Diagnostics and Treatment of Hepatocellular Carcinoma in 2016: Standards and Developments

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Introduction

Hepatocellular carcinoma (HCC) is one of the most common tumors worldwide, with more than 30 cases per 100,000 people annually in Southeast Asia and central Africa. Incidence rates of the disease are increasing globally. In the United States, the incidence has doubled over the past 20 years [1]. A similar trend has been observed in Germany [2], with currently around 6,000 new male and 2,500 new female patients per year (fig. 1).

The risk for HCC is influenced by etiology, activity, and stage of underlying liver disease. Patients with liver cirrhosis due to chronic infection with hepatitis C virus (HCV) or hepatitis B virus (HBV) have the highest risk. The rising incidence in Western countries is mainly due to the still high prevalence of HCV-associated liver cirrhosis and the increasing number of patients with advanced steatohepatitis as hepatic manifestation of the metabolic syndrome (e.g. due to diabetes and obesity) [1]. Several reports indicate that HCC in the setting of metabolic syndrome may also arise in the absence of cirrhosis; however, the exact proportion of this condition or its risk factors are still under investigation [3].

Non-Invasive and Invasive Diagnosis of HCC

HCC is typically a highly vascular tumor, which can be utilized regarding the non-invasive diagnostics of HCC. Non-invasive diagnosis of HCC in patients with liver cirrhosis is established by one imaging technique (contrast-enhanced computed tomography (CT) or magnetic resonance imaging (MRI)) in nodules larger than 2 cm showing the HCC radiological hallmark of contrast uptake in the arterial phase and wash-out in the venous/late phase (fig. 2). In nodules between 1 and 2 cm in size, two coincidental techniques (contrast-enhanced CT, MRI, or ultrasound) are necessary to establish the diagnosis. Pathological diagnosis of HCC is...
recommended for all nodules occurring in non-cirrhotic livers as well as for those cases with inconclusive or atypical imaging appearance in patients with liver cirrhosis. The histopathological diagnosis is based upon morphological criteria and immunohistochemistry. The marker glypican 3 (GPC3) has a sensitivity of approximately 70% and a specificity of more than 90%. Further helpful tissue markers are HSP70 and GS. The International Consensus Group of Hepatocellular Neoplasia recommends that a pathological diagnosis of HCC is established if at least two of these markers are positive. Additional staining can be considered to assess neo-vascularization (CD34) or potential progenitor cell origin (keratin 19, EpCAM) [4]. At the molecular level, HCC is characterized by high-level DNA amplifications in chromosome 6p21 (VEGFA) and 11q13 (FGF19/CNND1) as well as homozygous deletions in chromosome 9 (CDKN2A). The most frequent mutations affect the TERT promoter (60%), which is associated with an increased telomerase expression. The TERT promoter can also be affected by copy number variations and hepatitis B DNA insertions; moreover, mutations of it can be found in preneoplastic lesions. TP53 and CTNNB1 are the next most prevalent mutations, affecting 25–30% of tumors [5]. Some of these changes may cause activation of angiogenesis, which results in the typically seen vascularization of the tumor.

Staging and Therapeutic Algorithm

HCC includes a wide spectrum of tumors behaving biologically differently. Moreover, allocation of the available treatment options is influenced not only by tumor stage but also by the degree of liver dysfunction. Therefore, accurate staging and classification of the underlying liver disease is crucial for patient management. Several staging systems are well validated and widely used. The Barcelona Clinic Liver Cancer (BCLC) staging system is recommended by the American Association for the Study of Liver Diseases (AASLD) and the European Association for the Study of the Liver (EASL) [4, 6]. Further algorithms are the Hong Kong Liver Cancer (HKLC), the Cancer of the Liver Italian Program (CLIP), the Chinese University Prognostic Index (CUPI), the Japan Integrated Staging (JIS), the Liver Cancer Study Group of Japan (LCSGJ), the Taipei Integrated Scoring System (TIS), the Okuda, the Chinese University Prognostic Index (CUPI), and the French systems [7].

Using the BCLC algorithm, HCC is categorized as early- (BCLC 0 and BCLC A), intermediate- (BCLC B), advanced- (BCLC C), or end-stage (BCLC D) (fig. 3). Accordingly, surgical (resection or transplantation) or percutaneous thermal therapies (radiofrequency ablation (RFA) or microwave ablation (MWA)) are mainly considered suitable for early stages, while interventional therapies (transarterial chemoembolization (TACE) or radioembolization) are applied in patients with intermediate-stage HCC. Systemic treatment with the tyrosine kinase inhibitor sorafenib is considered as the treatment of choice for patients with advanced-stage HCC. Patients with BCLC stage D do not benefit from cancer treatment, and thus are being considered for best supportive care only.

Liver Resection and Transplantation

In patients with HCC without liver cirrhosis, surgical resection is the treatment of choice. In patients with cirrhosis, normal measurements of bilirubin and the absence of portal hypertension (no obvious signs of collaterals, platelets > 100/μl, and normal-sized spleen) have been found to be good predictors for a sufficient functional hepatic reserve after resection. Assessing the functional capacity of the remaining liver volume after resection can raise difficulties in the individual case. As a rule, only patients with Child–Pugh stage A cirrhosis can undergo resection. Intraoperative ultrasonography has become a standard intraoperative procedure to plan the resection with precision while taking anatomical structures into consideration. A safety margin is desirable, especially in view of existing satellite foci; however, in most cases, only limited resections are possible since the functional hepatic reserve is impaired. The 5-year survival rate after partial liver resection in Western countries is 20–50%. The risk of a tumor recurrence and/or dissemination is highest in the first 3 years postoperatively. In specialized centers, TACE is initially performed in borderline resectable tumors in neoadjuvant intention. However, this approach has not been validated in a randomized fashion yet. Adjuvant treatment after R0 resection is not established [8]. A complication associated with surgical treatment – in addition to the usual risks as-

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Fig. 1. Annual incidence rates of liver cancer in Germany from 1999 till 2012 for men (blue) and females (red). The numbers given indicate the estimated annual number of patients with newly diagnosed primary liver cancer (ICD-10 code C22) (Zentrum für Krebsregisterdaten im Robert Koch-Institut, www.krebsdaten.de; accessed 10.01.2016).
associated with hepatobiliary procedures – is hepatic decompensation, especially in patients with a low functional hepatic reserve.

In liver cirrhosis, liver transplantation is the best therapeutic option, because removal of the entire organ eradicates the tumor and precancerous lesions, and the impaired liver function is restored. Early experiences with liver transplants date back to the 1980s and were mostly disappointing, with 5-year survival rates of less than 40%. After the Milan criteria were introduced, the results of liver transplantation in this indication improved notably. However, this limits organ allocation to patients with a solitary HCC nodule of less than 5 cm in size or 3 nodules of less than 3 cm in size each, and no infiltration of the portal vein or extrahepatic manifestations. By selecting patients carefully, 5-year survival rates of more than 70% can now be achieved [9]. Currently, the criteria cannot be extended to patients with larger tumors because of the shortage of organs. The Eurotransplant organ allocation criteria have been changed in December 2006, and since that time transplant waiting times for patients with HCC are much shorter. Bridging treatment such as TACE is widely used in transplant centers. Another option is living liver donation.

**Thermal Ablation in Early-Stage HCC**

Thermal ablation is an alternative treatment for cirrhotic patients with early-stage HCC and contraindications for surgical treatment. In this setting, procedures such as RFA or MWA are potentially curative therapeutic options. The complication rate of RFA is low, and in suitable patients with good liver function a 5-year survival rate of up to 50% may be achieved [4]. The complications of RFA include liver abscess, biloma, and right-sided pleural effusion. Recent prospective and retrospective data showed a significant survival benefit for the combination of RFA and TACE [10, 11]. In a randomized trial comparing TACE followed by RFA versus RFA in patients with HCC < 7 cm in size, the combination resulted in 1-, 3-, and 4-year overall survival rates of 93, 67, and 62%, respectively, compared to 85, 59, and 45% in the RFA-only arm, respectively [10]. The corresponding recurrence-free survivals were 79, 61, and 55% as well as 67, 44, and 39%, respectively.

**Transarterial Chemoembolization**

For intermediate-stage tumors (BCLC stage B) TACE is the treatment of choice [4, 6]. TACE is based on the simultaneous application of a chemotherapeutic agent and embolization with occluding particles. Prerequisite for successful TACE is the presence of a hypervascularized tumor. Selective administration of the mixture of chemotherapeutic agent and occluding particles results in a high local concentration of the chemotherapeutic agent in the tumor with low systemic distribution. Due to occlusion of the tumor vessels, the chemotherapeutic agents remain in the tumor region and the resulting hypoxia improves the effect of the chemotherapeutic agent. Procedures for TACE are not standardized. Commonly used chemotherapeutic agents are doxorubicin, mitomycin C, and cisplatin. Lipiodol as an oily suspension has an affinity to HCC and acts as a carrier for the chemotherapeutic agent. A favorable prognostic factor for successful TACE besides tumor size of less than 4 cm and increased vascularization is also uptake of lipiodol in at least 75% of the tumor volume. TACE is either performed as ‘on demand’ (repeated in case of persistent vascularization) or as ‘continuous’ (repeated every 4–6 weeks until devascularization) schedule. An alternative approach is to use doxorubicin-eluting beads for TACE. An older meta-analysis showed a survival benefit for patients with TACE [12]. Overall, 545 patients were evaluated in this study. TACE improved the 2-year survival rate with an odds ratio of 0.53 (95% confidence interval (CI): 0.32–0.89; p < 0.017) in comparison to not treated control patients. Objective response could be achieved in 35% of patients.

Since both TACE and the oral tyrosine kinase inhibitor sorafenib are active treatments for HCC, there is a rationale to combine these two treatment modalities. Liu et al. [13] performed a meta-analysis on the subject and concluded that the combination seems to be
Radioembolization

Radioembolization with yttrium-90 labelled microspheres, also known as selective intra-arterial radiotherapy (SIRT), is increasingly offered for patients with unresectable HCC who are no candidate for transplantation or ablation. Yttrium-90 labelled microspheres preferentially lodge into the tumor microvasculature if injected into the tumor-feeding vessels and lead to locally limited tumor radiation, thus minimizing radiation hazard to surrounding normal liver parenchyma. The reported median survival ranges between 7 and 42 months in cohorts of patients with different stages of HCC. However, these data are based on mainly retrospective cohort analyses, and randomized data using SIRT for HCC treatment are currently not available [15]. However, data from the randomized SORAMIC trial comparing radioembolization in combination with sorafenib versus sorafenib are awaited in the near future [16].

Systemic Treatment

Sorafenib is an oral multi-tyrosine kinase and angiogenesis inhibitor with activity against vascular endothelial growth factor receptor (VEGFR)-2, platelet-derived growth factor (PDGFR), c-Kit receptors, BRAF, and p38 signaling pathways. It is the only approved systemic treatment option in advanced-stage HCC, resulting in a relevant survival benefit compared to placebo (10.7 vs. 7.9 months) [17]. In clinical practice, these results have been confirmed by several retrospective and prospective cohort studies [18]. Underlying liver dysfunction and low tolerance of treatment toxicity may adversely affect outcomes and increase susceptibility to adverse events during treatment with sorafenib. These adverse events include fatigue (up to 40%), diarrhea, hand-foot skin reactions, anorexia, and others. In clinical trials as well as in daily practice, they often lead to dose reduction, interruption, and early discontinuation of treatment and therefore warrant close monitoring.

Sorafenib is not eligible as adjuvant treatment for HCC. This has been recently reported in the STORM trial, which studied adjuvant treatment with sorafenib versus placebo after complete resection or ablation of HCC [19]. Sorafenib did not show an improvement in progression-free survival in this phase III trial. Therefore, adjuvant treatment with sorafenib should not be recommended in patients after complete resection or ablation.

Within the last years, several phase III trials of investigational agents in the first-line treatment of patients with advanced HCC have been reported. However, none of these drugs, e.g. sunitinib, brivanib, linifanib, or the combination of sorafenib and erlotinib, were superior to sorafenib in terms of overall survival or toxicity. Furthermore, until today there is no established systemic second-line treatment option in patients who have progressed or cannot tolerate sorafenib treatment. Currently, several agents with different targets are investigated in clinical trials in patients with advanced HCC. The most promising agents in advanced clinical development target angiogenesis (ramucirumab, lenvatinib, and regorafenib), the MET pathway (tivantinib, caboctanzinib, and tepotinib), anti-PD1 antibodies (nivolumab and pembrolizumab), and oncolytic viruses (Pexa-Vec, T-Vec) [20].

Disclosure Statement

J. Trojan is serving on advisory boards for Amgen, Bayer Healthcare, Bristol Myers Squibb, Daiichi Sankyo, Lilly, and Merck Serono.
References


