Future Treatment Option for Hepatocellular Carcinoma: A Focus on Brivanib

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Abstract
Hepatocellular carcinoma (HCC), one of the most common cancers worldwide, is particularly prevalent in the Asia-Pacific region. Guidelines on the treatment of HCC in Japan come from both consensus-based and evidence-based treatment algorithms. However, patients with extensive liver damage and/or more advanced disease (major vascular invasion and/or extrahepatic spread) are currently ineligible for any treatment. Recent knowledge of hepatocarcinogenesis has led to the targeting of new pathways, particularly the angiogenic pathway, with a specific focus on the vascular endothelial growth factor receptor (VEGFR). Apparently the most studied systemic antiangiogenic agent for HCC is sorafenib. An updated version of the aforementioned treatment algorithms recommends sorafenib therapy for advanced HCC patients with Child-Pugh A liver function and extrahepatic spread or major vascular invasion. Moreover, sorafenib is recommended for use in HCC patients who are refractory or intolerant to transarterial chemoembolization (TACE) with well-preserved liver function (Child-Pugh A). However, one of the unresolved issues is anti-VEGF resistance. It is speculated that novel antiangiogenic agents that combine inhibition of other pathways such as fibroblast growth factor receptor signaling in addition to VEGFR signaling might provide a potential mechanism to overcome anti-VEGF resistance in HCC. Brivanib inhibits both VEGF and fibroblast growth factor receptor signaling. To further investigate the benefits of brivanib for advanced HCC, a broad-spectrum, global, phase III development plan, the Brivanib studies in HCC patients at RISK (BRISK) clinical program, has been initiated. Clinical benefits seen with brivanib in the first-line setting, and following the failure of sorafenib therapy, highlight the potential to improve the clinical course of patients with advanced HCC, and this agent may provide a novel therapeutic option for the growing population of patients for whom no other treatment choice exists.

Introduction
Hepatocellular carcinoma (HCC), one of the most common cancers worldwide, is particularly prevalent in the Asia-Pacific region, with more than two thirds of global cases occurring in Asia-Pacific countries [1, 2]. In Japan, HCC is now the third leading cause of cancer death among males and females, and is responsible for the death of more than 33,000 Japanese citizens every year [3]. Throughout the Asia-Pacific region, the most important etiologic factors related to HCC are hepatitis B virus (HBV) and hepatitis C virus (HCV). Among Japanese
HCC patients, the primary etiology is HCV, with approximately 70–80% chronically infected with HCV and only a small proportion with HBV (<16%) [4, 5].

**Treatment Algorithm and Unmet Medical Needs**

Guidance on the treatment of HCC in Japan comes from both consensus-based and evidence-based treatment algorithms [6, 7]. As nationwide HCC screening programs are common in Japan, most patients present in the early stages of the disease and are eligible for potentially ‘curative’ treatments, such as surgical resection or local ablation (radiofrequency ablation or percutaneous ethanol injection) [6, 7]. If resection or ablation is contraindicated, or if the disease has progressed, then transarterial chemoembolization (TACE) or hepatic infusion chemotherapy may be recommended [6, 7]; however, patients with extensive liver damage and/or more advanced disease (major vascular invasion and/or extrahepatic spread) are currently ineligible for these treatments [6]. As such, there remains a significant unmet medical need for patients with advanced HCC in Japan.

**Present Status of Molecular Targeted Therapy**

Recent knowledge of hepatocarcinogenesis has led to the targeting of new pathways, particularly the angiogenic pathway, with a specific focus on the vascular endothelial growth factor receptor (VEGFR). Indeed, agents that inhibit angiogenesis via blockade of the VEGFR have seen some success in the treatment of HCC. Moreover, recent research data suggest the potential for an additional synergistic role for antiangiogenic agents whereby they might be used following TACE therapy to increase response rates [8]. Apparently the most studied systemic antiangiogenic agent for HCC is sorafenib. This is an oral multikinase inhibitor that targets the tyrosine kinase activity of VEGFRs 1, 2, and 3, as well as platelet-derived growth factor receptor-β, and has recently demonstrated some efficacy over placebo in Child-Pugh A patients with advanced HCC [9]. Similar results have been observed in a study of sorafenib for patients with advanced HCC conducted in the Asia-Pacific region [10], and a recent phase 1 study has indicated favorable safety/tolerability and promising antitumor activity in a Japanese population [11]. On the basis of these results, sorafenib has been approved in Japan for the treatment of advanced HCC since May 2009.

**Indication of Sorafenib in Treatment Algorithm**

There are, however, unresolved issues regarding the optimal use of sorafenib for HCC. To date, survival benefits in clinical trials have been modest, and a relatively high incidence of hand-foot syndrome (all-grade events reported in ~20–45% of patients) [9, 10] and an increased risk of bleeding events have been reported in the international literature [12]. In Japan, primarily due to the design of the pivotal trials and the available data in HCC patients, sorafenib use has been strictly regulated and limited to patients with Child-Pugh A cirrhosis who are not candidates for resection, ablation, or TACE. Moreover, post-marketing surveillance of sorafenib in Japan has raised safety concerns regarding interstitial pneumonia, hepatic coma, and hepatic failure, which has led to revision of the Japanese package insert. Updated version of the aforementioned treatment algorithms recommend sorafenib therapy for advanced HCC patients with Child-Pugh A liver function and extrahepatic spread or major vascular invasion. Moreover, sorafenib is recommended for use in HCC patients who are refractory or intolerant to TACE with well-preserved liver function (Child-Pugh A) (for details, see Kudo, fig. 7, p. 299) [13–15].

**Anti-VEGF Resistance**

Recent studies suggest that tumor progression following treatment with antiangiogenic agents that target the VEGF signaling pathway alone may result from either evasive or intrinsic resistance [16]. Furthermore, there is strong evidence to support the hypothesis that evasive resistance to anti-VEGF blockade is associated with reactivation of tumor angiogenesis by alternative signaling pathways, one such mechanism of resistance being activation of the fibroblast growth factor (FGF) signaling pathway [17, 18]. Basic FGF (FGF2) is a potent angiogenic factor. Indeed, expression of FGF2 enhances growth, invasion, and angiogenesis of many tumor types [19, 20]. Moreover, recent evidence has shown that FGF is overexpressed and activated in HCC and that high FGF2 levels may predict for a poor clinical outcome among patients with HCC [20].

**Importance of FGF Signaling**

Considering the proposed importance of FGF signaling in HCC angiogenesis, it is clear that novel antiangiogenic agents that combine inhibition of FGF receptor sig-
naling with inhibition of VEGFR signaling might pro-
vide a potential mechanism to overcome anti-VEGF
resistance in HCC (fig. 1). With this in mind, it is worth-
while considering the potential future impact of brivanib
on the treatment of advanced HCC. Brivanib, a small-
molecule tyrosine kinase inhibitor, is the first oral selec-
tive dual inhibitor of FGF and VEGF signaling. In mul-
tiple preclinical models of human xenograft tumors, in-
cluding patient-derived HCC xenografts, brivanib has
shown potent antitumor activity and no overt toxicity
when dosed orally [21, 22]. Furthermore, brivanib has
demonstrated promising antitumor activity and accept-
able tolerability in a phase 2, open-label study in patients
with unresectable locally advanced or metastatic HCC
[23, 24]. Crucially, within this trial, brivanib showed
activity both as first-line therapy (overall survival: 10
months) or as second-line therapy in patients who had
failed prior antiangiogenic treatment, primarily with
sorafenib (overall survival 9.5 months) [24]. Of note, the
incidence of all-grade hand-foot syndrome was only 8%
in this study.

**Phase I and II Data of Brivanib**

Additional retrospective studies and subanalyses have
also confirmed that brivanib is effective in patients from
the Asia-Pacific region. In a subanalysis performed to
evaluate the effects of brivanib among Asian versus non-
Asian patients enrolled in the aforementioned phase II
study, median overall survival was 10.6 months among
Asian patients treated with first-line brivanib (versus 5.7
months in non-Asian patients) and 9.8 months among
Asian patients receiving brivanib as second-line therapy
(versus 9.4 months in non-Asian patients) [25]. Another
subanalysis, this time including only patients who re-
ceived first-line brivanib therapy in the phase 2 study,
indicated that overall tolerability was similar or slightly
better in the Asian population versus non-Asian patients
[26]. A further subanalysis comparing 125 Asian and
non-Asian patients enrolled in separate phase I and II
studies [23, 27] confirmed that exposures in these patient
subpopulations were similar following brivanib doses of
800 mg daily [28]. Finally, a phase 1 study of brivanib in
Japanese patients with advanced or metastatic solid tu-
mors, including HCC, has shown manageable tolerability
and a similar safety profile at the same 800-mg once-dai-
ly dose as used in Caucasian patients [29]. Moreover, the
study provided evidence of antitumor activity in this
uniquely Japanese population, with 8 of 13 patients (62%)
showing stable disease.

**Design of Phase III Global Study**

To further investigate the benefits of brivanib for ad-
vanced HCC, a broad-spectrum, global, phase III devel-
opment plan, the Brivanib studies in HCC patients at
RISK (BRISK) clinical program, has been initiated. The global BRISK program will enroll patients from countries in Africa, Asia (including Japan), Australasia, Europe, and North, South, and Central America, and will include investigations of brivanib in a variety of clinically relevant settings, including first-line head to head with sorafenib, second-line post-sorafenib, and TACE adjuvant settings. In addition, it is noteworthy that the BRISK study of brivanib as adjuvant treatment to TACE therapy is being led by Japanese investigators and is one of the first global registration programs to be led from Japan.

Conclusion

HCC continues to be a major healthcare burden in Japan. Although it is detected in the early stages in most Japanese patients and treated accordingly, there remains a population of patients with advanced HCC who have limited therapeutic choices. With the recent approval of the antiangiogenic agent sorafenib, options for these patients have improved, but clinical studies to date suggest only a modest survival benefit with sorafenib and there is potential for safety/tolerability issues and the development of resistance to the anti-VEGF blockade. Clinical benefits seen with brivanib in the first-line setting, and following the failure of sorafenib therapy, highlight the potential to improve the clinical course of patients with advanced HCC, and this agent may provide a novel therapeutic option for the growing population of patients for whom no other treatment choice exists.

Disclosure Statement

The author has no conflict of interest to declare.


