Interventional Radiological Management of Hepatocellular Carcinoma

Hepatocellular carcinoma (HCC) is the sixth-most common cancer worldwide and the fourth-most common cause of death due to cancers.[1] Its incidence is steadily increasing, and the common etiological factors include hepatitis B and C, alcohol-induced chronic liver disease, and nonalcoholic fatty liver disease.[2,3] Multiple staging systems have been proposed for classifying and guiding treatment decisions based on various factors which include tumor burden, background liver function, and performance status of the patients. The Barcelona Clinic Liver Cancer (BCLC) system [Figure 1] is the most widely accepted and followed worldwide.[4,5] However, certain authors have suggested that BCLC is too conservative in treatment recommendations and thus have advocated the usage of the Hong Kong Liver Cancer system [Figure 2].[6]

HCC recurrences are frequently seen in patients who have undergone curative treatments.[7] Moreover, many patients in our country are no longer surgical candidates at the time of presentation.[8] Hence, a multidisciplinary approach is required for the management of HCC. With the development and emergence of interventional radiological (IR) treatments, the management of HCC has changed dramatically.[9,10] IR treatments are curative in patients with early-stage disease, while they play a major role in controlling disease, improving the quality of life, and improving survival in patients with advanced stages.[10,11]

The following IR treatment options are available for the treatment of HCC:

- **Ablation techniques**
  - Chemical
  - Thermal
  - Electrical.

- **Transarterial therapies**
  - Bland embolization
  - Chemoembolization (Trans-arterial Chemoembolization [TACE]).
    - Conventional
    - Drug-Eluting Beads (DEB) based
  - Radioembolization (transarterial radioembolization [TARE]).
  - Other adjunct procedures such as portal vein embolization (PVE).

**Ablation Techniques**

Thermal ablation systems are indicated in early-stage patients when surgical options are precluded or the patient refuses for surgery. Ablation can replace resection in selected cases as it spares more liver parenchyma than anatomical resection. Thermal ablation systems have proven advantages and survival benefit over percutaneous ethanol injection (chemical ablation) and are currently established as the standard ablation method for local tumor control.[12]

Currently, most of the generated data are on radiofrequency ablation (RFA), which is the most commonly used ablative technique worldwide. It works on the principle of generating heat to kill the tumor cells using electrical current. Newer ablative techniques, such as microwave ablation (MWA), cryoablation, and irreversible electroporation, overcome some of the limitations of RFA. MWA produces higher intratumoral temperatures than RFA in a shorter duration of time and is capable of ablating larger tumors with less effect from adjacent vasculature.[13] Early-stage HCC patients, with preserved liver functions (Child–Pugh Class A or B), with tumor size <3 cm, can be treated by RFA, and lesions up to 5 cm can be treated by MWA.

**Transarterial Therapies**

Transarterial therapies are suitable for unresectable and advanced-stage HCC, Child–Pugh class A or B. They can be combined with ablation/surgical resection or systemic therapy and act as a bridge to transplantation or help in downstaging before transplantation.[14,15]

Transarterial therapies can be performed in the following ways:

- Bland embolization performed with only embolic agents
- Chemoembolization (TACE) is the standard of care for patients with intermediate-stage HCC and relatively preserved liver functions.[16,17] It is performed with chemotherapeutic agents mixed along with embolic agents. It can be performed as follows:
  - cTACE (Conventional TACE) – ethiodized oil mixed with chemotherapeutic agents (doxorubicin or cisplatin)
  - DEB-TACE – DEBs mixed with chemotherapeutic agents.

There is still debate as to which one of the above is better, and both methods show similar clinical outcomes.[18] However, DEB-TACE has lesser drug-related adverse events and liver toxicity.[18] The dose of doxorubicin typically ranges from 30 mg to 75 mg mixed with 5–15 cc of lipiodol in cTACE; however, up to 150 mg of doxorubicin can be injected when combined with DEB-TACE for larger tumors.[19,20] It is generally a well-tolerated procedure, and patients might experience postembolization syndrome which includes nausea/vomiting, fever, and abdominal pain. Acute hepatic decompensation is a rarer event and can be prevented by appropriate patient selection and performing technique. Although there are no defined number of TACE treatments.
and consensus regarding retreatment schedule, TACE can be repeated after ascertaining the radiologic response and patient’s tolerance. Superselective TACE can be performed for patients with portal vein branch thrombus and is not an absolute contraindication.\(^\text{[21]}\) The absolute contraindications of TACE include decompensated advanced cirrhosis, extensive tumor with massive replacement of both entire lobes, and main portal vein thrombosis with a high risk of posttreatment failure.\(^\text{[22]}\)

- Radioembolization (TARE) is performed with selective intra-arterial administration of microspheres loaded with a radioactive element, with yttrium-90 (Y-90) being the most commonly used. It does not exert any macro-embolic effect and hence can be used even for patients with advanced portal venous thrombosis.\(^\text{[23]}\) The aim is to deliver a tumoricidal dose of approximately 120 Gy to the tumor while preventing damage to the surrounding liver parenchyma. It is done as a two-staged
procedure, whereby a planning macroaggregated albumin (MAA) scan is done initially by injecting 99m-Tc-labeled MAA. It is used to assess the lung shunt fraction and splanchnic shunting, based on which dose modifications can be made.

Newer modifications of TARE techniques include radiation segmentectomy and radiation lobectomy which have further expanded its clinical utility. Patients might experience a postradioembolization syndrome which is usually less as compared to TACE. The rarer adverse events including radiation-induced liver disease, radiation pneumonitis, and gastrointestinal ulcers can be avoided by meticulous planning on a MAA scan. There are limited data with respect to the comparison of radioembolization with sorafenib. A study by Gramenzi et al. showed similar survival data of both these treatments in cirrhotic patients with intermediate–advanced-stage HCC; however, downstaging allowing liver transplantation only occurred after radioembolization.²⁴

**Portal Vein Embolization**

In PVE, selective blockage of hepatic portal venous supply is done so that contralateral lobe hypertrophy can be induced by directing the portal flow to the segments of the future liver remnant. The patient can then undergo a resection with reduced chances of a complicated postoperative course.

With the availability of the abovementioned IR treatment options across our country, HCC patients, now, have multiple options. Combinations of these IR treatment options with systemic therapy including sorafenib and other newer agents have shown encouraging data in terms of improved disease control and survival.²⁵ However, a recent phase 3 trial (TACE 2) showed contradictory data with a lack of survival benefit when combining antiangiogenic drugs with TACE.²⁶ A combination of TACE with radiotherapy has also been studied in HCC patients with portal vein tumor thrombus, and few studies have shown improved patient outcomes.²⁷,²⁸ Similarly, for larger and oligonodular HCC, resection and liver transplant remain the curative options, and IR treatments can act as a bridge to these surgical options. Therefore, a multidisciplinary approach to the management of HCC that includes the IR treatment options would lead to better patient outcomes.

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