**Name of Journal:** *World Journal of Transplantation*

**Manuscript NO:** 88833

**Manuscript Type:** ORIGINAL ARTICLE

***Observational Study***

**Liver transplantation for hepatocellular carcinoma in India: Are we ready for 2040?**

Pahari *et al*. Liver transplant for HCC in India

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**Received:** October 11, 2023

**Revised:** December 21, 2023

**Accepted:** January 22, 2024

**Published online:**

**Abstract**

BACKGROUND

Liver transplantation (LT) for hepatocellular carcinoma (HCC) has been widely researched and is well established worldwide. The cornerstone of this treatment lies in the various criteria formulated by expert consensus and experience. The variations among the criteria are staggering, and the short- and long-term outcomes are controversial.

AIM

To study the differences in the current practices of LT for HCC at different centers in India and discuss their clinical implications in the future.

METHODS

We conducted a survey of major centers in India that performed LT in December 2022. A total of 23 responses were received. The centers were classified as high- and low-volume, and the current trend of care for patients undergoing LT for HCC was noted.

RESULTS

Of the 23 centers, 35% were high volume center (> 500 Liver transplants) while 52% were high-volume centers that performed more than 50 transplants/year. Approximately 39% of centers had performed > 50 LT for HCC while the percent distribution for HCC in LT patients was 5%–15% in approximately 73% of the patients. Barring a few, most centers were divided equally between University of California, San Francisco (UCSF) and center-specific criteria when choosing patients with HCC for LT, and most (65%) did not have separate transplant criteria for deceased donor LT and living donor LT (LDLT). Most centers (56%) preferred surgical resection over LT for a Child A cirrhosis patient with a resectable 4 cm HCC lesion. Positron-emission tomography-computed tomography (CT) was the modality of choice for metastatic workup in the majority of centers (74%). Downstaging was the preferred option for over 90% of the centers and included transarterial chemoembolization, transarterial radioembolization, stereotactic body radiotherapy and atezolizumab/bevacizumab with varied indications. The alpha-fetoprotein (AFP) cut-off was used by 74% of centers to decide on transplantation as well as to downstage tumors, even if they met the criteria. The criteria for successful downstaging varied, but most centers conformed to the UCSF or their center-specific criteria for LT, along with the AFP cutoff values. The wait time for LT from downstaging was at least 4–6 wk in all centers. Contrast-enhanced CT was the preferred imaging modality for post-LT surveillance in 52% of the centers. Approximately 65% of the centers preferred to start everolimus between 1 and 3 months post-LT.

CONCLUSION

The current predicted 5-year survival rate of HCC patients in India is less than 15%. The aim of transplantation is to achieve at least a 60% 5-year disease free survival rate, which will provide relief to the prediction of an HCC surge over the next 20 years. The current worldwide criteria (Milan/UCSF) may have a higher 5-year survival (> 70%); however, the majority of patients still do not fit these criteria and are dependent on other suboptimal modes of treatment, with much lower survival rates. To make predictions for 2040, we must prepare to arm ourselves with less stringent selection criteria to widen the pool of patients who may undergo transplantation and have a chance of a better outcome. With more advanced technology and better donor outcomes, LDLT will provide a cutting edge in the fight against liver cancer over the next two decades.

**Key Words:** Hepatocellular carcinoma; Liver transplant; India; Downstaging; Survey; Milan; University of California, San Francisco; Portal vein tumor thrombus; Expanded criteria

Pahari H, Raj A, Sawant A, Ahire DS, Rathod R, Rathi C, Sankalecha T, Palnitkar S, Raut V. Liver transplantation for hepatocellular carcinoma in India: Are we ready for 2040? *World J Transplant* 2024; In press

**Core Tip:** The current predicted 5-year survival rate of hepatocellular carcinoma (HCC) patients in India is less than 15%. The aim of transplantation is to achieve at least a 60% 5-year disease free survival which will truly provide a relief to the predictions of HCC surge over the next 20 years. The current worldwide criteria (Milan/University of California, San Francisco) may have a higher 5-year survival (> 70%) but the majority of patients still do not fit these criteria and are dependent on other sub-optimal modes of treatment with much lower survival rates. In order to face predictions for 2040, we must prepare to arm ourselves with less stringent selection criteria to widen the pool of patients who may avail transplant and have a chance at a better outcome.

**INTRODUCTION**

Hepatocellular carcinoma (HCC) comprises for approximately 75%–80% of all liver cancer types in most countries[1]. HCC is the sixth most common cancer worldwide, comprising approximately 5% of the total cancer incidence, and causes approximately six deaths per 100000 people annually[2,3]. In 2020, liver cancer was the third most common cause of cancer-related deaths worldwide (830000)[4]. There is a lack of statistical data from India, with the number of deaths estimated to be approximately 6.8 per 100000 people, with a total of approximately 14000 deaths annually in 2010[5,6].

The burden of HCC has been increasing worldwide, and India is no exception[7,8]. Asian countries have reported the highest global liver cancer incidence (73%) and liver cancer deaths in 2020[9]. Between 1978 and 2012, there was a steady increase in the number of HCC cases in India[10,11]. In the United States, a recent study predicted a continued increase in HCC rates through 2030[12]. At present, India contributes to approximately 18% of the incidence and 4% of the mortality. By 2040, the global burden of new cases and deaths from liver cancer may increase by up to 55% (an estimated 1.3 million cases and 1.4 million deaths)[13,14]. However, India still has a low 5-year survival rate for HCC (< 15%) despite the advancement of curative and palliative treatment options over the last two decades[15,16].

Liver transplantation (LT) for HCC in patients with cirrhosis has been widely researched and is now well established worldwide[17–19]. The cornerstone of this treatment lies in the various criteria formulated by expert consensus and experience over the years. The Milan criteria was established by Mazzaferro *et al*[20] in 1996 to improve the outcomes of LT for HCC in the initial aftermath of low survival and high recurrence rates[20]. Subsequent studies by Yao *et al*[21] and Mazzaferro *et al*[22] indicated the restrictive nature of these criteria, and slightly more liberal criteria, called the University of California, San Francisco (UCSF) criteria, were introduced in 2001[21,22]. These mainly included the number and size of HCC nodules, vascular invasion, and extrahepatic spread. Since then, several other criteria have been introduced, each with its own justification and outcomes. The variations among the criteria are staggering, and the short- and long-term outcomes are controversial[19,23,24]. Another factor is the evolution of living donor LT (LDLT) as a treatment option, which has led us to accept less stringent guidelines for LT in patients with HCC, as it does not affect the LT waitlist. However, the survival of HCC-LT recipients outside the standard criteria must be comparable to that of the expanded criteria to mitigate the additional risks to live donors. The incorporation of tumor markers into downstaging protocols has also contributed to improved outcomes and overall survival rates. We aimed to study the differences in the current practices of LT for HCC at different centers in India and discuss their clinical implications in the future.

**MATERIALS AND METHODS**

We created an electronic survey form using Google Docs. It included several multiple-choice and short-answer questions to elaborate on specific choices or topics. Data were collected regarding the name of each center, their overall experience, and their LT practices with respect to HCC. In total, 54 questions were included (Supplementary Figure 1). The survey was reviewed and acknowledged as exempt from the Institutional Review Board at Medicover Hospitals, Navi Mumbai.

The survey was conducted in 42 transplant centers in India. Each center communicated *via* a transplant surgeon or physician. Responses were obtained over a 3-month period between January 2023 and April 2023. No incentives or honorariums were provided for completing the survey. Participation in the study was voluntary. Any duplicate or doubtful responses were clarified by the concerned center, and only one complete response was included in the final assessment. Eventually, 23 responses were received, which were tabulated and analyzed using standard software.

**RESULTS**

Overall, 23 of 41 (56%) transplant centers across India responded voluntarily to our survey. Almost all centers perform LDLT rather than deceased donor LT (DDLT). High-volume centers were defined as those that had performed more than 50 Liver transplants/year in the last 3 years, whereas low-volume centers were defined as those that had performed less than 50 Liver transplants/year in the last 3 years. Centers with more than 500 Liver transplants were referred to as experienced centers for discussion. Among the 23 centers, eight centers (34.8%) were identified as experienced LT centers, with two centers performing more than 2,000 Liver transplants to date. More than 50% (12/23) of the centers were high-volume centers (Figure 1).Approximately 39% (nine centers) of the centers had performed over 50 cases of liver transplant for patients with HCC (Figure 2).

Among the centers, the majority (17/23) responded that HCC was present in 5%–15% of LT recipients(Figure 2). Only one center followed the Milan criteria for LT, whereas the remaining centers were equally divided (11 each) between the UCSF and center-specific criteria for the eligibility of patients with HCC for LT. Apart from one center, all other centers (21/22 responses; 95%) replied that the percentage of patients with HCC within the Milan criteria undergoing LT was < 5%. Thirteen out of 23 centers (56.5%) preferred surgical resection in a 43 year-old Child A cirrhosis patient with a 4 cm solitary HCC and good performance status over LT directly. Nine centers specified the criteria for liver transplant in patients with HCC. The different center-specific criteria at the time of transplantation (either primary or after downstaging) used by various institutes are outlined in Table 1.

The majority of centers (17/23; 74%) preferred positron-emission tomography (PET)-computed tomography (CT) as their modality of choice for metastatic work-up in HCC patients with chronic liver disease (CLD) planned for LT*.* The remaining centers (26%) opted for a combination of contrast-enhanced CT (CECT) of the abdomen and pelvis, chest CT, and bone scan (Figure 3).Approximately 65% of the centers did not have different criteria for LDLT and DDLT with respect to HCC–CLD patients. Six of the eight centers that had different criteria explained that they would list patients only under the UCSF criteria for DDLT, while they would opt for center-specific criteria to proceed with LDLT. One center mentioned that downstaged portal vein tumor thrombus (PVTT) with transarterial radioembolization (TARE) or stereotactic body radiotherapy (SBRT) would not be a candidate for DDLT at their center but would be a candidate for LDLT.

Most of the centers (21/23; 91%) used downstaging as a bridge to LT when the center-specific criteria were not fulfilled, but there was no absolute contraindication to LT. Of them, 18 centers (overall 18/23; 78%) would consider branch PVTT for downstaging prior to transplantation. Transarterial chemoembolization (TACE), TARE, and SBRT are common modalities used to downstage tumors for various indications. The indications for TACE, TARE, and/or SBRT as downstaging tools received eight responses, as outlined in Table 2.TARE was preferred over TACE in the presence of PVTT (12 responses), large or multiple tumors (six responses), and in all cases, when financially feasible (three responses), with some overlap in the responses. TACE was preferred mostly for large tumors without PVTT, in cases of financial restrictions, and when TARE was unavailable in some centers. The use of atezolizumab/bevacizumab combination in HCC patients awaiting transplantation was advocated by six centers, of which five would use it universally and one would use it when TACE/TARE was not feasible. Six other centers responded that they had no experience using atezolizumab or bevacizumab as part of the downstaging protocol.

Alpha-fetoprotein (AFP) was used as a marker for downstaging at most centers (17/23; 74%). The cut-off AFP value for transplant was 1000 ng/mL in most (10/17; 59%) centers, 400 ng/mL in four centers, and 2000, 500, and 200 ng/mL in one center each. All 17 centers considered AFP as a criterion for downstaging based on their set cut-off levels. Sixteen centers (70%) used protein induced by vitamin K absence or antagonist II (PIVKA-II) as a biomarker for HCC surveillance. All centers (19 responses) considered successful downstaging when their center-specific criteria or transplant listing criteria, including the AFP cutoff, were met. The most common determinants were decreased tumor size, clearance of PVTT, reduced AFP/PIVKA-II, loss of PET avidity or CT enhancement, and non-progression of tumor status. Opinions were divided among centers regarding when transplants should be performed after downstaging. Nine centers (9/23; 39%) thought it should be more than 6 wk, whereas six (26%) and seven centers (30%) thought it should be 4 wk and 6 wk, respectively. For post-operative surveillance, CECT-abdomen was the preferred imaging of choice (52%), followed by PET-CT (35%). The remaining few centers opted for CT + Bone Scan on follow-up (Figure 4).Everolimus was preferred by 22 of the 23 centers at different times post-transplant, with only one center not using it routinely (Figure 4)**.**

**DISCUSSION**

This survey covered a wide range of transplant centers across India, with an overall experience of over 8000 Liver transplants. Based on these results, we derived an idea of the distinct practices around the country regarding HCC leading to LT and its subsequent follow-up. Despite certain clear-cut agreements, many corresponding answers have highlighted gray areas where judgments and opinions differ and are of utmost importance in different settings.

***Selection criteria for HCC***

The selection criteria for HCC in LT have always been debated. From the early days of the Milan criteria to UCSFand, more recently, the Expanded Selection Criteria, it has been well established that cancer-free survival is dependent largely on extrahepatic spread and the level of vascular invasion, as compared to that on the size and number of tumors[20,21,25]. There is increasing evidence that outcomes outside the age-old criteria, such as the Milan criteria, are near-equivalent or at least good, as shown in Table 3[21,25–33].In a country like India, where the burden of cirrhosis patients is huge and most patients are from the lower socioeconomic status, it is most usual for HCC to present in a late-stage with a background of CLD where they are often beyond Milan or UCSF criteria[10,11,15,16]. The diagnosis of these patients is often delayed owing to the unavailability of facilities or a lack of awareness in rural/semi-urban centers. The 5-year survival rate of these patients is extremely low[15,16]. In this situation, external criteria from predominantly Western or other developed countries may not be suitable for Indian patients in the current scenario. The availability and use of direct-acting antivirals did not have any impact on the incidence or recurrence of HCC; however, extensive data are lacking in this regard[34–36].

In our survey, 5%–15% of patients undergoing LT in India were diagnosed with HCC annually. Of these, only 5% belonged to the Milan category. Since the advent of the Milan criteria, advancements in radiological techniques have made it possible to achieve extremely accurate staging. LDLT, with a high degree of donor safety, has mitigated organ availability issues. Hence, the expansion of recipient criteria has become possible with LDLT, even with slightly inferior outcomes compared to those in Milan[37]. In our opinion, any treatment that offers at least a chance of 60% 5-year disease-free survival should be acceptable and offered to a patient and their donor for LDLT and should not be outrightly rejected[38].

Regarding the listing of patients with HCC–CLD, there has been considerable debate on whether the same criteria used for LDLT are applicable for DDLT. More recently, expanded criteria have been shown to have comparable outcomes, and this dilemma has intensified. In general, DDLT listing has been reserved for those patients who have a similar 5 year survival as compared to non HCC patients (*e.g.* Milan or UCSF criteria)[37,38]. This reservation is due to the potential impact of this listing on other patients on the liver waitlist. It has also been suggested that DDLT listings should be subject to regional listing criteria for patients with HCC, whereas LDLT can be pursued with more liberal center-specific criteria, providing a full disclosure of risks and outcome benefits[37]. Our survey sheds light on the fact that up to 65% of centers preferred to use the same criteria for LDLT and DDLT listing. Of the eight experienced centers, three opted for separate listing criteria, while five opted for the same criteria.

***Metastatic work-up***

The current diagnostic tools for HCC include ultrasonography, CT, magnetic resonance imaging (MRI), and biopsy[39]. Biopsy confirmation is usually not required for a diagnosis[40]. Triple CT or MRI is the best imaging modality to diagnose HCC in patients with CLD. Current literature on the best imaging method for the evaluation of HCC metastasis is scarce. CT is the most accurate technique; however, it has limitations with respect to bone lesions, small vascular tumors, and difficulty in distinguishing between scarring and metastases[41–43]. The 18-Fluoro-deoxy-glucose-PET-CT has become increasingly established for the evaluation and treatment of metastatic HCC, with an average sensitivity of 60%–80% in most studies[44–46]. Other programs use a combination of dynamic CECT or MRI, chest CT and bone scintigraphy[47]. In our survey, 74% of centers chose PET-CT, whereas the remaining opted for the latter as a metastatic work-up prior to transplantation. AFP is considered an important biomarker for the diagnosis, treatment, and follow-up of patients with HCC before and after treatment[48]. It has also been implicated in the development and progression of HCC along with drug resistance in HCC cells[49]. However, only 60%–70% of HCC cases show elevated AFP levels, while 30%–40% of patients have normal values[50,51]. Newer biomarkers and models such as lens culinaris agglutinin-reactive fraction of AFP, des-carboxy-prothrombin, and GALAD scores (gender, age, AFP-L3, AFP, and DCP) are being increasingly used by various centers around the world[52,53]. In our study, AFP was universally followed, whereas PIVKA II was followed up in nearly 70% of the centers.

***Downstaging for HCC***

In our survey, more than 90% of the centers considered downstaging of HCC either as a bridge to transplantation or to fit the respective listing criteria or center-specific criteria for LDLT. The various indications mentioned by the surveyed participants, along with their corresponding modalities, are listed in Table 3. TACE and TARE were the most popular choices depending on availability and feasibility, whereas SBRT was mostly reserved for branch PVTT. A recent meta-analysis found that down-staged HCC–CLD patients who were initially beyond the listing criteria and who underwent transplantation had much better 3- and 5-year survival rates than non-transplanted patients[54]. They also noted that patients with downstaged HCC–CLD did not have inferior outcomes to transplant recipients who met the listing criteria[54]. Although the current European Association for the Study of the Liver and American Association for the Study of Liver Diseases guidelines suggest LT for downstaging to the Milan criteria, while the United Network for Organ Sharing (UNOS) adopted the UCSF criteria, the Indian perspective is different from the point of view of its socio-economics, advanced stage at diagnosis, and overall poor 5-year survival[55–57]. Mazzaferro *et al*[58] demonstrated that patients with downstaged HCC–CLD (to Milan) had a 77% 5-year overall survival rate compared to that of 31% with conventional anticancer therapies[58]. In this survey, TARE was preferred in many centers when available and affordable, especially in the presence of PVTT or multifocal HCC. An international systematic review of TARE as a downstaging tool before LT in 178 patients concluded that TARE is safer and better than TACE, with a 79% success rate[59]. Radunz *et al*[60] performed TARE downstaging in 40 pre-transplant patients and demonstrated an 87% tumor response (both complete and partial)[60]. However, another comparative meta-analysis indicated that TACE may have a better overall outcome than TARE when indicated with an approximately 60% tumor response[61–63]. Soin *et al*[64] demonstrated that after successful downstaging of PVTT (Vp1-3), a 5-year overall survival rate of 57% was obtained, which was comparable to that of patients without PVTT (65%)[64]. Regardless of the preference, downstaging with TACE or TARE is widely used throughout the country, with comparable results to those within the respective criteria for LDLT or DDLT.

SBRT is less frequently used but has been established as a safe alternative to conventional bridging therapies such as radiofrequency ablation (RFA), TACE, and TARE[64–67]. Patients with contraindications to TACE, especially those with PVTT, may receive SBRT[68]. Compared to other forms of treatment for PVTT like 3D-chemoradiation therapy, hepatic artery infusion chemotherapy, and molecular targeted drugs for HCC, SBRT offers a higher biologically effective dose in a shorter duration[69]. Retrospective studies of SBRT as a downstaging tool have indicated a good response and overall 5-year survival post-LT. In India, most centers select SBRT when TACE/TARE is not feasible or in the presence of branch PVTT (Vp1-2). However, the use of AFP in downstaging protocols remains controversial. There is no consensus among centers around the globe regarding the incorporation of biological (tumor markers, such as AFP) and morphological features for downstaging prior to transplantation. When adopting the UCSF criteria, the UNOS also suggested that a significant drop in AFP (< 500 ng/mL) along with stable disease at 6 months would be acceptable for DDLT listing[21,57]. Other studies have proposed various cutoffs for initial listing and downstaging endpoints ranging from < 100 to < 1000 ng/mL, while a few criteria have no cutoff and would accept any AFP if morphological variables were acceptable[25,30,48,70]. In our study, the majority of centers used 1000 ng/mL as a cut-off for AFP either at primary listing or after downstaging to proceed with LT. It is universally agreed that higher AFP levels impact the risk of recurrence and have worse outcomes than lower AFP levels. Finally, a combination of atezolizumab and bevacizumab was used by six centers as a bridge to transplantation. Several worldwide reports have suggested successful downstaging of advanced HCC with combination immunotherapy[71–72]. There is significant concern regarding the safety of using immunotherapy in patients with HCC who may later undergo liver transplant, especially given the risk of immune-related adverse events. In the IMBrave 150 trial, grade 3 to 4 toxicities were reported in 38% of patients receiving combination therapy with atezolizumab and bevacizumab[73]. In our study, many other centers did not use it because of a lack of experience, controversial nature or affordability issues.

The downstaging criteria for most centers were similar to their respective criteria for LT. The overall goal of downstaging is to give the opportunity for higher survival through LT to patients with HCC–CLD who would otherwise not fall into the LT criteria. Clavien *et al*[37] recommended that downstaging should only be performed when the 5-year survival rate after LT is comparable to those that fit the criteria without downstaging[37]. Our survey provided varying opinions on this aspect. Morphological and biological tumor responses were the main aspects, while the non-progression of tumors was also an important factor to consider. The modified Response Evaluation Criteria in Solid Tumors was also used by several centers[74,75]. Notably, all transplant centers waited at least 4 wk, with nearly 70% preferring to wait 6 wk after successful downstaging to ensure disease stability.

***Post-operative care and follow-up***

There is no international consensus on the post-transplant surveillance of HCC patients. The National Comprehensive Cancer Network guidelines suggest imaging and AFP every 3–6 months initially, followed annually thereafter[76]. We have a similar protocol for HCC surveillance after LT. Patients with hepatitis B usually continue antiviral therapy. In this survey, more than 50% respondents opted for CECT abdomen alone as their imaging of choice, while the remaining picked PET-CT or CECT abdomen with bone scintigraphy. Many pre-transplant factors are implicated in the risk of HCC recurrence, such as the number and size of nodules, vascular invasion, AFP level, neutrophil-to-lymphocyte ratio, bridging therapy prior to transplantation, presence of metabolic syndrome, viral infections, and time to transplant[77]. In the post-transplant period, immunosuppression with calcineurin inhibitors at higher levels has been implicated in recurrence but has not yet been established[78]. However, it is well established that most HCC recurrences occur within 2 years post-LT[79–81]. Regardless of the type of imaging or cause of recurrence, early diagnosis and treatment by RFA or resection offer the only hope for long-term survival. The use of mammalian target of rapamycin inhibitors in post-transplant period is not routinely recommended according to International Liver Transplant Society guidelines[78]. However, in the current context, everolimus was routinely used by 22 of the 23 centers listed in this study.

***Expansion of current criteria***

HCC is one of the leading causes of cancer-related deaths worldwide, with an annual global mortality rate of more than 800000[4]. An increase of up to 55% in the global burden of HCC is expected by 2040 (an estimated 1.3 million cases and 1.4 million deaths)[12–14]. LT offers hope to patients with HCC–CLD without extrahepatic disease for a better chance of survival[15–19]. It has already been established as the best treatment option for patients, with the highest survival rate. However, for long, LT was not considered an option for patients with HCC–CLD. This was followed by an era in which stringent criteria for sufficiently good outcomes were used to justify the use of deceased donor livers for other patients on waitlists[20,21,37]. Over the years, this has been accepted as the benchmark for new and upcoming guidelines and their corresponding results. The use of living donor grafts has mitigated the concern of the use of deceased donor livers for HCC patients; however, it has raised issues over overall survival rates compared to the risk of living liver donation. The benchmark of survival is highly debatable, but in a country like India, where the non-transplant survival of HCC–CLD patients is extremely low, any chance of a 5-year success beyond 50% warrants sufficient discussion[37,64]. Markov models and other recent downstaging studies suggest that a 5-year survival rate of 60% is worth the minimal risk of living donations and deceased donor candidacy[38]. However, other guidelines have suggested deceased donor candidacy at outcomes comparable to those of patients with CLD without HCC, whereas LDLT can be pursued with lower outcomes in the setting of full disclosure of risks and benefits[37].

***Summary***

Based on our survey, we summarize the following trends across liver transplant programs in India:

(1) Approximately 10% of CLD patients in India undergoing LT are diagnosed with HCC; however, only 5% of these patients fall within Milan criteria;

(2) Most centers follow the expanded center-specific criteria for LDLT, with comparable outcomes to those who fall within the Milan criteria. However, further validation is required through national collaborations and multicenter studies;

(3) PET-CT is the most preferred modality of metastatic work-up in HCC–CLD patients. AFP is the biological marker of choice; however, many centers opt for PIVKA-II surveillance;

(4) All centers opted for downstaging as a bridge to LT or to fit center-specific criteria if no extrahepatic metastasis or major vascular invasion was present. TACE, TARE, and SBRT are the therapies of choice with varying indications, whereas atezolizumab/bevacizumab combination immunotherapy is infrequently used. Downstaging was confirmed using both morphological and biological markers according to either international or center-specific guidelines;

And (5) Post-transplant surveillance was mostly guided by CECT abdomen and tumor markers, while some centers opted for PET-CT or CECT and bone scintigraphy. Despite the lack of concrete evidence, almost all centers started administering everolimus in the post-transplant period for HCC–LT patients.

**CONCLUSION**

The current predicted 5-year survival rate of HCC patients in India is less than 15%. The aim of transplantation is to achieve at least a 60% 5-year disease free survival rate, which will provide relief to the prediction of an HCC surge over the next 20 years. The current worldwide criteria (Milan/UCSF) may have a higher 5-year survival (> 70%); however, the majority of patients still do not fit these criteria and are dependent on other suboptimal modes of treatment, with much lower survival rates. To make predictions for 2040, we must prepare to arm ourselves with less stringent selection criteria to widen the pool of patients who may undergo transplantation and have a chance of a better outcome. With more advanced technology and better donor outcomes, LDLT will provide a cutting edge in the fight against liver cancer over the next two decades.

**ARTICLE HIGHLIGHTS**

***Research background***

Hepatocellular carcinoma (HCC) with chronic liver disease (CLD) is an indication for liver transplantation (LT). However, the overall survival for this condition is low in India, especially due to late presentation.

***Research motivation***

The various criteria that are established worldwide may lead to comparable outcomes compared to non-HCC patients, but significantly limit the number of patients that can avail this treatment option.

***Research objectives***

The aim of our study was to establish the current trends and give our opinion as to how to improve the donor pool or increase the access of patients to this life saving treatment option by relaxing stringent criteria while maintaining at least significant survival benefit.

***Research methods***

We conducted a survey to see the current trend of practices in India with regards to HCC-CLD patients undergoing LT.

***Research results***

In this survey, we were able to ascertain trends of practice in HCC-CLD patients with respect to LT. We were also able to identify possible pathways to improve access of LT to these patients and improve the overall survival rates of HCC patients in India to make it comparable to other cancers.

***Research conclusions***

This study shows that majority of patients are still dependent on sub optimal modes of treatment, and less stringent criteria may need to be followed with acceptable outcomes so that we may be able to match the increasing burden on HCC predicted over next 2 decades.

***Research perspectives***

To make predictions for 2040, we must prepare to arm ourselves with less stringent selection criteria to widen the pool of patients who may undergo transplantation and have a chance of a better outcome.

**ACKNOWLEDGEMENTS**

We would like to acknowledge the following hospitals for their invaluable contributions to this survey. In alphabetical order, they are: Aakash Hospital, New Delhi; AIG Hospital, Hyderabad; Amrita Institute, Kochi; Apollo Hospital, Chennai; Apollo Multispecialty Hospital, Kolkata; Aster CMI Hospital, Bengaluru; BGS Gleneagles Global Hospital, Bengaluru; Center for Liver & Biliary Sciences, New Delhi; Deenanath Mangeshkar Hospital, Pune; Gem Hospital, Coimbatore; Gleneagles Global Hospital, Chennai; Global Hospital, Hyderabad; Global Hospital, Mumbai; Indraprastha Apollo Hospital, New Delhi; Jaypee Hospital, Noida; Kokilaben Dhirubhai Ambani Hospital, Mumbai; Medicover Hospitals, Navi Mumbai; MGM Healthcare, Chennai; Narayana Health; New Era Hospital & Research Institute, Nagpur; Sahyadri Hospital, Pune; Sir HN Reliance Foundation Hospital, Mumbai; Zydus Hospital, Ahmedabad.

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**Footnotes**

**Institutional review board statement:** This is a survey of various institutions and a review of literature with authors opinion and directly involving any patients. It was reviewed by institutional board and exempted from review.

**Informed consent statement:** There is no patient information in the article.

**Conflict-of-interest statement:** All the authors report no relevant conflicts of interest for this article.

**Data sharing statement:** There is no patient information in the article and data of the survey is available on request.

**STROBE statement:** The authors have read the STROBE Statement—checklist of items, and the manuscript was prepared and revised according to the STROBE Statement—checklist of items.

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**Provenance and peer review:** Invited article; Externally peer reviewed.

**Peer-review model:** Single blind

**Corresponding Author's Membership in Professional Societies:** International Liver Transplant Society; Liver Transplant Society of India, 303; Indian Society of Organ Transplantation, LM 1427.

**Peer-review started:** October 11, 2023

**First decision:** November 21, 2023

**Article in press:**

**Specialty type:** Transplantation

**Country/Territory of origin:** India

**Peer-review report’s scientific quality classification**

Grade A (Excellent): 0

Grade B (Very good): B

Grade C (Good): C

Grade D (Fair): 0

Grade E (Poor): 0

**P-Reviewer:** Dabbous H, Egypt **S-Editor:** Li L **L-Editor:** A **P-Editor:** Li L

**Figure Legends**

图表, 饼图

描述已自动生成

**Figure 1 Total number and yearly volume of liver transplants at the participating centers.**

图表, 饼图

描述已自动生成

**Figure 2 Total number of liver transplants performed in patients with hepatocellular carcinoma (center-wise) and percentage of transplant patients with hepatocellular carcinoma.** HCC: Hepatocellular carcinoma; LT: Liver transplantation.

**图表, 条形图

描述已自动生成**

**Figure 3 Preferred metastatic work-up imaging modality in patients with hepatocellular carcinoma planned for transplant.** PET: Positron-emission tomography; CT: Computed tomography; HRCT: High resolution computed tomography; CECT: Contrast-enhanced computed tomography.

图表, 饼图

描述已自动生成

**Figure 4 Post-operative imaging and everolimus use preference in centers across India.** PET: Positron-emission tomography; CT: Computed tomography; HRCT: High resolution computed tomography; CECT: Contrast-enhanced computed tomography; LT: Liver transplantation.

**Table 1 Various center-specific criteria for hepatocellular carcinoma used at the time of liver transplantation across India**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **No. of centers** | **Center-specific criteria** | | | |
| **Size/No. of tumor** | **Invasion** | **Extrahepatic** | **AFP/markers** |
| 4 | Any size/any No. | No macrovascular | No | Any |
| 2 | Any size/any No. | No macrovascular | No | < 1000 |
| 1 | Encapsulated, any size, < 10 | No macrovascular | No | < 400 |
| 1 | Within UCSF size/No. | Vp1-vp3 invasion | No | < 400 |
| 1 | Any size/any No. | Vp1-vp2 invasion | No | Any |

UCSF: University of California, San Francisco criteria; AFP: Alpha-fetoprotein.

**Table 2 Indications of transarterial chemoembolization, transarterial radioembolization and stereotactic body radiotherapy in hepatocellular carcinoma–chronic liver disease patients awaiting liver transplantation**

|  |  |  |  |
| --- | --- | --- | --- |
| **Modality** | **TACE** | **TARE** | **SBRT** |
| Indications  (No. Of centers preferred) | HCC patients on waitlist[12] | PVTT[12] | Vp1-3 PVTT[12] |
| > Milan[4] | Large/multiple HCC[6] | Vp2 PVTT[2] |
| > UCSF[2] | All affordable cases[3] | TACE/TARE not possible[4] |
| Large tumor size[13] |  | Exophytic HCC[1] |
| Awaiting donor fitness/logistical delay in transplant[2] |  | Diaphragm involved or local infiltration[1] |
| High AFP[5] |  | Presence of shunt[1] |
| Absence of PVTT[2] |  | Not preferred[3] |
| TARE unaffordable/unavailable[4] |  |  |

There is overlap among the respondents for the indications of either modalities. HCC: Hepatocellular carcinoma; PVTT: Portal vein tumor thrombosis; UCSF: University of California, San Francisco criteria; TACE: Trans-arterial chemo embolization; TARE: Trans-arterial radio embolization; SBRT: Stereotactic body radiotherapy; AFP: Alpha fetoprotein.

**Table 3 Different criteria for liver transplantation in hepatocellular carcinoma patients**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Criteria name (yr)** | **Size of tumor (cm)** | **No. of tumors** | **Additional criteria** | **Overall 5-year survival** |
| Milan criteria (1996) | ≤ 5; ≤ 3 | 1; 3 | None | 75% |
| UCSF criteria (2001) | 6.5; ≤ 4.5 (total ≤ 8) | 1; 3 | None | 75.2% |
| Up-to-7 criteria (2001) | Size (cm) + No. ≤ 7 |  | None |  |
| Navarro criteria (2001) | ≤ 6; ≤ 5 | 1; 3 | None | 79% |
| Tokyo criteria (2007) | ≤ 5 | ≤ 5 | None | 75% |
| Asan criteria (2008) | ≤ 5 | ≤ 6 | None | 82% |
| Hangzhou criteria (2008) | < 8 (total) | Any No. | AFP < 400 ng/mL | 72% |
| Chang Gung criteria (2008) | ≤ 6.5; ≤ 4.5 | 1; ≤ 3 | None | 90% |
| Hong Kong criteria (2008) | ≤ 6.5; ≤ 4.5 | 1; ≤ 3 | None | 66% |
| Kyushu criteria (2009) | ≤ 5 | Any No. | PIVKA-II < 300 mAU/mL | 83% |
| Kyoto criteria (2010) | ≤ 5 | ≤ 10 | PIVKA-II < 400 mAU/mL | 87% |
| Toronto criteria (2011) | Any Size | Any No. | Poorly differentiated HCC excluded | 72% |
| Japanese National Expanded criteria (2019) | ≤ 5 | ≤ 5 | AFP < 500 ng/mL | 75.8% |

All the criteria exclude any vascular invasion or any extra-hepatic spread[25–33]. HCC: Hepatocellular carcinoma.