Laser Thermal Ablation in the Treatment of Small Hepatocellular Carcinoma: Results in 74 Patients

PURPOSE: To evaluate the safety, local effectiveness, and long-term results of laser thermal ablation (LTA) in the treatment of small hepatocellular carcinoma (HCC).

MATERIALS AND METHODS: Ninety-two biopsies proved small HCCs (range, 0.8–4.0 cm) in 74 patients who were treated percutaneously with LTA in an outpatient clinic. A laser at a power of 5.0 W was coupled with one to four fibers that were advanced through 21-gauge needle(s) for 6–12 minutes. All lesions were evaluated with computed tomography (CT) for changes in size and vascular pattern, recurrence rates, and cumulative survival rates. Patients were examined for complications.

RESULTS: No major complications occurred in 117 LTA sessions, with an average of 1.3 sessions per tumor. At 3 months, CT scans showed a nonenhancing area (complete necrosis) in 89 (97%) of 92 lesions. During follow-up (range, 6–66 months; mean, 25.3 months), 84 tumors (91%) decreased in size. The local recurrence rates (range, 1–5 years) ranged from 1.6% to 6.0%. Recurrence rates (range, 12–60 months) in other liver segments ranged from 24% to 73%. Cancer-free survival rates (range, 1–4 years) ranged from 73% to 24%. Overall survival rates were 99%, 68%, and 15% at 1, 3, and 5 years, respectively. Twenty-one patients (28%) died.

CONCLUSION: LTA is a safe and effective treatment for small HCC.

Early detection strategies for hepatocellular carcinoma (HCC) have increased the number of small HCCs that are treatable with resection in countries where this disease is endemic (1,2). The survival rates reported (3,4) after partial resection or subsegmentectomy have improved in the past decade, mainly because of the ever smaller tumors undergoing resection. However, resection is possible in only a small proportion of patients because of the underlying advanced cirrhosis (4). Furthermore, it has been observed that HCC frequently recurs in the liver remnant after resection (5). Orthotopic liver transplantation is also an effective treatment for small unresectable HCCs when donors are available for potential recipients (6).

These problems in the management of HCC prompted the development of other potentially therapeutic modalities, such as transcatheter arterial chemoembolization (7); transcatheter arterial chemoembolization (8); percutaneous cryotherapy (9); and ultrasonographically (US) guided percutaneous injection of hot saline (10), acetic acid (11), or ethanol (12), all of which destroy cancer cells. Transcatheter arterial chemoembolization does not eliminate all malignant cells, and although extensive tumor necrosis can be observed, viable residual tumor cells remain, particularly beneath the tumor capsule, in intracapsular invasion, in daughter nodules, and in portal venous tumor thrombi (8,13). Percutaneous ethanol injection (PEI) is now widely used for small HCC (12,14). The effect of PEI is roughly comparable to that of surgical resection (5,12). However, a larger number of treatment sessions is required (12), and PEI does not always produce complete necrosis (13,14). Furthermore, recurrence after treatment is very common (12,15–17), and needle track implantation is not unusual, especially when HCCs are 2 cm or greater at their largest diameter (18); in addition, sclerosing HCCs respond poorly to PEI therapy (19).
To improve the results obtained with PEI, other percutaneous techniques have been developed. The most promising minimally invasive techniques are hyperthermic methods of tissue destruction, such as radio-frequency electrocoagulation (20–22), microwave coagulation (23), and laser thermal ablation (LTA) (24–29). Each of these techniques generates intralesional heat by a different mechanism, but the local hyperthermia is the unifying mechanism underlying their effectiveness. To date, all of these new percutaneous lesion-heating techniques are less well established than PEI.

The purpose of our work was (a) to evaluate the safety of LTA in the treatment of small HCCs, (b) to investigate the local effectiveness of this technique, and (c) to analyze the long-term results in a series of 76 patients.

MATERIALS AND METHODS

Between March 1994 and December 1999, we enrolled 74 patients (92 lesions) with a single tumor 4 cm or smaller or with one to three nodules 3 cm or smaller, among 308 patients with HCC diagnosed and staged at the Regina Apostolorum hospital. All patients were poor surgical candidates or had unresectable lesions. Exclusion criteria were age older than 80 years, evidence of extrahepatic metastases (based on the coincident findings of at least two imaging techniques) and/or lobar and local portal venous thrombosis, uncontrolled liver disease decompensation (gastrointestinal bleeding, encephalopathy, refractory ascites, bacterial infection), clotting impairment (platelet count less than 40 × 10^9/L or prothrombin time with international normalized ratio, or INR, greater than 1.99 seconds) (22), renal failure, or Child-Pugh class C cirrhosis. Patients who fulfilled the criteria gave their written consent to enter the study, which had been approved by the investigation and ethics committee of the hospital. The patients’ baseline characteristics are summarized in the Table.

Baseline Characteristics of Patients

<table>
<thead>
<tr>
<th>Patient Characteristic</th>
<th>Value</th>
</tr>
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<tbody>
<tr>
<td>No. of patients</td>
<td>74</td>
</tr>
<tr>
<td>No. of lesions</td>
<td>92</td>
</tr>
<tr>
<td>Age range (y)</td>
<td>52–80</td>
</tr>
<tr>
<td>Mean age (y) ± SD</td>
<td>67.5 ± 7.7</td>
</tr>
<tr>
<td>Sex (M:F)</td>
<td>47:27</td>
</tr>
<tr>
<td>Origin of disease (VHC, VHB, VHC + VHB, cryptogenetic, or alcohol-related)</td>
<td>48:14:3:2:7</td>
</tr>
<tr>
<td>Mean serum bilirubin (μmol/L) ± SD</td>
<td>18.8 ± 10.3</td>
</tr>
<tr>
<td>Mean serum albumin (g/L) ± SD</td>
<td>32 ± 5</td>
</tr>
<tr>
<td>Mean prothrombin activity (INR) ± SD</td>
<td>1.17 ± 0.74</td>
</tr>
<tr>
<td>Child-Pugh score A or B</td>
<td>54:20</td>
</tr>
<tr>
<td>Tumor stage solitary or multinodular</td>
<td>58:16</td>
</tr>
<tr>
<td>Ratio of tumor of 0.8–1.0, 1.1–2.0, 2.1–3.0, 3.1–4.0 cm</td>
<td>5:24:51:12</td>
</tr>
<tr>
<td>Diameter of tumor (cm)</td>
<td>0.8–4.0</td>
</tr>
<tr>
<td>Mean ± SD</td>
<td>2.4 ± 0.7</td>
</tr>
<tr>
<td>Volume of tumor (mL)</td>
<td>Range 0.5–30</td>
</tr>
<tr>
<td>Mean ± SD</td>
<td>8.2 ± 6.1</td>
</tr>
<tr>
<td>Tumor stain present, absent, or mixed</td>
<td>78:7:7</td>
</tr>
<tr>
<td>Tumor location unilobar or bilobar</td>
<td>87.5</td>
</tr>
<tr>
<td>Tumor segment location 2, 3, 4, 5, 6, 7, or 8</td>
<td>4:7:13:9:30:9:20</td>
</tr>
<tr>
<td>US pattern hypoechoic, hyperechoic, isoechoic with hypoechoic rim, or mixed</td>
<td>74:5:7:6</td>
</tr>
<tr>
<td>Tumor capsule present, absent, or irregular</td>
<td>38:46:8</td>
</tr>
<tr>
<td>Histologic grade of tumor differentiation well differentiated, moderately differentiated, poorly differentiated, undifferentiated, or mixed</td>
<td>43:39:8:1:1</td>
</tr>
<tr>
<td>No. of patients with serum AFP value &lt;20, 20–200, or &gt;200 (μg/L)</td>
<td>54:15:5</td>
</tr>
</tbody>
</table>

Note.—AFP = α-fetoprotein, INR = international normalized ratio, VHB = virus hepatitis B, VHC = virus hepatitis C.

LTA Technique

Preparation for LTA was performed with the same guidelines as those used for large-core percutaneous biopsy. All patients were treated in our outpatient clinic. Percutaneous treatment is performed in the angiography suite by one radiologist in the simpler cases and by two radiologists in more complex and harder-to-reach cases, with one nurse who is experienced in dealing with con-
scious sedation during interventional procedures (30). Local anesthesia was achieved by using 5 mL of 2% xilocaine (Xilocaine 2%; AstraZeneca, Basiglio, Italy). Just before needle insertion, both 3–7 mg of midazolam maleate (Ipnovel 15; Roche, Milan, Italy) and 4 mL of fentanyl citrate (Fentanyl; Pharmacia & Upjohn, Milan, Italy) were administered intravenously while the cardiovascular and respiratory systems were continuously monitored. With US guidance and a 3.5-MHz convex transducer (CAB411A-EZU-PA3B4-1; Es- aote Biomedica), the needles were positioned within the lesion in the area to be treated, and the tip positions were confirmed with two-plane US images.

LTA was performed with a continuous-wave neodymium yttrium-aluminium-garnet, or Nd:YAG, laser (DEKA-M.E.L.A.; Florence, Italy) that operated at a wave-length of 1,064 μm. This laser source was used with an optical beam-splitting device (SMART 1064 HCC; DEKA-M.E.L.A.) with four separate fibers that were always illuminated concurrently (simultaneous or multiple-fiber application) (31). A 21-gauge modified Chiba needle (Hospital Service) and a plane-cut tip fiberoptic with a 300-μm quartz core (Medical Energy, Pensacola, Fla) were used. The need- dles were placed within the tumor prior to starting treatment, and each stylet was replaced by a sterile optical fiber, which was advanced until at least 1 cm of the bare fiber tip was in direct contact with the deepest part of the tumor. Both the number of needles (one to four) and the arrangement were chosen in accordance with the size, shape, and location of the lesions.

To achieve regional coagulation with necrosis of the main tumor, capsular in- vasion, and neighboring intrahepatic metastases (peripheral satellite nodules and portal vein thrombi), up to four need- dles were simultaneously placed in a square configuration with interneedle spacing of 1.0–1.5 cm, encompassing the whole tumor and surrounding noncan- cerous parenchyma, with the intention of creating a volume of necrosis larger (a rim at least 5–10 mm of the rim of nor- mal liver around the tumor) than the original volume of the neoplasm. Once the laser was turned on and the sched- uled energy was delivered for the single treatment, if warranted by the size of the lesion, another laser treatment was per- formed by withdrawing the needles and fiber tips 2.0 cm from the treated area and carefully repositioning them in the contiguous, still untreated cranial area within the lesion. All laser illuminations were monitored continuously with real- time US. At the end of the treatment, the optic fiber was slowly pulled out with the laser still on. Once the needles were pulled out, US was performed to reveal intra- and/or extrahepatic complications. LTA sessions were performed by one or two of five radiologists (C.M.P., G.B., F.M., P.C., B.C.) with at least 5 years experience in interventional abdominal procedures and fellowship training in our hospital.

After the procedure, patients were observed for 6 hours in the outpatient recovery room and, if no complications occurred, were discharged. Before the procedure and at 6 and 24–48 hours and 1 month after treatment, the following serologic values were measured: levels of transaminases, alkaline phosphatase, bil- irubin, electrolytes, hemoglobin, fibrino- gen, haptoglobin, and creatinine; pro- thrombin activity; and complete blood cell count. The degree of liver function impairment was estimated by means of routine biochemical parameters within 2 months after LTA. In addition, recorded data included Child-Pugh classification, growth of tumor, and occurrence of ex- trahepatic disease.

Study Design and Assessment of Treatment Effectiveness

Our aim was to achieve a rapid, safe, and complete destruction of small HCCs by producing a large area of necrosis while markedly reducing the number of sessions required to carry out a complete ablation. On the basis of our experience (32) and that reported by other authors (33), we used 5 W per fiber with an expo- sure time of 6 minutes to reach 1,800 J per fiber for each single tumor treatment. A laser illumination that lasted 6 minutes was considered a single tumor treatment. Up to two treatments were scheduled during any one session. We scheduled no more than two irradiation cycles for any one lesion, depending on the size of the tumor. The first irradiation cycle was composed of a maximum of two sessions and a maximum of two tumor treatments per session; and the second irradiation cycle, with the same design as the first cycle, took place after 3 months if there was incomplete necrosis.

We used one fiber for lesions less than or equal to 1 cm (n = 5), two fibers for lesions less than or equal to 2 cm (n = 24), and four fibers for lesions with a diameter of 2.1–4.0 cm (n = 63). Thus, we determined the number of LTA tumor treatments and sessions, the total energy used per session and per tumor, the mean volume of necrosis obtained and the en- ergy delivered per milliliter of tumor, and the mean total procedure time.

During and at 5–6 hours and 24–48 hours after treatment, US was performed to evaluate any changes in the echotex- ture of the lesions treated and in ther- mally induced necrosis. After 1–2 weeks and every 6 months thereafter, US was performed to monitor changes in size and lesion detectability during follow-up.

CT was performed 24–48 hours after the first irradiation cycle to evaluate the short-term effects of the procedure, assess the extent of necrosis, and plan further intervention. Still-enhancing areas were assumed to indicate residual viable tu- mor, and an additional irradiation cycle (ie, one or two sessions of LTA) was per- formed. Thus, tumor necrosis was con- sidered complete when no areas of en- hancement were seen in the tumor or at the periphery on CT scans obtained 3 months after the first irradiation cycle or 24–48 hours after a possible second irra- diation cycle. If CT scans obtained after the second irradiation cycle showed res-idual viable tissue, LTA was considered to have failed and was discontinued; in our hospital, patients then underwent segmental transcatheter arterial chemo-embolization within 30–60 days after LTA failure.

On the basis of the findings of this imaging technique, the response to LTA was defined in accordance with World Health Organization criteria as complete response, no evidence of neoplastic disease; partial response, reduction greater than 50%; no change, reduction less than 50%; and progressive disease, increase of greater than 25% (34). Local recurrence was diagnosed when enhancing foci were seen close to, along the edge of, or within the treated area. CT was performed 3 months after the end of treatment and every 6 months thereafter. All CT scans were analyzed in consensus by two expe- rienced radiologists (C.M.P., D.V.). Ther- apeutic effectiveness was also assessed with an α-fetoprotein assay performed af- ter the end of LTA and every 6 months thereafter. During follow-up, hepatic se- rum functional indexes were measured every 6 months.

Statistical Analyses

Statistical analysis was conducted ev- ery 6 months. The patients’ baseline char- acteristics and follow-up results are pre- sented as means plus or minus SDs. Rates of local recurrence, occurrence of new le- sions, death, cancer-free survival, and cu-
The mean volume of necrosis obtained was 3,600–44,600 J (median, 12,500 J). The energy used per tumor was 1.3). The total energy used at each session was 2,1) were performed during 117 sessions (range, one to six treatments; median, two; mean, 2.1) were performed during 117 sessions (range, one to six treatments; median, two; mean, 2.1). The total energy used at each session was 1,800–15,900 J (median, 8,000 J). The energy used per tumor was 3,600–44,600 J (median, 12,500 J). The mean volume of necrosis obtained was 15.2 mL ± 9.5 (SD) (range, 2–44 mL), with a total energy deposition of 800 J/mL of tumor (range, 350–1,600 J/mL; median, 700 J/mL). The mean total procedure time was 25.5 minutes ± 7.9 per session (range, 15–45 minutes).

**Imaging**

Laser illuminations were monitored continuously with real-time US. In every case, once LTA treatment had begun, continuous US monitoring revealed, after a delay of about 30–60 seconds, a slowly enlarging and coalescing hyperechoic zone around each fiber tip resulting from vaporization of fluid and formation of microbubbles of gas. This irregular and poorly defined hyperechoic area then progressively increased to envelop the entire tumor. Vapor appeared as echogenic linear foci radiating from the treatment area along tissue planes and/or hepatic vessels or coming up through the needle sheath and becoming visible as smoke in the needle hub. US performed after treatment depicted a decrease in the echogenicity of the treated area, and after 5–6 hours, a heterogeneous lesion was surrounded by a 3–5-mm hypoechogenic rim. At US performed after 24–48 hours, the thermal lesion was characterized by a small central hyperechoic area (zone of vaporization), which was surrounded by a thin hyperechoic rim (zone of carbonization) and a thick outer hypoechogenic area (zone of coagulation) with associated acoustic shadowing (Fig 1). This US appearance was nearly identical to that on 1–2-week and 6- and 12-month follow-up images, apart from a decrease in size of both the central cavity and the entire heated area. These changes preceded precise assessment of the coagulation size after LTA.

On precontrast CT scans, the entire treated tumor was replaced by an irregular area of hyperattenuation that was surrounded by a thin rim of lower attenuation (Fig 2). On contrast-enhanced CT scans obtained 24–36 months after treatment, the coagulation areas were no longer detectable at follow-up CT and on US images obtained 36–48 months after treatment. In our series, two lesions were still detectable after 48 and 60 months and had the characteristics just described.

**RESULTS**

A total of 196 tumor treatments (range, one to six treatments; median, two; mean, 2.1) were performed during 117 sessions (range, one to three sessions; median, one; mean, 1.3). The total energy used at each session was 1,800–15,900 J (median, 8,000 J). The energy used per tumor was 3,600–44,600 J (median, 12,500 J). The mean volume of necrosis obtained was 15.2 mL ± 9.5 (SD) (range, 2–44 mL), with a total energy deposition of 800 J/mL of tumor (range, 350–1,600 J/mL; median, 700 J/mL). The mean total procedure time was 25.5 minutes ± 7.9 per session (range, 15–45 minutes).

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On precontrast CT scans, the entire treated tumor was replaced by an irregular area of hyperattenuation that was surrounded by a thin rim of lower attenuation (the “ghost” of the original neoplasm) (Fig 2). On contrast-enhanced CT scans obtained after treatment, the whole heated region appeared as a uniform area of low attenuation that was surrounded by a rim of higher attenuation (Fig 2), which disappeared gradually during follow-up. In five (5%) of 92 cases, the area of hyperattenuation that surrounded the coagulative necrosis had, after 24 hours, a segmental configuration (Fig 3).

All 84 (91%) successfully treated tumors had diminished in size by 65% and 87% at 6 and 12 months, respectively, after LTA. The nonenhancing areas at the site of the treated HCC remained detectable as small hyperechoic areas with acoustic shadowing at US follow-up examination (Fig 1) and as small areas of hypointensity during the arterial (Fig 2) and equilibrium phases on follow-up CT scans obtained 24–36 months after treatment. The coagulation areas were no longer detectable at follow-up CT and on US images obtained 36–48 months after treatment. In our series, two lesions were still detectable after 48 and 60 months and had the characteristics just described.

**Therapeutic Responses and Follow-up Findings**

A single irradiation cycle was performed in 84 (91%) of 92 lesions. In eight (9%) tumors, a second irradiation cycle that consisted of only one session was performed (Fig 4). We obtained a volume of necrosis larger than that of the original lesion in 77 (84%) of 92 tumors; in 12 (13%) tumors, we achieved a volume equal to the initial volume and corresponding to the original lesion in size and shape (Fig 4); in the remaining three (3%) tumors, we observed evidence of tumor persistence (partial response). Complete response (complete necrosis) was achieved in 89 (97%) of 92 tumors. Of the three tumors (two patients) in which necrosis was incomplete, one was of grade 2 and located in the fifth segment and two were of grade 3 and located in the eighth segment. In both tumors, transcatheter arterial chemoembolization was performed; in one after 32 and in the other 55 days after the last LTA treatment.

During follow-up (range, 6–66 months; mean, 25.3 months), five (6%) of 89 successfully treated lesions showed a local recurrence. The 1-, 2-, and 5-year local recurrence rates were 1.6%, 6.0%, and 6.0%, respectively, and all local recurrences were observed within 2 years after LTA (range, 8–22 months; mean, 14.8 months) (Fig 5). A new HCC (one or more lesions with diameter of 0.8–3.0 cm; mean, 2.1 cm) occurred elsewhere in the liver in 36 (49%) of 74 patients within 3–64 months (mean, 17.7 months) after LTA. The 12-, 24-, 36-, 48-, and 60-month recurrence rates in segments other than that of the original site were 24%, 45%, 62%, 73%, and 73%, re-
Side Effects and Complications

No major complications requiring intensive care were encountered during or after the procedure. Three minor complications in the course of 120 LTA sessions occurred in three patients during the first 6–24 hours after the procedure: self-remitting variceal bleeding (reduction of hemoglobin level by 8 g/dl [80 g/L]), subcapsular hemorrhage, and perirenal inflammation. The first two complications required no treatment, although discharge from the outpatient recovery room was delayed, and hospital admission was indicated. Neither transfusion nor surgical repair was necessary for either the patient with variceal bleeding, since the hematocrit level returned to baseline values within 5 days, or the patient with intraparenchymal bleeding. In the third patient in whom the lesion was located peripherally, adjacent to the Glisson capsule in the sixth segment, we observed perirenal inflammation (hypoechogenic irregular area at US and ill-defined area of hypotenuation at contrast-enhanced CT), which cleared up in 20 days; the perirenal disease required the administration of 1 g of amoxicillin (Zimox; Pharmacia & Upjohn) once a day for 7 days but not hospitalization.

All patients experienced mild to moderate pain at the insertion site or the epigastic level, or diaphragmatic irritation with right shoulder tip pain during the procedure, especially when the tumor was superficial or contiguous to the porta hepatitis. Sixty-one (82%) of 74 patients experienced moderate pain or slight discomfort in the right hypochondrium during the next few days. In only 36 (59%) of these cases did the pain require the administration of 500 mg of acetaminophen (Tachipirina 500; Angelini, Rome, Italy) twice a day; in five (8%) cases, 10 drops of ketorolac tromethamine (Toradol; Recordati, Rome, Italy) was required three times a day in the fol-

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**Figure 2.** CT scans show complete response after LTA (energy delivered, 28,800 J). (a) Transverse arterial phase contrast-enhanced CT scan obtained before LTA shows a hyperattenuating 5.7-cm-diameter nodular lesion (arrow) in segment 8. (b) Transverse precontrast CT scan obtained 24 hours after LTA shows that the entire tumor has been replaced by an irregular and ill-defined area of hypotenuation (the “ghost” of the original neoplasm) (arrowhead) surrounded by a rim of lower attenuation (arrows). The small zones of vacuolation resulted from vaporization of fluid and formation of microbubbles of gas. (c) Transverse arterial phase contrast-enhanced CT scan obtained 24 hours after LTA shows the entire heated region characterized by a homogeneous area of hypotenuation, due to coagulation, surrounded by a thin rim of hypotenuation (arrowheads) due to hyperemia. The area of necrosis is larger than the original nodular lesion. (d) Transverse arterial phase CT scan obtained 24 months after LTA shows a marked reduction in size of the area of necrosis (arrow).
blood cell count, or platelet count were observed. At the time this article was written, no evidence of needle-route seeding had been seen.

DISCUSSION

To our knowledge, our patient population was the first series of patients with small HCC to be treated percutaneously with LTA. Our study findings demonstrate that treatment with LTA can result in complete tumor necrosis in a high percentage of tumors. We hypothesized that in the three nodules with incomplete response, treatment failure could be explained by improper placement of the needles in the lesion. When simultaneous multiple fiber application is used, the distance at which the fibers are positioned determines whether the synergistic thermal effect between the fibers will occur (31,35). Even a small deviation from the linearity of the insertion tract can cause either diverging fibers with insufficient tissue coagulation or converging channels with an accumulation of light and heat, causing subsequent tissue carbonization and fiber breakdown. If the optical fibers are correctly placed within the tumor, a thermal lesion that is larger than that obtained with PEI (5,12,17,22) or with other heating percutaneous techniques (22,23) invariably occurs.

An advantage of LTA is that it requires fewer treatment sessions for a complete treatment than does PEI (12,22). Our results are comparable to those achieved with radio-frequency electrocoagulation (22), but with the latter method, the investigators excluded patients who had lesions adjacent to the hepatic hilum or confluence of the hepatic veins (22). The use of thin fibers and fine needles with LTA makes it possible to reach the entire lesion safely, even if it is very deep, difficult to get at, near the gallbladder or the hepatic hilum, or superficial. Moreover, LTA treatment seems to be extremely well tolerated and produces no severe side effects or major complications.

Our study findings show that local recurrence (Figs 5, 8) does occur in small HCCs treated with LTA, but this drawback is infrequent. Other authors (5,11,14–16), by using PEI, report variable values that range from 4% to 30%. In our opinion, LTA may be equivalent to limited hepatic resection and may influence long-term survival, achieving results comparable to those of segmentectomies. Confirmation can come only from controlled prospective trials in which both treatments are compared and the patients are followed up for at least 5 years.

It is difficult to compare reliably radiofrequency ablation with other heating techniques, as different approaches have been used to increase the diameter of coagulation. Moreover, the relatively short follow-up time reported for some of the systems does not facilitate helpful comparison. However, some preliminary considerations should be stressed. Radio-frequency ablation, like PEI, demonstrated less effectiveness in the treatment of contiguous multinodular HCC, probably because of the fibrosis surrounding the small neoplastic nodules, which may influence heat diffusion and thereby limit treatment effectiveness (19,22). Moreover, by using this method in neoplastic tissue of the most common type (ie, nodular), one can generally reproduce the
size and shape of the necrotic area so that it is similar to that of the tumor, as if a mold of the tumor had been made (oven effect) (22). The same authors (38) confirmed these results in another study, explaining that with HCC it is generally sufficient to treat just the tumor itself, and that the oven effect may protect surrounding structures.

More recently, Rossi et al (39), by using the radio-frequency interstitial tumor ablation, or RITA, with a four-pronged umbrella needle system (RITA Medical Systems, Mountain View, Calif) after occlusion of the tumor blood supply, confirmed again that lethal temperatures are reached just inside the HCC nodule, whereas nonlethal temperatures are detected in the hepatic tissue immediately surrounding the nodule, and therefore, the resultant necrosis selectively involves only the HCC nodule and reproduces the nodule shape. The same authors (39) reported that in their study, the local recurrences may have been due to the unsatisfactory diffusion of heat within the HCC nodule, as well as extracapsular invasion or daughter nodules outside the original nodule. These results seem to indicate that radiofrequency ablation is unable to induce a volume of necrosis larger than the original lesion and thus obtain the complete destruction of neighboring neoplastic or preneoplastic nests. However, a longer follow-up is needed for a better understanding of the outcomes of these techniques, and more valid data are needed to compare the different options.

Recurrence of new HCCs in locations far from the original tumor was frequent during follow-up; this was probably due to multiple metachronous or unnoticed synchronous lesions or to intrahepatic metastases that were not detected at imaging (37). Such new lesions are part of the natural history of HCC in cirrhosis. The entire cirrhotic liver contains regenerative nodules, and hepatocarcinogenesis is probably a stepwise process from a regenerative nodule to an adenomatous hyperplasia and then to HCC (40). Thus, the entire liver is ready for the development of multiple HCCs, and the first lesion to be detected is only the beginning of a multifocal disease with several tumor clones (40,41).

The survival rates in our study were comparable to those of other studies (12,17). However, we stress that some reports involved patients who received diagnosis and were staged a few years ago when imaging techniques were less accurate. In our view, the use of these less accurate imaging techniques to verify local control of the disease may have caused an underestimation of the rates of local recurrence and of the short-, mid-, and long-term results with PEI. Hence, this limitation must also be taken into account when our results are compared with those of other authors.
Thus, there is a need for prospective, randomized, controlled trials with a greater number of patients with predefined characteristics to facilitate the analysis and comparison of the different treatment options offered to patients with HCC.

The best way to ascertain the effectiveness of LTA is to assess the presence of vascular enhancement after contrast material administration at CT, considering that nonenhancement represents absence of viable tissue and thus successful necrosis of the tumor (20–22,25–27). The use of triple-phase helical CT is, in our opinion, essential for assessing treatment effectiveness and completeness and for early detection of tumor recurrence (42). The relationship between the enhancing peripheral halo and LTA treatment deserves some comment, and in our view, the attenuation behavior, with its segmental appearance after bolus injection, can be explained. The hyperattenuating halo around the coagulated tissue that is thought to be related to hyperemia is probably due to a rapid change in hepatic arterial flow in response to thermal injury. Thermal ablation induces a marked increase in arterial perfusion in the surrounding liver within a few minutes after the start of treatment. The heated area of necrosis can cause hepatic segmental venous occlusion, and in this condition, the portal veins become draining veins and the occluded region is supplied with arterial blood alone (43). This phenomenon causes hyperattenuation in the occluded area at CT since arterial flow itself increases and the contrast medium is less diluted there. CT acquisition with rapid sequential sections makes it possible to evaluate these changes.

Determination of the volume of the area to be heated is complex and depends on the laser power, the laser irradiation time and wavelength of light involved, how the energy reaches the tissue, and such tissue factors as optical and thermal properties (22–27,29,44). Since different types of tissue have different optical characteristics, the choice of a specific laser treatment must be related to the tissue type involved (44). Because of these factors, according to a few authors (27), it is not practical to establish a fixed deposition of laser energy for any given tumor size; LTA of tumors of the same size may result in variable necrosis, despite the use of the same amount of laser energy. On the basis of our experience, however, we believe that the therapeutic parameters are 5 W of laser power administered so as to reach simultaneously a total energy of 7,200 J to an ellipsoid volume ranging from 8 to 18 mL ± 3.9 (mean, 13.4 mL) for a single tumor treatment and ranging from 12 to 40 mL ± 11.5 (mean, 21.2 mL) for two tumor treatments with a total energy deposition of 14,400 J (32) and a procedure time of 6 and 12 minutes, respectively.

Although no formal cost-effectiveness analysis was performed as part of this work, it should be pointed out that for LTA, the laser generator costs were $40,000–$45,000 and the costs for a set of fibers were only $1,000–$1,500, but this set can be used to treat up to 50 patients (45).

In conclusion, LTA is a safe and effective percutaneous technique for the treatment of primary liver tumors. However, further research in this field is needed to elucidate (a) the exact correlation between histopathologic structure and laser energy and (b) the correlation between hepatic blood flow and the actual extent of the area of necrosis (46).

Acknowledgment: We thank Manuela Latini and Laura Morelli for their help in the care of patients during conscious sedation.

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