Hepatocellular Carcinoma in Adults

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Abstract: Hepatocellular carcinomas are the most common primary neoplasia of the liver. The global distribution of hepatocellular carcinoma is related to the prevalence of hepatitis C in the population. Other major etiologic factors include alcoholic liver disease and nonalcoholic steatohepatitis. The majority of cases are discovered when screening patients with either chronic hepatitis or cirrhosis, but occasional incidental cases have been reported. Molecular markers and associated gene alterations are a work in progress. Serological markers and radiology are used to detect the disease in high-risk populations, and to monitor response to therapy in the affected patients. Even though radiologic features are specific, tissue diagnosis may be required, particularly for atypical and smaller lesions. Ancillary studies including reticulin stain and immunohistochemistry are important for confirmatory diagnosis. Liver transplantation is curative for hepatocellular carcinoma, but due to limitations in organ availability, palliative care is required, which mostly includes chemoembolization and radio-ablation.

Keywords: alpha fetoprotein; BCLC staging; CLIP staging; hepatocellular carcinoma; Okuda staging
INTRODUCTION

Malignancies that arise from hepatocytes are denominated hepatocellular carcinomas. Hepatoma is another term; however, this old terminology implies a benign process and therefore, to avoid confusion, this term should not be used. Hepatocellular carcinomas are malignant tumors and have the capacity to metastasize and cause death. The most common sites of metastases are lungs, abdominal lymph nodes and bones (1). Since hepatocytes are the most abundant cells within the liver, hepatocellular carcinoma is the most common primary liver disease in adults (85% of the malignancies of the liver). It is the 6th most common malignancy worldwide with 841,080 cases in 2018 and predicted to be 1,361,836 in 2040 with significant mortality and morbidity (2). The incidence of hepatocellular carcinoma is directly related to geography and prevalence of infectious diseases. The highest rates of disease are found in Asia and southern Africa (up to 150 per 100,000), mostly due to the high prevalence of infectious hepatitis. In the USA, the prevalence is much less but still significant (6 per 100,000), typically affecting Asian men (3). The epidemiology is similar in Canada with a reported prevalence of 6.8 per 100,000; this significant rise (3x) since the 1980s is likely related to the expansion of intravenous drug abuse and consequent increase in cases of hepatitis C (4). In Europe, the rate ranges from 3 to 6 per 100,000 individuals depending on the availability of treatment and prevention of infectious hepatitis, and abuse of alcohol (5). Hepatocellular carcinoma is the second leading cause of cancer mortality worldwide. Screening programs and curative therapy for infectious hepatitis have contributed to a decrease in mortality but alcohol abuse and metabolic diseases including diabetes and obesity have surpassed the initial benefit (6, 7).

ETIOLOGY AND PATHOGENESIS

The hepatocytes are active metabolic cells involved in many cell functions and are subject to a large number of insults that may result in abnormal proliferation (8). Exhaustion of their regenerative capacity is typically exemplified by cirrhosis. Thus, the etiologic factors involved in the cirrhotic process are also involved in tumorigenesis of hepatocellular carcinoma. Chronic hepatitis B and C are by far the most common causes of hepatocellular carcinoma, followed by alcoholic liver disease and nonalcoholic steatohepatitis. (9–11). Hepatitis B DNA levels in excess of 200,000 IU/mL (1,000,000 copies/mL) have been reported to increase the incidence of hepatocellular carcinoma to 1,152 per 100,000 individuals (12, 13). Other etiologic factors in adults include aflatoxin-contaminated food, diabetes, obesity, and hemochromatosis (14, 15). In some instances, hepatocellular carcinoma is detected incidentally during routine image examination. These tumors are considered sporadic and are result of gene mutations that occur during a person’s lifetime.

Genetic susceptibility to hepatocellular carcinoma has been demonstrated in animal models, but not established in humans. Family clustering has been
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reported in some Chinese families and Alaskan natives, but infectious hepatitis B is a requirement for HCC development in this particular population (16, 17).

Other etiologic factors include conditions associated with congenital, genetic, and metabolic conditions and are directly related to the capacity of these pathologies to develop cirrhosis. Such examples include hemochromatosis, Wilson’s disease, alpha-1-antitrypsin deficiency and Budd-Chiari syndrome.

Alcoholic liver disease is a risk factor that illustrates regeneration as the key factor in tumorigenesis. Active intake of alcohol causes hepatocyte injury with minimal regeneration, which clinically is a low risk factor for hepatocellular carcinoma. In contrast, alcohol abstinence results in significant regeneration and a higher risk of hepatocellular carcinoma (18).

Once the hepatitis B virus is integrated into the hepatocytes, transactivation of proto-oncogenes, activation of growth factors, and inactivation of tumor suppressor genes may result in abnormal proliferation (19). The hepatitis C virus damages double-stranded DNA, increasing the frequency of mutations in genes such as immunoglobulin genes, BCL-6, TP53, and β-catenin, causing abnormal proliferation (20). Other common mutations encountered in hepatocellular carcinoma include: telomerase promoter mutations (30 to 60% of HCC) (21); TP53 mutations (18 to 50%) (22) classically associated to exposure to aflatoxin; beta-catenin (18 to 40%); AXIN1 and AXIN2 genes; chromatin remodeling pathway (ARID1A and ARID2); Janus kinase (JAK)/signal transducers and activators of transcription (STAT) pathway (JAK1, IL6R, and IL6ST); genes involved in ubiquitination (KEAP1); genes involved in RAS/MAPK signaling (RPS6KA3); and genes involved the oxidative stress pathway (NFE2L2) (23).

NATURAL HISTORY AND PRE-NEOPLASTIC LESIONS

At the cellular level, hepatocellular carcinomas are thought to arise from a dysplastic focus (less than 1 mm) that develops into either a low-grade or a high-grade dysplastic nodule. The high-grade dysplastic nodule, more frequently than the low-grade dysplastic nodule, progresses to hepatocellular carcinoma (24). Since the primordial lesion (dysplastic focus) is less than 1 mm, it is not identifiable by radiologic studies and usually observed in hepatectomy specimens as focal cytological atypia within the hepatocytes. Morphologically, the atypia within the hepatocytes is described as small and large cell changes. The small cell change is characterized by increased nuclear to cytoplasmic ratio, hyperchromasia and cytoplasmic basophilia, giving the impression of crowding. Small change atypia is associated with high-grade dysplastic nodules and a higher risk of developing hepatocellular carcinoma. The small cell change is characterized by enlarged cells with larger nuclei and cytoplasm, but the nuclear to cytoplasmic ratio is preserved; frequent multinucleation, nuclear polymorphism and hyperchromasia are common. Large cell changes are the predominant alteration in low grade dysplastic nodules and indicate a benign feature (25). The biggest diagnostic challenge is to differentiate a high-grade dysplastic nodule from a well-differentiated hepatocellular carcinoma. Reticulin stain with delineation of the hepatic plates is a reliable ancillary technique in this context. Preservation of the hepatic plate is typical
of a high-grade dysplastic nodule, whereas expansion of the hepatic plates is typical of hepatocellular carcinoma. Other morphologic criteria for hepatocellular carcinoma include increased cell nuclei with consequent increased nuclear to cytoplasmic ratio, pseudogland formation and unpaired arteries with absence of portal triads (Figure 1), and sinusoidal capillarization appreciated with CD34 immunohistochemistry and stromal invasion without ductular reaction at the periphery of the nodules (26). The size of the lesion may also be helpful, especially radiologically, with lesions larger than 1 cm and less than 3 cm usually classified as dysplastic nodules, and lesions more than 3 cm as hepatocellular carcinoma (27).

Other preneoplastic lesions include clonal proliferation of hepatocytes forming hepatocellular adenomas. These lesions are characterized morphologically by well circumscribed proliferation of bland hepatocytes with intact hepatic plates and absence of portal triads. The real differential in this situation includes mostly a well differentiated hepatocellular carcinoma. Ancillary studies such as immunohistochemistry for CD34 demonstrating complete capillarization of the sinusoids, immunohistochemistry for Glypican-3 and reticulin stain (Figure 2) are helpful for differential. Hepatocellular adenomas are occasionally linked to oral contraceptives (28). Other possible etiologies include the use of clomiphene (used as hormonal treatment for infertility issues) (29), methyltestosterone (anabolic steroid) (30)

![Figure 1. Hematoxylin and Eosin stain 200X. On the left, normal hepatocytes with normal nuclear cytoplasmic ratio as compared to hepatocellular carcinoma (on the right) with increased nuclear cytoplasmic ratio, organized into thick trabeculae and few pseudoglandular structures with bile.](image-url)
and danazol (synthetic androgen used for treatment of endometriosis) (31). At the molecular level, hepatocellular adenomas are subdivided based on HNF1a mutations (steatotic adenomas), IL-6/STAT3 mutations (inflammatory adenomas), and β-catenin mutations (subgroup of inflammatory adenomas and unclassified adenomas with cytological atypia) (32). Prognostic implications justify such subtyping with inflammatory adenomas being frequently associated with metabolic syndrome, liver steatosis and alcohol exposure (33), and the β-catenin mutated adenomas immunophenotypically represented by nuclear staining associated with a higher risk of malignancy. Extensive literature and classification of hepatocellular adenomas is available, but the key is the identification of the lesions with great risk of progression to hepatocellular carcinoma.

**CLINICAL CHARACTERISTICS, DIAGNOSIS, AND DIAGNOSTIC TESTS**

Hepatocellular carcinoma usually affects the younger population. This is most likely related to significantly higher viral infection rate in individuals of reproductive age. Males are more affected than females, and males are also more frequently involved with alcohol abuse (34). Most patients are asymptomatic.
However, symptoms related to predisposing risk factors (cirrhosis and viral infections) are frequent, such as ascites and esophageal varices. Paraneoplastic syndromes may be occasionally associated with hepatocellular carcinoma, such as hypoglycemia (35), erythrocytosis (36), hypercalcemia (37) diarrhea (38) and cutaneous lesions (39). Other less common presentations include fever, infections, obstructive jaundice, and hemorrhage due to tumor rupture. Lymph node involvement is not common, and 10 to 15% of the cases present with advanced disease and metastases to lung, bone, and adrenal gland (40, 41).

Serological markers of the disease include alpha fetoprotein (AFP), heat shock protein, human cervical cancer oncogene, human telomerase reverse transcriptase mRNA, certain cytokines and microRNAs. AFP is by far the most used due to its availability, but screening limitations are significant due to its low sensitivity (41 to 65%). Pregnancy and primary liver and gastrointestinal disease also raise the levels of AFP, giving false positive results (42). The most accepted cut-off value is a serum concentration of 20 ng/mL (43). However, the most valuable indication of AFP is in the follow-up of previously treated hepatocellular carcinoma to detect the risk of recurrent disease (44). Three different glycoforms of AFP have been identified: AFP-L1, AFP-L2 and AFP-L3. The latter has been reported to be associated with more aggressive liver disease and consequently worse prognosis (45, 46). Glypican-3, a plasma membrane protein involved in the interaction with growth factors, has been suggested as an adjuvant serologic marker in addition to AFP to increase the sensitivity of detecting hepatocellular carcinoma (47, 48). Other liver enzymes have also been proposed in association with AFP to promote screening for hepatocellular carcinoma such as gamma-glutamyl transferase, Alpha-l-fucosidase and Des-gamma-carboxyprothrombin. Alternatively, instead of protein levels, mRNA levels of AFP and GGT may also be helpful prognostic factor after initial therapy for HCC (49, 50).

Additional serological markers in conjunction with AFP have been proposed to increase the sensitivity of detection such as alpha-L-fucosidase (AFU) and transforming growth factors alpha and beta (TGF-α and TGF-β) (51)

Radiologic diagnosis of hepatocellular carcinoma has significant implications for treatment options, with dysplastic nodules conservatively followed up based on the fact that one third of these lesions may progress to hepatocellular carcinoma; and early hepatocellular carcinoma should be treated with possible curative options such as ablation, resection and transplant (52). The image modality typically used is ultrasonography, likely due to the availability, but the sensitivity of 65% and specificity of 90% justify the use of more advanced image studies (53). In referral centers, the image protocol includes four-phase multidetector computed tomography (CT) or dynamic contrast enhanced magnetic resonance imaging (MRI), complementing each other, when suspicious features are noted (54). The distinction and detection of hepatocellular carcinoma is based on changes of vascularization including the presence of unpaired arteries and capillarization of the sinusoids that can be observed by image studies (55, 56). The definite radiologic criteria for the diagnosis of hepatocellular carcinoma are defined by contrast hyperenhancement at the arterial phase and hypoenhancement at the venous phase for lesions larger than 2 cm (57, 58). This criterion is highly specific and has replaced biopsy since 2000 (59). For lesions less than 2 cm, there is intense debate regarding the need of a biopsy, or not to define the lesion. Lesions less than 1 cm tend to be biopsied but there is variation in practice among centers (60).
When atypical nondiagnostic inconclusive features are noted by radiology, a biopsy is indicated. Most are image-guided targeted biopsies of the lesions, but a biopsy away from the lesion may be useful to assess other conditions, for example cirrhosis, that may increase the risk of hepatocellular carcinoma. Biopsy is preferred over fine needle aspiration due to the low sensitivity (67%) of the cytology specimen, although complications such as seeding is obviously higher in the more invasive biopsy procedure (61). The risk of seeding is small but considerable, being estimated at 2.7% (62). The hallmark of a neoplastic proliferation within a biopsy is the lack of portal triads. Features of hepatocellular differentiation such as bile production and canaliculi are important when distinguishing a metastatic lesion (63). Reticulin stain as previously commented highlights thickening of the hepatic plates (more than 3 cell thick) and the cytologic features will define the histologic grade of hepatocellular carcinoma. The best differential between hepatocellular carcinoma and hepatic adenoma is immunohistochemical stains for glypican-3 or AFP. However, AFP stain in some laboratories has a lot of background, making it very difficult to interpret. Glypican-3 may also be focal within the lesion and the representative core biopsy may be completely negative. By morphologic analysis, hepatocellular carcinoma may be classified as well-, moderately, and poorly differentiated, based on resemblance to normal hepatocytes. Well-differentiated tumors have slightly enlarged nuclei and increased nuclear to cytoplasmic ratio and may be very difficult to differentiate from normal hepatocytes. Moderately differentiated tumors have slightly thicker hepatic plates and larger cells. Poorly differentiated tumors are heterogeneous and have characteristics of an immature tumor with no typical differentiation or maturation, requiring immunohistochemical markers to obtain adequate and conclusive diagnosis (64).

Immunohistochemical markers are widely used in hepatobiliary pathology. The most useful marker is HEP PAR-1, a marker of hepatocellular differentiation but not useful for distinction between benign and malignant hepatocytes. Glypican-3 (Figure 3) and arginase-1 are specific markers for malignancy transformation (65). Supportive markers such as CD34 display typical capillarization of the sinusoids (Figure 4), and CD10 and pCEA demonstrate typical canalicular pattern. In addition, as an initial panel to exclude other possibilities, cytokeratins such as CK7 and CK20 are routinely used with the typical immunophenotype for HCC consisting of negativity for both markers.

Many histologic variants of hepatocellular carcinoma have been described and some have no clinical significance and may be confused with other entities. Such examples include clear cell hepatocellular carcinoma mimicking a metastatic clear cell renal cell carcinoma, pseudoglandular hepatocellular carcinoma mimicking a metastatic adenocarcinoma and scirrhous hepatocellular carcinoma mimicking a cholangiocarcinoma (66, 67). In other situations, recognition of a particular histologic variant is significant as in the case of diffuse cirrhosis-like hepatocellular carcinoma that radiologically mimics cirrhosis, and for that reason is usually missed (68). The same may be said of other variants such as giant cell variant and combined hepatocellular carcinoma and cholangiocarcinoma recognized as tumors with worse prognosis (66). Other variants are presumed to have better prognosis such as pedunculated hepatocellular carcinoma and fibrolamellar hepatocellular carcinoma (69). Finally, some variants are related to adjuvant therapy as ablated HCC causing significant necrosis (70).
Figure 3. Immunohistochemistry for Glypican-3 200X. On the left, normal hepatocytes with no staining (brown pigmentation) as compared to hepatocellular carcinoma (on the right) with cytoplasmic and few nuclei staining positive (brown discoloration).

Figure 4. Immunohistochemistry for CD34 200X. On the left, normal hepatocytes with no staining (brown disoloration) of the sinusoids as compared to hepatocellular carcinoma (on the right) with positivity within the sinusoids indicating capillarization.
STAGING

After diagnosis, staging of disease is required for adequate prognostication and subsequent therapy. The current staging systems include the pathologic tumor-node-metastasis (pTNM) (Table 1), the Okuda and the Cancer of the Liver Italian Program (CLIP) (Table 2), and the Barcelona Clinic Liver Cancer (BCLC) (71) (Table 3). The pTNM staging system is purely morphology-based and the other systems are mixed, using both morphology and clinical parameters to predict prognosis. The pTNM system fails to consider the residual function of the liver tissue and therefore does not predict behavior and prognosis especially in patients treated with partial resection (72). In contrast, the other staging systems take into

| Table 1 | The pTNM classification and staging (2017) |
|---------|------------------------------------------|
| **Tumor** |                                           |
| T0      | No tumor                                 |
| T1      | Solitary tumor smaller or larger than 2 cm without vascular invasion |
| T1a     | Solitary tumor smaller or equal to 2 cm   |
| T1b     | Solitary tumor larger than 2 cm without vascular invasion |
| T2      | Solitary tumor larger than 2 cm with vascular invasion; or multiple tumors, none larger than 5 cm |
| T3      | Multiple tumors, at least one of which is larger than 5 cm |
| T4      | Single tumor or multiple tumors of any size involving a major branch of the portal vein or hepatic vein; Tumor(s) with direct invasion of adjacent organs other than the gallbladder; Perforation of visceral peritoneum |
| **Node** |                                           |
| N0      | No nodal metastasis                      |
| N1      | Regional lymph node metastasis           |
| **Metastasis** |                                      |
| M0      | No distant metastasis                    |
| M1      | Distant metastasis                       |
| **Stage** |                                         |
| IA      | T1AN0M0                                  |
| Stage IB | T1BN0M0                                  |
| Stage II | T2N0M0                                   |
| Stage IIIA | T3N0M0                         |
| Stage IIIB | T4N0M0                                    |
| Stage IVA | Any T, N1, M0                        |
| Stage IVB | Any T, Any N, M1                      |
consideration of the background liver function, predicting prognosis considerably better, but are not suitable for orientating therapy. The Okuda staging system is efficient in identifying patients who have very poor prognosis (Stage III) but fails to distinguish different prognosis in patients grouped as Stage I and II (73, 74). The CLIP staging system predicts survival more accurately in those heterogeneous group staged as I and II in the Okuda staging system but also fails in orienting adequate therapy (75). Finally, the BCLC staging system identifies early HCC for aggressive therapy but fails again to orientate adequate therapy in the more advanced stages. As illustrated in Tables 2 and 3, both CLIP and the BCLC staging systems use Child-Pugh class when staging HCC. This scoring system is based on laboratory tests such as serum albumin, serum bilirubin and prothrombin time as well as the presence or absence of ascites and hepatic encephalopathy (76, 77).

**TABLE 2 The Okuda and the CLIP staging systems**

| Parameters                        | Okuda                         | Points<sup>a</sup> |
|-----------------------------------|-------------------------------|--------------------|
| Tumor size by imaging            | >50%                          | 1                  |
| Cross-sectional area             | <50%                          | 0                  |
| Ascites                           | Present                       | 1                  |
|                                  | Absent                        | 0                  |
| Serum Albumin (mg/dl)            | >3                            | 0                  |
|                                  | <3                            | 1                  |
| Serum total bilirubin (mg/dl)    | <3                            | 0                  |
|                                  | >3                            | 1                  |

**CLIP**

| Parameters                        | Points<sup>a</sup> |
|-----------------------------------|--------------------|
| Child-Pugh class                  | A 0                |
|                                  | B 1                |
|                                  | C 2                |
| Tumor morphology                  | Single nodule <50% area | 0 |
|                                  | Multiple nodules <50% area | 1 |
|                                  | Massive >50% area   | 2 |
| AFP (ng/mL)                       | <400               | 0                  |
|                                  | ≥400               | 1                  |
| Portal vein thrombosis            | No                 | 0                  |
|                                  | Yes                | 1                  |

<sup>a</sup>Total points = Stage.
TREATMENT

The therapeutic options for HCC consist of liver transplantation, hepatic resection, ablation, chemoembolization and chemotherapy. The best available treatment is transplantation removing the entire tumor and replacing the diseased liver (usually cirrhotic) by a new functional healthy organ. However, due to limitations regarding availability of donor organs other alternatives should be considered. Living donor transplantation is an alternative option that has been reported to have similar results in terms of survival rate (78). Contraindications for liver transplant include extrahepatic spread, multiple tumors with one of the lesions larger than 3 cm and a single tumor larger than 5 cm emphasizing the need for proper staging (79–82). Up to 75% of the patients that received a transplant, previously had received transarterial chemoembolization (TACE) and one third of this same population had previous radiofrequency ablation (RFA) preventing progression of disease and winning time until a donor is available (83). If no evidence of medical liver disease and/or cirrhosis is documented, the patient may be offered partial hepatectomy as curative therapy (84). The three-year disease-free survival in cirrhotic patients treated with partial hepatectomy is 18%, whereas it is 83% in patients treated by transplantation (85).

CONCLUSION

HCC is a frequent tumor endemic in areas of frequent infectious hepatitis. Patients with confirmed diagnosis of infectious hepatitis should be screened for early hepatocellular carcinoma and possible therapy. Early detection involves

| Stage       | Performance Status | Tumor Stage   | Liver Function                      |
|-------------|--------------------|---------------|-------------------------------------|
| A= Early    | A1 0               | Single, <5 cm | No portal HTN and normal bilirubin  |
|             | A2 0               | Single, <5 cm | Portal HTN and normal bilirubin     |
|             | A3 0               | Single, <5 cm | Portal HTN and elevated bilirubin   |
|             | A4 0               | 3 tumors <3 cm| Child-Pugh class A-B                |
| B = Intermediate | 0          | Large multinodular | Child-Pugh class A-B                 |
| C = Advanced| 1 and 2            | vascular invasion extrahepatic spread | Child-Pugh class A-B                 |
| D = End-stage| 3 and 4           | Any           | Child-Pugh class C                  |
serologic markers, mostly AFP and radiologic studies. Staging systems attempt to subdivide the cases of hepatocellular carcinoma in subgroups to predict behavior and prognosis as well as orientate therapy. Curative therapy may be achieved with either resection and/or transplant. Adjuvant therapy such and TACE and RFA are offered as palliative care in attempt to win some time until a donor liver is available.

Conflict of interest: The author declares no potential conflict of interest with respect to research, authorship and/or publication of this chapter.

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