Thermal ablation for unresectable liver tumours, time to move forward?

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INTRODUCTION

Advances in liver surgery and anaesthesia have expanded their potential applications and a fifth of liver tumours are now resectable. Nevertheless, in the majority of patients surgery is not possible and chemotherapy and palliative care remain the mainstays of treatment. In these patients ablative techniques are an attractive option and offer an opportunity to increase survival. Experimental research over the last 20 years has seen the development of a number of ablative modalities and several of these have been applied clinically. This has provided hepatobiliary surgeons with techniques which can be used palliatively but also in combination with resection for multiple and awkwardly positioned lesions. Techniques which achieve local destruction by modification of the tissue temperatures adjacent to the treatment probe have been increasingly favoured over the “chemical” procedures (alcohol injection and tissue chemoembolization)\[1\]. However ablative techniques which utilise temperature changes are potentially associated with significant complications. In fact, although microwave (MW) and radiofrequency (RF) are used in clinical practice, cryotherapy (CRYO) has been largely abandoned and laser ablation was never widely employed.

Examining the literature regarding thermal ablation (TA) reveal many similarities but also demonstrate significant differences among techniques. These comparisons are valuable, identifying those properties that are essential for the development of a safe and reliable treatment. In addition, the data is important when designing research strategies to facilitate further technical developments. The purpose of this editorial is to examine some of these common critical factors that influence local failures and to propose approaches to overcoming the current limitations.
RECURRENCES RATES AFTER THERMAL ABLATIVE PROCEDURES AND RISK FACTORS

Technical advances have allowed modifications of the equipment used for thermal ablative techniques. Different approaches have been used in an attempt to augment the advantages of the individual techniques and allow wider clinical application. Nevertheless, data from clinical studies does not confirm the superiority of any particular modality especially in respect of recurrence rates. The rate of local recurrences ranged from 0% to 50% after MW ablation[24, 25], 2% to 67% after RF[7-10], and 9% to 44% after CRYO[11-13]. Multivariate analysis of the risk factors that could potentially influence local failure after ablation found only a few which were important in determining outcome. The tumor size, number of nodules, and Child-Pugh classification had an influence after MW[14], tumor size and the ablative approach used (surgical versus percutaneous) after RFA[15] and tumor size, the presurgical level of carcinoembryonic antigen and tumor grading after CRYO[15,16].

REASONS FOR LOCAL FAILURE

Tumor size

With all ablative techniques, it is clear that tumor size is crucial and influences the short and long term results. Tumours of 3 cm or larger are most likely to recur and this is related to difficulties with achieving a complete ablation at the initial treatment. The ablation must encompass the lesion with a significant degree of overlap to ensure complete treatment and this can be difficult with large tumours due to the proximity of major vascular structures. One centimeter is considered a safe macroscopic margin and a 3 cm wide lesion therefore requires a 5 cm wide ablation to avoid local recurrence. Treatment zones which can be achieved with MW range between 3.5 and 5 cm[17,18], although newer devices have produced ablations with diameters up to 6.5 cm[19-22]. RF ablation is able to create lesions which range from 0.5 to 5 cm[23,24], although ablations of up to 7 cm have been reported[25]. In experimental CRYO studies most of the lesions produced were between 2 and 3 cm[26,27], with only one report described larger lesions with diameters up to 9 cm[28].

Different methods have been used to increase the diameters of ablated lesions and treat tumours greater than 3cm. Modifications of the shape and number of RFA electrodes and MW antennae lead to different shapes of the ablated zones and achieved larger diameters compared to the single straight configuration[19,21,22,29-31]. Another approach employs cooling of the RF-ablated tissue to avoid desiccation. The increase in tissue charring which occurs at the metal electrode–tissue interface during RF ablation results in elevated circuit impedance and markedly reduced RF output, limiting the heat diffusion. RF electrodes with internal cooling, those that release normal saline around the probe (wet electrodes) or where saline is added from an external source (saline-enhanced RFA) have all been investigated with promising results[29]. Finally, the temporary exclusion of vascular inflow (using a Pringle manoeuvre) almost doubled the diameters of the ablated lesions in all TA procedures and could potentially achieve volumes which are up to six-times greater[28,32,38].

Proximity to large vessels

Although tumour size is important both in respect of eligibility for initial surgery and the potential for treatment by an ablative technique some small liver tumours are still inoperable due to their anatomical position. Tumours adjacent to major vessels are often unresectable and at the same time at higher risks of recurrence and complications following TA. Large blood vessels (greater than 3 mm), and to a lesser degree bile ducts, are relatively protected from thermal injuries as the inflowing warm blood conducts the excessive temperature from the vessel wall, but the blood flow also prevents an adequate heating/freezing of perivascular tissues with consequent reduced destruction of tumour cells. The phenomenon is called “heat (or cold)-sink effect” and is the basis for incomplete ablations.

No specific methods have presently been developed to overcome this problem, which still remains a major contraindication to TA. However, there are other modalities which do not rely on temperature changes and are consequently not affected by local blood flow. Electrolytic ablation, also known as low-level direct current therapy, is an alternative method which produces no change in temperature and exerts its effects through electrochemical and pH modifications and the production of free radicals. These changes render the microenvironment close to the probe extremely cytotoxic, produce tissue necrosis and are not influenced to any large degree by the presence of blood vessels or bile ducts[39,40]. Due to the almost complete absence of thermal energy developed, the procedure has proved particularly safe in experimental studies where ablations were conducted close or even within major vessels (Figure 1)[41-43]. This characteristic makes this technique particularly useful for the treatment of lesions positioned close to major vessels[44]. The presence of the “electric-sink” effect, if any, does not modify significantly the shape of the ablated lesion which is able to include the vessel wall without damaging it[45]. If appropriate clinical trials confirm a clear survival advantage in this subgroup of patients with unresectable lesions close to major vessels, this could become a specific indication for electrolytic ablation that would negate the relative disadvantage of long ablation times compared to those of the more common TA procedures.

Histological zones

Although minor differences exist in descriptions of the pathological changes following different TA procedures, the results are generally very similar and independent of the method employed. The ablated region is composed predominantly of two concentric zones with a central area of complete coagulative necrosis and a peripheral transitional rim of inflammation, congestion,
hemorrhage and thrombosis. While the coagulative zone is occupied entirely by dead cells and amorphous material, the transitional rim still contain viable cells that may survive the adverse microconditions produced by the inflammatory environment and ultimately give rise to tumor recurrences\cite{31,46,47}. In this area cells express heat shock proteins and die by apoptosis, the effect of which peaks 6 h following the ablation and expands the zone of definitive necrosis\cite{46,51}, and a combination of capillary microthrombosis and vasoconstriction further contributes to the enlargement of the necrotic area\cite{52}.

Problems resulting from the production of a transitional zone have to be addressed if recurrences in this rim of tissue are to be eliminated. Experimental studies have provided considerable details about the process and mechanisms, but to date have failed to identify a reliable strategy to overcome the problem. Nevertheless, there are a number of hypotheses that may allow the development of future strategies. Firstly, techniques that selectively enhance the transitional rim may better differentiate tumor recurrences from the physiological repair that occurs normally following ablation. The recent advent of targeted ultrasound (US) contrast media has been achieved by linking specific antibodies against a wide variety of antigens with US echogenic microbubbles\cite{53,54}. In this way, inflammatory molecules (ICAM-1, VCAM-1 and E-selectin) have been successfully targeted by selective microbubbles able to visualise and identify different areas of inflammation\cite{58}. A similar approach has been employed for magnetic resonance imaging contrasts media, in which oxide nanoparticles have been rendered selective for AvB3 Integrin and successfully tested in-vitro\cite{59}.

Secondly, the creation of US microbubbles specific for the transitional inflammatory rim and linked with chemotherapeutic agents would guarantee a selective delivery of the anti-tumor therapy to the zone where recurrences are most likely and they would also be able to visualize the zone at the same time to ensure it was effectively treated. Preliminary studies have already been conducted with targeted microbubbles loaded with chemotherapy for the treatment of breast, ovarian and pancreatic tumours\cite{57,58}. In liver tumours a combined therapeutic approach can be envisaged which would consist of a TA procedures for the ablation of the majority of the tumour, and following the ablation selective delivery of chemotherapy to sterilize the remaining cells that may survive in the transitional inflammatory rim. Chemotherapy-loaded contrast agents could target not only inflammatory antigens but also tumor-specific molecules (i.e. CEA), and the simultaneous administering of both (inflammatory and tumor-specific loaded) could further help in the emergence prevention of resistant cells.

CONCLUSION

Although the best treatment for resectable liver tumour remains surgical resection\cite{10}, survival rates achieved by TA have been sufficiently promising that some centers have started to advocate trials comparing resection with ablation\cite{16}. Limitations that favour tumour recurrence following TA are common to all techniques, but some of those that did not enter clinical trials due to the disadvantage of long ablation times could help in particular clinical settings. Furthermore, newer techniques like MW have intra-operative advantages over the older RF in terms of faster execution times clearly demonstrated in dose-response studies\cite{19,25,29}. Unfortunately, modification of delivery probes and techniques have failed to reduce recurrence rates and a more integrated approach is required, possibly combining TA techniques with adjuvant therapies tailored to identify and treat specific areas in the transition zone.

REFERENCES

1. Cho YK, Kim JK, Kim MY, Rhim H, Han JK. Systematic review of randomized trials for hepatocellular carcinoma treated with percutaneous ablation therapies. *Hepatology* 2009; 49: 453-459
2. Kuang M, Lu MD, Xie XY, Xu HX, Mo LQ, Liu GJ, Xu ZF, Zheng YL, Liang JY. Liver cancer: increased microwave delivery to ablation zone with coiled-shaft antenna--experimental and clinical studies. *Radiology* 2007; 242: 914-924
3. Ohmoto K, Yoshioka N, Tomiyama Y, Shibata N, Kawase T, Yoshida K, Kuboki M, Yamamoto S. Radiofrequency ablation versus percutaneous microwave coagulation therapy for small hepatocellular carcinomas: a retrospective comparative study. *Hepatogastroenterology* 2007; 54: 985-989
4. Kawamoto C, Ido K, Isoda N, Hozumi M, Nagamine N, Ono K, Sato Y, Kobayashi Y, Nagae G, Sugano K. Long-term outcomes for patients with solitary hepatocellular carcinoma treated by laparoscopic microwave coagulation. *Cancer* 2005; 105: 965-993
5. Dong B, Liang P, Yu X, Su L, Yu D, Cheng Z, Zhang J. Percutaneous sonographically guided microwave coagulation therapy for hepatocellular carcinoma: results in 234 patients. *Am J Roentgenol* 2003; 180: 1547-1555
6. Ong SL, Gravante G, Metcalfe MS, Strickland AD, Dennison AR, Lloyd DM. Efficacy and safety of microwave ablation for primary and secondary liver malignancies: a systematic review. *Eur J Gastroenterol Hepatol* 2009; 21: 599-608
7. Sutherland LM, Williams JA, Padbury RT, Gotley DC, Stokes B, Maddern GJ. Radiofrequency ablation of liver tumours: a systematic review. *Arch Surg* 2006; 141: 181-190
8. Mulier S, Ni Y, Jamart J, Ruers T, Marchal G, Michel L. Local recurrence after hepatic radiofrequency coagulation: multivariate meta-analysis and review of contributing factors.
Gravante G. Liver thermal ablation

Ann Surg 2005; 242: 158-171

9 Wong SL, Mangub PB, Choti MA, Crocenzi TS, Dodd GD 3rd, Dorfman GS, Eng C, Fong Y, Giusti AF, Lu D, Marsland TA, Michelson R, Poston GJ, Schrag D, Seidenfeld J, Benson AB 3rd, American Society of Clinical Oncology 2009 Clinical Evidence Review on Radiofrequency Ablation of Hepatic Metastases From Colorectal Cancer. J Clin Oncol 2010; 28: 493-508

10 Mulier S, Ruers T, Jamart J, Michel L, Marchal G, Ni Y. Radiofrequency ablation versus resection for resectable colorectal liver metastases: time for a randomized trial? An update. Dig Surg 2008; 25: 445-460

11 Adam R, Akpinar E, Johann M, Kunstlinger F, Majno P, Bismuth H. Place of cryosurgery in the treatment of malignant liver tumors. Ann Surg 1997; 225: 39-48; discussion 48-50

12 Onik G, Rubinsky B, Zemel R, Weaver L, Diamond D, Cobb C, Porterfield B. Ultrasound-guided hepatic cryosurgery in the treatment of metastatic colon carcinoma. Preliminary results. Cancer 1991; 67: 901-907

13 Mala T, Edwin B, Mathisen G, Tillung T, Fosse E, Bergan A, Søreide O, Gladhaug I. Cryoaoblation of colorectal liver metastases: minimally invasive tumour control. Scand J Gastroenterol 2004; 39: 571-578

14 Liang P, Dong B, Yu X, Yu D, Wang Y, Feng L, Xiao Q. Prognostic factors for survival in patients with hepatocellular carcinoma after percutaneous microwave ablation. Radiology 2005; 235: 299-307

15 Seifert JK, Morris DL. Prognostic factors after cryotherapy for hepatic metastases from colorectal cancer. Ann Surg 1998; 228: 201-208

16 Seifert JK, Morris DL. Indicators of recurrence following cryotherapy for hepatic metastases from colorectal cancer. Br J Surg 1999; 86: 234-240

17 Yamashiki N, Kato T, Bejarano PA, Berho M, Montalvo B, Shebert RT, Goodman ZD, Schiffrer ER, Tzakis AG. Histopathological changes after microwave coagulation therapy for patients with hepatocellular carcinoma: review of 15 explanted livers. Am J Gastroenterol 2003; 98: 2052-2059

18 Shen P, Geisinger KR, Zagoria R, Levine EA. Pathologic correlation study of microwave coagulation therapy for hepatic malignancies using a three-ring probe. J Gastrointest Surg 2007; 11: 603-611

19 Hines-Peralta AU, Pirani N, Clegg P, Cronin N, Ryan TP, Liu Z, Goldberg SN. Microwave ablation: results with a 2.45-GHz applicator in ex vivo bovine and in vivo porcine liver. Radiology 2006; 239: 94-100

20 Awad MM, Devgan L, Kamel IR, Torbensen M, Choti MA. Microwave ablation in a hepatic porcine model: correlation of CT and histopathologic findings. HPB (Oxford) 2007; 9: 357-362

21 Simon CJ, Dupuy DE, Ianniiti DA, Lu DS, Yu NC, Aswad BI, Busuttil RW, Lassman C. Intraoperative triple antenna hepatic microwave ablation. AJR Am J Roentgenol 2006; 187: W333-W340

22 Yu NC, Lu DS, Raman SS, Dupuy DE, Simon CJ, Lassman C, Aswad BI, Ianniiti D, Busuttil RW. Hepatocellular carcinoma: microwave ablation with multiple straight and loop antenna clusters—pilot comparison with pathologic findings. Radiology 2006; 239: 269-275

23 Goldberg SN, Gazelle GS, Compton CC, Mueller PR, Tanabe TF. Characterization of intrahepatic malignancy with radiofrequency ablation: radiologic-pathologic correlation. Cancer 2000; 88: 2452-2463

24 Goldberg SN, Walovitch RC, Straub JA, Shore MT, Gazelle GS. Radio-frequency-induced coagulation necrosis in rabbits: immediate detection at US with a synthetic microsphere contrast agent. Radiology 1999; 213: 438-444

25 Goldberg SN, Solbiati L, Hahn PF, Cosman E, Conrad JE, Fogle R, Gazelle GS. Large-volume tissue ablation with radio frequency by using a clustered, internally cooled electrode technique: laboratory and clinical experience in liver metastases. Radiology 1998; 209: 371-379

26 Matsuzono R, Selig AM, Colucci VM, Jolesz FA. MR monitoring during cryotherapy in the liver: predictability of histologic outcome. J Magn Reson Imaging 1993; 3: 770-776

27 Tacke J, Adam G, Haage P, Sellhaus B, Grosskortenhaus S, Günther RW. MR-guided percutaneous cryotherapy of the liver: in vivo evaluation with histologic correlation in an animal model. J Magn Reson Imaging 2001; 13: 50-56

28 Seifert JK, Gerharz CD, Mattes F, Nassir F, Fachinger K, Bell C, Junginger T. A pig model of hepatic cryotherapy. In vivo temperature distribution during freezing and histopathological changes. Cryotherapy 2003; 47: 214-226

29 Mulier S, Miao Y, Muller P, Dupas B, Pereira P, de Baere T, Lencioni R, Leveillee R, Marchal G, Michel L, Ni Y. Electrodes and multiple electrode systems for radiofrequency ablation: a proposal for updated terminology. Eur Radiol 2005; 15: 798-808

30 Mulier S, Ni Y, Frich L, Burdjo D, Denys AL, De Wispelaere JF, Dupas B, Habib N, Hoey M, Jansen MC, Lacrosse M, Leveillee R, Miao Y, Muller P, Mutter D, Ng KK, Santambrogio R, Stippel D, Tamaki K, van Guldik TM, Marchal G, Michel L. Experimental and clinical radiofrequency ablation: proposal for standardized description of coagulation size and geometry. Ann Surg Oncol 2007; 14: 1381-1396

31 Gravante G, Ong SL, Metcalfe MS, Strickland A, Dennison AR, Lloyd DM. Hepatic microwave ablation: a review of the histological changes following thermal damage. Liver Int 2008; 28: 911-921

32 Patterson EJ, Scudamore CH, Owen DA, Nagy AG, Buczko­owski AK. Radiofrequency ablation of porcine liver in vivo: effects of blood flow and treatment time on lesion size. Ann Surg 1998; 227: 559-565

33 Chinn SB, Lee FT Jr, Kennedy GD, Chinn C, Johnson CD, Winter TC 3rd, Warner TF, Malvhi DM. Effect of vascular occlusion on radiofrequency ablation of the liver: results in a porcine model. Am J Roentgenol 2001; 176: 789-795

34 Chang CK, Hendy MP, Smith JM, Recht MH, Wellin RE. Radiofrequency ablation of the porcine liver with complete hepatic vascular occlusion. Ann Surg Oncol 2002; 9: 594-598

35 Shen P, Fleming S, Westcott C, Challia V. Laparoscopic radiofrequency ablation of the liver in proximity to major vascularule: effect of the Pringle maneuver. J Surg Oncol 2003; 83: 36-41

36 Shibata T, Murakami T, Ogata N. Percutaneous microwave coagulation therapy for patients with primary and metastatic hepatic tumors during interruption of hepatic blood flow. Cancer 2000; 88: 302-311

37 Shibata T, Niinobu T, Ogata N. Comparison of the effects of in-vivo thermal ablation of pig liver by microwave and radiofrequency coagulation. J Hepatobiliary Pancreat Surg 2000; 7: 592-598

38 Mala T, Frich L, Aurdal L, Clausen OP, Edwin B, Søreide O, Gladhaug IP. Hepatic vascular inflow occlusion enhances tissue destruction during cryoablation of porcine liver. J Surg Res 2003; 115: 265-271

39 Hagedorn R, Fuhr G. Steady state electrolysis and isoelectric focussing. Electrophoresis 1990; 11: 281-289

40 Baxter PS, Wemmyss-Holden SA, Dennison AR, Maddern GJ. Electrochemically induced hepatic necrosis: the next step forward in patients with unresectable liver tumours? Aust N Z J Surg 1998; 68: 637-640

41 Wemmyss-Holden SA, Dennison AR, Finch GJ, Hall PD Pde L, Maddern GJ, Electrolytic ablation as an adjunct to liver resection: experimental studies of predictability and safety. Br J Surg 2002; 89: 579-585

42 Wemmyss-Holden SA, de la M Hall P, Robertson GS, Dennison AR, Vanderzon FS, Maddern GJ. The safety of electrolytically induced hepatic necrosis in a pig model. Aust N Z J Surg 2000; 70: 607-612
43 Metcalfe MS, Mullin EJ, Texler M, Berry DP, Dennison AR, Maddern GJ. The safety and efficacy of radiofrequency and electrolytic ablation created adjacent to large hepatic veins in a porcine model. *Eur J Surg Oncol* 2007; 33: 662-667

44 Wemyss-Holden SA, Dennison AR, Berry DP, Maddern GJ. Local ablation for unresectable liver tumors: is thermal best? *J Hepatobiliary Pancreat Surg* 2004; 11: 97-106

45 Finch JG, Fosh B, Anthony A, Slimani E, Texler M, Berry DP, Dennison AR, Maddern GJ. Liver electrolysis: pH can reliably monitor the extent of hepatic ablation in pigs. *Clin Sci (Lond)* 2002; 102: 389-395

46 Fraser J, Gill W. Observations on ultra-frozen tissue. *Br J Surg* 1967; 54: 770-776

47 Bhardwaj N, Strickland AD, Ahmad F, Atanesyan L, West K, Lloyd DM. A comparative histological evaluation of the ablations produced by microwave, cryotherapy and radiofrequency in the liver. *Pathology* 2009; 41: 168-172

48 Ohno T, Kawano K, Sasaki A, Aramaki M, Yoshida T, Kitano S. Expansion of an ablated site and induction of apoptosis after microwave coagulation therapy in rat liver. *J Hepatobiliary Pancreat Surg* 2001; 8: 360-366

49 Nikfarjam M, Muralidharan V, Su K, Malcontenti-Wilson C, Christophi C. Patterns of heat shock protein (HSP70) expression and Kupffer cell activity following thermal ablation of liver and colorectal liver metastases. *Int J Hyperthermia* 2005; 21: 319-332

50 Hoffman AL, Wu SS, Ohaid AK, French SW, Lois J, McMonigle M, Ramos HC, Sher LS, Lopez R. I. Histological evaluation and treatment outcome after sequential radiofrequency ablation and hepatic resection for primary and metastatic tumors. *Am Surg* 2002; 68: 1038-1043

51 Rai R, Richardson C, Flecknell P, Robertson H, Burt A, Manas DM. Study of apoptosis and heat shock protein (HSP) expression in hepatocytes following radiofrequency ablation (RFA). *J Surg Res* 2005; 129: 147-151

52 Strickland AD, Clegg PJ, Cronin NJ, Swift B, Festing M, West KP, Robertson GS, Lloyd DM. Experimental study of large-volume microwave ablation in the liver. *Br J Surg* 2002; 89: 1003-1007

53 Klibanov AL. Microbubble contrast agents: targeted ultrasound imaging and ultrasound-assisted drug-delivery applications. *Invest Radiol* 2006; 41: 354-362

54 Klibanov AL. Ultrasound molecular imaging with targeted microbubble contrast agents. *J Nucl Cardiol* 2007; 14: 876-884

55 Lindner JR. Microbubbles in medical imaging: current applications and future directions. *Nat Rev Drug Discov* 2004; 3: 527-532

56 Chen K, Xie J, Xu H, Behera D, Michalski MH, Biswal S, Wang A, Chen X. Triblock copolymer coated iron oxide nanoparticle conjugate for tumor integrin targeting. *Biomaterials* 2009; 30: 6912-6919

57 Gao Z, Kennedy AM, Christensen DA, Rapoport NY. Drug-loaded nano/microbubbles for combining ultrasonography and targeted chemotherapy. *Ultrasonics* 2008; 48: 260-270

58 Rapoport NY, Kennedy AM, Shea JE, Scaife CL, Nam KH. Controlled and targeted tumor chemotherapy by ultrasound-activated nanoemulsions/microbubbles. *J Control Release* 2009; 138: 268-276

59 Lorentzen T, Christensen NE, Nolsle CP, Torp-Pedersen ST. Radiofrequency tissue ablation with a cooled needle in vitro: ultrasonography, dose response, and lesion temperature. *Acad Radiol* 1997; 4: 292-297

S- Editor Li LF  E- Editor Yang C