Continuous versus pulsed microwave ablation in the liver: any difference in intraoperative pain scores?

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Abstract

Background This study prospectively compared intraoperative pain scores during percutaneous microwave ablation of the liver in patients randomized between continuous and pulsed energy delivery algorithms.

Methods During a 12-month period, 20 patients who underwent microwave liver ablation were prospectively randomized between 2 different energy delivery modes: “continuous mode” (CM, n=10) and "pulsed mode” (PM, n=10). All ablation sessions were performed using the same microwave ablation platform under computed tomographic guidance and intravenous analgesia. Within 30 min post ablation, all patients completed a questionnaire assigning a numeric pain intensity score from 0 (no pain) to 10.

Results Mean pain scores were 8.17±1.850 in the CM group and 4.50±1.567 in the PM group, with a statistically significant difference of 3.667±2.807 pain units (P=0.001). The mean procedure time was 53.5±20.90 min in the PM group vs. 58.5±17.44 min in the CM group (P=0.279). The mean size of the lesions was 2.81±0.95 cm in the PM group and 2.81±0.85 cm in the CM group (P=0.984). On a per-lesion basis, technical success was achieved in all evaluable tumors in both groups. No difference was noted in the local tumor control on the 6-month imaging evaluation. No complications were observed in the CM arm, while small perihepatic hemorrhagic fluid collections were reported in the PM group.

Conclusions Both algorithms for microwave energy delivery have comparable treatment effects in terms of 6-month local tumor control for liver lesions <3 cm in diameter. PM treatments compared to CM appear to induce significantly less pain in patients undergoing percutaneous liver ablation under intravenous analgesia.

Keywords Microwave ablation, continuous, pulsed, percutaneous, liver

Introduction

Microwave (MW) ablation is an increasingly applied technique for the treatment of malignant tumors, achieving high local tumor control rates in both primary and secondary lesions, irrespectively of the target's histology; the resultant larger ablation volumes may be achieved in shorter treatment times compared to any other thermal ablation modality [1-8]. The typical MW ablation zone obtained through a single energy delivery of a single probe, dwelling in a given position for the entire treatment duration, is an elongated ellipsoid with rotational symmetry around the probe’s axis, whose aspect ratio (defined as: S=D/L, where L is the maximum ablation size along the probe axis and D is the maximum ablation size perpendicular to it) generally ranges from 0.55-0.75 [9]. Over
the last decade, in an attempt to enhance ablation sphericity (i.e., taking S closer to 1), several MW vendors have deployed a variety of technical solutions, from new antenna designs to multi-probe configurations or modification of the standard MW delivery scheme (i.e., continuous provision of a fixed power P for the entire treatment duration T) [10-12].

Anesthesiology and analgesia protocols in the routine practice of percutaneous ablation are widely variable across centers and clinicians, ranging from intravenous analgesia to conscious or deep sedation and even up to general anesthesia. Significant factors include, but are not limited to, availability of the anesthesiologist, personal preferences of both the operator and the anesthesiologist, as well as patient’s comorbidities and performance status.

The purpose of this study was to prospectively evaluate pain tolerability during liver MW ablation with continuous or pulsed energy delivery modes, by comparing the patients’ self-reported pain scores immediately upon completion of the ablation session.

**Patients and methods**

All patients were informed about the interventional technique to be used, with its possible benefits and complications, and signed a written informed consent form prior to the procedure and the study. The study protocol conformed to the ethical guidelines of the 1975 Declaration of Helsinki and received approval by the institution’s human research committee prior to patients’ enrollment and treatment.

**Patient selection and evaluation**

Prior to percutaneous MW ablation, a cohort of 20 patients was randomized on a 1:1 basis between continuous (CM group) and pulsed (PM group) treatment. The study duration was 12 months. Lesions were diagnosed based on either contrast-enhanced magnetic resonance imaging (MRI) or positron emission tomography findings. Patients were selected for MW ablation by a multidisciplinary team of medical, radiation, surgical and interventional oncologists. Each patient underwent laboratory tests (including renal function and coagulation tests) at least 24 h prior to the percutaneous ablation session. Patient and tumor characteristics, treatment algorithm and parameters, intra- and postoperative complications, and intraoperative pain scores were recorded for each procedure. All ablation sessions were performed by 2 interventional radiologists (one with 20 and one with 10 years of experience); neither was blind to the energy delivery mode applied. Pain score questions were asked by a nurse within the first 30 min post ablation; the nurse was blind to the energy delivery mode applied.

The primary objective of the study was to compare the intraoperative pain between continuous and pulsed energy delivery modes; secondary objectives included local tumor control on follow-up imaging checks with MRI (including diffusion-weighted sequences and sequences after i.v. gadolinium injection).

Inclusion criteria included patients ≥18 years old with primary or metastatic hepatic disease, confirmed either by prior biopsy or through imaging (defined as new or growing nodules in cases of histological proven primary cancer); up to 3 hepatic lesions with a maximum size of 3 cm; Karnofsky Performance Scale score ≥60, coagulation parameters within normal limits and a life expectancy of >3 months. All included patients and lesions had to be evaluable over the 6-month follow up. All patients were referred to the study by the multidisciplinary tumor board. Exclusion criteria included a baseline pain level ≥2 prior to the ablation procedure, uncontrollable primary or metastatic disease outside the liver, patient’s non-compliance, uncontrollable international normalized ratio, systemic or local infection, poorly controlled ascites, expected survival less than 3 months, Eastern Cooperative Oncology Group score less than 3, and the presence of a medical or psychiatric illness that would preclude informed consent or follow up.

For the randomization process, we used the following procedure: when a patient met the inclusion criteria, he or she was assigned a number (the first patient referred to our department was assigned number 1, the second patient, number 2, the third patient, number 3, etc.). Odd-numbered patients were assigned to the CM group and even-numbered patients to the PM group.

**Percutaneous ablation procedure**

MW ablation was always performed in an inpatient setting. Computed tomographic (CT) guidance with sequential scanning (120 Kv peak, 240 mAs wavelength and 2 mm slice thickness) was used for planning, targeting and intraprocedural modification during the ablation session. The analgesia protocol was the same throughout the cohort: 30 min prior to ablation tramadol was injected intravenously, diluted in 100 mL normal saline, whilst acetaminophen was administered during the ablation session to treat intraprocedural pain (amounts of both analgesics were based on each patient’s weight) [13]. Under local sterility, MW ablation was performed via a percutaneous approach in all cases. After the initial CT scan, a skin entry point was selected. All treatments were performed using the same MW ablation equipment (HS AMICA, HS Hospital Service SpA, Rome, Italy), comprising a 2450 MHz solid-state generator with integrated peristaltic pump (AMICA-GEN) and internally cooled MW applicators (AMICA-PROBE). The MW probe was inserted in the lesion of interest through a MW probe (AMICA-PROBE). The MW probe was inserted in the lesion of interest through a MW probe generator with integrated peristaltic pump (AMICA-GEN) and internally cooled MW applicators (AMICA-PROBE).
lab experiments on *ex vivo* bovine liver, provide for the ablation dimensions (L × D) to be expected—with all the inherent limitations of an *ex vivo* animal model—for given settings of the energy delivery scheme (either continuous or pulsed, with \( t_{\text{ON}} = 4 \text{ sec} \) and \( t_{\text{OFF}} = 6 \text{ sec} \)), the output power (up to 100 W in continuous mode and up to 140 W in pulsed mode) and the overall treatment time (up to 15 min). Depending on the size of the lesion to be treated, the ablation protocols in the CM group provided for 40 or 60 W for either 5 or 10 min, while in the PM group all sessions lasted 10 min at either 100 or 120 W. In either trial arm, whenever deemed necessary, the MW antenna was repositioned and a second ablation session was performed, so as to ensure that the final ablation completely encompassed both the target tumor and an annular safety zone around it at least 5 mm thick.

### Statistical analysis

Study outcomes were numerically presented as mean±standard deviation. Pain scores in the 2 trial arms were compared using the paired-samples Student’s *t*-test. \( P \)-values <0.05 were considered to indicate a statistically significant difference. Multivariate analysis was performed in order to evaluate the role of confounding factors. Sample size calculation using a pain difference of 3 numeric visual scale (NVS) Units (considered clinically significant) was also performed, suggesting a sample of 14 patients for a power of 80%. Statistical analyses were performed using SPSS Statistics 22 (IBM Corp. Released 2013. IBM SPSS Statistics for Windows, Version 22.0. Armonk, NY: IBM Corp).

### Outcome measures

CT (contrast enhanced, in the arterial and portal venous phase of enhancement) was used to assess both the ablation zone size and the potential immediate complications at the end of the ablation treatment (Fig. 2). Exactly 30 min after the completion of ablation sessions, all patients completed a questionnaire in which they assigned a numeric pain intensity score from 0 (no pain) to 10. Patients were kept under observation for 24 h and eventually discharged. We evaluated technical success, complication rates and intraoperative pain scores. Pain score was evaluated using a brief pain inventory containing questions about pain intensity; answers were provided in terms of NVS [14]. Technical success was defined as complete coverage of the lesion by the ablation zone immediately after the procedure, plus a safety margin of at least 5 mm (as depicted by CT scan before and after iodinated contrast medium injection in the arterial and portal venous phases of enhancement). Complications were defined according to the Cardiovascular and Interventional Radiological Society of Europe (CIRSE) classification system [15].

### Results

The male:female ratio in both groups was 8:2. The mean age was 69.60±9.55 years in the CM group and 63.30±11.40 years in the PM group (\( P=0.197 \)). The PM group included 10 patients with 11 lesions: 5 hepatocellular carcinomas (HCCs) and 6 metastases (colon \( n=5 \) and breast \( n=1 \)); 4/11 (36.6%) tumors were subcapsular. The CM group included 10 patients with 10 hepatic lesions: 5 HCCs and 5 metastases (colon \( n=2 \), pancreas \( n=1 \), breast \( n=1 \), bronchogenic \( n=1 \)); 3/10 (30%) tumors were sub-capsular (i.e., ≤2 cm from the liver capsule). Five of 10 patients in the CM group and 6/10 in the PM group received systematic therapy during the ablation and study period. All patients remained hospitalized for the night following ablation and exited the hospital next morning.

The mean procedure time, including local anesthesia, placement of MW antenna, ablation and post-procedural CT evaluation was 53.5±20.90 min in the PM group vs. 58.5±17.44 min in the CM group (\( P=0.279 \)) (Fig. 2). The mean size of the lesions was 2.81±0.95 cm in the PM group and 2.81±0.85 cm in the CM group (\( P=0.984 \)).

On a per-lesion basis, technical success was achieved in all evaluable tumors in both groups. There was no repeat treatment of an index tumor. Local recurrence-free response (local tumor efficacy) of the treated lesions at 6 months was 90.90% (10/11) in the PM group and 100% in the CM group. The recurrent tumor in the PM group was depicted in a lesion next to a vessel and was retreated with continuous-mode MW ablation. On a per patient basis, at 6-month follow up in the...
Figure 2 Chart illustrating mean duration values of the ablation session between the 2 groups. There was no statistically significant difference (P=0.279)

PM Group overall treatment response was complete for 90% (9/10) in the PM group and 100% (10/10) in the CM group.

The mean pain score was 8.17±1.850 in the CM group and 4.56±1.567 pain units in the PM group (Fig. 3). There was a statistically significant difference of 3.667±2.807 pain units (P=0.001). No complications were noted in the CM group, whereas in the PM group there were 2 grade 1 complications according to the CIRSE classification system (small, perihepatic hemorrhagic fluid collections). During the follow-up period there were no deaths related to the procedure or to disease progression.

Discussion

A newer approach to sphericity enhancement involves the modification of the standard MW delivery scheme (i.e., continuous provision of a fixed power P for the entire treatment duration T). A few authors have proposed MW energy pulsing, i.e., intermittent energy administration characterized by a succession of active delivery periods, each of duration t_{acq} and rest periods, each of duration t_{off} (Fig. 1A) [12]. After a given treatment time T, the amount of energy deposited in the patient during a pulsed ablation procedure is: E=P*P*T*DC, where P is the power intensity delivered during active periods and DC = t_{acq}(t_{on}+t_{off}) is the percentage active time (i.e., duty cycle); therefore, as DC<1, higher P values are needed to compensate for rest periods and to warrant the same final energy deposition, compared to a continuous delivery scheme of equal duration. The ability of pulsing algorithms to generate more spherical ablations relies on the possibility of differentiating the temperature profiles parallel and perpendicular to the MW probe axis: for internally cooled probes, during rest periods tissues lying closer to the probe shaft are cooled faster than tissues at the ablation periphery farther from the probe; for appropriate choices of t_{on} and t_{off} (and provided shaft cooling is adequately intensive and effective), this can be exploited to reduce the ablation length while minimally affecting its transversal diameter D, so as to yield an improved S=D/L aspect ratio (Fig. 1B).

Pulsing algorithms are a valid means for creating more uniform temperature gradients and less anisotropic thermal profiles within tissues. Since postprocedural pain at the ablation site is certainly correlated with inflammatory reactions, which in turn are triggered by the treatment-induced thermal profile, the less steep and more uniform temperature gradient associated with pulsing rather than continuous MW delivery schemes may indeed result in less painful procedures [16,17]. A paper by Haugh et al nicely highlights the different temperature profiles obtained using the 2 MW delivery techniques under investigation: pulsed ablations resulted in much more uniform temperature maps around the emitting probe compared to continuous ablations [18]. In other words, the continuous mode allowed for higher temperatures closer to the emitting antenna but ended up with very similar temperatures compared to the pulsed mode at a given distance from the emitting tip: as the overall ablation zone is the ellipsoidal tissue volume contained within the 55°C isotherm induced by MW energy delivery during the treatment, both modalities proved equally capable of ablating similar tissue volumes, but pulsed ablation achieved this through a much more uniform temperature profile. One may infer (though not conclusively) that pulsed ablations provide for a “milder” treatment compared to continuous ablations, while depositing an equal amount of energy into tissues: this may account for a local reduction in temperature-triggered biological mechanisms (such as inflammation, nervous stimulation, production of cytokines, etc.), which also correlate with pain response. Another potential explanation might associate pain with the ablation shape: since the ablation zone is much longer than it is wide using continuous ablation mode, one would expect that the ablation zone more often extended into the capsule and/or perivascular areas (which are the sensitive areas in the liver) using continuous vs. pulsed ablation mode.

There are several limitations to our study that should be taken into account. Pain scores are inherently subjective and may be biased by patients’ concurrent disabilities.
Additionally, the number of patients enrolled for the study was certainly small considering the wide histological variety of the treated tumors, potentially introducing further biasing factors. Furthermore, there are no long-term follow-up results concerning the efficacy and survival rates associated with the 2 different energy delivery modes. Finally, this paper considers only a small subset of the many available treatment protocols (in terms of power and time settings), for both the continuous and the pulsed energy delivery modes.

Our results provide preliminary evidence of significant intraoperative pain reduction in CT-guided MW liver ablation procedures compared to conventional continuous delivery schemes, with no difference in the outcome (i.e., complete tumor necrosis) for liver lesions <3 cm in diameter. Hence, pulsed MW ablations appear to be an attractive option for patients with small to medium sized lesions undergoing liver ablation under i.v. analgesia, or whenever anesthesiologists are not available for the procedure. Future larger scale prospective trials with more circumscribed inclusion criteria are warranted in order to accumulate more robust and disease-specific data.

Summary Box

What is already known:

- Continuous-mode microwave (MW) ablation is an increasingly applied technique for the treatment of malignant tumors
- In an attempt to enhance ablation zone sphericity, a pulsed mode of MW ablation has been developed

What the new findings are:

- A pulsed mode of MW ablation can be an attractive alternative for percutaneous ablation of liver lesions <3 cm under i.v. analgesia or mild sedation
- The pulsed mode of MW ablation seems to induce less intraoperative pain, while having similar safety rates compared to continuous mode for liver lesions <3 cm
- Both algorithms for MW energy delivery have comparable treatment effects in terms of 6-month local tumor control for liver lesions <3 cm

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