Subclinical Breast Cancer: Minimally Invasive Approaches. Our Experience with Percutaneous Radiofrequency Ablation vs. Cryotherapy

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Keywords
Cryoablation · Radiofrequency ablation · Breast cancer · Percutaneous ablation · MRI · Minimally invasive techniques

Summary
Background: The aim of this study was to compare the efficacy of radiofrequency ablation vs. cryoablation in the treatment of early breast cancer. Patients and Methods: 80 women (mean age 73 ± 5 years) with early breast cancer were retrospectively evaluated. 40 patients underwent cryoablation and 40 patients underwent radiofrequency ablation, both with sentinel lymph node excision. Tumor volume and histopathological data were compared by means of postprocedural 3.0-T magnetic resonance imaging (MRI). 30–45 days after the percutaneous ablation, all patients underwent surgical resection of the tumor. The mean follow-up was 18 months without any local recurrences. Results: Both techniques allow good correlation with histopathological data. In 75 patients (93.8%) we observed complete necrosis; in 5 cases there was residual disease in the postprocedural MRI and postoperative histological examination. There was a good correlation between MRI volume and histologic samples. Cosmetic results were good in all patients but 2. Conclusion: Both percutaneous radiofrequency ablation and cryotherapy are minimally invasive techniques with a good clinical and cosmetic outcome in selected cases. MRI examination is an ideal method to assess breast neoplasms in terms of quality and quantity as well as residual tumor extent after percutaneous ablation. Cryotherapy is the preferred method because of the analgesic effect of freezing with better patients compliance.

Schlüsselwörter
Cryoablation · Radiofrequenzablation · Mammakarzinom · Perkutane Ablation · MRT · Minimalinvasive Techniken

Zusammenfassung
Introduction

In the past decades, advanced techniques for breast-conserving surgery have been developed which provide good oncological and cosmetic results without altering the survival rate of patients [1]. As a standard of care, surgical biopsy has been replaced by percutaneous core needle biopsy, and axillary dissection has been replaced by sentinel lymph node mapping. Percutaneous excision without major surgery of a single, subclinical, invasive cancer in selected patients is a new goal. Being a superficial structure, the breast is a suitable organ for percutaneous treatment.

Breast cancer is a cause of morbidity and mortality also in older women, affecting up to 30% of the over-70s. With the increase in life expectancy, this figure is likely to exceed 30% in the next decade [2, 3]. Due to the common presence of comorbidities, elderly patients are often treated with less aggressive approaches. Radiofrequency ablation (RFA), cryotherapy, interstitial laser ablation, focused ultrasound ablation, and focused microwave thermotherapy represent valid alternatives to open surgery with less psychological impact for the patient and good clinical and cosmetic outcome. Furthermore, these techniques are characterized by a low grade of complications, require shorter periods of hospitalization, and incur lower health care costs [4–7]. We aimed to compare in vivo the efficacy of RFA and cryotherapy in the treatment of small invasive breast cancers in terms of tumor necrosis, pathological and cosmetic outcome, and cosmetic outcome.

Patients and Methods

Patients

This study was approved by our institutional review board. We retrospectively reviewed all the minimally invasive procedures performed on breast cancer patients in our Department between October 2008 and March 2011. 80 postmenopausal women (mean age 73 ± 5 years (standard deviation, SD); range 64–82 years) were enrolled (table 1). Inclusion criteria were biopsy-proved ductal invasive unifocal breast cancer 2 cm or smaller (T1), well differentiated tumor (G1 and G2) visible in both ultrasound (US) and magnetic resonance imaging (MRI) studies, and tumor located at least 1 cm from the skin and 1 cm from the chest wall at US examination. We excluded patients with multifocal or multicentric neoplasia, lobular neoplasia, intraductal carcinoma, retro-areolar tumors, and other non-ductal infiltrating cancers. All patients provided written informed consent. 40 patients underwent cryotherapy and 40 patients underwent RFA. In all patients, vacuum-assisted biopsy (VAB) was performed for histopathological analysis. After the percutaneous procedure, US-guided strand reperage and sentinel lymph-node biopsy were performed. A preoperative MRI 1 and 4 weeks after the procedure was performed as follow-up. 30–45 days after the percutaneous ablation, all patients received definitive surgery.

Breast US

Ultrasonographic examinations were performed using a 5–12 MHz transducer and a US unit (ATL HDI 5000, Philips Medical Systems, Best, The Netherlands).

Breast MRI

Breast MRI examinations were performed with the patient placed in a prone position in a 3.0-T system (Achieva, Philips Medical Systems). A dedicated sensitivity-encoding breast coil was used for radiofrequency signal reception. Axial T2-weighted and T1-weighted images were obtained. The dynamic series consisted of a T1-weighted 3-dimensional fast-field echo sequence (repetition time/echo time, 5.1/2.4 ms; flip angle 20°). A total of 8 dynamic acquisitions with a temporal resolution of 70 s for a single dynamic acquisition were performed. Each dynamic acquisition consisted of 60 2.5-mm thick sections (gap, 0) with a 256×512 matrix and a field of view of 330 mm (adjusted to the size of the breast). Contrast agent bolus injection consisted of 0.2 mmol gadopentetate dimeglumine (Magnevist®, Schering, Berlin, Germany) per kilogram of body weight administered at an injection rate of 2.5 ml/s. Patients underwent 3 MRI examinations: 1 week before the procedure, 1 week after the procedure, and 4 weeks after the procedure.

Vacuum-Assisted Biopsy

Small invasive cancers were characterized based on histological samples obtained with the Mammotome® breast biopsy system (Ethicon Endo-Surgery, Cincinnati, OH, USA) or the Vacora® vacuum biopsy system (Bard Biopsy Systems, Tempe, AZ, USA) under US guidance. Specimens were subjected to standard histological and immunohistochemical evaluation (receptors for estrogen (ER), progesterone (PR), and HER2/neu) (table 1). VAB involves the use of US guidance and needles of different calibers with a coaxial system of collection/ aspiration that allows multiple samples to be taken through a single percutaneous insertion.

Cryotherapy

The procedure was performed in the angiography room. The devices were placed percutaneously under ‘real-time’ US guidance using a US probe at 5–12 MHz (ATL 5000, Philips Medical Systems). After disinfecting the skin of the breast, 10 ml of xilocaïne 2% were administered subcutaneously, along the path of the probe and in the target area. After skin incision with a scalpel (3 mm), 1 cryoprobe (1.47 mm diameter, 17-gauge isotherms, IcRod® model, Galil Medical, Yokneam, Israel) and 2 thermocouples for temperature monitoring were positioned. The device was set with the tip of the exposed needle at 2–5 mm from the distal margin of the lesion. Light intravenous sedation was always given, but patients stayed awake during the procedure. If the distance between the lesion and the skin profile and/or muscle was equal to 1 cm, sterile saline solution heated at 40 °C was injected to avoid surrounding skin and chest wall damage. A cooling double-cycle (each 10 min) separated by 5 min of passive heating (stop of argon coolant flow), and active heating with helium of the cryoprobe after the second cooling cycle, determined complete cell necrosis at −40 °C. Further heating was then necessary to dissolve the iceball. At the end of the procedure, the probe and thermocouples were removed without the need for stitches, and a sterile patch and compression bandage were put in place. In the same session, surgical excision of the sentinel lymph node for subsequent histological evaluation was performed. Patients were strongly advised to report postprocedural symptoms at the clinical follow-up 1 week later. The appearance of the skin, the size of the lesion, and the level of comfort of the procedure for the patient were

Table 1. Patient characteristics and hormone status pattern

<table>
<thead>
<tr>
<th>Patients, n</th>
<th>80</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age ± SD (range), years</td>
<td>73 ± 5 (64–82)</td>
</tr>
<tr>
<td>Hormone status, n</td>
<td></td>
</tr>
<tr>
<td>ER/PR/HER2</td>
<td>72</td>
</tr>
<tr>
<td>ER–/PR–/HER2</td>
<td>5</td>
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<tr>
<td>ER–/PR–/HER2</td>
<td>3</td>
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ER = Estrogen receptor; PR = progesterone receptor; HER2 = human epidermal grow factor receptor 2.

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Breast Care 2013;8:356–360
carefully recorded at the end of the session. Oral broad-spectrum antibiotic therapy was prescribed for the 5 days following the procedure.

**Radiofrequency Ablation**

An RFA system using a 460-KHz generator (TAG100 RF Generator, Fogazzi, Concesio, Italy) capable of 100 W maximum output at 200 Ω and a cool-tip single-needle electrode (Miras PTV, University Hospital Policlinico Tor Vergata, Rome, Italy, in collaboration with INVATEC ITALIA, Roncadelle, Brescia, Italy) were employed for the breast RFA procedures. The Miras PTV is a 15-gauge monopolar 200-mm double-lumen single electrode with active stainless steel tip exposure of 25 mm. The procedure was performed with patients in supine position and under general anesthesia, and the neoplasm was identified using US guidance. A small incision in the breast skin facilitated needle insertion. The cool-tip single-needle electrode was inserted into the core of the tumor. RF current was emitted, starting with the generator set to deliver energy according to the initial impedance value for a mean time of 5 min. The power was increased in 10-W intervals every 5 min until the readings on the external thermocouples indicated that the target temperature (90 °C) had been reached. Then, the power output was adjusted as needed to maintain the final temperature of 90 °C for 12 min. At the end of the procedure, a cool-down period of 60 s was applied to ensure necrosis of the tissue surrounding the needle tip. Upon completion of the procedure, the needle was withdrawn. Sterile skin closure strips were used to close the skin. Oral broad-spectrum antibiotic therapy was prescribed for the 5 days following the procedure.

**Surgery**

Lumpectomy was performed in all patients after a mean time of 34 days from percutaneous ablation (range 30–45 days). Axillary lymph node dissection was performed in 15 patients because of positivity of the sentinel lymph node biopsy.

**Adjunctive Treatment**

In accordance with the St Gallen oncologic criteria [8], patients underwent breast irradiation or adjuvant systemic chemotherapy. Hormonal therapy was administered according to grade, and ER, PR, and HER2/neu status determined at the pretreatment biopsy (table 1).

**Histopathology**

VAB was performed to histologically characterize the lesions before treatment. Samples were formalin-fixed and paraffin-embedded. Histological sections were stained with hematoxylin and eosin for tumor staging. Predictive and prognostic markers were studied by immunohistochemistry, including ER (CONFIRM Anti-ER SP1, Ventana Medical Systems, Tucson, AZ, USA), PR (CONFIRM anti-PR 1E2, Ventana), and HER2 (PATHWAY HER2, Ventana); the proliferative activity of tumor cells was evaluated using the Ki-67 index (CONFIRM Ki-67, Ventana).

**Cosmetic Evaluation**

Cosmetic evaluation was performed at the end of ablation (Time 0) and at 4 weeks from the procedure (Time 1), prior to surgical excision. All patients were evaluated based on the following signs: appreciable nodule on palpation, skin bruising, skin rash, and skin hyperpigmentation. The absence of skin pigmentation was considered excellent cosmetic outcome (grade 1), breast with slight texture changes or mild pigmentation was considered good cosmetic outcome (grade 2), breast with moderate texture changes or pigmentation was regarded as acceptable outcome (grade 3), and the presence of marked texture or pigmentation changes was considered a poor cosmetic outcome (grade 4).

**Follow-Up**

All patients underwent a weekly US examination from the 1st to the 4th week after the ablation procedure. MRI was performed at the 1st and 4th week after cryotherapy or RFA.

**Statistical Analysis**

All data analysis was performed using the Statistical Package for the Social Sciences, version 15.0 (SPSS Inc., Chicago, IL, USA). Descriptive statistics consisted of the mean ± SD for variables with Gaussian distributions or median (min-max) for categorical parametric and non-parametric data. Comparisons of paired data were carried out with the t-test or Wilcoxon test for non-parametric data. Correlation analysis was carried out by calculating the Pearson coefficient (r). Statistical significance was set at p < 0.05.

**Results**

Biopsies revealed all tumors to be invasive ductal carcinomas, of which 72 were hormone-responsive (ER, PR, HER2–), 5 were not endocrine responsive (ER–, PR–, HER2), and 3 were triple-negative (ER–, PR–, HER2–). 73 were well differentiated (G1), 7 were moderately differentiated (G2).

**Cryotherapy**

All 40 patients undergoing cryotherapy were treated with lumpectomy. At the end of the second cooling cycle, the iceball around the probe at –40 °C was a mean 16 mm
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Time 1 MRI contained partial degenerative changes (2 for cryotherapy and 3 for RFA) with NADH diaphorase activity revealing vital cells.

Cosmetic Evaluation
Cosmetic outcome was evaluated immediately after percutaneous ablation and 4 weeks after the procedure. At Time 0, it was grade 1 in 49 patients (26 for cryotherapy and 23 for RFA), grade 2 in 18 (8 for cryotherapy and 10 for RFA), grade 3 in 12 (7 for cryotherapy and 5 for RFA), and grade 4 in 1 (0 for cryotherapy and 1 for RFA) (fig. 1). At Time 1, cosmetic evaluation was grade 1 in 71 patients (37 for cryotherapy and 34 for RFA), grade 2 in 5 (2 for cryotherapy and 3 for RFA), grade 3 in 2 (1 for cryotherapy and 1 for RFA), and grade 4 in 2 patients (0 for cryotherapy and 2 for RFA). The comparison between the 2 series did not reveal a significant prevalence of esthetic complications in any group.

Discussion
Minimally invasive ablative techniques for small breast cancer may be a desirable goal in breast-conserving therapy. These kinds of treatment could improve patient comfort and cosmetic results, and provide a treatment window for patients who are not surgical candidates.

To our knowledge, this is the first study comparing the effects of RFA and cryotherapy in a homogeneous population. In the past decade, some studies have been published showing the efficacy of both techniques in independent series. The most important achievement is real-time imaging with which complete ablation of the lesion can be ascertained. US guidance is useful for needle and thermocouple placement within the tumor in both techniques, even though it is not accurate for depicting the ablation volume. In US monitoring during RFA, the appearance of gas bubbles and a hyperechoic halo with a posterior acoustic shadow in the treated area indicates the thermal effect of the procedure. Moreover, it produces an increasing echogenicity, with a shadow effect (‘fog effect’) that hides the deep tissue layers and the mass itself [4]. In the future, it would be desirable to use MRI real-time monitoring which allows visualization of the entire boundaries of the ablation area. In 2008, van der Ploeg et al. [9] published the first case of MRI-guided breast RFA in 3 patients with invasive ductal carcinoma and no artifacts around the needle electrode [9, 10]. Although the iceball produced during cryoablation reduces the visibility in deeper tissues, it is much more regular, definite, and symmetric, allowing an easier comparison between the margins of the lesion and the ablation volume [11]. All RFAs were performed successfully, with no tissue impedance rising above 20 Ω from baseline during the procedure, independent of the breast composition pattern. Spherical shapes were obtained in all cases with the type of RFA needle used [6]. In our series, some cases with MRI and histopathology showed residual disease. In 38 lesions, there were no vital neoplastic cells. No local recurrence was registered.

Table 2. Magnetic resonance imaging (MRI) evaluation: 1-week post-ablation MRI showed no residual enhancement in 71 (88.7%) of 80 lesions (36/38); 4-week post-ablation MRI showed increasing enhancement in 5 (6.25%) of the 9 areas of residual enhancement (2/3) suggesting residual tumor tissue which was confirmed by NADH diaphorase staining

<table>
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<th>Cryotherapy (n = 40)</th>
<th>RFA (n = 40)</th>
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<tbody>
<tr>
<td></td>
<td>positive</td>
<td>negative</td>
</tr>
<tr>
<td>MRI at 1 week</td>
<td>4</td>
<td>36</td>
</tr>
<tr>
<td>MRI at 4 weeks</td>
<td>2</td>
<td>38</td>
</tr>
<tr>
<td>Histology</td>
<td>2</td>
<td>38</td>
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</table>

All 40 patients undergoing RFA were treated with lumpectomy. The target temperature (90 °C) was reached after 15 ± 3.7 min (range 12–23 min). In 37 patients, we observed complete necrosis of the ablated tumor in both post-procedural MRI and after surgery, without enhancing lesions. In 5 cases, postprocedural MRI showed residual enhancement which in 2 patients was no longer visible at the presurgical MRI (expression of granulation tissue around avascular areas); in the other 2 cases both MRI and histopathology showed residual disease. In 38 lesions, there were no vital neoplastic cells. No local recurrence was registered.

Radiofrequency
All 40 cases of RFA were performed successfully, with no tissue impedance rising above 20 Ω from baseline during the procedure, independent of the breast composition pattern. Spherical shapes were obtained in all cases with the type of RFA needle used [6]. In our series, some cases with MRI and histopathology showed residual disease. In 38 lesions, there were no vital neoplastic cells. No local recurrence was registered.

### MRI Evaluation

After both procedures, MRI showed altered signal intensity and a minor degree of peripheral enhancement in the ablation areas. 1-week post-ablation MRI images displayed no suspicious residual enhancement in 71 (88.7%) of 80 lesions (36 for cryotherapy and 35 for RFA). The 4-week post-ablation MRI showed that 4 (5%) (2 for cryotherapy and 2 for RFA) of the 9 areas of residual enhancement were less conspicuous, suggesting reactive granulation tissue around the ablated areas. 5 (6.25%) of the 9 areas of residual enhancement increased in intensity on the 4-week postprocedural MRI images, suggesting residual tumor tissue which was confirmed by NADH diaphorase staining (table 2). There was a strong correlation between MRI volume and pathologic examination values (r = 0.896, p < 0.0001, 95% confidence interval (CI) 0.847–0.944).

Histopathology
Specimen examination showed only necrosis or degenerative changes in 77 cases. The 5 enhancing lesions revealed by Time 1 MRI contained partial degenerative changes (2 for cryotherapy and 3 for RFA) with NADH diaphorase activity revealing vital cells.

### Cosmetic Evaluation
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the first post-ablation MRI was performed to assess the short-
term effect on tumor and tissues. In clinical practice, a post-
ablation MRI may support the decision on further treatment
if incomplete tumor ablation is suspected. As a matter of fact,
50% of focal enhancement at the first MRI after cryotherapy
and 60% of positive cases after RFA were confirmed at the
4-week MRI and postsurgical histological evaluation. The
4-week post-ablation MRI images were helpful for further
evaluation of residual tumor enhancement, without the bias
related to inflammatory peripheral ring enhancement. We
achieved 38 disease-free specimens after cryotherapy
(94.84%) and 37 disease-free specimens after RFA (92.90%).
There was no statistically significant difference between
the cryotherapy and RFA groups (p = 0.082) so the results are
substantially equivalent. A total of 5 failures occurred in
terms of persistence of viable tumor cells. In both procedures,
this failure can be ascribed to incorrect positioning of the
device and insufficient size of the area of necrosis. In all failures,
the necrosis area was displaced from the center of the tumor.
Several studies showed that the absence of positive surgical
margins does not guarantee complete removal of disease,
and some studies have associated neoadjuvant endocrine therapy
with improved treatment outcome in elderly women [12, 13].
In our series, a strong correlation was shown between post-
procedural MRI and pathologic data. Both percutaneous ab-
lation techniques provided good results in terms of cosmetic
outcome and patient compliance. This is in accordance with
the literature [10, 14], although cryotherapy is preferred be-
cause of the analgesic effect of freezing. Furthermore, cryo-
therapy only requires mild sedation compared to general
anesthesia with RFA, which is more popular with patients
[6, 10, 14–16].

In all cases but 2, the cosmetic results were acceptable for
the patients. 1 RFA case with grade 4 outcome was complica-
ted by a 1-cm² skin necrosis, which was, however, remedied
during lumpectomy. Another grade 4 RFA case progressed
from grade 3 between Time 0 and Time 1 due to a retraction
of the skin caused by fibrosis in a small-sized breast. We
finally obtained 71 excellent results and good-mild results in
8 patients, with hyperpigmentation and/or subcutaneous
thickening. The difference between cryotherapy and RFA
regarding grade 4 cosmetic outcome may be related to a huge
energy deposition in the tissue during RFA.

This study is limited by some bias; furthermore, it is a
retrospective study without patient randomization, and the
study population was small and highly selected for low-grade
tumors in elderly patients. However, we can state that both
RFA and cryotherapy resulted in a good level of patient satis-
faction, although the analgesic effect of coldness is better en-
dured than the heat from RFA. Minimally invasive ablation
techniques may be useful in the treatment of single small
breast cancers with complete necrosis of the lesion, and good
esthetic outcome and patient satisfaction are achieved in
most cases.

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