Review

Treatment Strategy of Transarterial Chemoembolization for Hepatocellular Carcinoma

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Abstract: Transarterial chemoembolization (TACE) is a first-line treatment for patients with hepatocellular carcinoma (HCC) in Barcelona Clinic Liver Cancer stage B (BCLC-B). There are two major techniques of TACE: conventional TACE (cTACE) using iodized oil and gelatin sponge particles, and TACE using drug-eluting beads (DEB-TACE). The latest randomized controlled trial proved the superiority of cTACE regarding local effects over DEB-TACE; however, cTACE also damages the liver more severely. Therefore, cTACE should be performed for localized HCCs as selectively as possible. On the other hand, DEB-TACE has less liver toxicity and is favorable for patients with an advanced age, large and/or bilobar tumors, or a poor liver function. However, some BCLC-B HCCs are TACE-resistant and the concept of TACE unsuitability (mainly up-to-7 criteria out) has been proposed by Asia-Pacific Primary Liver Cancer Expert Meeting. Systemic therapy is recommended for patients with TACE-unsuitable HCC; however, the condition of TACE-unsuitable HCC does not always rule out TACE monotherapy and some up-to-7 criteria out tumors may also be good candidates for superselective cTACE when localized in limited liver segments. The sequential therapy of an antiangiogenic and TACE is also a novel option for patients with TACE-unsuitable HCC, antiangiogenic-refractory HCC, or even down-staged HCC.

Keywords: hepatocellular carcinoma; transarterial chemoembolization; iodized oil; gelatin sponge particle; drug-eluting bead; chemotherapeutic; systemic therapy; antiangiogenic; treatment strategy

1. Introduction

Since the first report by Yamada et al. [1], transarterial chemoembolization (TACE) has been widely performed for inoperable hepatocellular carcinoma (HCC). In 2002, two randomized controlled trials (RCT) demonstrated that TACE for inoperable HCC showed a survival benefit compared with best supportive care (BSC) [2,3]. Two meta-analyses also proved the clinical usefulness of TACE in prolonging HCC patients’ survival [4,5]. Therefore, TACE has been recognized as an effective treatment option for inoperable HCC worldwide, and as a first-line treatment for HCC in Barcelona Clinic Liver Cancer stage B (BCLC-B) (Figure 1) [6].

Now, two major techniques of TACE have been performed: conventional TACE (cTACE) with iodized oil (Lipiodol 480, Guerbet Japan, Tokyo, Japan) and gelatin sponge (GS) particles; and TACE with drug-eluting beads (DEB-TACE). cTACE has been mainly developed in Asian regions, whereas DEB-TACE has been focused in Western countries. The concepts and rationales for both techniques, as well as adverse effects, are quite different; therefore, we should use them appropriately according to the patient and tumor condition. Thus, the purpose of this article is to describe the TACE strategy for BCLC-B HCC, mainly based on the conceptualistic position and techniques of TACE treatment in Japan.
2. Hemodynamics in the Liver and HCC

Understanding of the hemodynamics in the liver and HCC is very important to perform TACE procedures safely and effectively.

2.1. Microvasculature of the Normal Liver

Two major branches are arising from the terminal hepatic artery: one terminates within the portal tract supplying the bile duct (peribiliary vascular plexus: PBP), portal tract interstitium, and portal vein wall (portal vein vasa vasorum); and the other one penetrates the liver parenchyma unaccompanied by the portal vein or bile duct, named the isolated artery. Most terminal hepatic arterioles connect with the PBP, and blood from the PBP directly drains into the portal venules and hepatic sinusoids. Therefore, bile duct injury can develop following TACE when the PBP is severely damaged. The incidences of bile duct injury are usually high in the normal liver compared with the cirrhotic liver because the PBP is more hypertrophied in the cirrhotic liver. The isolated artery communicates with the hepatic capsular arterial plexus, as well as with extrahepatic arteries, such as the internal mammary and inferior phrenic arteries, through the ligaments in the bare area of the liver. This communication can help the tumor mainly in the bare area of the liver to survive after TACE and promote the tumor recurrence.

The normal liver parenchyma is supplied by a dual blood supply, including the portal venous system and the hepatic arterial system; thus, embolization of the hepatic artery can theoretically achieve selective ischemic necrosis of the tumor tissues alone. However, hepatic necrosis can also develop when superselective cTACE is performed.

2.2. Hemodynamics in Hypervascular HCC

Hemodynamics in the hepatocytic nodules can change according to the grades of malignancy of the nodules. During multistep hepatocarcinogenesis, the intranodular arterial supply initially decreases, but then acutely increases because of the increase in abnormal arterial supply. On the other hand, the intranodular portal supply gradually decreases, and finally disappears in moderately to poorly differentiated HCC tissue. Therefore, overt HCC is predominantly supplied by arterial blood; however, some tumor tissues are still partially supplied by portal blood, such as well-differentiated tumor portions and tumors invading the surrounding liver through the capsule (capsular/extracapsular invasion) mainly located in the periphery of the tumor, as well as microsatellite lesions.
Simultaneously, the hepatic veins surrounding HCC are also occluded by tumor invasion and the tumor blood drains into the peritumoral area through the portal vein remnant in the capsule, depicted as corona enhancement on computed tomography (CT) during hepatic arteriography (CTHA) [16,17].

3. Limitation of TACE and Necessity of Curative TACE

3.1. Limitation of TACE

As mentioned above, well-differentiated tumor portions, capsular and/or extracapsular tumor invasions, and microsatellites are usually supplied by a dual blood supply; therefore, they can survive after simple embolization of the arterial side, particularly when it is performed non-selectively [15,18]. Additionally, there are two possible causes of local tumor recurrence after TACE. One important cause is portal blood supply to tumors via the portal venules and surrounding hepatic sinusoids due to the reversal of portal flow through the tumor drainage appearing during blockage of the hepatic arterial flow [18,19]. The other cause is arterial collateral supply to tumors from the neighboring arteries, sometimes from the extrahepatic arteries through the communication between the hepatic capsular arterial plexus and isolated artery [10].

3.2. Necessity of Curative TACE

TACE is recognized as a palliative treatment option for HCC; however, it is reported that a complete response (CR) after the initial TACE is the most robust predictor of long-term survival [20]. In other words, ineffective TACE may shorten a patient’s lifespan. TACE loads hypoxic and chemotherapeutic stress on HCC, and some surviving tumors frequently change to more aggressive and TACE-resistant tumor tissues, such as sarcomatous appearances or mixed hepatocchalangiocellular phenotypes [21,22]. Hypoxia induced by TACE also stimulates vascular endothelial growth factor (VEGF) production by the residual tumor cells, which can promote the tumor progression following TACE [23,24]. Furthermore, some surviving tumors can be nourished by the portal venous system if the arterial branches are markedly attenuated by TACE and extrahepatic collaterals cannot develop [25]. These suggest that ineffective TACE can also produce uncontrollable tumors. In our recent analysis, the incidences of intrahepatic distant recurrence (IDR) of HCC in patients with local tumor progression (LTP) were significantly higher than in patients without LTP (p = 0.0004) [26]. This indicates that some IDR may be associated with LTP: metastases may be from locally progressed tumors or may be from newly developed tumors promoted mainly by high expression of VEGF production from the viable tumor cells. In addition, TACE damages the liver and bile ducts [8,27,28] (Figure 2). Therefore, it is important to perform curative TACE when the tumor is localized and controllable.

Figure 2. Schematic representation of negative aspects of TACE. Abbreviations: VEGF, vascular endothelial growth factor.
4. Rationale for Each TACE Technique

4.1. Rationale for Bland Embolization and DEB-TACE

Various kinds of microspheres are available. Experimental studies show that smaller-sized microspheres induce more necrosis of the hepatic parenchyma. Doxorubicin-loaded DEBs can also cause a coagulative necrosis of the adjacent liver parenchyma; therefore, smaller-sized DEBs induce more marked necrosis of the hepatic parenchyma than the same-sized bland beads or larger-sized DEBs [29]. For tumor tissues, smaller DEBs are likely to occlude distal tumor vessels, resulting in more localized chemotherapeutic delivery and a good imaging response [30], although the risk of unexpected complications, such as biliary necrosis and systemic embolization, may increase [28,31,32]. However, one of the important technical problems of particle embolization is that microspheres cannot embolize the tumor drainage. Therefore, the reversed flow from the surrounding hepatic sinusoids and portal venules into the peripheral tumor portion following arterial blockage cannot be blocked. Micro-arterial communications may also be unblockable. As a result, tumor tissues supplied not only by both arterial and portal blood but also by collateral blood including portal blood may survive mainly at the periphery (Figure 3) [33,34]. In DEB-TACE, it is expected that eluted chemotherapeutics may necrotize the surviving tumor cells; however, it is uncertain how the chemotherapeutics reach the viable tumors.

![Figure 3](image-url). Schematic representation of rationale for superselective cTACE and DEB-TACE/bland embolization. Reproduced with permission from Ref. [33]. An Open Access article (CC BY 4.0). Abbreviations: W-D, well-differentiated; PB, peribiliary vascular plexus, PV, portal vein.

In addition, DEB-TACE damages the hepatic artery more severely than cTACE (per branch, \( p < 0.001 \) and per patient, \( p = 0.005 \)) [35] and the incidences of arterioporal shunt formation are also significantly higher following DEB-TACE than cTACE (48.7 vs. 8.1%, respectively, \( p < 0.001 \)) [36].
4.2. Rationale for cTACE

Iodized oil is a semi-fluid embolic agent that can change its shape in vessels. Two types of iodized oil emulsion can be prepared: water-in-oil emulsion (WOE) and oil-in-water emulsion (OWE). WOE should be properly prepared because WOE has stronger embolic effects on tumor vessels than OWE [37,38]. Iodized oil injected into the hepatic artery is initially retained in the tumor sinusoids. When more than a certain amount of iodized oil is pooled in the tumor sinusoids, some iodized oil overflows into the portal veins through the drainage route of the tumor [39,40]. Iodized oil also enters the portal veins through the PBP in the surrounding liver [40]. As a result, the peritumoral portal blood flow, including reversed portal blood flow into the tumor, can be temporarily blocked by viscosity of iodized oil. Moreover, some iodized oil can pass through the arterial communications and reach the neighboring hepatic arterial branches and/or extrahepatic arteries [10,13,41]. This allows embolization or identification of an occult tumor feeder and may prevent the arterial collateral supply to the tumor following TACE [13]. By adding arterial blockage with GS particles after injection of iodized oil, both the hepatic artery and portal vein surrounding the tumor can be embolized (Figure 3). Small-sized GS particles (slurry, approximately 0.2–0.5 mm in diameter) are favorable to achieve more necrosis because the reported mean diameter of tumor feeders in tumors of 7 to 63 mm (mean, 20.3 ± 12.7) in diameter was 0.41 ± 0.32 mm (range, 0.12–1.79) [42]. This leads not only to ischemic necrosis of the hypervascular tumor portions but also to peritumoral necrosis including tumor portions supplied by a dual blood supply and induces shrinkage of the surrounding liver parenchyma [11–13,33,40,43]. This also indicates that cTACE damages the normal liver; therefore, selective catheterization is essential to minimize the liver toxicity. It is also important to minimize the total dose of iodized oil, and the maximum dose in one cTACE procedure is generally 10 mL in Japan and 15 mL in Western countries [44].

4.3. Therapeutic Effects of Bland Embolization with Particles

In the recent RCT conducted by Malagari et al. [45], TACE using doxorubicin-loaded DEB presented a better local response, fewer recurrences, and a longer time to progression than bland embolization with same-sized microspheres (p = 0.008). However, no significant benefit to survival of DEB-TACE was found, partly because of a short follow-up. On the other hand, the latest RCT conducted by Brown et al. [46] showed no significant differences in tumor response (p = 0.48), median progression-free survival (p = 0.11), and overall survival (p = 0.64) between the bland embolization and doxorubicin-loaded DEB-TACE groups.

It is obvious that bland embolization has lesser damage to the liver than cTACE and DEB-TACE because non-drug-loaded microspheres can occlude the vessels with mild foreign body reaction [47], mostly non-necrotic vascular changes without hepatic necrosis [29]. However, the role of bland embolization with microspheres has not yet been established in the treatment strategy for HCC. In a report by Osuga et al. [48], the 1- and 2-year survival rates of 59 naïve HCC patients with a mean largest tumor diameter of 4 cm (range, 1.2–14.7 cm) treated with bland embolization using HepaSphere (Merit Medical, Jordan, UT, USA) of 53 to 212 µm in a dry state were 100% and 83%, respectively, and the median survival time (MST) was 30 months. Maluccio et al. [32] reported the outcomes of bland embolization with small particles (50-µm polyvinyl alcohol particles and 40–120-µm Embosphere (Merit Medical) for 322 HCC patients. Among them, 109 patients without portal vein invasion or extrahepatic disease had an MST of 40 months and 1-, 2-, and 3-year survival rates of 84%, 66%, and 51%, respectively. However, four patients died of acute hepatic failure and two died of pulmonary embolism. Small particles have strong therapeutic effects because these can impart terminal vessel blockage within the tumor; however, the risk of unexpected complications caused by migrated particles may increase [31,32]. Therefore, a superselective approach and careful observation of pulmonary shunting is recommended in the use of small-sized particles. In a report by Lanza et al. [49], 40- or 100-µm Embozene (Varian, Austin, TX, USA) could be safely used in the superselective bland embolization procedure.
Bland embolization with gelatin sponge particles can also be effective for achieving hemostasis of ruptured HCC in patients in a hemodynamically unstable condition with minimal invasiveness [50].

4.4. Proper Use of Chemotherapeutics in TACE

One to three chemotherapeutics are usually used in TACE, although the necessity of chemotherapeutics is still controversial [3–5,45,46].

cTACE has a long history in Japan; however, the most effective and least toxic regime for cTACE has not yet been established. Some retrospective studies showed the superiority of cisplatin over doxorubicin/epirubicin in terms of overall survival [51–53], although a prospective study showed no significant differences [54]. However, cisplatin requires pre- and post-hydration because of its renal toxicity and also has severe side effects, including thrombocytopenia, hepatic failure, and hypersensitivity reactions. The incidence of hypersensitivity reactions increases when the number of treatment cycles using cisplatin is increased [55]. Therefore, cisplatin is mainly used for doxorubicin/epirubicin refractory tumors and/or advanced tumors [56]. Miriplatin is a lipophilic platinum chemotherapeutic that is available only in Japan. It has less toxicity; however, the RCT showed no significant differences in therapeutic efficacies over epirubicin [57]. Some retrospective studies also showed a significantly higher local recurrence rate in the miriplatin TACE group compared with the epirubicin/epirubicin plus mitomycin C TACE group [58,59]. Very slow release of active platinum (5.9% at 28 days) and less damage to the artery are considered to be the main causes of high frequent local tumor recurrence [58]. Now, miriplatin is mainly used in balloon-occluded TACE because of its association with less arterial damage [60].

In the clinical setting, switching the chemotherapeutics may be a feasible option for advanced HCC, especially when the tumor becomes resistant to the initial form of cTACE, although there is not enough scientific evidence [54,56,57].

In DEB-TACE, DC Bead (Boston Scientific, Marlborough, MA, USA) and Embozene-Tandem (Varian) can load positively charged chemotherapeutics, such as doxorubicin or epirubicin. HepaSphere (Merit Medical) can also load cisplatin by its reservoir effect, besides positively charged chemotherapeutics, but it is released within 24 h [61]. No comparisons are available of the long-term prognosis after DEB-TACE between each embolic agent, because of the heterogeneity in patient selection, etiology of the underlying cirrhosis, embolization technique and particle size, and relatively short-term follow-up.

4.5. Comparison of Therapeutic Effects between DEB-TACE and cTACE

Table 1 shows the results of RCT studies regarding therapeutic effects between DEB-TACE and cTACE. Three RCTs conducted in Europe showed no significant differences in local tumor control and/or survival between the DEB-TACE and cTACE groups [62–64]. Supplementary analyses of the PRECISON V study showed that in 67% of patients with more advanced disease (Child-Pugh B class, Eastern Cooperative Oncology Group [ECOG] performance status 1, bilobar, or recurrent disease), the incidences of an objective response and disease control were significantly higher in the DEB-TACE than cTACE group ($p = 0.038$ and $p = 0.026$, respectively) [62]. Regarding the systemic side effects of doxorubicin, such as alopecia, skin discoloration, mucositis, and bone marrow suppression, there was a significant benefit of DEB-TACE over cTACE ($p = 0.012$) [62]. In the PRECISION Italia study, postprocedural pain was more frequent and severe after cTACE ($p < 0.001$) [64].
Table 1. Comparison of therapeutic effects between DEB-TACE and cTACE.

| Author Group/Technique | N | Bead Size (µm) | CR (%) | PR (%) | OR (%) | P-Value | OS (%) 1 y | OS (%) 2 y | P-Value |
|------------------------|---|----------------|--------|--------|--------|---------|-----------|-----------|---------|
| Lammer Ref. [62] (2010) Advanced disease * | DEB cTACE | 93 | mainly, 300–500 | 26.9 | 22.2 | 51.6 | 0.11 | 90 | 87 | 0.96 |
| | DEB cTACE | 63 | | | | | | | | |
| | cTACE | 108 | | | | | | | | |
| Sacco Ref. [63] (2011) | DEB cTACE | 33 | 100–300 | 51.5 | 48.5 | 100 | 0.1 | 90 | 87 | 0.96 |
| | cTACE | 34 | | | | | | | | |
| Golfieri Ref. [64] (2014) | DEB cTACE | 89 | 100–300 | 52.2 | 70.6 | 100 | 0.1 | 84 | 45 | 0.949 |
| | cTACE | 88 | | | | | | | | |
| Ikeda Ref. [65] (2020) | DEB | 99 | 100–300 | 35.7 at 1 month | 27.6 at 3 months | 84 at 1 month | <0.0001 | 86 | 57 | 0.949 |
| | cTACE | 101 | | | | | | | | |

* At least, one of Child-Pugh B, ECOG 1, recurrent disease, bilobar disease was present. Abbreviations: CR, complete response; PR, partial response; OR, objective response; OS, overall survival.

On the other hand, the latest RCT conducted in Japan proved significant differences in CR rates between cTACE and DEB-TACE groups (84.2 vs. 35.7% at 1 month and 75.2 vs. 27.6% at 3 months, respectively [both p < 0.0001]), although the rates of grade 3–4 of elevation of serum aspartate aminotransferase and alanine aminotransferase were significantly higher in the cTACE group (48.5 vs. 5.1% and 38.7 vs. 3.1%, respectively [both p < 0.0001]) [65]. This study indicates that cTACE can achieve complete tumor necrosis by sacrificing the surrounding liver, and the discrepancy of outcomes of cTACE between Europe and Japan might be caused by different devices and techniques because superselective cTACE using a smaller (1.5- to 2-F tip) microcatheter under intraprocedural monitoring by CT or cone-beam CT (CBCT) is a standard technique in Japan [13]. This study did not evaluate the survival benefit of superselective cTACE over DEB-TACE because of a short follow-up period; however, it is expected the extension of the patient’s lifespan in the cTACE group, because a CR after the initial TACE is the most significant prognostic factor in HCC patients [20].

Theoretically and clinically, liver toxicity of DEB-TACE is mild compared with cTACE [62,64] because of a lower dose of active chemotherapeutics and proximal embolization for liver and tumors, although sudden death caused by systemic embolization infrequently develops [31,32] and the rate of biliary complication is significantly higher compared with that of cTACE (36.8 vs. 14.4%, respectively, p < 0.001) [28]. Therefore, DEB-TACE is favorable for patients with an advanced age, large and/or bilobar tumors, or a poor liver function. On the other hand, cTACE should be indicated for patients with localized tumors and a preserved liver function. Adverse effects of bland embolization with microspheres are milder than those of DEB-TACE [47]; therefore, it is suitable for elderly patients with large HCCs, Child-Pugh score 9 patients, or patients with ruptured HCC.

4.6. Technical Advantages of Selective cTACE Compared with Non-Selective cTACE

Non-selective cTACE using a large volume of iodized oil is not only less effective for tumors but also more harmful for normal liver parenchyma; therefore, catheter position is very important to reduce the total dose of iodized oil and minimize the liver damage. cTACE also damages the hepatic artery by arteritis due to chemotherapeutics and allergic reaction to gelatin [66,67], and attenuation of the hepatic arterial circulation reduces the hepatic function and causes the extrahepatic collaterals to develop [68,69]. Therefore, damage to the hepatic artery by cTACE should be minimized so as to preserve the liver function and prolong the duration of TACE treatment. Generally, selective cTACE is defined as cTACE at the segmental hepatic artery, whereas superselective cTACE is defined as cTACE at the subsegmental hepatic artery. cTACE at the most distal level of the subsubsegmental hepatic artery is termed ultraselective cTACE [13,40].
Most microsatellite lesions, which are usually non-selective cTACE-resistant [15], are located in the corona enhancement area [70]. The corona area should be included in the treatment area because cancer cells first enter there before forming distant metastatic lesions. However, the width of the corona is not uniform and is thick in places [17]; therefore, the safety margin should be normally determined individually before curative treatment, such as for hepatic resection or radiofrequency ablation (RFA), based on CTHA findings [70]. In TACE, we conveniently determine the safety margin according to the tumor size: at least 5 mm wide for tumors <25 mm and 10 mm wide for tumors ≥25 mm, corresponding to the width of corona enhancement [26,71,72] based on a histological study [73]. As mentioned above, superselective cTACE can simultaneously treat the drainage area in most tumors because the corona area is usually supplied by the tumor feeder [39]; however, in some cases, a part of the corona area is supplied by another arterial branch. In such cases, the arterial branch should be subsequently embolized, although iodized oil has already been distributed there via the portal vein connecting with tumor drainage.

5. Technical Tips for Performing Effective Superselective cTACE

5.1. Achievement of Marked Portal Vein Visualization with Iodized Oil During cTACE

There is a relationship between the grades of portal vein visualization with iodized oil and local tumor recurrence. We divided the grades of portal vein visualization with iodized oil into three grades on spot radiographs obtained during the procedure: grade 0, no visualization; grade 1, visualization adjacent to the tumor; and grade 2, visualization in the whole or extended embolized area, and the local recurrence rates in the grade 2 group were significantly lower than those in the grades 1 and 0 groups (p < 0.0485 and p < 0.0001, respectively) (Figure 4) [40].

![Figure 4](image_url)

**Figure 4.** Classification of grades of portal vein visualization with iodized oil and relationship between local tumor recurrence. (a) Schematic representation of grades of portal vein visualization with iodized oil; (b) Graph shows the local recurrence rates calculated with the Kaplan-Meier method for each portal vein visualization group. The local recurrence rates for tumors in the grade 2 group were significantly lower than those for tumors in the grades 1 and 0 groups (p = 0.0485 and p < 0.0001, respectively). The local recurrence rates for tumors in the grade 1 group were also significantly lower than those in the grade 0 group (p < 0.0001). Reprinted with permission from Ref. [40].

In superselective cTACE, iodized oil should be slowly injected to avoid “oil cast formation” in the arteries. The endpoint of iodized oil injection is grade 1 portal vein visualization, not grade 2, because iodized oil in the tumor and surrounding liver is pushed into the portal vein by GS slurry injection and widely distributes into the whole embolized area, and frequently beyond the embolized area. As a result, grade 2 visualization can be obtained [74]. When the flow of the tumor-feeding branch unexpectedly stops before adequate portal vein visualization, we can adopt two measures: injection of 0.5 μg of prostaglandin E1 or 0.5 mL of 2% lidocaine through the catheter to increase
arterial flow; and more distal advancement of a microcatheter to achieve a semi-wedged condition [74]. The latter technique should be attempted first, if possible, because it is usually more effective. When the microcatheter is advanced more distally, iodized oil can be re-injected. In such a condition, iodized oil can flow forward due to the injection force, because stretching the catheterized artery and the mass effect of the microcatheter itself can prevent backflow of iodized oil. As a result, the volume of iodized oil passing through the arteriportal and arterial communications can increase and embolic effects can be enhanced.

5.2. Importance of the Order of Embolization of Each Tumor Feeder

When multiple feeding branches are present, the order of embolization of each feeder is key to improve the therapeutic effect of cTACE. The main tumor feeder should be embolized last because dense retention of iodized oil and contrast material disturbs the depiction of the residual tumor stain and other small feeders on digital subtraction angiography performed immediately after cTACE. Additionally, embolic materials injected into the main tumor feeder are sometimes pushed back by the reversed flow through the minor tumor feeder when the flow of the main tumor feeder becomes near stasis [74]. In tumors partially fed by an extrahepatic collateral supply, the feeding branch of the extrahepatic artery should also be embolized first, if possible, because arterial blood flow to tumors through the extrahepatic artery immediately increases when the hepatic arterial flow is stopped [75]. Moreover, each tumor feeder should also be embolized from distal to proximal in order to avoid inadvertent occlusion of the distal tumor feeders with overflowing embolic materials during embolization of the proximal tumor feeder [74]. However, feeders that are considered difficult to catheterize, such as the caudate artery or tiny tumor feeders arising from extrahepatic arteries, should be embolized first, because the performance of a microcatheter-microwire system is gradually deteriorated during the procedure [74].

5.3. Usefulness of Stepwise cTACE for Large HCCs

cTACE is also widely performed for large HCCs; however, the previously reported 5-year overall survival rate was only 7–9.7% [76,77]. In addition, the use of a large volume of iodized oil for large tumors carries the risk of catastrophic complications, such as systemic embolization of the lung or brain [78,79] and acute tumor-lysis syndrome [80]. Therefore, we typically perform stepwise superselective cTACE for localized tumors >7 cm [81,82]. Two to three cTACE sessions are scheduled depending on the vascular anatomy, and each session is performed in a superselective fashion using ≤10 mL of iodized oil at 3- to 10-week intervals according to the patient and tumor conditions. In our recent analysis of 25 patients with HCC ≥10 cm (mean, 130 ± 27.6 mm [range, 101–193], single [n = 12], 2–9 [n = 6, mean, 4.2 ± 2.6 ], and ≥10 tumors [n = 7, including 2 with ≥50 tumors], with vascular invasion [n = 5]), the 5-year survival rate was 23.1%, and that of patients with tumors ≤3 lesions was 38.9% [82]. This indicates that cTACE is also effective for localized huge HCCs. However, cTACE of extrahepatic collateral arteries was required in 84% of patients during the treatment course [82]. Therefore, the outcomes of cTACE for huge HCC may be more strongly influenced by the TACE technique compared with cTACE for small HCC. Bland embolization with gelatin sponge particles followed by cTACE is another alternative option for huge HCC to safely embolize the tumor [83].

5.4. Usefulness of CT/CBCT and TACE Guidance Software

A unified CT and angiography system that has been developed in Japan can improve the technical success rate of superselective cTACE [84]. However, it is relatively expensive and is not widely distributed. CBCT is a new technology to obtain CT images using a flat panel detector of an angiography unit. It has been reported that CBCT can improve not only local tumor control but also the prognosis of HCC patients [71,85]. TACE guidance software, including automated tumor feeder detection (AFD), is also a novel software using CBCTHA data that can reduce the physician’s workload and improve the treatment accuracy [26,72,86–89]. This software can provide
useful information regarding the order of embolization and division of chemotherapeutics for each tumor feeder. The curability and invasiveness can also be controlled by changing the virtual target size including the safety margin. If curative TACE is expected, a wider safety margin should be created to embolize all potential tumor feeders. If palliative TACE is recommended, the safety margin should be narrower to reduce the adverse effects [90]. In our previous analysis of 155 HCCs ≤5 cm (mean diameter, 17.4 ± 7.4 mm [range, 10–48]) including 18 angiographically occult tumors, AFD software using CBCT/THA data could detect 85.4% of tumor-feeding subsubsegmental hepatic arteries, 90.3% in new tumors and 75.3% in recurrent tumors. Regarding the technical success, complete ultraselective cTACE could be performed in 82.6% of tumors [72].

AFD software can also be useful to identify a tumor feeder arising from extrahepatic arteries and expand the indication and safety of TACE through extrahepatic collaterals. However, if the tumors are fed by the cystic, gastric, or colonic artery, the safety margin should not be included in the target to avoid non-target embolization [90,91].

In our recent analysis, the 5-year survival rate of 102 patients with tumors ≤7 cm and ≤5 lesions (mean largest tumor diameter, 25.3 ± 12.3 mm [range, 6–69]; mean tumor number, 1.9 ± 1.1 [range, 1–5]) treated with superselective cTACE using TACE guidance software was 60%, and about half (13/27) of patients died of HCC-unrelated causes [26]. This indicates that TACE guidance software can prolong a patient’s lifespan.

6. Concept of TACE Unsuitability

However, TACE-resistant HCCs are also present. In 2019, Asia-Pacific Primary Liver Cancer Expert (APPLE) Meeting proposed the concept of TACE unsuitability. TACE unsuitability is defined as each one of the following three clinical conditions that prevent a survival benefit from TACE or conditions that TACE is even harmful: (i) unlikely to respond to TACE: confluent multinodular type, massive or infiltrative type, simple nodular type with extranodular growth, poorly differentiated type, intrahepatic multiple disseminated nodules, or sarcomatous changes after TACE; (ii) likely to develop TACE failure/refractoriness: up to seven criteria out nodules; and (iii) likely to become Child-Pugh B or C class after TACE: up-to-7 criteria out nodules (especially, bilobar multifocal HCCs) or modified albumin bilirubin (ALBI) grade 2b [92].

In patients with TACE-unsuitable HCC, systemic therapy is generally recommended. However, TACE-unsuitable HCC is not always contraindicated for TACE monotherapy. Because up-to-7 criteria out tumors include various tumor conditions, like BCLC-B HCC, and some up-to-7 criteria out tumors may also be good candidates for superselective cTACE when tumors are located in limited segments of the liver [92]. Complete necrosis can also be achieved in some confluent multinodular and simple nodular type with extranodular growth HCCs when a sufficient safety margin can be embolized. A recent report indicated that age ≥75 years (p = 0.022), AFP ≥200 ng/mL (p = 0.010), tumor counts ≥11 (p = 0.008), and heterogenous enhancement (p = 0.024) were significant prognostic factors in BCLC-B HCC patients treated with superselective cTACE [93]. This indicates that TACE is also effective for some up-to-7 criteria out tumors. Therefore, the indication of TACE for unsuitable patients should be discussed according to the ability of TACE in own institution and a tailor-made treatment strategy is recommended. When TACE is performed first and the tumors may rapidly recur and/or the tumor features may change to unfavorable forms (such as hypovascular/ring-like enhancement, infiltrative tumors around the previously treated tumor that iodized oil is densely accumulated, hyperintensity on diffusion-weighted magnetic resonance imaging, and development of vascular invasion and/or lymph node metastases) following TACE, additional TACE should be abandoned and systemic therapy should be immediately stated.

The combination of an antiangiogenic and TACE is a promising treatment option for TACE-unsuitable HCC because pre-administration of an antiangiogenic can enhance the therapeutic effects of TACE considered to be facilitated by promoting vascular normalization and improving the distribution of iodized oil mixed with chemotherapeutics [94,95]. Therefore, TACE or hepatic arterial
infusion chemotherapy should be promptly performed for patients with antiangiogenic-refractory HCC, if possible.

7. TACE Strategy According to the Number and Size of HCC

BCLC-B HCC includes various tumor conditions and the goal of TACE treatment is not only to control the tumors but also to maintain the liver function [92]. Therefore, it is essential to change the technique and magnitude of TACE in individual cases, although there is no consensus on how to change the TACE techniques according to the tumor spread.

In a report by Yamakado et al. [96], a patient subgroup with Child-Pugh A class and tumors ≤ 7 cm and ≤ 4 lesions showed significantly favorable outcomes among BCLC-B HCC patients. The Child-Pugh score was also a significant prognostic factor, and the prognosis of Child-Pugh score 9 patients treated with cTACE was worse than that of the BSC group in the previous RCT [3,97]. Yamakado et al. [98] also reported that cTACE techniques has an impact on patient survival. In those with HCC ≤ 7 cm and ≤ 5 lesions, the prognosis of patients who underwent selective/superselective cTACE was significantly better than that of patients treated with non-selective cTACE ($p = 0.033$). These studies suggest that superselective cTACE is recommended for patients with Child-Pugh scores of 5–8 and HCC ≤ 7 cm and ≤ 5 lesions.

Now, we perform cTACE for almost all HCC patients with Child-Pugh scores ≤ 8 because it has a stronger therapeutic effect compared with other techniques. For tumors ≤ 7 cm and ≤ 3 lesions, superselective cTACE, mainly ultraselective cTACE, is performed. For tumors > 7 cm but ≤ 3 lesions, stepwise superselective cTACE is also performed with curative intent [81,82]. Bland embolization, DEB-TACE, or systemic therapy is another option, especially for elderly patients. Bland embolization followed by cTACE is also another option to safely embolize the huge tumor [83]. For bilobar multiple lesions ≥ 4, superselective cTACE is not always indicated but the embolized area should be minimized to reduce liver toxicity. In this subgroup, DEB-TACE or systemic therapy is also an alternative option. For far advanced (large and bilobar) tumors, palliative cTACE, DEB-TACE, or bland embolization is indicated to aim at mass reduction or prevention of tumor rupture; however, systemic therapy is recommended first when the patient has a preserved hepatic function, although there is a risk of tumor bleeding in antiangiogenic therapy (Figure 5) [99]. In HCC patients with Child-Pugh score 9, DEB-TACE or bland embolization is recommended.

![Figure 5. Our TACE strategy according to the patient and tumor conditions. Abbreviation: C-P, Child-Pugh; n, number; RFA, radiofrequency ablation.](image-url)
According to APPLE Consensus Statements, tumors ≤6 cm and ≤6 lesions are classified into the good response to cTACE subgroup, tumors >6 cm and ≤6 lesion and tumors ≥7 lesions are classified into the poor response to cTACE subgroup. DEB-TACE and systemic therapy is another option for these tumor conditions (Figure 6) [92]. In particular, antiangiogenic therapy is recommended for TACE-unsuitable tumors. In addition, the sequential therapy of an antiangiogenic and TACE is a novel option for antiangiogenic-refractory HCCs or even for down-staged HCCs by antiangiogenic therapy [92].

Figure 6. Heterogeneity of intermediate-stage HCC and grade of response to cTACE in each subgroup. Reproduced with permission from Ref. [92]. An Open Access article (CC BY 4.0).

8. Further Directions

With the potential of ultraselective cTACE to cure small HCC including hypovascular tumor portions [11–13,100], it can replace surgical resection and RFA in selected patients with BCLC 0-A HCC. In the eUpdated European Society of Medical Oncology (ESMO) guideline on hepatocellular carcinoma algorithm on 19 June 2020, indication of TACE has been expanded to BCLC 0-A HCC (Figure 7) [101]. This treatment strategy will become more popular, if TACE guidance software and a small-sized microcatheter system become widely available. Additionally, the sequential therapy of an antiangiogenic and TACE is a promising treatment strategy for BCLC-C HCC patients with a preserved hepatic function [102]. Indeed, in the fourth version of clinical practice guidelines for HCC proposed by the Japan Society of Hepatology in 2017, TACE has also been recommended for not only a solitary tumor >3 cm but also tumors with vascular invasion, in addition to BCLC-B HCC [103]. We believe that TACE still plays an important role in the treatment of HCC patients. We should keep in mind that TACE is not a simple local chemotherapy, and the outcomes are markedly influenced by TACE techniques.
Figure 7. Updated European Society of Medical Oncology (ESMO) guideline on hepatocellular carcinoma algorithm on 19 June 2020. Reprinted from Ref. [101]. An Open Access article (CC BY 4.0).

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