Lymphoedema therapists: a national and international survey

Elizabeth A Anderson, Allison B Anbari, Nathan C Armer and Jane M Armer

Key words
Lymphoedema therapists, survey

Abstract
The American Lymphedema Framework Project (AFLP) surveyed lymphoedema therapists in the US in 2009 to describe their preparation, patient population and care practices. In the autumn of 2018, the survey was expanded to trained therapists worldwide to describe and compare current and past therapist characteristics and practices. The updated 2009 survey was distributed via Qualtrics to US and international therapists. The current analysis includes over 950 completed surveys. Preliminary results showed: country: US (n=672/922 [73%]; Canada (n=92 [10%]); United Kingdom (n=42 [5%]), Australia (n=28 [3%]); gender: identifying as female (n=633/676 [93%]); mean age: 47 yrs (range 21–76); discipline: physical therapist [45%], occupational therapist [31%], massage therapist [24%]); mean practice years: 10.7 yrs (range 0–41); and practice setting: hospital out-patient clinic (47%); private practice (38%); hospital in-patient (13%); home care/hospice (9%). Further 2009–2018 comparative analyses will be shared. Understanding characteristics and practices of lymphoedema therapists and patients will help stakeholders meet under- and unmet needs of this population.

Declaration of interest: None.

Over 200 million people around the world have or are at risk of developing lymphoedema (Grada and Phillips, 2017). Lymphoedema is a failure of the lymphatic transport system, resulting from cancer treatment, infection, trauma, and/or genetic/familial structural malformations leading to distressing and debilitating swelling of the affected area (International Society of Lymphology [ISL], 2016; Armer et al, 2018). Volume reduction and symptom management by a trained lymphoedema therapist is critical to improving symptoms and maintaining quality of life (ISL, 2016). Certification training involves licensed healthcare professionals completing a 135-hour didactic course and 1 year of clinical practice in lymphoedema management (Lymphology Association of North America [LANA], 2017; North American Lymphoedema Education Association [NALEA], 2017). LANA certification is voluntary and available worldwide. The goal of this updated survey by the American Lymphoedema Framework Project (ALFP) was to determine the current state of lymphoedema care and practice characteristics worldwide as reported by the therapists.

The ALFP is a national, United States-based collaboration of healthcare providers, researchers, patients, advocates, educators, industry representatives and third-party payers led by recognised clinical experts and investigators in lymphoedema care (ALFP, 2019). Since 2008, its mission has been to evaluate appropriate healthcare services for patients with all forms of lymphoedema and advance the quality of lymphoedema care both in the US and worldwide. The partnership with the International Lymphoedema Framework (ILF) has resulted in increased global awareness and research advancement towards improving functional, physical, and quality of life outcomes for patients with lymphoedema (Armer et al, 2010; International Lymphoedema Framework, 2019).

Between 2008 and 2014, five ‘open-space’ stakeholder meetings were held to ensure focus on priority issues remained current: Chicago, IL; Columbus, OH; Atlanta, GA; Columbia, MO; and Cape
Table 1. Characteristics of lymphoedema therapists.

<table>
<thead>
<tr>
<th>Gender identification</th>
<th>Age</th>
<th>Years of treating lymphoedema</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female = 94%</td>
<td>Mean = 47.4 years (SD ± 11 years)</td>
<td>Mean = 10.7 years (SD ± 7.7 years)</td>
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<tr>
<td></td>
<td>Minimum = 21 years</td>
<td>Minimum = 1 month</td>
</tr>
<tr>
<td>Male = 6%</td>
<td>Maximum = 76 years</td>
<td>Maximum = 41 years</td>
</tr>
</tbody>
</table>

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Town, South Africa (Armer et al, 2017): The international meeting held in Cape Town contributed to the eventual formation of the Lymphoedema Association of South Africa (LAOSA) (Lymphedema Association of South Africa, 2019). The 2008 issues were confirmed at each meeting and continue to be priorities, with awareness and education ranked first:

- Increase awareness of lymphoedema and related lymphatic system disorders
- Improve patient education, support, and self-management
- Establish criteria for health provider education
- Continue to build the credibility of the ALFP
- Develop and implement research to refine diagnostic standards and provide evidence for effective treatments
- Promote evidence-based practice for lymphoedema management
- Improve reimbursement for lymphoedema care and resources.

ALFP goals of building a minimum dataset (MDS) to support outcomes research and defining best practice for lymphoedema care have matured since 2008. The MDS contains over 1,300 patients with data points including volume measurements, symptoms, and longitudinal visit information (Armer et al, 2017). Data-mining tools and a 3-D mobile imaging platform allow more research questions to be explored and increase accuracy and frequency of lymphoedema measurements. Best practice aims fostered the completion of 11 systematic reviews addressing lymphoedema care outcomes, providing healthcare professionals with information to support clinical practice. In addition, the ALFP Therapist Directory ‘Look4LE’ continues to expand, registering information on over 1,200 LANA-certified US and international therapists (Armer et al, 2017).

The priority of increasing lymphoedema awareness and education motivated the ALFP national survey of lymphoedema therapists in 2009, with a follow-up survey encouraged by NALEA training schools in 2011. Continued growth of the therapist directory, new lymphoedema management research results, and the continued recognition of gaps in provider education that affect the care of patients with lymphoedema (Ng et al, 2015; Armer et al, 2017) was a catalyst for the authors to explore current practice environments and educational frameworks of therapists in both the US and worldwide.

Methods

Between June and July 2018, the 2009 ALFP survey was updated. Online searches reviewed treatment types, referral sources, measurement methods, payment methods, patient educational resources, licensure processes, and sources of licensure training to determine changes since 2009. The survey questions were reviewed by research team members and edited through electronic review. The final survey included 56 items that were imported into Qualtrics™ (Qualtrics, Provo: UT). The items queried information about therapists’ demographics, practice location, patient population, therapy modalities, training processes, treatment payment sources and practice setting descriptions. This study was approved as an exempt study by the University of Missouri Institutional Review Board.

Lymphoedema therapists were invited to complete the survey by email invitation sent from the ALFP stakeholder database. Snowball-sampling techniques were used, such as inviting recipients to forward the survey link to eligible colleagues. Additional network members and partnership organisations were invited to forward the survey link to contacts and therapists on their membership lists. The survey was available for online completion for 8 weeks from October through December 2018.

Results

Demographics

Data were submitted by 950 therapists from all 50 states of the United States (US) (n=662) and all seven Canadian provinces, along with 41 additional countries (n=288) (Figure 1). The majority of respondents self-identified as female (93%), with an average age of 47 years (range 21–76) (Table 1). The three most frequent disciplines reported were physical therapy (45%); occupational therapy (31%); and massage therapy (24%). Mean reported years in practice was 10.7 years (range 0–41). The majority of therapists (96%) self-reported they met the 135-hour training requirement to be recognised as a Certified Lymphedema Therapist (CLT) and 33% reported achieving LANA certification. The top four reported practice settings included: hospital outpatient clinic (47%); private practice (38%); hospital inpatient (13%); and home care/hospice (9%). A descriptive summary of the data is provided, with number of responses varying from 680–719 for each question because participants were not required to answer all questions.

Treatment

The most commonly-reported treatments offered by responding lymphoedema therapists were the various elements of comprehensive decongestive therapy (CDT), consisting of manual lymphatic drainage (MLD), compression bandaging and compression garments, exercise, movement, risk-reduction education, skin care and soft tissue mobilisation. Less than 15% of responding therapists reported offering single-phase pneumatic compression devices, aquatic treatment, low-level laser, vibrator treatment, compression bandage only, reflexology and other treatments. A majority (55%) of therapists offered seven or more treatment options.

On average, therapists reported that 80% of patients treated had secondary lymphoedema. Lymphoedema therapists also reported treating patients with the following areas of oncology-related lymphoedema: upper extremities (53%); lower extremities (30%); trunk (7%); head and neck (8%); and genitals (2%). Concerning the comparison between wound care and lymphoedema management, therapists, on average, reported that 81% of...
their patients required lymphoedema care only; 3% required wound care only; and 16% required both.

Further descriptive-comparative analyses will be performed to compare 2009 and 2018 findings. Overall, preliminary findings from the updated survey reveal modest variance from the 2009 survey, on average 0–4%. A companion manuscript detailing the comparative analysis is forthcoming.

Discussion
All 50 US states, all seven Canadian provinces, and an additional 41 countries had representation in this 2018 ALFP-sponsored lymphoedema therapist survey. The high level of training among therapists could be due to selection bias related to the method of survey dissemination and differential access to the online survey. It could also be that highly-prepared therapists are more likely to respond. One-third of the therapists reported they had representation in this 2018 ALFP-sponsored lymphoedema therapist survey. The largest percentage of therapists required lymphoedema care only; 3% required wound care only; and 16% required both.

Further descriptive-comparative analyses will be performed to compare 2009 and 2018 findings. Overall, preliminary findings from the updated survey reveal modest variance from the 2009 survey, on average 0–4%. A companion manuscript detailing the comparative analysis is forthcoming.

Conclusion
With this update to the 2009 survey, we were able to continue exploring the perspectives and practices of therapists from around the world for over a decade. Lymphoedema therapists are critical members of the health care team providing care to persons with and at risk of lymphoedema from all causes. Understanding the training, characteristics, and practices of lymphoedema therapists and their patients will help health professionals, educators, policymakers, and funders better meet the under- and unmet needs of this growing population.

Acknowledgements
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Figure 1. Framework countries responding to the therapist survey.