**Name of journal:** *World Journal of Gastroenterology*

**ESPS Manuscript NO: 13407**

**Columns: REVIEW**

**Current management of hepatocellular carcinoma: An** **Eastern perspective**

Yim HJ *et al*. Management of HCC

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**Author contributions**: Um SH designed the study, outlined the draft, and supervised the overall project; Yim HJ wrote and organized the manuscript; Suh SJ searched reference materials and contributed to the writing of the manuscript.

**Supported by** grants from Korea Healthcare Technology R&D Project, Ministry of Health and Welfare, South Korea, No. HI10C2020 (partly); and Korea University Research Grant (partly).

**Conflict-of-interest:** The authors do not have any conflicts to report.

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**Received:** August 21, 2014

**Peer-review started:** August 23, 2014

**First decision:** October 29, 2014

**Revised:** December 11, 2014

**Accepted:** February 12, 2015

**Article in press:**

**Published online:**

**Abstract**

Hepatocellular carcinoma (HCC) is one of the leading causes of cancer death, especially in Eastern areas. With advancements in diagnosis and treatment modalities for HCC, the survival and prognosis of HCC patients are improving. However, treatment patterns are not uniform between areas despite efforts to promote a common protocol. Although many hepatologists in Asian countries may adopt the principles of the Barcelona Clinic Liver Cancer staging system, they are also independently making an effort to expand the indications of each treatment and to combine therapies for better outcomes. Several expanded criteria for liver transplantation in HCC have been developed in Asian countries. Living donor liver transplantation is much more commonly performed in these countries than deceased donor liver transplantation, and it may be preceded by other treatments such as the down-staging of tumors. Local ablation therapies are often combined with transarterial chemoembolization (TACE) and the outcome is comparable to that of surgical resection. The indications of TACE are expanding, and there are new types of transarterial therapies. Although data on drug-eluting beads, TACE, and radioembolization in Asian countries are still relatively sparse compared with Western countries, these methods are gradually gaining popularity because of better tolerability and the possibility of improved response rates. Hepatic arterial infusion chemotherapy and radiotherapy are not included in Western guidelines, but are currently being used actively in several Asian countries. For more advanced HCCs, appropriate combinations of TACE, radiotherapy, and sorafenib can be considered, and emerging data indicate improved outcomes of combination therapies compared with single therapies. To include these paradigm shifts into newer treatment guidelines, more studies may be needed, but they are certainly in progress.

**Key words**: Hepatocellular carcinoma; Eastern; Treatment; Guidelines; Combination

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**Core tip:** This article describes the current status of the management of hepatocellular carcinoma, focusing on the changing trends of treatment modalities in Eastern countries. Newly adopted therapies as well as emerging combination strategies are discussed based on recent data.

Yim HJ, Suh SJ, Um SH. Current management of hepatocellular carcinoma: An Eastern perspective. *World J Gastroenterol* 2015; In press

**INTRODUCTION**

Worldwide, hepatocellular carcinoma (HCC) is the sixth most prevalent cancer[[1](#_ENREF_1)]. More than 600000 people are newly diagnosed every year and approximately the same number die due to HCC annually. The main etiology of HCC is liver cirrhosis caused by chronic hepatitis B or C, alcohol, fatty liver diseases, or less commonly, autoimmune or genetic metabolic liver diseases[[2](#_ENREF_2)]. The incidence, characteristics, and prognosis of HCC vary from region to region according to the prevalence of underlying chronic liver diseases as well as the screening and treatment strategies for HCC. Currently, efforts are being made to promote the use of common protocols, but the patterns of treatment are still not uniform as the therapeutic approach to HCC mainly depends on the availability of treatment modalities as well as the preferences of physicians[[2-7](#_ENREF_2)]. As three-quarters of HCC cases occur in East Asia, the experiences and data in this area should have been substantially accumulated, and the treatment trends would have characteristic features. This article aims to review the current status of the management of HCC from an Eastern perspective. The first section introduces the principles and current trends of different treatment modalities, and the second section summarizes the findings on multidisciplinary treatments based on recently available data.

**MAIN TREATMENT MODALITIES: EVOLVING ROLES AND CHANGING TRENDS**

For decisions regarding initial treatments, the Barcelona Clinic Liver Cancer (BCLC) staging system from Western guidelines is frequently applied[[3](#_ENREF_3),[4](#_ENREF_4)]. This system has very strict guidelines for treatments; only very early stage and early stage HCCs are indicated for the curative therapies, and only one treatment option is assigned to each of intermediate stage and advanced stage disease. Furthermore, no combination therapy is recommended according to the BCLC algorithm. Hence, despite the worldwide use of the BCLC guidelines, debates regarding their practicality are ongoing. The Asian Pacific Association for the Study of the Liver has guidelines for HCC treatment similar to those in the BCLC system[[5](#_ENREF_5)]; both consider hepatic function as well as tumor size, number, and its extent, and the treatment options are not much different from BCLC. However, indications of pre-existing therapies are expanding and newly emerging therapies are currently being implemented (Figure 1). In addition, alternative therapies or combination therapies for each stage are available as determined by clinical situations in real practice. In this section, current status of surgical, interventional, medical, and radiation therapies are reviewed with newly available data, particularly, from Asian countries.

***Surgical therapies***

**Liver transplantation:** Transplanting a healthy liver provides the most favorable survival outcomes in HCC patients[[8](#_ENREF_8)]. If the patients have underlying decompensated liver cirrhosis, no other option exists. However, the availability of organs limits access to this best curative therapy. The annual incidence of deceased organ donors does not exceed 5 per million in most Asian countries[[9](#_ENREF_9)]. Compared with those in Western countries, patients with HCC in Asia have a low probability of receiving a deceased donor liver transplantation (DDLT) in a timely manner and thus have a higher risk of drop-out because of tumor progression[[10](#_ENREF_10)]. For this reason, living donor liver transplantation (LDLT) has been promoted. In Korea, the proportion of adult LDLT recipients with HCC has increased to 30%–40% of all HCC liver transplant recipients[[11](#_ENREF_11)]. Despite technical complexity, LDLT is replacing DDLT in other Asian countries as well[[8](#_ENREF_8)]. Donor mortality and morbidity rates of LDLT were 0.2% and 24%, respectively, according to a report of a worldwide survey[[12](#_ENREF_12)]. Most LDLT centers develop their own criteria for maximizing donor safety[[13](#_ENREF_13)]. Although the right lobe is the most suitable graft for the recipient, its procurement is limited by size of donor liver. When the right lobe cannot be used alone, a dual graft from 2 donors containing the left lobe can be utilized[[14](#_ENREF_14)]. Despite this method, the donor pool has not significantly expanded because of the technical complexity of the surgery and ethical concerns. To further overcome organ shortage, ABO-incompatible LDLT was attempted and became successful after the implementation of rituximab, which decreased antibody-mediated rejection rates from 23.5% to 6.3%, as shown in a Japanese multicenter study[[15](#_ENREF_15)].

The Milan criteria (solitary tumor < 5 cm, 2 or 3 tumors < 3 cm each, and absence of vascular invasion and extrahepatic metastasis) have been applied for the selection of candidates for liver transplantation[[16-18](#_ENREF_16)]. However, these criteria have been criticized because many patients missed opportunities for transplants because of the strictness of the criteria. Therefore, Yao *et al*[[19](#_ENREF_19)] proposed their own set of criteria, permitting the listing of patients with somewhat larger-sized tumors. In Asia, several independent criteria have also been proposed, expanding indications without increasing the risk of HCC recurrence significantly[[20-25](#_ENREF_20)] (Table 1). Five-year survival rates were as high as 80% after transplantation using these criteria. However, these criteria should be applied very carefully to DDLT candidates until a consensus is achieved.

Current issues related to “bridging therapy” and “downstaging” are discussed in the section on multidisciplinary treatments.

**Hepatic resection:** As hepatic resection is a potentially curative therapy, it has been considered a first-line option for HCC patients with well-preserved hepatic function, especially when there is only one tumor or when tumors are confined to a single lobe. To assess hepatic function, a Japanese group measured the indocyanine green retention rate at 15 min (ICG15)[[26](#_ENREF_26)]. Feasibility and the extent of the resection are decided according to the degree of retention of the dye[[27](#_ENREF_27)]. Although the BCLC algorithm mandates Child-Pugh A liver function without portal hypertension for hepatic resection, selective resection has been attempted in HCC patients exhibiting upper Child-Pugh B liver function or mild portal hypertension in Asian countries, with reference to the ICG15 value[[6](#_ENREF_6),[26](#_ENREF_26)].

Prognosis after hepatic resection is determined by number and size of tumor, vascular invasion, and level of alpha-fetoprotein[[28-30](#_ENREF_28)]. Five-year survival rates are > 50% after the resection of solitary tumors, whereas rates of 20%–30% have been reported for 3 or more nodules[[28-30](#_ENREF_28)]. With respect to tumor size, 5-year survival rates for patients with HCCs < 2 cm, 2–5 cm, and > 5 cm are 66%, 52%, and 37%, respectively[[28-30](#_ENREF_28)]. However, in selected cases with proper hepatic function, large single HCCs can be surgically removed with favorable long-term survival outcomes[[29](#_ENREF_29)]. More advanced stages of HCCs have been resected in 511 Chinese patients, yielding a 5-year survival rate of 30.5%[[31](#_ENREF_31)]. The presence of vascular invasion or extrahepatic metastasis resulted in poor outcomes[[31](#_ENREF_31)].

Recently, laparoscopic liver resection has been implemented for the treatment of HCC. This is a minimally invasive surgery, so postoperative morbidity and duration of hospitalization are reduced with no changes in surgical margin status, tumor recurrence, and overall survival[[32](#_ENREF_32)]. This technique is successfully being applied for the resection of large tumors between 5 and 10 cm and lesions at difficult-to-approach locations[[33-35](#_ENREF_33)] as well as intra-abdominal metastatic HCCs in Asian countries[[36](#_ENREF_36)].

***Interventional therapies***

**Local ablative therapies:** Local ablation can be categorized as chemical or thermal. Chemical ablation includes percutaneous ethanol injection (PEI) and acetic acid injection, whereas thermal ablation includes radiofrequency ablation (RFA), the use of microwaves, cryotherapy, and high-intensity focused ultrasound[[3](#_ENREF_3),[5](#_ENREF_5)]. As these are considered potentially curative therapies, patients with early stage HCCs are the candidates, especially when surgical treatments are not available. Among these modalities, RFA is currently the most commonly used. Excellent long-term results of RFA, up to 10 years, were reported in Korean HCC patients meeting the Milan criteria[[37](#_ENREF_37)].The results at 5 and 10 years were as follows: cumulative local tumor progression rates, 27.0% and 36.9%; cumulative intrahepatic distant recurrence rates, 73.1% and 88.5%; and overall survival rates, 59.7% and 32.3%, respectively[[37](#_ENREF_37)]. Comparison of the efficacy of RFA with other local therapies showed that RFA was substantially superior to PEI, especially in tumors with a diameter > 2 cm[[38](#_ENREF_38),[39](#_ENREF_39)]. Nevertheless, PEI is associated with a necrosis rate of 90%–100% in tumors < 2 cm and is still useful in selected patients when RFA is not technically feasible[[40-42](#_ENREF_40)]. Recently, it was reported that in cases where the tumor is located under the diaphragm or near the surface of the liver, creating artificial ascites or pleural effusion is helpful in performing RFA and avoiding burns on adjacent organs[[43](#_ENREF_43),[44](#_ENREF_44)]. This technique is being applied in several Asian countries with good results[[43](#_ENREF_43),[44](#_ENREF_44)].

Several randomized controlled trials compared the efficacy of RFA with that of resection in Asian patients with HCC meeting the Milan criteria[[45](#_ENREF_45),[46](#_ENREF_46)]. Pooled data demonstrated no significant differences in overall survival or recurrence-free survival between the treatments at 1 and 3 years. The 5-year overall survival [relative risk (RR), 0.72; 95% confidence interval (CI): 0.60–0.88] and recurrence-free survival (RR = 0.56; 95%CI: 0.40–0.78) rates were higher in the resection group[[46](#_ENREF_46)]; however, the 5-year data were provided by only one study, which advocated surgery. Complication rates were lower and hospitalization period shorter in patients who received RFA rather than resection[[46](#_ENREF_46)]. Although the efficacy of RFA appears to be comparable to that of hepatic resection with lower complication rates, additional data may be needed and the need for long-term surveillance should be re-enforced.

**Transarterial chemoembolization:** Patients with either large tumors or multinodular tumors and a good performance status are candidates for transarterial chemoembolization(TACE). However, the presence of decompensated liver disease, severe hepatic dysfunction, portal vein thrombosis, or extrahepatic tumor spread precludes TACE. Although TACE is associated with a complete response rate of only 40%, it improved survival compared with supportive treatment in 2 independently performed randomized controlled trials in Eastern and Western countries[[47](#_ENREF_47),[48](#_ENREF_48)]. A meta-analysis of 7 trials that included 545 HCC patients showed similar results [odds ratio (OR), 0.42; 95%CI: 0.20-0.88][[49](#_ENREF_49)]. Importantly, when the tumor size is ≤ 2 cm, prognosis is even better; a Korean study of TACE in small HCCs reported cumulative survival rates of 93.4%, 75.4%, 63.1%, and 51.1% at 1, 3, 5, and 8 years, respectively, for TACE, which were not significantly different from those of 97.6%, 86.7%, 74.5%, and 60.0%, respectively, for RFA[[50](#_ENREF_50)]. Therefore, TACE may have a potential role as a curative therapy for small HCCs when surgical or local ablative therapies are not feasible.

Although TACE has been contraindicated in cases of HCC with portal vein invasion, multiple studies reported that it can be safely performed and may have better survival benefits than supportive care in patients with compensated liver function[[51-54](#_ENREF_51)]. Notably, when a tumor is nodular and restricted to 1 lobe or 1–2 segments and hepatic function is classified as Child-Pugh class A, median survival after TACE is as long as 22–30 mo even in the presence of main portal vein tumor thrombosis[[51](#_ENREF_51),[52](#_ENREF_52)]. When compared with sorafenib, which is a current standard treatment for advanced HCC, median overall survival rates for TACE were not significantly different from those of sorafenib (9.2 and 7.4 mo, respectively, *P* = 0.377)[[55](#_ENREF_55)]. Therefore, TACE could be an alternative therapeutic option for advanced HCC.

TACE was originally intended to maintain intratumoral concentrations of chemotherapeutic agents by transiently obstructing supply vessels and thus minimizing systemic exposure. The strategy for TACE was recently refined after the introduction of microspheres that can increase the duration of drug retention in the tumor without blocking blood flow, which reduces hepatic derangement and systemic toxicity[[56](#_ENREF_56)]. In a multicenter phase II randomized study of 201 HCC patients, TACE with drug-eluting beads (DEB) was compared with conventional TACE, and hepatic toxicity and drug-related adverse events were significantly less observed in the DEB-TACE arm[[57](#_ENREF_57)]. Although this study showed a nonsignificant trend toward better antitumoral effects with DEB-TACE, a case-control study conducted in Korea reported a significantly better objective response rate with DEB-TACE (85%) than with conventional TACE (30%, *P* < 0.01) as assessed by modified Response Evaluation Criteria in Solid Tumors. A systemic review of the published data demonstrated the superiority of DEB over conventional TACE in terms of overall disease control, especially in patients with more advanced stage disease[[58](#_ENREF_58)]. To summarize, the indications of TACE are expanding, and new types of transarterial therapy are currently available in Eastern areas.

***Medical therapies***

**Cytotoxic chemotherapies:** Cytotoxic chemotherapy has been attempted continuously since treatment of HCC began but has failed to improve overall survival in most clinical trials to date[[59](#_ENREF_59),[60](#_ENREF_60)]. The main problem of cytotoxic chemotherapy in HCC is the co-existence of liver cirrhosis. Cirrhosis can delay the metabolism of chemotherapeutic agents and may enhance their toxicity[[61](#_ENREF_61)]. In addition, HCC is relatively chemoresistant to most cytotoxic anticancer drugs. An early randomized trial of doxorubicin conducted in Hong Kong showed a tumor response of less than 10% and borderline improvement in overall survival (10.6 weeks) compared with no treatment (7.5 wk, *P* = 0.036)[[61](#_ENREF_61)]. Notably, 25% of patients died due to doxorubicin-related complications, including septicemia and cardiotoxicity. The antitumor activity of other cytotoxic agents such as gemcitabine[[62](#_ENREF_62),[63](#_ENREF_63)], oxaliplatin[[64](#_ENREF_64)], and capecitabine[[65](#_ENREF_65)] in clinical and retrospective studies was modest with objective responses of < 20%. In randomized controlled trials, combination therapies such as PIAF (cisplatin, interferon, adriamycin, fluorouracil) and FOLFOX (5-fluorouracil, folic acid, and oxaliplatin) did not significantly improve survival compared with doxorubicin[[59](#_ENREF_59),[60](#_ENREF_60)]. Moreover, a high rate of myelotoxicity was reported in the PIAF group[[59](#_ENREF_59)]. Therefore, no cytotoxic chemotherapy regimen has provided strong evidence of improving the survival of HCC patients, and regular practice of chemotherapy is not advised. Nonetheless, a current retrospective study in Korea indicated that ECF (epirubicin, cisplatin, and 5-fluorouracil) combination therapy prolonged overall survival in sorafenib-refractory patients with metastatic HCC if a tumor response was observed; overall survival periods were 20.4 mo in responders and 4.9 mo in nonresponders (*P* < 0.001)[[66](#_ENREF_66)]. Thus, ECF may be an alternative or rescue therapy for patients who failed sorafenib therapy, but further prospective evaluations will be needed.

Hepatic arterial infusion chemotherapy (HAIC) has been used for treatment of advanced HCC with portal vein tumor thrombosis in Asian countries[[67-72](#_ENREF_67)]. Traditionally, the presence of tumor thrombus is assumed to aggravate ischemic injuries after TACE, so alternative modalities were sought. HAIC does not use embolic material, and the chemotherapeutic agent is infused into the hepatic artery *via* an implanted catheter, which reduces systemic side effects by first-pass effects and maximizes drug delivery to the tumor. Although this is considered an experimental treatment modality and is not recommended for treatment of HCC in Western countries, a large amount of clinical data on HAIC have been accumulated in Eastern countries[[67-72](#_ENREF_67)]. A small retrospective study showed survival benefits of HAIC using low doses of cisplatin and 5-fluorouracil compared with systemic cytotoxic chemotherapy or supportive care (median survival, 6, 4, and 2 mo, respectively, *P* = 0.003) in cases of advanced HCC with portal vein tumor thrombosis[[73](#_ENREF_73)]. A subsequent prospective study showed better efficacy of HAIC when a higher dose of cisplatin was used[[74](#_ENREF_74)]. Importantly, a recent retrospective study by the same group in Korea compared HAIC and sorafenib in advanced HCC patients with portal vein tumor thrombosis and showed better overall survival (7.1 and 5.5 mo, respectively, *P* = 0.011) and longer median time to progression (3.3 and 2.1 mo, respectively, *P* = 0.034) in the HAIC group[[75](#_ENREF_75)]. These findings are consistent with those of a Japanese study[[76](#_ENREF_76)]. Although well-designed prospective studies are warranted to confirm these results, HAIC at least appears to be an alternative therapy for patients with portal vein tumor thrombosis when sorafenib is not available or is intolerable. Further research is also needed regarding the use of HAIC as salvage therapy in patients with advanced HCC who do not respond to standard therapy.

**Molecular target therapies:** Sorafenib is the only approved systemic agent for the treatment of advanced HCC. It is a multikinase inhibitor whose targets include Raf-1 and B-Raf serine/threonine kinases, vascular endothelial growth factor receptor (VEGFR) and platelet-derived growth factor receptor (PDGFR) tyrosine kinases, and c-kit receptors[[77](#_ENREF_77)]. The Sorafenib Hepatocellular Carcinoma Assessment Randomized Protocol trial, which enrolled 602 patients with advanced stage HCC, showed improved median overall survival in the sorafenib group compared with the placebo group (10.7 and 7.9 mo, respectively, *P* < 0.001)[[78](#_ENREF_78)]. A subsequent study conducted in the Asia-Pacific region showed a similar trend (overall survival of 6.5 and 4.2 mo in the sorafenib and placebo groups, respectively, *P* = 0.014)[[79](#_ENREF_79)]. On the basis of the results of these trials, sorafenib became the standard treatment for advanced HCC with well-preserved liver function. The most significant adverse effects were diarrhea and hand-foot skin reactions[[78](#_ENREF_78),[79](#_ENREF_79)]. Interestingly, these toxicities were associated with better survival in patients receiving sorafenib[[80](#_ENREF_80),[81](#_ENREF_81)]. Therefore, despite the occurrence of adverse reactions, the use of sorafenib should not be discouraged when tolerable.

Sorafenib had not been compared with other treatment modalities before its approval. Currently, its efficacy in a real-life setting was compared with the efficacy of other treatments (TACE, radiation, and cytotoxic chemotherapy) in Korean patients with advanced HCC[[82](#_ENREF_82),[83](#_ENREF_83)]. Overall survival times were 8.4 and 8.2 mo for sorafenib and other treatments, respectively, and the difference was not significant[[82](#_ENREF_82),[83](#_ENREF_83)]. To improve the efficacy of sorafenib, combination therapy or a multidisciplinary approach may be needed[[84](#_ENREF_84)].

Several newer molecular target therapeutic agents were evaluated in clinical trials. Sunitinib, an orally administered multikinase inhibitor of receptor tyrosine kinases, showed modest activity against HCC. Although an overall survival time of 9.8 mo was observed in a phase II study[[85](#_ENREF_85)], sunitinib did not outperform sorafenib in a phase III randomized study (overall survival, 8.1 and 10.0 mo, respectively, *P* = 0.0019)[[86](#_ENREF_86)]. Brivanib, a selective dual inhibitor of fibroblast growth factor (FGF) and vascular endothelial growth factor (VEGF) signaling, was associated with a median overall survival of 10 mo in a phase II trial[[87](#_ENREF_87)] and was considered a promising new drug for advanced HCC. However, the primary endpoint of brivanib not being non-inferior to sorafenib was not met in a subsequent phase III trial (overall survival, 9.5 and 9.9 mo, respectively, *P* value is nonsignificant)[[88](#_ENREF_88)]. The efficacy of brivanib in advanced HCC patients who were intolerant to sorafenib or failed to respond to sorafenib previously was also tested. The results of this study showed no significant improvement in overall survival compared with placebo (9.4 and 8.2 mo for brivanib and placebo, respectively)[[89](#_ENREF_89)]. Linifanib (ABT-869), a receptor tyrosine kinase inhibitor targeting VEGFRs, also failed to significantly improve survival compared with sorafenib in a phase III trial (overall survival, 9.1 and 9.8 mo, respectively)[[90](#_ENREF_90)]. The reasons for most of these novel agents failing to improve survival may be diverse and include lack of understanding of critical drivers of cancer progression, unpredicted toxicity, and marginal antitumor effects[[91](#_ENREF_91)]. To overcome these obstacles, the clinical trial design should be modified to focus on biomarker-based subpopulation targeting strategies, and thereby, personalized therapies should be pursued in the future. In addition, efficacy and toxicity need to be evaluated in detail in phase I and II studies before moving to phase III studies[[91](#_ENREF_91)]. Currently, several novel molecular targeting agents are being evaluated in phase III trials as first-line or second-line therapies including lenvatinib [VEGFR1–3, FGF receptor (FGFR) 1–3, PDGFR-β, RET, KIT], ramucirumab (VEGFR2), regorafenib (VEGFR, TIE-2, PDGFR-β, FGFR, KIT, RET, RAF), cabozantinib (MET, VEGFR-2), and tivantinib (MET)[[92](#_ENREF_92)]. Table 2 summarizes the current status of the randomized controlled trials of molecular target therapies.

In summary, there are no currently available first-line molecular targeted agents other than sorafenib and no standard second-line treatments for patients intolerant or nonresponsive to sorafenib. If underlying liver function is well preserved, novel molecular target therapies, HAIC, or systemic cytotoxic chemotherapy may have a role as second-line treatment, but further studies are warranted. If a patient with advanced HCC has poor hepatic function, aggressive anticancer treatments are not indicated.

***Radiotherapies and emerging therapies***

**External radiation therapies:** Radiotherapy techniques for the treatment of HCC have substantially evolved over the past decades. Delivery of radiation energy became more precise, which enabled the exposure of tumors to higher doses of radiation, while saving non-tumorous liver parenchyma[[93](#_ENREF_93)]. In the past, the role of radiation therapy was limited to alleviation of bone pain due to bone metastasis and to emergency use in spine and brain metastasis[[94-96](#_ENREF_94)]. Radiation therapy has currently been adopted as a definitive therapy with curative intent if the tumor is at an early stage. Particularly, stereotactic body radiation therapy can achieve high rates of locoregional tumor control as it can deliver high doses of radiation in a single treatment session or in a small number of fractions[[97](#_ENREF_97),[98](#_ENREF_98)]. In locally advanced HCCs, radiation therapy can be used to relieve obstruction and improve portal blood flow if the tumor invades the biliary tree or portal vein[[99](#_ENREF_99),[100](#_ENREF_100)]. A large multicenter study in Korea of 994 HCC patients with portal vein tumor thrombosis showed a median survival of 9.2 mo[[101](#_ENREF_101)]. This was a relatively longer survival time than that of advanced HCC patients who did not receive any treatment in previous trials[[78](#_ENREF_78),[79](#_ENREF_79)]. Studies from Japan and China also reported the efficacy of radiotherapy for HCC with portal vein thrombosis, and overall survival was significantly better in patients receiving radiotherapy than in patients receiving sorafenib (10.9 and 4.8 mo, respectively, *P* = 0.025) or undergoing surgery (12.3 and 10.3 mo, respectively, *P* = 0.029)[[102](#_ENREF_102),[103](#_ENREF_103)]. Although these studies were retrospective, they suggest the usefulness of radiotherapy in advanced HCC. However, radiotherapy has not been incorporated into the international guidelines for HCC despite its efficacy. This may be attributed to the paucity of well-designed randomized controlled studies, which are urgently needed. In addition, guidelines for optimal dose fractionation and protocols for avoiding radiation toxicity should be further established[[93](#_ENREF_93)].

Proton beam therapy (PBT) can dramatically reduce damage to surrounding liver tissue by modulation of the Bragg peak of protons in energy and time, and thereby, maximizes the effects of radiation on the tumor. In Eastern areas, studies of PBT in HCC patients have been reported mainly by Japanese groups[[104-106](#_ENREF_104)]. A retrospective study of PBT in 162 surgically unresectable patients reported a local control rate of 89% and an overall survival rate of 23.5% at 5 years[[106](#_ENREF_106)]. Although the tumor stages of the patients were diverse and TACE or PEI may have also been administered, the overall efficacy seems quite favorable. PBT showed a good response rate even for large tumors (> 10 cm) and HCCs with main portal tumor thrombosis[[107](#_ENREF_107),[108](#_ENREF_108)].

**Radioembolization:** Radioembolization is a modality involving the use of a transarterial approach to the hepatic tumor and subsequent infusion of radioactive substances. The rationale for this approach is that the efficacy of external beam radiation therapy is limited by the low tolerability of cirrhotic livers leading to radiation hepatitis or decompensation. To avoid exposure of non-tumorous parenchyma to radiation, microspheres emitting high-energy and low-penetration radiation are selectively delivered to the tumor[[109](#_ENREF_109)]. The most commonly used radioembolic agents are iodine-131 and yttrium-90 glass beads, both of which showed favorable antitumoral effects with an acceptable safety profile[[109](#_ENREF_109),[110](#_ENREF_110)]. The benefits of radioembolization over the other types of transarterial therapies still need to be validated. A retrospective analysis showed no significant differences in efficacy between radioembolization and TACE for intermediate stage HCC; median survival times were 15.0 and 14.4 mo, respectively[[111](#_ENREF_111)]. However, patients receiving radioembolization needed less hospitalization and fewer treatments. Fewer treatment sessions should improve quality of life and reduce the possibility of liver derangement; therefore, in these respects, radioembolization is considered better than conventional TACE. The efficacy of radioembolization in patients with advanced HCC patients has also been evaluated. Sixty- three patients with portal vein thrombosis were analyzed from an European HCC cohort according to underlying liver function[[112](#_ENREF_112)]. Median overall survival and time to progression were 13.8 and 5.6 mo, respectively, for Child-Pugh A patients and 6.5 and 4.9 mo, respectively, for Child-Pugh B patients[[112](#_ENREF_112)]. Although these data appear very promising, there are still no randomized controlled trials comparing radioembolization with standard treatments for each stage. Data from Asian countries are limited, but a multicenter prospective study in Korea showed a median time to progression of 18 mo and a 3-year survival rate of 75%[[113](#_ENREF_113)]. This is an improved result compared with data from Western countries[[114](#_ENREF_114),[115](#_ENREF_115)], but future well-designed studies are needed.

**Emerging therapies:** Recently, the oncolytic and immunotherapeutic vaccinia virus has been reported to induce antibody-mediated, complement-dependent cancer cell lysis in humans[[116](#_ENREF_116)]. Immunotherapy may benefit patients with advanced stage HCC who do not have further treatment options. The results of a phase III trial need to be confirmed.

Currently, several target delivery systems has been exploited for the treatment of HCC. New formulations including polymetric nanoparticles, nanocapsules, liposomes, nanoemulsions, microsphere, and polymeric micelles have been reported[[117](#_ENREF_117),[118](#_ENREF_118)]. Novel drug delivery systems are expected to improve treatment efficacy and to decrease toxicity by drug targeting to the specific site of action[[118](#_ENREF_118)]. For example, the asialoglycoprotein (ASPG) receptor is expressed on hepatocyte, and a synthetic ligand, lactosylated liposomes can be used for effective delivery vehicles of doxorubicin in HCC therapy[[119](#_ENREF_119)]. In a previous report, lactosylated liposomes encapsulating doxorubicin showed stronger anti-tumor response than the non-targeted liposomal doxorubicin and free doxorubicin. A galactose ligand with chitosan modifications, galactosylated chitosan, is also a promising carrier of chemotherapeutic agent, such as 5-fluorouracil, to the ASPG receptor, and its *in* *vitro* and *in vivo* efficacy was well described[[120](#_ENREF_120)]. It is thought that efficacy of anticancer therapy utilizing target delivery system will be more synergized by combination of molecular target therapy. Further studies are warrantied.

**MULTIDISCIPLINARY TREATMENT BEYOND TREATMENT ALGORITHMS**

As treatment modalities for HCC are very diverse, not only hepatologists but also surgeons, intervention radiologists, medical oncologists, and radiation oncologists should jointly discuss the best treatment options for HCC patients. Treatment may not necessarily be a sole modality; combinations of multiple treatments can be considered. Although current guidelines do not recommend multiple treatments, emerging data indicate better outcomes with multidisciplinary treatments for HCC. Furthermore, several newer clinical trials aim to properly evaluate such strategies. In this context, multimodality treatment options based on currently available evidence, especially from Eastern countries, are described in this section, according to HCC stage.

***Very early or early stage***

Guidelines recommend liver transplantation, hepatic resection, or RFA/PEI for very early or early stage HCC[[3-5](#_ENREF_3)]. However, treatment may be diversified according to the status of the patient or the tumor. In addition to the single treatments described above, the following treatments can be considered as an adjuvant or a combination.

**Bridging therapy for liver transplantation:** Although the best outcome can be achieved with liver transplantation, HCCs may progress while patients are on the waiting list. To avoid dropout, the rate of which approaches 20%, bridging therapies may be needed[[121](#_ENREF_121)]. Most commonly applied therapies are RFA, TACE, and surgical resection, although data from Asian countries are rather sparse. If the tumor is within the Milan criteria and liver function is not decompensated, RFA should be the first bridging therapy attempted because its post-procedural intratumoral necrosis rate is higher than that of other locoregional therapies, and it is associated with the lowest drop-out rate[[122](#_ENREF_122)]. PEI appears to be less efficacious than RFA, but can be chosen if the lesions are close to adjacent organs, where RFA is dangerous to perform. If the tumor size is > 3 cm, TACE or TACE plus RFA may be favored as tumors become more vascularized and the effect of RFA may be diminished[[121](#_ENREF_121)]. Surgical resection can precede liver transplantation, and salvage transplantation can be performed in the event of recurrence, without a decrease in overall post-transplant survival[[123](#_ENREF_123)]. However, most data on bridging therapy data are uncontrolled, so it is difficult to strongly recommend this therapeutic strategy, especially if patients are eligible for LDLT.

**Adjuvant therapy after resection:** Hepatic resection is the preferred treatment for patients with early stage tumors and well-compensated liver function, but recurrence is the main obstacle to improving long-term prognosis. To reduce the recurrence rate, various neoadjuvant and adjuvant therapies were evaluated, but they failed to demonstrate any benefits to recurrence-free survival[[124-126](#_ENREF_124)]. Recently, sorafenib was tested for prevention of recurrence after curative therapy, including resection and ablation (the STORM study), but again no benefits to recurrence-free survival were observed[[127](#_ENREF_127)]. A Japanese group previously showed positive effects of acyclic retinoids and vitamin K analogues on recurrence-free survival, but overall survival was not improved and large-scale studies were not performed appropriately[[128](#_ENREF_128),[129](#_ENREF_129)]. Interferon has been suggested as an adjuvant therapy after resection[[130](#_ENREF_130)]. According to a large current database in Taiwan, antiviral therapies reduce the recurrence of HCC after surgery in patients with chronic hepatitis B or C[[131](#_ENREF_131),[132](#_ENREF_132)]. In agreement, a randomized controlled trial conducted in China showed better recurrence-free survival (RR = 0.651; 95%CI: 0.451–0.938) and overall survival (RR = 0.420; 95%CI: 0.271–0.651) in patients receiving antiviral therapy, especially in terms of prevention of late recurrence[[133](#_ENREF_133)]. It would be reasonable to recommend nucleoside or nucleotide analogues or interferon therapy to patients with hepatitis B and pegylated interferon-based therapy to patients with hepatitis C after curative hepatic resection[[130-133](#_ENREF_130)].

**RFA/PEI combined with TACE:** Local ablative therapies, which are curative modalities for HCC as mentioned previously[[3](#_ENREF_3),[5](#_ENREF_5)], have been very useful in the treatment of patients reluctant to undergo or ineligible for surgery because of issues other than liver diseases. As the size of the tumor limits the efficacy of RFA or PEI, the combination of RFA or PEI and vaso-occlusive therapies such as TACE has been attempted to overcome the limitations of interventional therapies and to maximize synergistic effects[[134](#_ENREF_134)]. A retrospective study conducted in Korea evaluated the therapeutic efficacy of RFA plus TACE in patients with medium-sized (3.1–5.0 cm) HCCs and found that it significantly lowered the local tumor progression rate compared with RFA alone (55% and 86% at 5 years, respectively, *P* < 0.001)[[135](#_ENREF_135)]. Subsequently, several randomized controlled trials compared RFA and RFA plus TACE in Japan and China[[136](#_ENREF_136),[137](#_ENREF_137)], and a meta-analysis of these studies showed that the combined treatment was significantly associated with higher overall survival (OR = 1.85; 95%CI: 1.26–2.71) and recurrence-free survival (OR = 2.13; 95%CI: 1.41–3.20) rates[[138](#_ENREF_138)]. The benefits of the combination therapy could be attributed to the avoidance of the heat sink effect and the subsequent increase in the size of the thermal coagulation zone. In addition, synergism between hyperthermia and high concentrations of chemotherapeutic agents may enhance the destruction of microscopic satellite lesions. Recently, a non-randomized controlled study compared RFA plus TACE and surgical resection for the treatment of single HCCs ranging in size from 2 to 5 cm[[139](#_ENREF_139)]. The study showed that the combination therapy was as effective as resection in terms of recurrence-free survival (69.4% and 65%, respectively, at 4 years, *P* value is nonsignificant) and overall survival (78.4% and 80.3%, respectively, at 4 years, *P* value is nonsignificant). Collectively, the available data suggest that RFA plus TACE provides better outcomes than RFA alone and may be as efficacious as surgical resection for medium-sized HCCs.

PEI may be also combined with TACE, thereby peripheral micrometastasis will be better controlled and diffusion of the ethanol can be more facilitated compared with PEI alone[[140](#_ENREF_140)]. This combination has been reported to be associated with superior efficacy in terms of local control, but no survival benefits compared with PEI monotherapy have been confirmed[[141](#_ENREF_141)].

***Intermediate stage***

Most HCCs beyond the Milan criteria correspond to intermediate stage HCCs if vascular invasion and distant metastasis are absent. TACE is the recommended therapy for this stage[[3](#_ENREF_3),[5](#_ENREF_5)], but the beneficial effects of TACE on long-term survival are limited. Therefore, further treatment would be necessary even in the presence of an initial tumor response.

**Liver transplantation after downstaging:** As liver transplantation is associated with the best treatment outcome of HCC, listing patients for transplantation should be considered whenever available; patients in the intermediate stage will be eligible if effective treatment was achieved and their HCC status was shifted to meet the Milan criteria[[142](#_ENREF_142)]. TACE is the most commonly used modality for downstaging, and local ablative therapies may be combined[[142](#_ENREF_142)]. The expected 5-year overall survival rate in patients who received liver transplants after downstaging is comparable to that of HCC patients who met the Milan criteria without downstaging[[143](#_ENREF_143)]. However, the 5-year disease-free survival rate is lower in the downstaged patients[[143](#_ENREF_143)]. Stricter follow-ups would be necessary for these patients.

**TACE combined with RFA or radiotherapy:** Combination therapy with RFA or PEI and TACE is being used to treat early stage HCC as mentioned earlier. This therapy can also be applied to intermediate stage HCC. However, tumor may be too extensive or multiple to combine ablative therapies in intermediate stages. For best results, modalities commonly used for more advanced stage HCC can be adopted in combination with TACE (*e.g.*, sorafenib and radiotherapy). The efficacy of TACE plus radiotherapy has been studied in 12 non-randomized and 5 randomized controlled trials in Korea, Japan, and China. A meta-analysis of these trials showed significantly improved survival at 1 year (OR = 2.23; 95%CI: 1.76–2.83) and 5 years (OR = 4.47; 95%CI: 2.08–9.61) and a better tumor response (OR = 2.58; 95%CI: 1.64–4.06) in patients receiving TACE plus radiotherapy compared with patients receiving TACE alone[[144](#_ENREF_144)]. In this analysis, most although not all patients in the individual studies had intermediate stage HCC. Therefore, the combination of TACE and radiotherapy should be beneficial for this stage, but consensus is needed for routine recommendation in practice guidelines.

**TACE combined with sorafenib:** TACE may upregulate circulating VEGF, which is associated with vascular invasion, tumor growth, metastasis, and poor survival. Therefore, control of VEGF signaling and of other tumor growth factors is necessary to prevent the progression and recurrence of HCC in patients receiving TACE[[145](#_ENREF_145)]. A randomized controlled trial was conducted in Japan and Korea to assess the effects of TACE plus sorafenib[[146](#_ENREF_146)]. In that study, time to progression was significantly longer in Korean patients receiving TACE plus sorafenib than in those receiving TACE alone, but not in Japanese patients. To clarify the clinical results, a meta-analysis of 6 studies was performed, and the pooled results showed that overall survival [hazard ratio (HR) = 0.65; 95%CI: 0.47–0.89] and time to progression (HR = 0.68; 95%CI:, 0.52–0.87) were significantly longer in patients who received TACE plus sorafenib than in patients who received TACE only[[147](#_ENREF_147)]. Another recent meta-analysis of 9 studies mostly from China reached the same conclusions[[148](#_ENREF_148)]. These results indicate that appropriate combination therapies will improve clinical outcomes in patients with unresectable HCCs.

***Advanced stage***

This stage encompasses locally advanced HCCs with vascular invasion and HCCs with extrahepatic metastasis. Whether the tumor has advanced locally or distantly, the BCLC guidelines uniformly recommend treatment with sorafenib. Although survival benefits were observed compared with no treatment, the efficacy of sorafenib is limited[[78](#_ENREF_78),[79](#_ENREF_79)]. Therefore, it would be appropriate to search for more effective methods. For instance, sorafenib may be combined with other type of therapies (*e.g*., TACE, radioembolization, and external radiation) and TACE or HAIC may be combined with radiotherapy in patients with advanced HCC.

**Sorafenib combined with TACE:** A relatively large retrospective study compared the efficacy of TACE plus sorafenib and sorafenib alone in 355 advanced stage HCC patients (164 and 191 patients, respectively) [[149](#_ENREF_149)]. Overall survival was significantly longer in the combination group than in the sorafenib monotherapy group (8.9 and 5.9 mo, respectively, *P* = 0.009) as was median time to progression (2.5 and 2.1 mo, respectively, *P* = 0.008). The difference in time to progression was still significant after propensity score matching, whereas the difference in overall survival was not. Another study compared the efficacy of TACE plus sorafenib and TACE alone in 246 advanced stage HCC patients (82 and 164 patients, respectively) after propensity score matching. Overall survival was significantly longer in the combination group than in the TACE monotherapy group (7.0 and 4.9 mo, respectively, *P* = 0.003) as was time to progression (2.6 and 1.9 mo, respectively, *P* = 0.001)[[150](#_ENREF_150)]. These data suggest that the combination of TACE and sorafenib is most likely more efficacious than either therapy alone in advanced HCC. Other types of transarterial therapies, including DEB-TACE or radioembolization, are emerging modalities for the treatment of advanced HCC as mentioned above[[112](#_ENREF_112),[151](#_ENREF_151)]. To potentially improve their efficacy, combining new modalities with sorafenib are being evaluated in clinical trials[[152](#_ENREF_152),[153](#_ENREF_153)]; a phase II study which combined DEB-TACE with sorafenib showed objective response rate of 58% and disease control rate of 100% in advanced HCC patients[[152](#_ENREF_152)]. The combination is a promising HCC treatment strategy considering the current data, but its benefits compared with monotherapy needs to be confirmed in a future phase III trial.

**Sorafenib combined with radiotherapy:** Sorafenib was reported to enhance the radiosensitivity of human HCC cell lines by inhibiting radiation-induced activation of VEGFRs, a downstream kinase (extracellular signal-regulated kinase), and nuclear factor-κB and by increasing radiation-induced apoptosis[[154](#_ENREF_154)]. Therefore, combining sorafenib and radiotherapy, in the form of either radioembolization or external beam radiation, is expected to be synergistic. A multicenter phase II study evaluated safety and efficacy of combining sorafenib therapy and radioembolization in several Asian-Pacific countries[[155](#_ENREF_155)]. Sorafenib was administered after radioembolization, and the median overall survival time was 8.6 mo in patients with advanced stage HCC[[155](#_ENREF_155)]. Most of toxicities were associated with sorafenib therapy. Considering phase III Asian-Pacific trial data of sorafenib which showed median survival time of 6.5 mo in advanced HCC, the data of radioembolization plus sorafenib combination therapy appears favorable[[79](#_ENREF_79)]. Data of sorafenib plus external beam radiation are emerging, recently. A phase II study of sorafenib therapy plus external beam radiation reported an initial complete or partial response rate of 55% and a 2-year overall survival rate of 32% in 40 Taiwanese patients with advanced HCC[[156](#_ENREF_156)]. These efficacy data seem encouraging, but further investigations are warranted.

**TACE combined with radiotherapy:** As mentioned in the intermediate stage section, TACE plus radiotherapy is an effective synergistic strategy. Most of the previous randomized and non-randomized clinical trials of TACE plus radiotherapy included both intermediate stage and advanced stage HCC[[100](#_ENREF_100),[144](#_ENREF_144)], whereas studies of advanced stage HCC only are few[[157](#_ENREF_157),[158](#_ENREF_158)]. A retrospective study assessed outcome of patients with locally advanced HCC; 27 patients who were treated with TACE plus radiotherapy and another 27 patients who received sorafenib alone were compared after propensity score matching. Interestingly, overall survival was better in the former group than in the latter one (6.7 and 3.1 mo, respectively, *P* < 0.001)[[158](#_ENREF_158)]. Although this was a small study, the results iterate that universal application of sorafenib for advanced stage patients may not be the best option. Further well-designed studies are warranted.

**HAIC combined with radiotherapy:** To facilitate the efficacy of HAIC for treatment of advanced HCC with portal vein thrombosis, radiotherapy may be combined. In a previous pilot clinical trial, infusion of 5-fluorouracil was performed at 1st and 5th weeks of radiotherapy, and then continued every 4 wk. Objective response rate was 45% and median survival time was 13.1 mo[[99](#_ENREF_99)]. More recently, HAIC combined with radiotherapy was compared with HAIC in advanced HCC patients, and the combination therapy was shown to be better than HAIC monotherapy in terms of time to progression (5.0 and 2.7 mo, respectively, *P* = 0.0024) and overall survival (8.6 and 5.0 mo, respectively; *P* = 0.0002), particularly, among the HAIC non-responders[[159](#_ENREF_159)]. Although, these studies are retrospectively performed, HAIC combined with radiotherapy appears to have more benefits than monotherapy, suggesting synergistic effects of the therapies.

**Sequential therapy with metastasectomy:** In cases of extrahepatic metastasis, radiotherapy is considered if the lesions cause severe symptoms[[160](#_ENREF_160)]. Metastatic lesions are often surgically removed if: (1) liver function is well preserved; (2) intrahepatic lesions are adequately controlled by surgery or locoregional therapy; and (3) extrahepatic metastatic lesions are confined to a single organ[[161-164](#_ENREF_161)]. Studies from Asian countries showed 5-year survival rates ranging from 26% to 37% in HCC patients with lung metastasis who underwent metastasectomy[[162-164](#_ENREF_162)], which are surprising survival data at the patients’ tumor stage despite the selection of surgical candidates. Although sorafenib became currently the first-line treatment for HCC with distant metastasis, uniform application of sorafenib monotherapy does not seem to be the best way because of its low objective response rate. Therefore, when possible, treating intrahepatic and metastatic lesions *via* metastasectomy, locoregional therapy or radiotherapy before the administration of sorafenib would be a reasonable plan[[165](#_ENREF_165)]. However, because no data exist regarding this strategy, it needs to be evaluated further in the future.

***Terminal stage***

At this stage, best supportive care (BSC) is recommended. BSC includes management of cirrhotic complications such as ascites, hepatic encephalopathy, variceal hemorrhage, and hepatorenal syndrome. Another important aspect would be management of cancer pain. Indeed, pain management has been frequently neglected at many tertiary hospitals in Asian countries. However, a systematic approach to cancer pain control is important.

Non-opioid drugs (paracetamol) and mild opioids (codeine, tramadol, and dihydrocodeine) may be useful for mild to moderate pain if administered on a regular basis[[166](#_ENREF_166)]. Nonsteroidal ant-inflammatory drugs, which can cause renal derangement, should be avoided. Transdermal patches (fentanyl and buprenorphine) are considered if patient’s requirements of opioid are stable[[166](#_ENREF_166)]. Breakthrough pain can be managed with rapid-acting rescue therapies administered via intravenous or subcutaneous routes. Strong opioids (morphine, oxycodone, hydromorphone, oxymorphone, and fentanyl) are used to control severe pain[[166](#_ENREF_166)], but expose the patient to the risk of developing hepatic encephalopathy. Hence, close monitoring is essential. Emotional and nutritional support is also important for terminal stage care, so collaboration between the hospice team and the clinical nutrition team would be helpful[[167](#_ENREF_167)]. Until the final round, a multidisciplinary approach should be maintained.

***Need of updated staging system***

With the advancement of therapeutic modalities and aggressive treatment by either mono- or combination therapy as reviewed so far, the prognosis of HCC has improved remarkably; survival benefits are better observed in more advanced stage diseases. In this regard, reevaluation of preexisting staging systems and refinement of the best-fit models have been performed in Korea, Japan, Taiwan, and China[[168-171](#_ENREF_168)]. Most recently, a newer staging system was pronounced from a single center in Hong Kong, reflecting recent improved survival outcomes in subsets of intermediate and advanced stage patients with more radical therapies[[172](#_ENREF_172)]. In addition, the Hong Kong Liver Cancer staging was better than BCLC staging in stratifying HCC patients with different prognostic groups. Although further validation may be needed in non-Asian patients, the system will be helpful for identifying patients who are suitable for more aggressive treatments than what BCLC staging system recommends.

**CONCLUSION**

There is an increasing demand that international HCC treatment guidelines should be updated properly. Still, combinations of treatment modalities have not been incorporated into recent guidelines, and there are several unmet needs. Treatment intervals, strategies in the event of recurrence, and the timing of retreatment have not been properly studied, and no established recommendations are available. Therefore, to further improve the outcomes of HCC patients, strategies for surveillance, diagnosis, initial treatment, recurrence monitoring, and treatment after recurrence should be more organized. Close collaboration between specialists in multiple fields is of utmost importance in achieving these aims.

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**P-Reviewer:** Decena Sollano JD, Yu CY **S-Editor:** Yu J **L-Editor:** **E-Editor:**

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**Figure 1 Expanding indications of treatment modalities for hepatocellular carcinoma.** Indications of each treatment are currently expanding and treatments may be combined as determined by clinical situations. RFA: Radiofrequency ablation; TACE: Transarterial chemoembolization; HAIC: Hepatic arterial infusion chemotherapy; BSC: Best supportive care.

**Table 1 Expanded criteria for liver transplantation proposed in Eastern countries**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Criteria (city, country, reference)** | **Tumor number** | **Tumor diameter** | **Additional criteria** |  | **Overall survival****within criteria** |
| Hong Kong, China[20] | 1≤ 3 | ≤ 6.5 cm≤ 4.5 cm | No diffuse type,no vascular invasion |  | 3 years5 years | 78%66% |
| Hangzhou, China[21] | NC | Total ≤ 8 cm | Histopathologic grade I or II with AFP ≤ 400 ng/dL if tumor > 8 cm |  | 3 years5 years | 70.7%70.7% |
| Seoul (AMC), Korea[22] | ≤ 6 | ≤ 5 cm | No gross vascular invasion |  | 3 years5 years | 87.5%81.6% |
| Seoul (CMC),Korea[23] | ≤ 7 | ≤ 7 cm | NC |  | 5 years | 86.3% |
| Tokyo, Japan[24] | ≤ 5 | ≤ 5 cm | NC |  | 3 years5 years | 82%75% |
| Kyoto, Japan[25] | ≤ 10 | ≤ 5 cm | PIVKA-II ≤400 mAU/mL |  | 5 years | 87% |

AMC: Asan Medical Center; CMC: Catholic Medical Center; AFP: alfa-fetoprotein; PIVKA II: Protein induced by vitamin K absence or antagonist-II; NC: Not commented.

**Table 2 Randomized controlled trials with molecular target therapies**[92]

|  |  |
| --- | --- |
| **First line** | **Status** |
| Comparison with placeboSorafenib (SHARP, Asian-Pacific)Sorafenib in Child B (BOOST)Comparison study between sorafenib and single agent (head to head) | Proven benefitPhase III Ongoing |
|  Sunitinib -> endpoint not met | Terminated |
|  Brivanib (BRISK-FL) -> endpoint not met | Failed |
| Linifanib -> endpoint not metLenvatinib  | TerminatedPhase III ongoing |
| Combination of sorafenib and another agent |  |
|  Sorafenib + Erlotinib (SEARCH) -> endpoint not met | Failed |
| Sorafenib + Doxorubicin (CALGB-80802)Sorafenib + Everolimus | Phase III ongoingR-Phase II; Failed |
| **Second line** |  |
| Sorafenib failure |  |
| Brivanib (BRISK-PS) -> endpoint not metBrivanib (BRISK-APS)Everolimus (EVOLVE-1) -> endpoint not metRamucirumab (REACH)Regorafenib (RESORCE)Cabozantinib (CELESTAL) Tivantinib (Metiv-HCC) | FailedTerminatedFailedPhase III ongoingPhase III ongoingPhase III ongoingPhase III ongoing |
| Combination or addition to standard therapies |  |
| Adjuvant setting after surgery or RFA: sorafenib (STORM) | Failed |
| Combination with TACE: sorafenib (SPACE) -> endpoint not metbrivanib (BRISK-TA) -> endpoint not metsorafenib (TACTICS)  | FailedFailedR-Phase II ongoing |

RFA: Radiofrequency ablation; TACE: Transarterial chemoembolization.