Gamma probe and ultrasound guided fine needle aspiration cytology of the sentinel node (GULF) trial - overview of the literature, pilot and study protocol

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Abstract

Background: Sentinel node (SN) biopsy (SNB) detects clinically occult metastases of breast cancer and melanoma in 20–30%. Wound infections, seroma and lymph edema occur in up to 10%. Targeted ultrasound (US) of the SN, (with fine needle aspiration cytology (FNAC) if appropriate) has been investigated as a minimally invasive alternative, but reported sensitivity rates are too low to replace SNB. Our hypothesis is that the use of a handheld gamma probe concomitant with US may improve sensitivity.

Our aim is to provide an overview of the current literature on preoperative nodal staging of clinical N0 melanoma patients, report on a pilot, and present a study protocol for a minimally invasive alternative to the SNB: Gamma probe and Ultrasound guided Fine needle aspiration cytology of the sentinel node (GULF trial).

Methods: The GULF trial is a multicenter open single arm observational trial. Newly diagnosed cT1b-4N0M0 cutaneous melanoma or cT1-3N0M0 breast cancer patients, aged >18 years, presenting for SNB are eligible. 120 patients will be included for preoperative targeted gamma probe guided US and FNAC of the SN. Afterwards all patients proceed to surgical SNB. Primary endpoint is the sensitivity of FNAC. Secondary endpoints include SN identification rate and the histopathological compatibility of Core Needle Biopsy and FNAC vs. SNB. Secondary endpoints were investigated in a pilot with 10 FNACs and marker placements, and 10 FNACs combined with Core Needle Biopsy.

Results: A pilot in 20 patients showed that SN identification rate was 90%, supporting the feasibility of this technique.

Discussion: There is broad experience with US (in combination with FNAC) prior to SNB, but sensitivity and specificity are too low to completely abandon SNB. Promising alternative techniques potentially will replace SNB in the future but more evidence is needed in the form of prospective studies. Accurate identification of the SN for US-FNAC has been proven feasible in our pilot. When adequate sensitivity can be reached, US-FNAC provides a minimally invasive alternative for the surgical SNB procedure.

Trial registration: The GULF trial is registered in the Netherlands Trial Registry (NTR), ID: NRT5193. May 1st 2015.

Keywords: Sentinel lymph node biopsy, Ultrasonography, Fine-needle biopsy, Melanoma, Breast Neoplasms, Minimally invasive surgical procedures
Background
Sentinel node biopsy
With the introduction of sentinel node (SN) biopsy (SNB) as a less invasive alternative to elective lymph node dissection for melanoma and breast cancer with clinically negative lymph nodes, this has become the gold standard for adequate staging. Although less invasive than an elective lymph node dissection, SNB is still associated with some potential morbidity. Morbidity occurs in up to 10% of patients; wound infections and seroma are the most frequently seen complications [1, 2]. Rarely lymph edema is seen after SNB. Around 70–80% of SNB’s are tumor negative after histological assessment, these patients cannot benefit from the SNB procedure. In that light the morbidity of a surgical SNB procedure is deemed considerable, and any less invasive procedure, if accurate enough, would be preferred.

The detection rate of submicrometastases has increased considerably in the past decades; adaptation of the melanoma and breast cancer SN sectioning protocols and use of standard immunohistochemistry staining enabled pathologists to detect even the smallest tumor deposits accurately [3–8]. This has clear clinical implications; more patients are diagnosed as SN positive and will be offered a completion lymphadenectomy (CLND) [6, 8]. It is questionable whether this morbid surgical procedure is justified in cases with minimal SN tumor burden [9], as several retrospective melanoma studies and recently the prospective DeCOG study have shown that survival for this group of melanoma patients is similar to SN negative patients [10–13]. In breast cancer, presence of isolated tumor cells (≤0.2 mm) or micrometastases (>0.2 ≤ 2.0 mm) is associated with a slightly worse prognosis [7, 14], but its clinical relevance is debated as well [15–17], and CLND is omitted in certain groups of patients with a positive SN [18].

Prospective studies currently investigating the therapeutic value of CLND in melanoma are the EORTC-1208MG (Minitub) [9], including patients with minimal SN tumor burden only, and the MSLT2, which included all SN positive patients [19]. Parallel to this, certain adjuvant therapy trials (EORTC 18071, EORTC 1325, Combi-AD) recruit stage IIIA patients only in case of ≥1 mm SN tumor burden [20–22].

Primary results from the EORTC 18071 show that SN positive patients benefit the most from adjuvant treatment measured as recurrence free survival at 3 years [21, 22]. In this light, it remains worthwhile to keep selecting patients for adjuvant therapy in trial setting and/or CLND based on nodal staging, and a cut-off for detection of (sub) micrometastases (<1.0 mm) may aid in prevention of overtreatment in low risk patients.

Ultrasound guided fine needle aspiration cytology
Ultrasound (US) guided fine needle aspiration cytology (FNAC) or core needle biopsy (CNB) may provide a good minimally invasive alternative to SNB. In breast cancer patients screening US of the regional lymph node basin is part of the preoperative staging process; this way up to 17% of patients undergo axillary lymph node dissection immediately and are spared a SNB [23, 24]. Melanoma patients do not routinely undergo a preoperative US of the regional lymph node basin, due to previously reported poor identification rates of occult lymph node metastases with US and FNAC [25, 26]. Several studies have been conducted in order to analyze if US (with FNAC or CNB) can replace SNB, but thus far reported sensitivity rates for US vary considerably, ranging between 9 and 94% [27, 28].

For the current study we will focus on melanoma, as the therapeutic value of both SNB and CLND are debated, and alternatives for SNB are more limited for this type of cancer.

In our search for a reliable and accurate minimally invasive alternative to SNB for staging of clinical N0 melanoma patients, we examined the current available literature and performed a systematic search of all major databases to explore whether other methods than US guided FNAC may have proven adequate alternatives to SNB.

Literature overview
All relevant studies on US imaging of regional lymph nodes in melanoma patients scheduled for SNB are displayed in Table 1 (search details are given in Additional file 1). Some of the studies mentioned in Table 1 are overlapping; the studies from Voit et al. [29–33] concern the same database with more inclusions over time. In the studies that performed US prior to lymphoscintigraphy, sensitivity rates were low, ranging from 4.7% to 39%, and specificity rates were high, ranging from 86% to 100%. Two studies did not mention the exact timing of US in relation to lymphoscintigraphy; Hocevar et al. reached a sensitivity and specificity of 71% and 84%, and Testori et al. reached a sensitivity of 94% and 90%. In the studies that performed a targeted US (i.e. US of the marked “SN” area on the skin after lymphoscintigraphy), sensitivity ranged from 22% to 100%, and specificity ranged from 62% to 100% (Table 1).

Besides US and targeted US with FNAC prior to SNB, several groups have focused on development of new imaging techniques for examination of the SN/lymph nodes and detection of SN tumor deposits, such as sonoelastography [34, 35], contrast enhanced US [36], and multispectral optoacoustic tomography (MSOT) [37] (Table 2). Sonoelastography measures tissue consistency; which can be visualized on top of US
| Author, year       | Study Design, monocentric | N     | US Setting     | FNAC/other technique              | Sens (%) | Spec (%) | PPV (%) | NPV (%) |
|-------------------|---------------------------|-------|----------------|-----------------------------------|----------|----------|---------|---------|
| Rossi [61], 2000  | Not mentioned             | 69    | Pre-lympho     | No                                | 33       | 100      | 100     | 86      |
| Rossi [25], 2003  | Prospective, monocentric | 125   | Pre-lympho     | FNAC if US suspicious             | US alone: 39 | 100      | 100     | 85      |
| Hocevar [62], 2004| Prospective, monocentric | 57    | unknown        | FNAC if US suspicious             | US alone: 71 | 84      | 59      | 90      |
| Testori [63], 2005| Retrospective, monocentric| 88    | unknown        | No                                | 94       | 90       | 64      | 99      |
| Starritt [64], 2005| Prospective, monocentric | 31    | all SN +       | No                                | NA       | NA       | NA      | NA      |
| Voit [65], 2006   | Prospective, monocentric | 127   | Post-lympho    | FNAC if US suspicious             | US alone: 79 | 72      | 53      | 90      |
| Van Rijk [26], 2006| Retrospective, monocentric| 107   | Pre-lympho     | FNAC if US suspicious             | US alone: 34 | 87      | -       | -       |
| Sibon [66], 2007  | Prospective, monocentric | 131   | Pre-lympho     | No                                | 9        | 96       | 43      | -       |
| Kunte [67], 2009  | Prospective, monocentric | 25    | Pre- and post-lympho | No                        | 33       | 100      | 100     | 88      |
| Voit [30], 2009   | Prospective, monocentric | 400   | Post-lympho    | FNAC if US suspicious             | 65       | 99       | 93      | 92      |
| Sanki [60], 2009  | Prospective, monocentric | 716   | Post-lympho    | No                                | 33       | 97       | 60      | 88      |
| De Giorgi [36], 2010| Prospective monocentric | 15    | Post-lympho    | Standard CEUS                     | CEUS: 100 | 62      | 55      | 100     |
| Voit [31], 2010   | Prospective monocentric | 400   | Post lympho    | FNAC if US suspicious             | All Berlin criteria combined: 82 | 80      | 52      | 94      |
| Hinz [58], 2011   | Prospective monocentric | 81    | Pre and post lympho | No                        | 22       | 100      | 100     | 96      |
| Chai [68], 2012   | Retrospective monocentric| 325   | Pre-lympho     | FNAC if US suspicious             | 34       | 86       | 37      | 84      |
| Marone [69], 2012 | Prospective monocentric | 623   | Pre-lympho     | No                                | 15       | 100      | 100     | 87      |
| Pilko [70], 2012  | Retrospective Monocentric| 405   | Pre-lympho     | FNAC if US suspicious             | Not mentioned | -     | -       | -       |
| Stoffels [71], 2012| Retrospective Monocentric| 221   | Pre-lympho     | FNAC if US suspicious             | 14       | 97       | 100     | 97      |
| Hinz [72], 2013   | Retrospective Monocentric| 20    | Pre-lympho & pre PET-CT | No if US malig, Direct LND | 12       | 100      | 100     | 74      |
| Ulrich [73], 2014 | Prospective monocentric | 800   | Post lympho    | FNAC if US suspicious             | US-FNAC: 56 | 99      | 92      | 89      |
| Voit [32], 2014   | Prospective monocentric | 1000  | Post lympho    | FNAC if US suspicious             | US alone: 71 | -      | -       | -       |
| Voit [33], 2016   | Prospective monocentric | 1000  | Post lympho    | FNAC if US suspicious             | US alone: 71 | -      | -       | -       |

Overview of studies reporting on ultrasound imaging of regional lymph nodes in melanoma patients prior to sentinel node biopsy. Abbreviations: US ultrasound, FNAC fine needle aspiration cytology, CEUS contrast enhanced ultrasound, Sens sensitivity, Spec specificity, PPV positive predictive value, NPV negative predictive value, lympho lymphoscintigraphy, NA not applicable.
images using different color shades; red indicating soft tissue, and blue indicating rigid tissue [34, 35]. As metastases tend to be more solid than normal lymph node tissue regions of interest for FNAC can be identified. For contrast enhanced US an intravenous contrast agent is applied to detect possible areas of hyperperfusion or hypoperfusion; indicating potential metastatic lesions [36]. These techniques reached a high sensitivity for identification of SN metastases (Table 2).

Two recently developed techniques for improved SN identification peri-operatively are SPECT-US [38, 39], and near infrared light fluorescence imaging [40–44]. SPECT-US displays the location of a radio-active SN in the US images; making it easier for the surgeon to locate SN’s in anatomically challenging area’s such as the cervical and occipital area; or to guide radiologists for FNAC [38, 39]. Near infrared light fluorescence imaging is conducted with Indocyanine green as tracer, which can be combined with 99Tc nano-colloids to form a hybrid tracer [41]. Intraoperative identification is similarly accurate to 99Tc-colloid, and particularly helpful for SN localization in the cervical and occipital area, where overprojection from the 99Tc-colloid injection site is a common obstacle. Preoperative (transcutaneous) SN identification has reached lower identification rates, due to the limited penetration depth of the fluorescent tracer [45].

Summarizing, few US imaging studies have used a method to accurately identify the SN prior to US examination and FNAC. This may have contributed to lower than expected sensitivity rates for detection of SN metastases in studies where this was not applied. It explains why to date no alternative method for SN staging has been adopted in daily clinical practice, and the need for such a method remains.

Rationale for a new trial

SN identification

To overcome the problem of suboptimal identification of the SN, we hypothesize that use of a handheld gamma probe (Geiger teller) to detect the SN post lymphoscintigraphy may further aid the radiologist in accurately identifying the SN for ultrasound guided FNAC. Several pilot studies have been performed using this technique in breast cancer patients; correct localization of the SN occurred in 75% - 100% [46–49]. This formed the rationale for the GULF Trial (Gamma probe and Ultrasound guided Fine needle aspiration cytology of the sentinel node).

Cytology or histology?

In order to reach the sample size needed for proof of concept with accurate power and within an acceptable term, both melanoma patients and breast cancer patients will be included in the GULF trial. The SN procedure is uniformly applied for both melanoma and breast cancer, and breast cancer patients may equally benefit from a minimally invasive alternative for the SN. All patients will undergo FNAC. Since metastatic size may have clinical implications for breast cancer patients [18], a subset of 10 breast cancer patients will undergo CNB additionally after FNAC. This allows for a comparison of results between CNB, FNAC and SNB.

Hypotheses GULF trial

We hypothesize that a sensitivity of 90% with a 95% confidence interval of 80% - 100% is achievable. Secondly, we expect that a SN identification rate of more than 75% is feasible.

Study aims

- To present a study protocol for a minimally invasive alternative to the sentinel node biopsy (GULF trial), with as primary objective to determine whether an acceptable sensitivity for US and gamma probe guided FNAC can be achieved.
- Secondary objective is 1) the identification rate of the SN and 2) the histological results of CNB versus FNAC and versus SNB.

Prior to starting the GULF-trial, we had to prove the concept of adequate identification of the SN. A pilot

| Author, yr | Topic | No. of patients | Technique | Sens (%) | Spec (%) |
|------------|-------|----------------|-----------|----------|---------|
| Hinz [72], 2013 | Elastography | 36 | US + power Doppler: | 81 | 76 |
| | | | Elastography: | 91 | 76 |
| | | | Combined: | 95 | 76 |
| Ogata [35], 2014 | Elastography | 12 | US: | 77 | 57 |
| | | | Elastography: | 100 | 71 |
| Stoffels [37], 2015 | MSOT and indocyanin green | 20 | MSOT | 100 | 48.6 |

Overview of pilot studies investigating non-invasive detection of melanoma lymph node metastases

Abbreviations: yr. year, Sens sensitivity, Spec specificity, US ultrasound, MSOT multispectral optoacoustic tomography
study focusing on the adequate detection rate of the SN was conducted.

Pilot
After approval of the Ethical Review Board a pilot was performed in 20 patients presenting at the Erasmus MC Cancer Institute. All patients underwent gamma probe guided US-FNAC after written informed consent. In the 10 first melanoma patients additional metallic marker placement (O-Twist-Marker, BIP) was performed after local infiltration of the skin and surrounding tissue with 1-10 mL lidocaine 2%. Correct identification of the SN was assessed by examining the excised SN (s) on presence of the marker. Separately, in the first 10 breast cancer patients CNB was performed after FNAC with a 14G needle, after local infiltration similar to marker placement. CNB was done for assessment of concordance with FNAC results and to detect potential superiority of either technique.

All patients proceeded to OR for SNB, which was performed according to the triple technique: preoperative 99Tc lymphoscintigraphy <24 h prior to surgery, intradermal injection of patent blue near the primary tumor site prior to first incision, and peroperative use of a handheld gamma-probe to locate SN (s) [50, 51]. Lymph nodes were considered SN when radioactive and/or blue. A marker was retrieved from the SN in 9 out of 10 patients; which meant the SN identification rate was 90%. CNB samples were investigated on presence of lymphoid tissue. This was present in 6 out of 10 patients. 40% of CNBs was not representative. In comparison: FNAC color staining was representative in 19 out of 20 patients (95%), and FNAC immunohistochemistry staining was representative in 14 out of 20 patients (70%).

During the pilot study no safety issues occurred. In the second enrolled study patient none of the 2 placed markers were found at histopathological examination of the SN and in another patient only 1 of 2 placed markers was found. A detailed shoulder X-ray confirmed the markers were still in situ in both patients. In the latter patient the X-ray images were suggestive of marker displacement towards mamma tissue; this was probably due to intraoperative displacement of the marker during SN removal.

Considering the positive results from this pilot, the study will be continued with an expansion of the pilot population in order to reach a sufficient sample size according to the presented study protocol.

Methods
GULF design
Patients with a newly diagnosed cT1b-4N0M0 cutaneous melanoma or cT1-3N0M0 breast cancer presenting at the outpatient clinic of the Erasmus MC Cancer Institute, and the Netherlands Cancer Institute – Antoni van Leeuwenhoek (only melanomas) will be assessed for inclusion. All patients will undergo US and gamma probe guided FNAC of the SN. The pilot patients received additional marker placement (n = 10) for identification purposes, or additional CNB (n = 10) for assessment of potential benefit of CNB (i.e. histology and size measurement possible) (Fig. 1).

Study population
Inclusion criteria
Age ≥ 18 years, new diagnosis of cT1b-4N0M0 cutaneous melanoma or cT1-3N0M0 breast cancer.

Prior to start of any study related procedure, written informed consent must be given according to ICH/GCP and national legislation.

Exclusion criteria
Clinically suspect lymph node, other known malignancy with potential to disseminate to axillary or groin lymph
node basins, prior lymph node biopsy, no SN visible at lymphoscintigraphy/not identifiable with gamma probe.

Study procedures

US-FNAC
All patients will be admitted to the surgical ward on the day of surgery. First, a lymphoscintigram <24 h prior to SNB will be performed, as is standard procedure. Following successful lymphoscintigraphy (i.e. SN(s) is/are visible) the study procedures can start. A dedicated radiologist will perform US imaging of the lymph node basin where a SN or multiple SNs was/were identified by lymphoscintigraphy. The exact location of the SN (s) will be determined using a handheld gamma probe, and by combination with US; the assumed SN (s) will be visualized (being a visible lymph node at the center of the hotspot found with the gamma probe). FNAC will be performed of all visualized assumed SN (s). In case of multiple SNs in one lymph node basin or multiple draining lymph node basins with an SN in every basin (for instance a melanoma on the back draining to both axilla and groin), FNAC will be performed of all lymph nodes pointed out as primary tier SN by the nuclear medicine specialist. For FNAC 1–4 cortical samples will be taken. Whenever additional clearly suspect lymph nodes are visible, the radiologist will perform FNAC from these nodes as well, as is standard of care. All samples will be transported to the pathology lab for analysis. US findings will be recorded according to the Berlin morphologic Criteria to create uniformity in recording per center [30]. After FNAC, all patients will proceed to the operating room for SNB according to standard procedure (as described in the Pilot section). Lymph nodes were considered SN when radioactive and/or blue. No diagnostic procedure or treatment is postponed or elongated. No additional visits to the outpatient clinic are required.

Endpoints

Primary endpoint: Primary outcome is the sensitivity of gamma probe and US guided CNB or FNAC.

Secondary endpoints: Secondary outcome is 1) the identification rate of the SN 2) the histological results of CNB versus FNAC and versus SNB.

Ad 1) an identification rate of at least 75% is deemed acceptable (concordant with literature). This has been proven feasible in the pilot study.

Statistical considerations

Sample size and accrual
Based on retrospective data, the prevalence of metastatic SNs is expected to be 30%. Our gold standard is the histological outcome of SNB (absence or presence and size of metastases in the SN). Submicrometastases (i.e. <0.1 mm at any site or 0.4 mm subcapsular) in melanoma patients, and isolated tumor cells (i.e. ≤0.2 mm) in breast cancer patients will be considered negative: a negative FNAC is accepted in these cases. Based on previous reports, we expect to find around 10% of these submicrometastases and isolated tumor cells in both melanoma and breast cancer patients [52, 53]. Considering this, the maximum achievable sensitivity of FNAC will be 90%. For this sensitivity, and a 95% confidence interval of 80–100% (With a two-sided significance level $\alpha = 0.05$ and power $1 – \beta = 0.8$), the required sample size is 116 considering a 30% prevalence of metastatic SNs. Around 3% of patients are expected to have a negative lymphoscintigram: the sample size will be increased to 120 patients.

With an average accrual rate of 60 patients per year, maximum accrual will be met at 2 years post start of study.

Statistical analysis plan

The main analysis addressing the primary endpoint will be performed after inclusion of all 120 patients. No interim analysis is planned for this endpoint.

Ethical considerations

This study has been approved by the Erasmus MC medical-ethical committee. The study will be conducted according to the principles of the Declaration of Helsinki and in accordance with national and regional legislation, guidelines, regulations and acts.

Discussion

Currently SNB is the most important staging procedure for clinically N0 melanoma patients, especially in the light of trial participation for adjuvant therapies based on N-status [20, 21, 54]. The therapeutic role of SNB for melanoma is still under debate [55–57]. Considering the fact that this is a surgical staging procedure associated with complications in up to 10% of patients, our group sought to investigate a more minimally invasive alternative.

The ongoing improvement of imaging techniques (i.e. more accurate and detailed US imaging) and increased experience with FNAC renders combined US-FNAC as a high potential minimally invasive alternative for surgical SNB [31, 32]. Correct transcutaneous identification of the SN forms the main obstacle for broad application of this technique as this is key in obtaining reliable FNAC.

The current study aims to give an overview of the current melanoma literature, report a pilot and present a study protocol for a minimally invasive technique to investigate the SN using gamma probe guided US-FNAC.
Overview of the literature
The studies presented provide evidence that it is difficult to detect clinically occult lymph node metastases in melanoma patients, and although some studies have achieved high sensitivity and specificity rates, these results have not been reproduced by other groups. There are many differences between the reported studies; namely retrospective vs. prospective study setting; US prior to lymphoscintigraphy vs. targeted US after lymphoscintigraphy; the number of persons performing US and their expertise; variation in US morphology criteria used to discriminate between benign and suspicious or malignant lymph nodes; and use of FNAC or not. All these factors will have contributed to the outcome of these studies. It is interesting to see that sensitivity rates are low in the studies that performed an US of the entire lymph node basin without knowing the location of the SN(s), but that even in the studies were targeted US of the SN area was applied, sensitivity rates could be as low as 22% [58] and as high as 82% [31] or even a perfect 100% [36] as well. Thompson et al. proposed a possible explanation for these disparate results; many of the micrometastases present in SNs are too small to be detected by the US-equipment used [59, 60]. However, Voit et al. demonstrated that it was possible to successfully perform a FNAC in a lesion as small as 0.4 mm. Nevertheless, most smaller SN metastases will be overlooked by US and/or missed by FNAC. The question is if this has any clinical implications.

As long as US-imaging is limited by a detection limit, and alternative imaging techniques are tested in pilot settings, the need for a reliable, minimal invasive easy to perform and replicate method to assess SN status remains. Hence the presentation of the GULF trial study protocol here.

Pilot
Our pilot results show that correct identification of the SN for FNAC was possible in 90%, and that the sampled material was representative in 95% of FNAC samples. CNB was representative in only 60%. This confirms that the described technique for targeted US-FNAC of the SN is feasible. CNB will not be added to the study procedure considering the low rate of representative tissue in the pilot phase.

If an acceptable sensitivity can be achieved for FNAC, patients can proceed to undergo radical lymph node dissection immediately in case of positive FNAC, bypassing the SNB procedure. When the FNAC sample is negative, surgeons can choose to perform a SNB or continue with only surgical excision of the primary tumor and monitoring of potential lymph node involvement at follow up visits. This way up to 80% of patients eligible for SNB can be spared this invasive procedure and the risk of morbidity related to this procedure. Furthermore, for melanoma patients this would mean that general anesthesia is no longer needed, as WLE can be performed under local anesthesia. Ultimately operative nodal staging may become completely obsolete.

Conclusions
The literature on pre-operative assessment of regional lymph nodes with US in clinically N0 melanoma patients is disparate. Targeted US of the SN area in combination with FNAC or other new techniques has potential to become a minimally invasive alternative for the SNB, however, findings need to be replicated in prospective clinical trials first. A pilot with gamma probe guided US-FNAC show that accurate SN identification in up to 90% of patients is feasible. Our group presents a study protocol of the Gamma probe and Ultrason sound guided Fine needle aspiration cytology of the sentinel node Trial (GULF trial) as a potential improvement to the reported US-FNAC techniques and ultimately even a possible replacement of the SNB.

Additional file

Additional file 1: Data gives an overview of the number of found articles per literature database, and specific entry terms are provided. (PDF 14 kb)

Abbreviations
CLND: Completion lymphadenectomy; CNB: Core needle biopsy; FNAC: Fine needle aspiration cytology; GULF: Gamma probe and ultrasound guided fine needle aspiration cytology of the sentinel node trial; ICH-GCP: International Conference on Harmonisation - Good Clinical Practice; MSOT: Multispectral optoacoustic tomography; SN: Sentinel node; SNB: Sentinel node biopsy; SPECT: Single photon emission computed tomography; US: Ultrasound

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Availability of data and material
The pilot and literature datasets analyzed during the current study are available from the corresponding author on reasonable request.

Authors’ contributions
DG, LK, COO, AvA and CV conceptualized the study design. All authors developed the study protocol and are responsible for implementation of the study. COO, LK, DG and CG are responsible for inclusion of patients for the pilot study. CdM is responsible for performance of the intervention. SK and CoO are responsible for analysis of study material. COO was responsible for drafting the manuscript and all authors contributed to the revision of the manuscript. All authors have read and approved the final version of the manuscript.
Competing interests
The authors declare that they have no competing interests.

Consent for publication
Informed consent was obtained for all participants.

Ethics approval and consent to participate
The ethics committee of the Erasmus Medical Center has approved this study (committee’s reference number: 078), informed consent was obtained for all participants.

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References
1. Wrightson WR, Wong SL, Edwards MJ, et al. Complications associated with sentinel lymph node biopsy for melanoma. Ann Surg Oncol Jul 2003; 10(6): 676-680.
2. Biver-Dalle C, Puzenat E, Puyraveau M, et al. Sentinel lymph node biopsy in melanoma: our 8-year clinical experience in a single French institute (2002-2009). BMC Dermatol. 2012;1:22.
3. Cook MG, Green MA, Anderson B, et al. The development of optimal pathological assessment of sentinel lymph nodes for melanoma. J Pathol Jun 2003; 200(3):314-319.
4. Chakera AH, Heese B, Burak Z, et al. EANM-EORTC general recommendations for sentinel node diagnostics in melanoma. Eur J Nucl Med Mol Imaging Oct 2009; 36(10):1713-1742.
5. Garbe C, Peris K, Hauschild A, et al. Diagnosis and treatment of melanoma. European consensus-based interdisciplinary guideline - update 2016. Eur J Cancer. Jun 28 2016;63:201-217.
6. van Akkooi AC, de Vilt WH, Verhoef C, Graveland WJ, van Geel AN, Kliffen M, Eggermont AM. High positive sentinel node identification rate by EORTC melanoma group protocol. Prognostic indicators of metastatic patterns after sentinel node biopsy in melanoma. Eur J Cancer Feb 2006;42(3):372-380.
7. Weaver DL, Ashikaga T, Krag DN, et al. Effect of occult metastases on survival in node-negative breast cancer. N Engl J Med. Feb 3 2011;364(5):412-421.
8. Weaver DL. Pathology evaluation of sentinel lymph nodes in breast cancer: protocol recommendations and rationale. Mod Pathol May 2010;23 Suppl 2: S26-S32.
9. van der Ploeg AP, van Akkooi AC, Verhoef C, Eggermont AM. Completion lymph node dissection after a positive sentinel node: no longer a must? Curr Opin Oncol Mar 2013;25(2):152-159.
10. van der Ploeg AP, van Akkooi AC, Schmitz PI, Koljenovic S, Verhoef C, Eggermont AM. EORTC melanoma group sentinel node protocol identifies high rate of submicrometastases according to Rotterdam criteria. Eur J Cancer Sep 2010;46(13):2414-2421.
11. van Akkooi AC, de Vilt WH, Verhoef C, Eggermont AM. Isolated tumor cells and long-term prognosis of patients with melanoma. Ann Surg Oncol May 2008;15(5):1547-1548.
12. de Vilt WH, van Akkooi AC, Verhoef C, Eggermont AM. Detection of melanoma micrometastases in sentinel nodes - the cons. Surg Oncol Sep 2008;17(3):175-181.
13. Leiter U, Stadler R, Mauch C, et al. Complete lymph node dissection versus no dissection in patients with sentinel lymph node biopsy positive melanoma (DeCDG-SLT): a multicentre, randomised, phase 3 trial. Lancet Oncol. Jun 2016;17(6):757-767.
14. de Boer M, van Deurzen CH, van Dijk JA, et al. Micrometastases or isolated tumor cells and the outcome of breast cancer. N Engl J Med. Aug 13 2009;361(7):653-663.
15. Hansen NM, Grube B, Ye X, Turner RR, Brenner RJ, Sim MS, Giuliano AE. Impact of micrometastases in the sentinel node of patients with invasive breast cancer. J Clin Oncol. Oct 1 2009;27(28):4679-4684.
16. Meittini I, Desideri I, Saleva C, et al. Impact of sentinel node tumor burden on outcome of invasive breast cancer patients. Eur J Surg Oncol Oct 2014; 40(10):1195-1202.
17. van der Heiden-van der Loo M, Schaapveld M, Ho VK, Siesling S, Rutgers EJ, Peeters PH. Outcomes of a population-based series of early breast cancer patients with micrometastases and isolated tumour cells in axillary lymph nodes. Ann Oncol Nov 2013; 24(11):2794-2801.
18. Mammacarcinoom W. Oncoline, Mammacarcinoom 2.0, Landelijke Richtlijn. 2012; http://www.oncoline.nl/mammacarcinoom. Accessed 15 Feb 2016.
19. Morton DL. Overview and update of the phase III multicenter selective Lymphadenectomy trials (MSLT-1 and MSLT-II) in melanoma. Clin Exp Metastasis Oct 2012;29(7):699-706.
20. Eggermont AM, Maio M, Robert C. Immune checkpoint inhibitors in melanoma provide the cornerstones for curative therapies. Semin Oncol Jun 2015;42(3):429-435.
21. Eggermont AM, Chiarion-Sileni V, Grob JJ, et al. Adjuvant ipilimumab versus placebo after complete resection of high-risk stage III melanoma (EORTC 18071): a randomised, double-blind, phase 3 trial. Lancet Oncol May 2015; 16(5):522-530.
22. Eggermont AM, Chiarion-Sileni V, Grob JJ. Correction to Lancet Oncol 2015; 16:522-30. Adjuvant ipilimumab versus placebo after complete resection of high-risk stage III melanoma (EORTC 18071): a randomised, double-blind, phase 3 trial. Lancet Oncol. Jun 2015; 16(6):e262.
23. Housami N, Ciatto S, Turner RM, Cody HS, 3rd, Macaskill P. Preoperative ultrasonoguided needle biopsy of axillary nodes in invasive breast cancer: meta-analysis of its accuracy and utility in staging the axilla. Ann Surg Aug 2011;254(2):243-251.
24. de Kanter AJ, van Eijck CH, van Geel AN, et al. Multicentre study of ultrasonographically guided auxiliary node biopsy in patients with breast cancer. Br J Surg Nov 1999;86(11):1459-1462.
25. Rossi CR, Mocellin S, Sgabetti N, et al. The role of preoperative ultrasonographic scan in detecting lymph node metastasis before sentinel node biopsy in melanoma patients. J Surg Oncol Jun 2003;83(2):280-84.
26. van Rijk MC, Teertstra HJ, Petersen JL, Nieweg OE, Dlimos RA, Hoefnagel CA, Kroon BB. Ultrasonography and fine-needle aspiration cytology in the preoperative evaluation of melanoma patients eligible for sentinel node biopsy. Ann Surg Oncol Nov 2006;13(1):1511-1516.
27. Catalano O. Critical analysis of the ultrasonographic criteria for diagnosing lymph node metastasis in patients with cutaneous melanoma: a systematic review. J Ultrasound Med. 2011;30(4):547-60.
28. Xing Y, Bronstein Y, Ross MI, et al. Contemporary diagnostic imaging modalities for the staging and surveillance of melanoma patients: a meta-analysis. J Natl Cancer Inst. Jan 19 2011;103(2):129-142.
29. Voit C, Kron M, Schafer G, et al. Ultrasound-guided fine needle aspiration cytology prior to sentinel lymph node biopsy in melanoma patients. Ann Surg Oncol Dec 2006;13(12):1682-1689.
30. Voit CA, van Akkooi AC, Schafer-Hesterberg G, Schoengen A, Schmitz PI, Sterry W, Eggermont AM. Rotterdam criteria for sentinel node (SN) tumor burden and the accuracy of ultrasound (US)-guided fine-needle aspiration cytology (FNAC): can US-guided FNAC replace SN staging in patients with melanoma? J Clin Oncol. Oct 20 2009;27(30):4994-5000.
31. Voit C, Van Akkooi AC, Schafer-Hesterberg G, et al. Ultrasound morphology criteria predict metastatic disease of the sentinel nodes in patients with melanoma. J Clin Oncol. Feb 10 2010;28(5):847-852.
32. Voit CA, Gooskens SLM, Siegel P, et al. Ultrasound-guided fine needle aspiration cytology as an addendum to sentinel lymph node biopsy can perfect the staging strategy in melanoma patients. Eur J Cancer. 2014;50(13):2280-8.
33. Voit CA, Oude Ophuis CMC, Ulrich J, van Akkooi ACJ, Eggermont AMM. Ultrasound of the sentinel node in melanoma patients: echo-free island is a discriminatory morphologic feature for node positivity. Melanoma Res. 2016; 26(3):267-71.
34. Hinze T, Hoeller T, Wenzel J, Biebert T, Schmid-Wendtner MH. Real-time tissue elastography as promising diagnostic tool for diagnosis of lymph node metastases in patients with malignant melanoma: a prospective single-center experience. Dermatology. 2013;226(1):81-90.
35. Ogata D, Uematsu T, Yoshikawa S, Kyoohara Y. Accuracy of real-time ultrasound elastography in the differential diagnosis of lymph nodes in cutaneous malignant melanoma (CMM): a pilot study. Int J Clin Oncol Aug 2014;19(4):716-721.
36. De Gori V, Gore A, Grazzini M, et al. Contrast-enhanced ultrasound: a filter role in AJCC stage III melanoma patients. Oncology (Switzerland). 2010;70(5-6):370-5.
37. Stoffels I, Marschner S, Heflich H, et al. Metastatic status of sentinel lymph nodes in melanoma determined noninvasively with multispectral optoacoustic imaging. Sci Transl Med. 2015;7:317.
38. Freeny C, Winkens T, Oppester T, Elsepier P, Runnebaum I, Darr A. Real-time ultrasound and freehand-SPECT: experiences with sentinel lymph node mapping. Nuklearmedizin. 2014;53(6):259-64.
39. Bluemel C, Safak G, Cramer A, et al. Fusion of freehand SPECT and ultrasound: first experience in preoperative localization of sentinel lymph nodes. Eur J Nucl Med Mol Imaging. 2016;43(13):2304-2312.
40. Fujiwara M, Mizukami T, Suzuki A, Fukamizu H. Sentinel lymph node detection in skin cancer patients using real-time fluorescence navigation with indocyanine green: preliminary experience. J Plast Reconstr Aesthet Surg Oct 2009;62(10):e173-e178.
41. Brouwer OR, Buckle T, Vermeersen L, et al. Comparing the hybrid fluorescent-radioactive tracer indocyanine green-99mTc-nanocolloid with 99mTc-nanocolloid for sentinel node identification: a validation study using lymphoscintigraphy and SPECT/CT. J Nucl Med Jul 2012;53(7):1034-1040.
42. Gillmore DM, Khullar OV, Gioux S, Stockdale A, Frangioni JV, Colson YL, Russell SE. Effective low-dose escalation of indocyanine green for near-infrared fluorescent sentinel lymph node mapping in melanoma. Ann Surg Oncol. Jul 2013;20(7):2357-2363.
43. Floyd JM, Wapnir IL, Read BM, Swetter S, Greco RS. Indocyanine green and fluorescence lymphangiography for sentinel lymph node identification in cutaneous melanoma. J Surg Oncol Dec 2014;110(7):888-892.
44. Kom JM, Tellez-Diaz A, Bartz-Kurych M, Gathman B. Indocyanine green SPY elite-assisted sentinel lymph node biopsy in cutaneous melanoma. Plast Reconstr Surg Apr 2014;133(4):914-922.
45. Namikawa K, Tsutsumida A, Tanaka R, Kato J, Yamazaki N. Limitation of indocyanine green fluorescence in identifying sentinel lymph node prior to skin incision in cutaneous melanoma. Int J Clin Oncol Feb 2014;19(1):198-203.
46. Hollerweger A, Macheiner P, Hubner E, Gritzmann N. Axillary sentinel lymph-node biopsy: gamma probe assisted sonographic localization. Ultraschall Med Feb 2006;27(1):34-39.
47. Wheelahan P, Vinnicombe SJ, Brown DC, McLean D, Evans A. Accuracy of non-operative identification of the sentinel lymph node using combined gamma and ultrasound scanning. Clin Radiol Aug 2014;69(8):849-852.
48. Testori A, Meroni S, Moscovici OC, et al. Surgical sentinel lymph node biopsy in early breast cancer. Could it be avoided by performing a preoperative staging procedure? A pilot study. Med Sci Monit Sep 2012;18(9):CR543-CR549.
49. Motomura K, Inaji H, Komolke Y, et al. Gamma probe and ultrasonographically-guided fine-needle aspiration biopsy of sentinel lymph nodes in breast cancer patients. Eur J Surg Oncol Mar 2012;37(2):141-145.
50. Boistock P, Eisdner R, Glass E, et al. Comparison of blue dye and probe-assisted intraoperative lymphatic mapping in melanoma to identify sentinel nodes in 100 lymphatic basins. Arch Surg Jan 1999;134(1):43-49.
51. Bluemel C, Herrmann K, Giannmane F, et al. EANM practice guidelines for lymphoscintigraphy and sentinel lymph node biopsy in melanoma. Eur J Nucl Med Mol Imaging. Oct 2015;42(11):1750-1766.
52. van der Ploeg AP, van Akkooi AC, Haydu LE, et al. The prognostic significance of sentinel node tumour burden in melanoma patients: an international, multicenter study of 1539 sentinel node-positive melanoma patients. Eur J Cancer Jan 2014;50(1):111-120.
53. Hovensnejghel G, Classe JM, Garbay JR, et al. Prognostic value of isolated tumor cells and micrometastases of lymph nodes in early-stage breast cancer: a French sentinel node multicenter cohort study. Breast Cancer Oct 2014;23(5):561-566.
54. Eggermont AM, Seduc S, Rutkowski P, et al. Long term follow up of the EORTC 18952 trial of adjuvant therapy in resected stage IB-II melanoma patients comparing intermediate doses of interferon-alpha-2b (IFN) with observation: ulceration of primary is key determinant for IFN-sensitivity. Eur J Cancer Mar 2016;55:111-121.
55. Morton DL, Thompson JF, Cochran AJ, et al. Final trial report of sentinel-node biopsy versus nodal observation in melanoma. N Engl J Med. Feb 13 2014;370(7):599-609.
56. van Akkooi AC. Sentinel node followed by completion lymph node dissection versus nodal observation: staging or therapeutic? Controversy continues despite final results of MSLT-I. Melanoma Res Aug 2014;24(4):291-294.
57. Thomas J, Lissmann-Thesis D, Buchner A, et al. High-resolution ultrasound combined with power Doppler sonography can reduce the number of sentinel lymph node biopsies in cutaneous melanoma. Dermatology 2011;222(2):180-188.
58. Thompson JF, Haydu LE, Sanki A, Uren RF. Ultrasound assessment of lymph nodes in the management of early-stage melanoma. J Surg Oncol Sep 2011;104(4):354-360.
59. Sanki A, Uren RF, Moncrieff M, Tran KL, Scolyer RA, Lin HY, Thompson JF. Targeted high-resolution ultrasound is not an effective substitute for sentinel lymph node biopsy in patients with primary cutaneous melanoma. J Clin Oncol Nov 20 2009;27(33):5614-5619.
60. Rossi CR, Scagnot B, Vecchiato A, et al. Sentinel node biopsy and ultrasound scanning in cutaneous melanomaclinical and technical considerations. Eur J Cancer. 2000;36(7):895-900.
61. Hocevar M, Bracko M, Pogacic K, Vidiger-Kralj B, Besic N, Zgajnar J, Music MW. The role of repetitive ultrasonography in reducing the number of sentinel lymph node procedures in melanoma. Melanoma Res Dec 2004;14(6):533-536.
62. Testori A, Lazzaro G, Baldini F, et al. The role of ultrasound of sentinel nodes in the pre- and post-operative evaluation of stage I melanoma patients. Melanoma Res 2005;15(3):191-198.
63. Starritt EC, Uren RF, Scolyer RA, Quinn MJ, Thompson JF. Ultrasonic examination of sentinel nodes in the initial assessment of patients with primary cutaneous melanoma. Ann Surg Oncol. 2005;12(1):18-23.
64. Voit CA, Schafer G, Scheongen A, et al. Ultrasound (US) and US-guided fine needle aspiration cytology (FNAC) prior to sentinel lymph node biopsy (SLNB) in melanoma patients: accuracy of US-FNAC and lack of further improvement by RT-PCR of the aspirate. J Clin Oncol Jun 2006;24(18):4675-4675.
65. Sibon C, Chagnon S, Thakariner A, et al. The contribution of high-resolution ultrasonography in preoperatively detecting sentinel-node metastases in melanoma patients. Melanoma Res Aug 2007;17(4):233-237.
66. Kunte C, Schuh T, Eberle JY, et al. The use of high-resolution ultrasonography for preoperative detection of metastases in sentinel lymph nodes of patients with cutaneous melanoma. Dermatol Surg Nov 2009;35(11):1757-1765.
67. Chai CY, Zager JS, Szabunio MM, Marzban SS, Chau A, Rossi RM, Sondak VK. Preoperative ultrasound is not useful for identifying nodal metastases in melanoma patients undergoing sentinel node biopsy: preoperative ultrasound in clinically node-negative melanoma. Ann Surg Oncol Apr 2012;19(4):1100-1106.
68. Marone U, Catalano O, Canaco C, et al. Can high-resolution ultrasound avoid the sentinel lymph-node biopsy procedure in the staging process of patients with stage III cutaneous melanoma? Ultraschall Med 2012;33(7):E179-85.
69. Pilko G, Zgajnar J, Music M, Hocevar M. Lower tumour burden and better overall survival in melanoma patients with regional lymph node metastases and negative preoperative ultrasound. Radiol Oncol. 2012;46(1):60–8.
70. Stoffels I, Dissendom J, Poppell T, et al. Advantages of preoperative ultrasound in conjunction with lymphoscintigraphy in detecting malignant melanoma metastases in sentinel lymph nodes: a retrospective analysis in 221 patients with malignant melanoma AJCC stages I and II. J Eur Acad Dermatol Venereol Jan 2012; 26(1):79-85.
71. Hinz T, Voith H, Ahmadzadehfar H, Hoeller T, Wenzel J, Bieber T, Schmid-Wendtner MH. Role of high-resolution ultrasound and PET/CT imaging for preoperative characterization of sentinel lymph nodes in cutaneous melanoma. Ultraschall Med Jan 2013;33(1):30-36.
72. Ulrich J, van Akkooi AC, Eggermont AM, Voit CA. [Sonographic criteria for diagnosing sentinel node metastases in melanoma patients]. Ultraschall Med Apr 2015; 36(2):149-153.