Use of chemotherapy to treat hepatocellular carcinoma

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SUMMARY Hepatic malignancies remain a global challenge. Hepatocellular carcinoma (HCC) accounts for around 90% of patients with liver cancer and is the sixth most common neoplasm worldwide and the fourth leading cause of cancer-related death. However, the long-term prognosis for HCC remains far from satisfactory, with a late diagnosis and limited treatment. DOX has served as conventional chemotherapy with the longest history of use. Although conventional chemotherapy is being challenged by molecular therapy and immune therapy, there is renewed optimism and interest in both systematic and locoregional therapy. Combined chemotherapy is widely used in clinical practice. In specific terms, FOLFOX can serve as a first-line (category 2B) option as recommended by the 2021 NCCN guidelines, while the efficacy of LTLD plus RFA has been confirmed in the phase III HEAT study. These approaches have challenged the dominant status of molecular therapy in terms of health economics and they have potential benefits in Asia, where HBV-related hepatocellular carcinoma is prevalent. Moreover, locoregional chemotherapy can be achieved with TACE and HAIC (possibly involving FOLFOX, DOX, mitomycin C, cisplatin, epirubicin, etc.). TACE was officially recommended by the 2021 NCCN guidelines for patients with Child-Pugh class B liver disease. In addition, HAIC has demonstrated a potential advantage in preliminary clinical practice, although it hasn’t been included in any guidelines. Hence, this review summarizes large-scale trials and studies examining the development and innovative use of chemotherapeutic agents. Mounting clinical evidence warrants an exploration of the efficacy of chemotherapy.

Keywords chemotherapy, hepatocellular carcinoma, immune therapy, molecular therapy

1. Introduction

Primary hepatic malignancies include hepatocellular carcinoma (HCC) and intrahepatic cholangiocarcinoma originating from the parenchyma. In addition, rare primary lesions originating from the mesenchyme develop into liver sarcoma. HCC is the sixth most common neoplasm worldwide and the fourth leading cause of cancer-related death, and it accounts for around 90% of patients with hepatic malignancies with an unfavorable prognosis due to its largely asymptomatic natural history (1), high recurrence, and ineffective therapeutic strategies for advanced HCC (2-5) (Figure 1). Hepatitis B virus (HBV) infection is an independent high risk factor for HCC among unvaccinated persons, mostly in Asia and sub-Saharan Africa (6). In addition, hepatitis C virus (HCV) infection, dietary exposure to aflatoxin B1, and alcoholic liver disease (ALD) and nonalcoholic fatty liver disease (NAFLD) have become the leading causes of HCC in developed countries (7,8).

Currently, there are various controversies regarding the therapeutic options for HCC. Since localized liver cancer is asymptomatic for much of its natural history, the major obstacles are a late diagnosis and a subsequently low resection rate, limiting treatment alternatives. Thus, a significant fraction of patients will eventually become eligible for chemotherapy. Chemotherapy has become a conventional option for HCC as a result of drug research and development. These drugs play an indispensable role in systematic therapy and are also being developed to act on locoregional targets through approaches such as transarterial chemoembolization (TACE) and hepatic artery infusion chemotherapy (HAIC). Here, chemotherapy is outlined and its corresponding role in treating HCC has been described.

The stage of HCC is identified using various staging systems, namely Barcelona Clinic Liver Cancer (BCLC) tumor staging, the Hong Kong Liver Cancer staging system (3,9), and the Cancer of the Liver Italian Program (10). The BCLC is most widely used and was introduced...
in 1999 (3,11). Since it provides a comprehensive evaluation based on liver function, performance status, and tumor burden, the BCLC system has been approved by the European Association for the Study of the Liver (4,10-12). The algorithm classifies patients into one of five stages and it provides treatment recommendations for each stage (3). (Figure 2) Based on this classification system, the use of chemotherapeutic agents for different stages of HCC has been described here (Figure 3).

2. Systemic chemotherapy

Systemic therapies are, along with transarterial therapies, recommended for patients with BCLC stage B or C HCC
DOX was first introduced in the early 1960s (17) and was widely used to treat solid tumors. A bold attempt has been made to use DOX to treat HCC. Nonetheless, controversies surrounding DOX’s mechanism of action abound. In a nutshell, DOX inhibits the replication and translation of DNA via various approaches including as a topoisomerase II poison (17) and targeting p53 (17).

Like other antitumor agents, DOX was soon approved, but its drawbacks have been serious cardiac adverse events, namely chronic cardiomyopathy and congestive heart failure (CHF), and progressive drug resistance after completion therapy within a year (18). These faults have limited the maximum recommended volume of DOX to 40 to 75 mg/m² according to successive clinical studies conducted from 1977 to 2007 (16). (Table 1) As a monotherapy for HCC, DOX only conferred a survival benefit of 3.0 to 4.1 months and an ORR of 19% (16,19). As diagnosis and technology have advanced, fortunate patients are more likely to receive systemic treatment in earlier stages (16), but DOX’s marginal survival benefit has not changed.

2.2. DOX derivatives

After DOX was introduced, DOX derivatives were examined as chemotherapeutic agents. Pegylated liposomal doxorubicin (PLD) has long-acting pegylated ‘stealth’ liposomes encapsulating a doxorubicin hydrochloride inner core for intravenous administration to target HCC lesions. PLD has a better permeability and liposolubility that delay its clearance from the circulation via leaky capillaries, resulting in an attenuated circulation time and a superior cardiac safety profile (20). However, a phase II study found that PLD has almost no effect in advanced HCC (20,21), with a response rate of 10-17% at best (20,22).

When liposomal doxorubicin (LD) is pegylated, it does not have significant systemic efficacy, but lysosphospholipid-sensitive LD (LTLD) locally releases a high concentration of doxorubicin and quickly diffuses into local tissues when heated to ≥ 40°C (23,24). The tumor concentration of doxorubicin increases 25-fold (23). Tumor microvasculature is more permeable than normal.
Table 1. Summary of outcomes of systemic use of DOX according to clinical trials from 1977 to 2007

<table>
<thead>
<tr>
<th>Years</th>
<th>Clinical Trials</th>
<th>Demographic Characteristics</th>
<th>Doses of DOX</th>
<th>Median OS</th>
<th>Response</th>
<th>Treatment-related Adverse Events</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Most Common Grade 3-4</td>
</tr>
<tr>
<td>1978</td>
<td>—</td>
<td>1. Patients were ineligible for surgery</td>
<td>60 mg/m²; 30 mg/m²: WBC &lt; 2,000 μL or PLT &lt; 100,000 μL; 15 mg/m²: WBC &lt; 1,000 μL or PLT &lt; 50,000 μL, Calculated maximum: 550 mg/m²</td>
<td>2-5 months</td>
<td>Objective response: 32%</td>
<td>NR</td>
</tr>
<tr>
<td>1984</td>
<td>NCZ vs. DOX</td>
<td>1. Patients from South Africa</td>
<td>60 mg/m² for anicteric patients; 40 mg/m² for icteric patients and AST &gt; 2 times normal</td>
<td>22 weeks</td>
<td>NR</td>
<td>NR</td>
</tr>
<tr>
<td>1984</td>
<td>Eastern Cooperative Oncology Group Trial</td>
<td>1. Patients from the US, Europe, and South Africa 2. Contraindications for patient eligibility included previous treatment and poor liver function</td>
<td>60 mg/m² for anicteric patients; 40 mg/m² for icteric patients</td>
<td>17 weeks</td>
<td>Objective response: 12%</td>
<td>Common adverse events: Leukopenia (leukocyte count: 2,000/mm³) and/or neutropenia (neutrophil count: 1,000/mm³); 21%</td>
</tr>
<tr>
<td>1985</td>
<td>Clinical trials of DOX in Italy</td>
<td>1. Patients from Italy 2. Grade A or B liver function 3. Unresectable HCC</td>
<td>60 mg/m²; 30 mg/m²: WBC &lt; 3,500 μL or PLT &lt; 80,000 μL; DB &gt; 2 mg/mL Calculated maximum: 550 mg/m²</td>
<td>4.1 months</td>
<td>Objective response: 24%</td>
<td>Common adverse events: Alopecia: 100%</td>
</tr>
<tr>
<td>1988</td>
<td>DOX vs. no antitumor therapy</td>
<td>1. Patients from China 2. Inoperable tumor 3. Liver function with acceptable therapy 4. Without detectable cardiac diseases</td>
<td>Initial dose 60 mg/m² with subsequently increased dose of 75 mg/m², NPC &lt; 3.5 × 10⁷, 50 mg/m²; NPC &lt; 1 × 10⁷, 37.5 mg/m²</td>
<td>10.6 weeks</td>
<td>Partial response: 3.3%</td>
<td>Cardiotoxicity; Neutropenia</td>
</tr>
<tr>
<td>2005</td>
<td>DOX vs. PIAF</td>
<td>1. Unresectable HCC 2. Liver function with acceptable therapy</td>
<td>60 mg/m²</td>
<td>6.83 months</td>
<td>Overall response: 10.5%</td>
<td>Neutropenia: 63%; Anemia: 28%; Thrombocytopenia: 24%</td>
</tr>
</tbody>
</table>

**Abbreviation:** NCZ, neocarzinostatin; ORR, objective response rate; NR, not report; OS, overall survival.
blood vessels, so LTLD is better able to reach tumors and reduce systemic toxicity (24-26). Given the significant correlation between LTLD and heat, the best approach might be to increase the local temperature using radiofrequency ablation (RFA). Recent studies involving a combination of intravenous administration of LD and RFA suggested that RF-induced thermal energy at 42°C in particular might yield better efficacy, improving the release of DOX from the long-circulating drug/liposome complex and resulted in accumulation of a higher concentration at the target lesion (27). In the recent phase III HEAT study, the initial complete response of multinodular intermediate-sized lesions (3-7 cm) to RFA + LTLD was > 94% and the therapeutic failure was < 5% (23) according to a sub-analysis of relatively large tumors (5-7 cm), and median PFS was 13.9 months and the median OS was 53 months (23,24). That said, there were no statistically significant differences between the RFA + LTLD arm and the RFA alone arm. However, a subsequent post hoc study suggested that prolonging the RF ablation of larger tumors would be more likely to increase efficacy and have a survival benefit (28).

2.3. Other cytotoxic agents

Nevertheless, DOX continues to be used as conventional chemotherapy. Continued innovation in chemotherapeutic agents is expected to result in the replacement of DOX, but studies have yielded conflicting results. For instance, fluoropyrimidine 5-fluorouracil (5-FU) has been widely used in a vast number of regimens and it has been used as an essential component of transarterial treatment as well (29-31). Like DOX, 5-FU suppresses DNA and RNA synthesis via the misincorporation of fluoronucleotides, and it inhibits the nucleotide-synthesizing enzyme thymidylate synthase (TS) as well. Tegafur-uracil is an oral prodrug metabolized to 5-FU mostly in the liver, and it has higher efficacy and is better tolerated (32). Nolatrexed (NOL) is a novel anticancer agent that also inhibits TS. NOL is taken up into cells without active transport, and it acts without polyglutamation (16). However, a large-scale randomized controlled trial (RCT) has compared the efficacy of DOX and NOL and found that NOL resulted in a negligible improvement in survival, with an OS of 20.7 weeks, compared to DOX (16). Thus, numerous novel cytotoxic agents have been examined, including gemcitabine, capecitabine, and oxaliplatin (33). These agents have demonstrated modest efficacy alone but considerable efficacy when used in combination (Table 2) (29,34). For instance, gemcitabine kills cells in the progress of DNA synthesis by inhibiting pyrimidine metabolism, so it is specific to certain cell phases. Moreover, it is attractive as a component of a combined strategy due to its favorable nonhematologic toxicity spectrum and mild and reversible hematological toxicity profile (35,36). Like gemcitabine, capecitabine targets fluoropyrimidine but via an oral protocol (37). Capecitabine resulted in an ORR of 11%, (including complete remission in 1 patient), and a disease control rate of 22% (29,38). The antitumor role of platinum was subsequently discovered by accident, and platinum-based drugs were approved in 1978. Platinum's mechanism of action, as confirmed by a number of highly reliable studies, differs from the mechanisms mentioned thus far since it interferes with DNA synthesis by covalently binding directly with DNA (39,40). Typical platinum-based chemotherapeutic agents include cisplatin, carboplatin, and oxaliplatin. Cisplatin was initially found to play a significant role in treating reproductive system tumors, namely testicular and ovarian cancers, with notable toxicity to the kidneys and gastrointestinal tract (39). Cisplatin resulted in a response rate of 16-27% when used in combination to treat advanced HCC (29,31), but it did provide a marginal survival benefit when used alone. Cisplatin is also used in intra-arterial strategies. Indeed, cisplatin has a marginal survival benefit because its interference with DNA binding and repair diminishes as HCC becomes resistant to the drug. Carboplatin, a second-generation platinum-based drug, has greatly reduced nephrotoxicity but it has an efficacy similar to that of cisplatin. Oxaliplatin (1R,2R-diaminocyclohexane oxalatoplatinum (II)) is actively antagonistic to tumors with acquired resistance to cisplatin (39,41,42), and it even overturned the previously accepted view that platinum-based drugs are insensitive to colorectal cancer (39). Interferons (IFNs) are a group of signaling cytokines secreted by immune cells. They display antitumor action by provoking antitumor immune responses and regulating the expression of proliferation-related genes (43).

2.4. Combination chemotherapy

Since single agents had limited efficacy against HCC, the question is whether those drugs would be efficacious when used in combination. A phase II study indicated that a gemcitabine plus PLD regimen resulted in a response rate of 24% and it increased opportunities for surgery, including resection and transplantation, for eligible patients with unresectable HCC (21,35,44,45). Due to its different mechanisms of action, it resulted in an acceptable toxicity and it prevented cross-resistance (35). In specific terms, gemcitabine promotes topoisomerase II expression, which is a process that PLD targets (35,46). Thus, better results are achieved when gemcitabine is administered before PLD. In addition, a gemcitabine plus oxaliplatin (GEMOX) regimen is likely to be well-tolerated when treating primary NAFLD according to a phase II study (47). The regimen results in an ORR of 18% and a disease control rate of 76%. Although these figures seem to indicate a marginal benefit, the regimen resulted in durable stabilization of HCC characterized by chemo-resistance (47). Combination chemotherapy with PIAF (cisplatin, interferon, doxorubicin, and...
Table 2. Examination of combination chemotherapies

<table>
<thead>
<tr>
<th>Combination Regimens</th>
<th>Pharmacological Mechanism</th>
<th>Administration</th>
<th>Median PFS</th>
<th>Median OS</th>
<th>ORR</th>
<th>Treatment-related Adverse Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gemcitabine + PLD</td>
<td>Gemcitabine: pyrimidine antimetabolite</td>
<td>Gemcitabine 1,000 mg/m² on Days 1 and 8, followed by pegylated liposomal doxorubicin 30 mg/m2 on Day 1.</td>
<td>5.8 months</td>
<td>22.5 months</td>
<td>25.00%</td>
<td>31.70%</td>
</tr>
<tr>
<td>Gemcitabine + Oxaliplatin (GEMOX)</td>
<td>Oxaliplatin: covalently binds directly with DNA to interfere with DNA synthesis</td>
<td>Gemcitabine 1,000 mg/m² on Day 1 and oxaliplatin 100 mg/m2 on Day 2</td>
<td>6.3 months</td>
<td>11.5 months</td>
<td>18.00%</td>
<td>42.00%</td>
</tr>
<tr>
<td>PIAF (cisplatin, interferon, doxorubicin, and 5-fluorouracil)</td>
<td>Cisplatin: covalently binds directly with DNA; 5-FU: misincorporation of fluoronucleotides into RNA &amp; DNA, inhibits thymidylate synthase; IFN: increases immune response</td>
<td>Cisplatin (20 mg/m²) on Days 1 through 4, interferon α-2b (5 MU/m²) on Days 1 through 4, doxorubicin (40 mg/m²) on Day 1, and 5-Fluorouracil (400 mg/m2) on Days 1 through 4 every 3 weeks for up to six cycles.</td>
<td>NR</td>
<td>8.67 months</td>
<td>20.90%</td>
<td>NR</td>
</tr>
<tr>
<td>FOLFOX4 (fluorouracil, leucovorin, and oxaliplatin)</td>
<td>Leucovorin: increases the binding affinity of a ternary complex mainly consisting of 5-Fluoro-2-deoxyuridine monophosphate (FdUMP) converted by 5-FU</td>
<td>OXA 85 mg/m² intravenously on Day 1; LV 200mg/m² IV from hour 0 to 2 on Days 1 and 2; and FU 400 mg/m2 IV bolus at hour 2, then 600 mg/m² over 22 hours on Days 1 and 2, once every 2 weeks</td>
<td>2.93 months</td>
<td>6.40 months</td>
<td>8.15%</td>
<td>55.74%</td>
</tr>
</tbody>
</table>

**Abbreviation:** PLD, pegylated liposomal doxorubicin; ORR, overall response rate; OXA, oxaliplatin; LV, leucovorin; IV, intravenously; NR, not report.
5-fluorouracil) has yielded positive outcomes in terms of a pathologic complete response (15) in a small but marked proportion of patients, and it resulted in a marginally prolonged median survival of 8.67 months (range = 6.36 to 12.00) versus DOX (6.83 months (range = 4.80 to 9.56)) (31). Attention has continued to focus on one combination chemotherapy. The drug 5-FU induces misincorporation of fluoronucleotides by replacing dUMP to form a ternary complex with a higher binding affinity and increased stabilization. The presence of leucovorin can boost ternary complex formation, and oxaliplatin can further increase efficacy. Combination chemotherapy with the FOLFOX (fluorouracil, leucovorin, and oxaliplatin) regimen has been widely used in metastatic colorectal cancer (mCRC) with demonstrable efficacy (48), and it has also been examined in advanced HCC. Although FOLFOX4 conferred a slight advantage as indicated by a median PFS of 2.93 months (2.43 to 3.53) versus DOX (1.63 to 2.30; P < 0.001), it was not effective at prolonging OS (6.4 months versus 4.97 months, P > 0.05) according to a phase III study (49). That said, FOLFOX4 offered a statistically significant benefit in terms of OS when used to treat metastatic HCC (49). Given its limited superiority in terms of a statistically significant survival benefit, the 2021 clinical practice guidelines of the NCCN recommend that FOLFOX serve as a first-line (category 2B) option (1).

Numerous studies have examined chemotherapy with various combinations of drugs, such as capecitabine plus cisplatin (XP) (29) and gemcitabine plus carboplatin (50). Nevertheless, these trials noted only a modest and inconsistent efficacy.

In summary, HCC is a multidrug-resistant tumor caused by a high level of MDR1 expression (51). Randomized trials of novel therapeutics have failed to note a significantly improved survival until recently, with a median OS of 6 to 8 months (35). Systematic chemotherapy is modestly effective in treating HCC (49).

2.5. Chemotherapy and molecular therapy

Examination of subsequent approaches to treating advanced HCC has ushered in a new era of molecularly targeted agents (MTAs) (Table 3) and immune therapy (32). Sorafenib was the first systemic molecular agent approved by the FDA, and it is rapidly replacing DOX as the drug of choice for frontline therapy (3). Sorafenib targets multi-kinases, including the serine-threone kinases Raf-1 and B-Raf, and tyrosine kinases, which are key substances that activate vascular endothelial growth factor receptors (VEGFRs) and platelet-derived growth factor receptors (PDGFR-ß). These receptors play a key role in cell proliferation and angiogenesis in the tumor microenvironment (TME) (53-55). The SHARP trial (a phase III RCT) initially noted an improved prognosis in cases of advanced HCC, with a median survival benefit of nearly 3 months and a median OS of 10.7 months versus the placebo group (52,53). The National Institute for Health and Clinical Excellence argued that sorafenib had limited cost-effectiveness as a first-line treatment for advanced HCC (35). Therefore, studies examined MTAs, including erlotinib (56), brivanib (57), sunitinib (58), linifanib (59), and everolimus (60), as a way to achieve greater efficacy at a lower cost. However, a global phase III trial found that these agents had efficacy no better than or on par with what of sorafenib (3). Studies on MTAs appeared to have reached an impasse, but lenvatinib subsequently appeared as an alternative for advanced HCC with a broader pharmacological mechanism profile against VEGFR, FGFR, PDGFR α, RET, and KIT (61). According to a phase III trial, lenvatinib resulted in an OS of 13.6 months (95% CI 12.1-14.9 months) similar to that of sorafenib (12.3 months, 95% CI 0.79-1.06 months) (61). However, all secondary efficacy endpoints were statistically superior, namely PFS, TTP, and OR (61). Moreover, a recent cost-utility analysis found that lenvatinib was superior in cost (62). Lenvatinib is reasonably given priority (62). Recently, the 2021 clinical practice guidelines of the NCCN recommended sorafenib and lenvatinib as a category 1 option for patients with Child-Pugh class A liver function (1). In addition, the later phase III RESORCE trial (63) and CELESTIAL trial confirmed the role of regorafenib and cabozantinib, both of which are oral multikinase inhibitors, as subsequent-line therapy in the event of disease progression after sorafenib administration (1).

Although clinical trials have demonstrated the benefits of sorafenib and lenvatinib, a retrospective study in South Korea reached the opposite view. In that study, the efficacy of conventional chemotherapy (fluorouracil plus doxorubicin and platinum) was not inferior to that of sorafenib (34); this finding is presumably due to the fact that trials included patients with Child-Pugh class B or C liver disease. Moreover, sorafenib alone had a limited benefit in select patients with extrahepatic disease (49). In addition, a pivotal phase III study in Asia demonstrated that sorafenib has modest efficacy, with an OS of 6.5 months (95% CI 5.56-7.56 months) versus 4.2 months in the placebo arm (95% CI, 3.75-5.46 months). Although the HRs were comparable between that study and the SHARP trial, the OS in Asia was inferior to that in the West (64). This is presumably due to the higher proportion of patients infected with HBV or poor screening in developing countries (64,65). A subsequent systemic review confirmed that sorafenib had superior efficacy in cases of non-metastatic HCC caused by HCV (5). The rapid emergence of sorafenib resistance in the majority of patients and the conflict between high costs and low incomes has limited its use in Asia (34). Use of MTAs and alternative chemotherapies is hotly contested in abound in certain countries.

Would the combination of sorafenib and chemotherapeutic agents result in a considerable
### Table 3. Examination of molecular therapy to treat advanced HCC

<table>
<thead>
<tr>
<th>Clinical Trial</th>
<th>Molecular Strategies</th>
<th>Median OS (months)</th>
<th>Median PFS (months)</th>
<th>ORR</th>
<th>Treatment-related adverse events</th>
<th>Leading to Death</th>
<th>Leading to Discontinuation</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>SHARP</td>
<td>Sorafenib</td>
<td>10.7</td>
<td>Symptomatic progression: 4.1</td>
<td>PR: 2%</td>
<td>DCR: 71%</td>
<td>Diarrhea: 8%</td>
<td>HFS: 8%</td>
<td>Fatigue: 4%</td>
</tr>
<tr>
<td>Sorafenib vs. placebo</td>
<td>Sorafenib</td>
<td>6.5</td>
<td>2.8</td>
<td>DCR 35.3%</td>
<td>NR</td>
<td>HFS: 10.7%</td>
<td>Diarrhea: 6.0%</td>
<td>Fatigue: 3.4%</td>
</tr>
<tr>
<td>SEARCH</td>
<td>Sorafenib + Erlotinib vs. Sorafenib + placebo</td>
<td>9.5 vs. 8.5</td>
<td>3.2 vs. 4.0,</td>
<td>6.6% vs. 3.9%,</td>
<td>87%</td>
<td>Diarrhea: 18.5%</td>
<td>Fatigue: 17.1%</td>
<td>NA</td>
</tr>
<tr>
<td>BRISK-FL Study</td>
<td>Brivanib</td>
<td>9.5</td>
<td>4.2</td>
<td>12%</td>
<td>11.70%</td>
<td>Hyponatremia: 4%</td>
<td>Fatigue: 2.5%</td>
<td>NA</td>
</tr>
<tr>
<td>Sunitinib vs. Sorafenib</td>
<td>Sunitinib</td>
<td>7.9</td>
<td>3.6</td>
<td>6.20%</td>
<td>NR</td>
<td>Thrombocytopenia: 29.7%</td>
<td>Neutropenia: 25.7%</td>
<td>NA</td>
</tr>
<tr>
<td>Linifanib vs. Sorafenib</td>
<td>Linifanib</td>
<td>9.1</td>
<td>5.4</td>
<td>13%</td>
<td>85.30%</td>
<td>Hypertension: 20.8%</td>
<td>NA</td>
<td>36.30%</td>
</tr>
<tr>
<td>EVOLVE-1</td>
<td>Everolimus</td>
<td>7.6</td>
<td>3</td>
<td>DCR 56.1%</td>
<td>Grade 3: 52.1%</td>
<td>Grade 4: 18.8%</td>
<td>Anemia: 7.8%</td>
<td>Asthma: 7.8%</td>
</tr>
<tr>
<td>REFLECT</td>
<td>Lenvatinib</td>
<td>13.6</td>
<td>7.4</td>
<td>24.1</td>
<td>75%</td>
<td>Hypertension: 23%</td>
<td>Decreased weight: 8%</td>
<td>NR</td>
</tr>
<tr>
<td>RESORCE</td>
<td>Regorafenib</td>
<td>10.6</td>
<td>3.1</td>
<td>7%</td>
<td>50%</td>
<td>Hypertension: 13%</td>
<td>HFS: 13%</td>
<td>Fatigue: 9%</td>
</tr>
<tr>
<td>CELESTIAL</td>
<td>Cabozantinib</td>
<td>10.2</td>
<td>5.2</td>
<td>4%</td>
<td>68%</td>
<td>HFS: 17%</td>
<td>Hypertension: 16%</td>
<td>Increased AST: 12%</td>
</tr>
<tr>
<td>REACH-2</td>
<td>Ramucirumab</td>
<td>8.5</td>
<td>2.8</td>
<td>5%</td>
<td>NR</td>
<td>Hypertension: 8%</td>
<td>Liver injury or failure: 4%</td>
<td>Proteinuria: 2%</td>
</tr>
</tbody>
</table>

**Abbreviation:** HCC, hepatocellular carcinoma; OS, overall survival; PFS, progression-free survival; ORR, objective response rate; PR, partial response; DCR, disease control rate; NA, not applicable; HFS, hand-foot skin reaction; NR, not report.
survival benefit? Sorafenib plus DOX (34), sorafenib plus GEMOX (66,67), sorafenib plus tegafur-uracil (UFT)(32), and other combinations have been studied in clinical trials, but they all demonstrated a moderate benefit in terms of PFS versus sorafenib alone.

2.6. Chemotherapy and immune therapy

Immune therapy’s mechanism of action is based on the TME around the malignant lesion and dysfunctional tumor-immune system interactions, which lead to immune evasion by reducing the recognition of tumor-associated antigens (TAAs) (68). HCC expresses immune checkpoint ligands, including co-inhibitory molecules (cytotoxic T lymphocyte-associated antigen 4 (CTLA4), PD1, T cell immuno-globulin and mucin domain containing molecule 3 (TIM3), and lymphocyte-activation gene 3 (LAG3)) (68). Immune checkpoints expressed by effector lymphocytes subsequently bind with the ligands to inhibit overwhelming activation (68). Interaction between receptors and their ligands needs to be blocked in order to sustain the activity of effector lymphocytes against the malignant proliferation of tumor cells. Immune checkpoint inhibitors (ICIs) and monoclonal antibodies can achieve this goal. PD1 and PD-L1 inhibitors have demonstrated promising results in preventing the proliferation of HCC cells (68).

The efficacy of atezolizumab (a PD-L1 inhibitor) plus bevacizumab (a VEGF inhibitor) was confirmed by the IMbrave150 trial; the combination resulted in a PFS of 6.8 months (95% CI, 5.7 to 8.3 months) and an ORR of 67.2% for one year among patients with unresectable HCC (69). The combination was also recommended by 2021 NCCN guidelines as first-line therapy (category 1) for patients with Child-Pugh class A liver disease. After the IMbrave150 trial, more attention was paid to patients with a high risk of recurrence after curative resection or ablation (70). Although the combination displayed a benefit to an extent and it was superior to current systemic therapy, a subgroup analysis of a phase III study noted lower efficacy in NAFLD and HCC with activated Wnt/β-catenin signaling (71). The incidence of NAFLD has been increasing, and NAFLD has become a main etiological risk factor for HCC in the US (71) due to imperfect screening systems, leading to advanced tumor stages. Mahipal et al. found that about 50% of HCC cases were accompanied by overactivation of the Wnt/β-catenin signaling pathway, promoting proliferation and metastasis as well as sorafenib resistance (72-74).

The combination of GEMOX and bevacizumab resulted in an OS of 9.6 months OS (95% CI, 8.0 months to not available) and a PFS of 5.3 months (95% CI, 3.7 to 8.7 months) according to a phase II study (33), but further evidence is lacking. Atezolizumab and other chemotherapy agents have displayed efficacy in triple-negative breast cancer, but there have been few studies on the combination of chemotherapeutic drugs and immune agents to treat HCC.

3. Locoregional use of chemotherapy

3.1. TACE

Although systemic chemotherapy has failed to play a major role in the treatment of HCC, local use of chemotherapeutic agents is still considerable, and approaches include TACE and HAIC. The theoretical basis for these approaches is the vascular shift where benign lesions are supplied by branches of the portal system, while malignant nodules are nourished by the hepatic artery (3,60) according to CT and MRI (3,60). Patients with BCLC stage B HCC are eligible for TACE (15) according to the 2021 NCCN guidelines, which specify a reasonable level of liver function (Child-Pugh class A or B) or a multinodular tumor without vascular invasion or extrahepatic metastasis (1,4,5).

In addition to embolization, TACE can deliver a chemotherapeutic agent in a high concentration to a target lesion; this means less of the drug in the circulation, thus reducing adverse events (1,4). TACE includes conventional TACE (cTACE) and drug-eluting bead TACE (DEB-TACE) (1,75). Doxorubicin, mitomycin C, and cisplatin are usually used in cTACE. These drugs are suspended in lipiodol for delivery to the target location, followed by embolization with gelatin sponge particles (75). DEB-TACE involves agents similar to those used in cTACE but different carriers. In DEB-TACE, beads are implanted in the tumor vasculature, where they remain for a prolonged period (75) to maximize the duration of their presence and to attenuate systemic toxicity (4,76).

A retrospective study initially indicated that DEB-TACE for unresectable HCC in all stages resulted in an OS of 610 days versus 284 days for cTACE (77). Moreover, a stratification analysis indicated that DEB-TACE resulted in a significant benefit for patients with Child-Pugh class A or B liver function since they potentially suffered liver failure when undergoing cTACE. Another retrospective study (78) and two prospective studies subsequently agreed with the earlier findings (4,79). However, the PRECISIONV trial suggested that the two procedures have equivalent efficacy and safety, possibly due to the inclusion of patients with all BCLC stages of liver disease (4,80).

Although widely used, chemoembolization remains highly controversial. A retrospective study (81) and 3 RCTs were optimistic about the benefits of TACE compared to symptomatic treatment of unresectable HCC (82,83). Moreover, a subsequent meta-analysis corroborated the survival advantage of TACE (15). However, two French studies reached the opposite conclusion, possibly because of the inclusion of a disproportionate number of patients with alcoholic cirrhosis (15). In addition, an RCT over 5 years compared the efficacy of TAE and transarterial DOX.
embolization (84). The study indicated that TAE resulted in a median PFS of 6.2 months and an OS of 19.6 months versus a PFS of 2.8 months and an OS of 20.8 months for the TACE arm, so the PFS and OS did not differ significantly. Does this mean that chemotherapy drugs have completely failed in both systemic and regional treatment of HCC? Fortunately, the answer is no. Despite those negative and disappointing results, the study in question did not stratify patients by BCLC stage, which led to a confounding bias. In addition, the study only used DOX and it did not consider other effective agents. Hence, a subsequent RCT compared the efficacy of cisplatin and epirubicin in TACE, and it has concluded that cisplatin was not significantly superior to epirubicin (85). Both cisplatin and mitomycin C have yielded consistent results according to a prospective study (86). One final aspect to consider is that the drug carrier may limit efficacy. HepaSpheres, which are vinyl alcohol-sodium acryl-ate microspheres, were the conventional carrier system, and they lasted almost 30 years in clinical practice. HepaSpheres provided superior absorption and release of a chemotherapeutic drug and they were pliant in blood vessels (87). However, micron-sized iron powder, barium ferrite (BaFe12O19), and carbon-coated iron nanocrystals (CCINs) are a novel carrier system (88). This new carrier enhances chemotherapy by maximizing the drug-loading capability and controlled-release, and this new carrier system has displayed great potential in animal experiments in vitro compared to the conventional carrier system (88). Together, these aspects play an essential role in the efficacy of chemotherapeutic drugs in TACE.

3.2. HAIC

HAIC has been widely used in Asia, and especially in Japan (1,89). HAIC’s mechanism of action is to deliver a high concentration of a chemotherapeutic agent to a targeted lesion via the hepatic artery without embolization (90). There are two approaches to HAIC: single administration and continuous infusion with a subcutaneously sited reservoir system (90). Different chemotherapeutic agents involve different approaches to targeting in order for them to be most effective. In specific terms, epirubicin hydrochloride and mitrplatin, which are concentration-dependent agents, can be used for bolus injection, while time-dependent agents, namely DOX and 5-FU, can be used for continuous HAIC. Notably, some agents, such as cisplatin and mitomycin C, are suitable for both bolus injection and continuous infusion (90). Although HAIC has been routinely used in Asia, HAIC is not mentioned as a common locoregional therapy for advanced HCC in the 2021 NCCN guidelines or by the Asian Pacific Association for the Study of the Liver (90). Therefore, attention has focused on obtaining highly reliable clinical evidence of HAIC’s efficacy. Although a systematic review and a retrospective study have indicated that HAIC conferred a survival benefit versus supportive therapy in advanced HCC (91), prolonging long-term survival was difficult due to the aggressive nature and rate of recurrence of HCC (92). Moreover, HAIC with 5-FU and cisplatin yielded a marginal survival benefit in patients with macrovascular invasion (MVI) but without extrahepatic metastasis (EHM) (93).

Would the combination of systemic therapy and HAIC result in increased efficacy? In theory, HAIC could compensate for MTAs failing to reach the expected dose at the targeted lesion because of PVTT (94), while systemic therapy could target extrahepatic lesions (52). However, the SIIUS study noted a similar OS of 11.8 months for combination therapy (HAIC (cisplatin)+sorafenib) vs. 11.5 months for monotherapy (sorafenib) but with a higher likelihood of Grade 3-4 adverse events (95). Therefore, subsequent studies have focused on alternative agents for use in HAIC. As mentioned before, oxaliplatin is superior to cisplatin (96), so the question is whether oxaliplatin could avoid the adverse effects caused by cisplatin. A phase II and III trial on FOLFOX plus sorafenib in HAIC yielded positive results in terms of the OS (13.37 months vs. 7.13 months, P < 0.001), a higher RR (40.8% vs. 2.46%; P < 0.001), and a longer PFS (7.03 [95% CI, 6.05-8.02 months] vs. 2.6 [95% CI, 2.15-3.05 months]; P < 0.001), and especially when HAIC was combined with PVTT (96). Hence, FOLFOX has been the mainstay of HAIC (97,98). Nevertheless, a cost-effectiveness analysis found that SoraHAIC was moderately cost-effective in developing areas (99). Immune therapy is recommended in the 2021 NCCN guidelines on advanced HCC, so further studies need to be conducted to determine if HAIC is cost-effective.

Compared to TACE consistently administered via the hepatic artery, HAIC with FOLFOX resulted in a median OS (23.1 vs. 16.1 months, P < 0.001) and PFS (9.6 vs. 5.4 months, P < 0.001) superior to those of TACE for large HCC according to a prospective non-randomized study and a randomized study (100,101). However, TACE for preoperative and postoperative resection of large HCC has been studied, and its use remains controversial (102,103). TACE can serve as a bridge therapy and downstaging therapy (1). That said, a point worth noting is that HAIC to treat HCC has yet to be fully examined.

4. Discussion and Conclusion

Conventional chemotherapy faces challenges from molecular therapy and immune therapy, but it plays an essential role in the treatment of hepatic malignancies. Over time, DOX was isolated from the pigment-producing Streptomyces peucetius and then DOX derivatives such as PLD and LTLD were examined. The efficacy of a combination of LTLD and RFA has been confirmed in the phase III HEAT study. As
chemotherapeutic drugs continue to advance, a variety of novel chemotherapeutic agents have been developed. In 2021, the NCCN recommended that FOLFOX serve as a first-line option (category 2B). Further advances in molecular therapy and immune therapy have challenged the dominance of conventional chemotherapy. Unlike the direct inhibition of DNA synthesis by chemotherapeutic agents, molecular strategies inhibit multi-kinases involved in cell proliferation. Molecularty targeted agents such as sorafenib and lenvatinib have demonstrated efficacy. That said, they are restricted to certain patients with Child-Pugh class A liver disease since they are not superior to conventional chemotherapy for patients with Child-Pugh class B or C liver disease. The appearance of immune agents has inaugurated a new era of systemic treatment of liver cancer, but immune therapy has a modest efficacy in circumstances involving overactivation of the Wnt/β-catenin signaling pathway, which occurs in 50% of HCC and which is related to a recurrence rate as high as 70% at 5 years. Immune therapy also offers a marginal survival benefit in hepatic virus infection-related HCC. Thus, alternatives to conventional chemotherapy, MTAs, and immune therapy are particularly controversial. What strategies should be adopted for patients with Child-Pugh class B or C liver disease? What is the nature of HCC recurrence? The current PFS for these strategies is less than one year according to clinical studies, and the current authors have justified questions about the costs and benefits of those strategies. All of the aforementioned topics need to be examined further.

Moreover, chemotherapeutic agents have been used to treat locoregional lesions through approaches such as TACE and HAIC. TACE includes cTACE and DEB-TACE. Various agents are used in TACE systems. In addition, HAIC is a consistent transcatheter arterial infusion strategy without embolization that is often used because it delivers a drug at a higher concentration. FOLFOX is a mainstay of HAIC. Moreover, HAIC is superior to TACE according to a recent prospective study. Although HAIC confers overwhelming advantages and is likely to have the most potential as a therapy, it is only used in Asia and is not mentioned in guidelines globally. Moreover, TACE can serve as a bridge therapy and downstaging therapy; whether HAIC can perform those roles is uncharted territory.

**Funding:** This work was supported by grants from the National Key Technologies R&D Program (2018YFC1106800), the Natural Science Foundation of China (82173124, 82173248, 82103533, 82002572, 82002967, 81972747, and 81872004), the National Program to Support Innovative Postdoctoral Personnel (BX20200225, BX20200227), China's Postdoctoral Science Foundation (2021M692278, 2020M673231), Sichuan Province's Program to Support Science and Technology (2021YJ0436), the Postdoctoral Science Foundation of Sichuan University (2021SCU12007), the 1.3.5 Project for Fields of Excellence, Sichuan University's West China Hospital (ZYSJC18008), and West China Hospital's Postdoctoral Science Foundation (2020HXBH075, 2020HXBH007).

**Conflict of Interest:** The authors have no conflicts of interest to disclose.

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Received January 12, 2022; Revised February 8, 2022; Accepted February 12, 2022.

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Released online in J-STAGE as advance publication February 15, 2022.