Improving the quality of life of patients with breast cancer-related lymphoedema by lymphaticovenous anastomosis (LVA): study protocol of a multicentre randomised controlled trial

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ABSTRACT

Introduction Early breast cancer detection and advancements in treatment options have resulted in an increase of breast cancer survivors. An increasing number of women are living with the long-term effects of breast cancer treatment, making the quality of survivorship an increasingly important goal. Breast cancer-related lymphoedema (BCRL) is one of the most underestimated complications of breast cancer treatment with a reported incidence of 20%. A microsurgical technique called lymphaticovenous anastomosis (LVA) might be a promising treatment modality for patients with BCRL. The main objective is to assess whether LVA is more effective than the current standard therapy (conservative treatment) in terms of improvement in quality of life and weather it is cost-effective.

Methods and analysis A multicentre, randomised controlled trial, carried out in two academic and two community hospitals in the Netherlands. The study population includes 120 women over the age of 18 who have undergone treatment for breast cancer including axillary treatment (sentinel lymph node biopsy or axillary lymph node dissection) and/or axillary radiotherapy, presenting with an early stage lymphoedema of the arm, viable lymphatic vessels and received at least 3 months conservative treatment. Sixty participants will undergo the LVA operation and the other sixty will continue their regular conservative treatment, both with a follow-up of 24 months. The primary outcome is the health-related quality of life. Secondary outcomes are societal costs, quality adjusted life years, cost-effectiveness ratio, discontinuation rate of conservative treatment and excess limb volume.

Ethics and dissemination The study was approved by the Ethics Committee of Maastricht University Medical Center (METC) on 19 December 2018 (NL67059.068.18). The results of this study will be disseminated in presentations at academic conferences, publications in peer-reviewed journals and other news media.

Trial registration number NCT02790021; Pre-results.

INTRODUCTION

An increasing number of women survive breast cancer due to advancements in treatment options. As a result, the number of women living with the long-term effects of breast cancer treatment grows, making the quality of survivorship more relevant. Between 8% and 56% of breast cancer survivors develop arm or shoulder problems such as restricted shoulder mobility, shoulder pain and lymphoedema, with one of the most underestimated and debilitating morbidities of them all being upper limb lymphoedema.

Up to 70% of the patients who develop breast cancer-related lymphoedema (BCRL) do so within the first 2 years post-treatment, however, cases have been described of women developing upper limb lymphoedema 20 years or later after initial treatment. In the Netherlands, between 7% and 30% of the 14 000 annual patients with invasive breast
cancer will develop lymphoedema depending on certain treatment and patient related risk factors.14 15

The following risk factors are associated with the development (and severity) of BCRL: the extent of breast/axillary surgery, adjuvant radiation, (neo-)adjuvant chemotherapy, the number of positive nodes, treatment in dominant limb and obesity.5 16–20 Limb swelling may present with symptoms of heaviness, tightness, pain and loss of normal arm function and range of motion. The negative psychological effects brought on by the impairments of activities in daily life and reduced limb aesthetics constitute an additional burden and decrease in health-related quality of life (HRQoL).1 7 11 13 21 Moreover, infections of the skin are regularly seen in a severe stadium of lymphoedema, such as erysipelas or cellulitis.2 8 21

Conservative therapy

Complex decongestive therapy (CDT), currently accepted as the standard treatment for lymphoedema, is initially aimed at alleviating symptoms without curative intent which for most patients means lifelong treatment and a constant reminder of the breast cancer period. CDT includes general skin care, patient education, compression therapy with compression bandages and garment, manual lymphatic drainage (MLD) and exercise therapy.14 22 23 A systematic review concluded that compression garment in combination with manual lymph drainage induces a significant limb volume reduction of 17% to 60%.24 Another randomised controlled trial (RCT) demonstrated a 29% reduction in excess limb volume with combined conservative therapy.25 However, after reaching maximum limb volume reduction, compression garment are lifelong necessary for the patients to maintain the volume reduction obtained.

Lymphaticovenous anastomosis

Connections can be made between the lymphatic and venous systems to divert static lymph fluid away from the obstruction site in a technique called lymphaticovenous anastomosis (LVA).26 Due to advancements, microvascular surgery is more developed, and anastomoses in vessels as small as 0.3 mm in diameter are made possible. Several studies on lymphatic super microsurgery performing LVA are available.26–41 Most of the studies describe results on both upper and lower limb lymphoedema and not only secondary lymphoedema.27 34 Nevertheless, studies mention a volume or circumference decrease between 30% and 61%, and positive results on subjective complaints with low incidence or no complications.26–29 31 36–39 41 Furthermore, more than half of the patients eventually were able to discontinue compression garment after an LVA procedure.27 42

Many studies have been performed, mostly reporting on a small study population. Furthermore, the majority were retrospective, few were prospective, yet none of them were randomised. Another disadvantage is the heterogeneity of the patient population, assessment modalities and inconsistent reporting of outcomes and complications.27 30 34

The aim of this multicentre RCT is to examine HRQoL and (cost-)effectiveness of LVA compared to CDT in a large homogenous group of patients with BCRL.

METHODS AND ANALYSIS

Study design

The LYMPH trial is a multicentre, non-blinded, RCT and will be conducted in the Maastricht University Medical Center, Radboud University Medical Center, Zuyderland Medical Center and Canisius-Wilhelmina Hospital in the Netherlands.

Enrolment will take place at the outpatient clinics of the participating hospitals. The inclusion and exclusion criteria are listed in box 1. A total of 120 women must be recruited after a period of 2 years. After inclusion and informed consent, participants will be randomly assigned to either the LVA or conservative (CDT) group with a 1:1 allocation as per a computer-generated randomisation schedule stratified by site using block randomisation. This computer-generated randomisation is done within the electronic Case Report Form in CASTOR EDC. Since only patients with early-stage lymphoedema are included and no large imbalances are expected, no stratification for other demographic data are applied.

Blinding is not possible in this study, since the operation scars on the arm are easily detectable during the study measurements. However, HRQoL is the primary outcome which is examined by a digital standardised questionnaire. The patients only have access to the questionnaires and the researcher has no influence on this data. The start date of the study is November 2018, and...
the estimated completion date of the study is November 2022. An overview of the study design is shown in figure 1.

**Interventions to be measured**

**Group A: conservative therapy**

The current standard treatment for BCRL is a combination of different methods of conservative therapy, also known as CDT.14 CDT incorporates two stages of treatment. The first treatment phase entails skincare, MLD, exercises aimed at improvement of mobility/range of motion in the shoulder, elbow or wrist joints, and compression therapy through bandaging. Most patients already underwent this phase short after the diagnosis of lymphoedema. CDT in the second treatment phase is aimed at maintenance of the achieved limb volume/circumference reduction through compression therapy with therapeutic elastic compression garment for the arm. Skincare, mobility exercises and MLD is continued in this phase if needed.14 24 Since CDT aim to alleviate symptoms without curative intent, this treatment is mostly lifelong needed. In this study, the patients are followed for 2 years during their regular conservative treatment.

**Complex decongestive therapy**

Patients allocated to group A will be referred to one of the following dedicated lymphoedema (physical/skin) therapy clinics, if not already treated by one, according to their place of residence for continuation of standard conservative therapy. Only standard conservative therapy, as they would have gotten if not participating in this study, will take place in these clinics, no study measurements. All women in this study group will be treated according to a protocol which is already in use for patients not participating in this study, since it is considered as the best available standard care. To be able to compare the outcomes for the conservative therapy group, a standardised treatment protocol using the standard lymphatic drainage methods applied in the Netherlands and Germany (‘Verdonkmethod’ and ‘Asdonkmethod’, respectively), will be used in this study. See the online supplementary data for the CDT protocol. Ongoing conservative treatment and the frequency is controlled by the skin therapist. All information regarding conservative treatment is noted in the patient diary.

**Group B: surgical treatment**

**Lymphaticovenous anastomosis**

LVA is a relative minimally invasive registered procedure which can be performed under local anaesthesia. The patient lies comfortable on the operation table, and a limb table is used. The limb is then prepared for surgery. Before making the incision, a mix of bupivacaine (Marcaine) and epinephrine (1:100 000) is injected at the site of incision to achieve local anaesthesia and optimal haemostasis.

The following steps of the operation are performed using a surgical microscope. Based on the ICG lymphography mapping, incisions of 1–2 cm are made at the predetermined sites. Lymphatic vessels are identified, and an anastomosis is performed with a similarly sized adjacent recipient vein in the subdermal plane. The anastomosis is usually performed in an end-to-end fashion in case both the lymphatic vessel and vein have approximately the same calibre (otherwise end-to-side). The end-to-end anastomosis is created with an 11–0 suture. The patency of the LVA is confirmed by direct visual examination under the microscope. On average, 1–4 anastomosis are performed in a lymphoedematous arm. The superficial wound is closed using 4–0 Ethilon covered by adhesive plasters and a bandage. The operation length is approximately 2–3 hours.29
Postoperative treatment

From 2 weeks after surgery, when the stitches are removed, patients will be treated with conservative therapy the same way and in the same frequency as preoperatively. The participants are treated by the same method as group A if needed, as described in phase 2 (maintenance phase) of the CDT protocol. After 3 months, the plastic surgeon will determine whether conservative therapy can be reduced or stopped, depending on the decrease of subjective complaints and swelling of the arm. The frequency of MLD will be controlled by the skin therapist and noted in the patient diary.

Follow-up moments for both groups will be at 3, 6, 12, 18 and 24 months. For group A, the follow-up starts from the day of the informed consent signing and for group B from the day of the surgery.

Sample size calculation

We made the following assumptions for the calculation of the sample size to show a statistically significant and clinically relevant difference in quality of life between treatment groups at 12 months’ follow-up as measured with the Lymphedema Functioning, Disability and Health (Lymph-ICF) questionnaire:

Comparing LVA to conservative treatment, the minimal difference in HRQoL that is considered as clinically relevant is 15 points (15% decrease on the 0 to 100 scale) on the Lymph-ICF questionnaire at 12 months’ follow-up. The EQ-5D-5L is a generic HRQoL measure that can be used to calculate QALYs to be used in the economic evaluation. The EQ-5D-5L is a questionnaire responsive to changes in health in breast cancer patients after conclusion of treatment.

The EQ-5D-5L examines a patient’s HRQoL on the day of the interview. It consist of the EQ-5D-5L descriptive system and a Visual Analog Scale (EQ VAS). The descriptive system comprises five dimensions: mobility, self-care, usual activities, pain/discomfort and anxiety/depression. Each dimension has five levels: no problems, slight problems, moderate problems, severe problems and extreme problems. Responses to the five items result in a patient’s health state that can be transformed into an index score representing a HRQoL-weight, ranging between 0 (death) and 1 (perfect health). These index scores are combined with length of life to calculate the QALYs. The EQ VAS records the patient’s self-rated health with endpoints labelled ‘the best health you can imagine’ at the top and ‘the worst health you can imagine’ at the bottom.

Discontinuation of conservative treatment will be assessed with a patient diary to record the frequency of treatments received (ie, skin therapy visits, number of compression garment, etc). Lastly, bilateral limb volume measurements will be done using VECTRA 3D imaging and the water displacement method. The excess limb volume is measured as the difference in volume between the affected and unaffected limb which is reported as a percentage of the volume of the unaffected limb. A relative volume reduction (relative to the unaffected arm) as well as an absolute volume reduction (volume reduction of the affected arm at next measurement) will be calculated. The calculated volume will be corrected for the body mass index and for volume differences between the dominant and non-dominant arm.

Secondary outcomes

Secondary outcomes are the societal costs, QALYs, incremental cost-effectiveness, discontinuation of conservative treatment and excess limb volume. Assessment will be done at baseline and 3, 6, 12, 18 and 24 months after informed consent (group A), or after surgery (group B). Costs include healthcare-related costs, costs to patients and family, and costs due to lost productivity. Complete individual-level hospital resource use data (eg, surgical intervention, diagnostic procedures, hospital admissions, outpatient clinic visits) will be measured using medical records and provider information systems. Resource use outside the hospital (eg, lymphoedema therapy, general practitioner visits, out-of-pocket expenses such as for therapeutic elastic garment and over-the-counter drugs, travel costs and quantities of lost paid work) will be determined by means of prospective cost diaries as kept by participants. The cost diary developed for this study is an adapted version of the MTA Medical Cost Questionnaire (MCQ) and MTA Productivity Cost Questionnaire (PCQ). The Dutch manual for costing research will be used to determine prices for each volume of resource use.

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Besides using the water displacement method, volume measurement will also be done by arm circumference measurement.
measurement using tape. Both arms will be measured during every visit at the level of the olecranon, 5 and 10 cm proximally, 5 and 10 cm distally, at the level of the wrist, and the dorsum of the hand.

In the outpatient clinic, a fluorescent marker, called indocyanine green (ICG) is injected intracutaneously into the second and fourth finger webspaces of the lymphoedematous limb and a so-called ICG lymphography is performed in search for viable lymphatic vessels. This is a technique using near-infrared fluorescence imaging (NIRF). After 0.05 mL of ICG (5 mg/mL) is injected per webspace, a near-infrared camera is used to visualise the lymphatic vessels. Proximal to the injection sites fluorescent stains are identified. When using the images as a guide, the lymphatic pathways and the sites for incisions for lymphaticovenous anastomoses are marked with a pen and a colour picture is taken. These colour pictures are used to identify the location when LVA will be performed. NIRF will be done at introduction visit and after 12 and 24 months.

Data analysis
For the HRQoL a paired Student’s t-test will be used to evaluate the changes in quality of life scores and in limb volume measurements between preinclusion and the different postinclusion periods of time within individuals from the same study group. For each of the follow-up moments (3, 6, 12, 18 and 24 months) the change in quality of life from baseline will be compared between groups using the two sample unpaired t-test, to evaluate short-term and long-term treatment effects. If baseline imbalance is present, assessed qualitatively, adjusted differences per follow-up moment will be computed using linear regression. In addition to statistical testing per follow-up measurement, a linear mixed-effects model will be used to test for an overall difference between the two groups. To account for clustering of measurements at the patient-level, a model with a random intercept and random slope will be used.

Economic evaluation
An economic evaluation will be performed alongside the clinical trial to determine the cost-effectiveness of LVA compared to CDT. The design of the economic evaluation follows the principles of a cost–utility analysis and adheres to the Dutch guideline for economic evaluations in healthcare and the Dutch manual for costing research.\(^{30,31}\) Outcome measures for the economic evaluations will be costs, HRQoL and QALYs. The trial-based evaluation adopts a societal perspective and has a time horizon of 2 years.

An incremental cost-effectiveness ratio (ICER), that is, cost per QALY gained, will be calculated by dividing the difference in costs between the two treatments with the difference in QALYs. Bootstrapping techniques will be used to summarise the uncertainty in estimates of incremental costs, effects and the ICER. In addition, cost-effectiveness acceptability curves will show the probability that LVA is cost-effective compared to conservative treatment, given the observed data, for a range of maximum monetary values that a decision-maker might be willing to pay for a QALY gained.

The impact of uncertainty surrounding deterministic parameters (eg, prices) on the ICER will be explored using one-way sensitivity analyses. Results, presented in a tornado diagram, can help determine which parameters are key drivers of the cost-effectiveness results. Predetermined subgroup analyses will address possible variation between patients (heterogeneity).

Missing values will be imputed using mean substitution or multiple imputation, as appropriate.

ETHICS AND DISSEMINATION
Data monitoring
Data will be handled confidentially. Source data will be stored by the investigator in a locked place. Data of all measurements during follow-up moments, (serious) adverse events (AEs) and digital questionnaires including patient cost diary are stored immediately in the online database of CASTOR EDC. The investigator and project leader only have access to this database with an account with password. Identifying data will be stored in coded form; the key to the form is known only to the supervisor, the investigator, the Dutch Health Care Inspectorate, the study monitors and the members of the review committee.

Harms
AEs are defined as any undesirable experience occurring to a subject during the study, whether or not considered related to the trial procedure. AEs related to the LVA operation or conservative therapy that have a possible impact on the lymphoedema and reported spontaneously by the subject or observed by the investigator or his staff will be recorded directly in CASTOR EDC.

The research team will report the serious AEs (SAEs) through the web portal ToetsingOnline to the accredited METC that approved the protocol, within 7 days of first knowledge for SAEs that result in death or are life threatening followed by a period of maximum of 8 days to complete the initial preliminary report. All other SAEs will be reported within a period of maximum 15 days after the research team has first knowledge of the SAEs.

Auditing
Monitoring of the conduct of the study will be done by the Clinical Trial Center Maastricht on a frequent basis following their protocol as is requested by the Board.

Protocol amendments
Any modifications to the protocol which may impact the study will be notified to the METC that gave a favourable opinion prior to implementation.

Patient and public involvement
The Dutch Network for Lymphedema and Lipedema, and the Patient Advocacy Group, a joint initiative from Wolfs J, et al. BMJ Open 2020;10:e035337. doi:10.1136/bmjopen-2019-035337

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the Breast Cancer Research Group of the Dutch breast cancer association, were consulted. They provided feedback from the patients’ perspective on our research protocol, patient participation and implementation plan, feasibility, patient information sheet, outcome parameters and the burden for the patients.

Ethical considerations
This study will be conducted according to the principles of the Declaration of Helsinki, recently changed in Fortaleza (2013) and in accordance with the Medical Research Involving Human Subjects Act (WMO).

DISSEMINATION
The results of this study will be disseminated in presentations at academic conferences, publications in peer-reviewed journals and other news media. Data will be kept confidential and will not be shared with the public. Requests for data sharing for appropriate research purposes will be considered on an individual basis after trial completion and after publication of primary manuscripts.

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Contributors
JB, AAPdG, EH, DU, RvdH and SSQ conceived the study and initiated the study design. MK provided statistical and cost-effectiveness expertise. JW, XK, HT, DU, RvdH and SSQ completed the study design and protocol. All authors contributed to refinement of the study protocol and approved the final version.

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Competing interests
None declared.

Patient consent for publication
Not required.

Ethics approval
The study was approved by the Ethics Committee of Maastricht University Medical Center on 19 December 2018 (NL67059.068.18).

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