR, Axelrod D, Guth AA, et al. A web- and mobile-based intervention for women treated for breast cancer to manage chronic pain and symptoms related to lymphedema: results of a randomized clinical trial. *JMIR Cancer.* 2022;8(1):e29485. <https://doi.org/10.2196/29485>. Published 2022 Jan 17.
- Fu MR, Axelrod D, Guth AA, et al. Proactive approach to lymphedema risk reduction: a prospective study. *Ann Surg Oncol.* 2014;21(11):3481–3489. <https://doi.org/10.1245/s10434-014-3761-z>.
- Fu MR, Axelrod D, Guth AA, et al. mHealth self-care interventions: managing symptoms following breast cancer treatment. *mHealth.* 2016;2:28. <https://doi.org/10.21037/mhealth.2016.07.03>.
- Fu MR, Axelrod D, Guth AA, et al. Usability and feasibility of health IT interventions to enhance self-care for lymphedema symptom management in breast cancer survivors. *Internet Interv.* 2016;5:56–64. <https://doi.org/10.1016/j.invent.2016.08.001>.
- Anderson KO, Dowds BN, Pelletz RE, Edwards WT, Peeters-Asdourian C. Development and initial validation of a scale to measure self-efficacy beliefs in patients with chronic pain. *Pain.* 1995 Oct;63(1):77–84.
- Porter LS, Keefe FJ, Garst J, McBride CM, Baucum D. Self-efficacy for managing pain, symptoms, and function in patients with lung cancer and their informal caregivers: associations with symptoms and distress. *Pain.* 2008 Jul 15;137(2):306–315. <https://doi.org/10.1016/j.pain.2008.04.011>.

- doi.org/10.1016/j.pain.2007.09.010. Epub 2007 Oct 17. PMID: 17942229; PMCID: PMC2522367.
43. Nahum JL, Fu MR, Scagliola J, et al. Real-time electronic patient evaluation of lymphedema symptoms, referral, and satisfaction: a cross-sectional study. *mHealth*. 2021;7:20. <https://doi.org/10.21037/mhealth-20-118>. Published 2021 Apr 20.
44. Bustos VP, Friedman R, Pardo JA, Granoff M, Fu MR, Singhal D. Tracking symptoms of patients with lymphedema before and after power-assisted liposuction surgery. *Ann Plast Surg*. 2023;90(6):616–620. <https://doi.org/10.1097/SAP.0000000000003430>.