CHIC Risk Stratification System for Predicting the Survival of Children With Hepatoblastoma: Data From Children With Hepatoblastoma in China

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Objective: The aim of this study was to compare the accuracy of the Children’s Oncology Group (COG) risk stratification system to the Children’s Hepatic tumor International Collaboration (CHIC) risk stratification system for predicting the prognosis of Chinese children with hepatoblastoma (HB).

Methods: Clinicopathological data of 86 patients diagnosed with HB between January 2014 and December 2017 were retrieved. The study endpoints were the 1- and 3-year overall survival (OS) and disease-free survival (DFS) were analyzed to evaluate the predictive value.

Results: The 1-, 3-year OS and DFS of the 86 patients were 86.0%, 76.3%, and 74.4%, 54.0%, respectively. Univariate analyses revealed that age at diagnosis had a significant role in prognosis for both OS and DFS, along with PRETEXT staging and metastasis at diagnosis. Multivariate analysis showed that metastasis at diagnosis (HR 3.628, 95% CI 1.404-9.378, P=0.008), PRETEXT staging system (HR 2.176, 95% CI 1.230-3.849, P=0.008) and age at diagnosis (HR 2.268, 95% CI 1.018-4.982, P=0.043) were independent factors for OS. For DFS, the independent factors were the PRETEXT staging system (HR 2.241, 95% CI 1.533-3.277, P<0.001) and age at diagnosis (HR 1.792, 95% CI 1.018-3.154, P=0.043). Both COG and CHIC risk stratification systems could effectively predict the prognosis of children with HB for OS. For DFS, the CHIC risk stratification system was more effective. In addition, the CHIC risk stratification system had a higher c-index (OS 0.743, DFS 0.730), compared to the COG risk stratification system (OS 0.726, DFS 0.594).
INTRODUCTION

HB is a rare childhood tumor with an annual incidence of 1.5 cases per million children, occurring predominantly in the first 2 years of life (1). Its overall survival in the past three decades has increased from ~30% to ~70% (2, 3); largely contributed due to improvement in imaging technology, surgical methods, timely surgeries, and chemotherapy prescription (4–6). However, milestone developments for the risk stratification of HB still lag behind more common pediatric cancers (7).

In the past decades, four separate cooperative multicenter trial groups have performed systematic treatment studies in HB, namely, the International Childhood Liver Tumor Strategy Group (SIOPEL); the Children’s Oncology Group (COG), and its legacy groups the Children’s Cancer Group (CCG) and the Pediatric Oncology Group (POG); the German Society for Pediatric Oncology and Haematology (GPOH); and the Japanese Study Group for Pediatric Liver Tumors (JPLT) (8). After accumulating and analyzing large amounts of data, these organizations proposed that the small cell undifferentiated (SCU) histology and low serum AFP at diagnosis could be associated with adverse outcomes in survival (9, 10) and have been constantly investigating novel prognostic factors (11–14). However, as the stratification criteria used in each group are different, it is not possible to compare the prognosis of each group clearly.

The small number of patients and the use of multiple different staging systems in the four main trial groups hindered the comparative evaluation of treatment outcomes in the pediatric HB trial. To solve this problem, CHIC has established a new international risk-stratified staging system by analyzing the database that includes comprehensive data from 1605 children treated in eight multicenter HB trials over 25 years (15). The new risk-stratified staging system—the Children’s Hepatic tumors International Collaboration—Hepatoblastoma Stratification (CHIC-HS) is shown as stratification trees based on four clinically familiar PRETEXT groups. CHIC-HS has not only confirmed the prognostic importance of factors such as metastatic disease, low AFP concentration (≤100 ng/ml), and PRETEXT group, that were originally proposed but also suggested additional new prognostic factors and more detailed grouping criteria (7).

This study aimed to compare the prognostic ability of the COG risk stratification system (Table S1) and the CHIC risk stratification system for prognosticating the survival of Chinese children with HB (Figure S1).

Conclusion: Age at diagnosis played a significant role in prognosis. Compared to the COG risk stratification system, the CHIC risk stratification system was superior in predicting the survival of Chinese children with HB.

Keywords: hepatoblastoma, pediatric, risk stratification, prognostic factors, Childhood Hepatic Tumor International Consortium

PATIENTS AND METHODS

Ethics Statement

The research was approved by the institutional review board (IRB) of Sun Yat-sen University Cancer Center (SYSUCC; Guangzhou, China), the IRB number is B2019-232-01, and written informed consent was obtained from each patients’ parents involved in the study. All the original data was deposited on http://www.researchdata.org.cn (RDD number RDDA2020001362).

Patients

Patients who were diagnosed as HB from January 2014 to December 2017 at the SYSUCC and the First Affiliated Hospital of Sun Yat-sen University (Guangzhou, China) were identified from our prospective database.

Patients who met all of the following criteria were included in the study analyses: 1) newly diagnosed patients; 2) age less than 18 years old; 3) histologic confirmation of HB. Since the current study aimed to analyze the prognostic factors for long-term survival, we excluded patients who died <30 days following surgery or died of chemotherapy side effects.

At the time of initial diagnosis, all the parameters mentioned by risk-stratified staging system were recorded and evaluated as possible predictors of survivals including age, alpha-fetoprotein (AFP), metastatic disease (M), macrovascular involvement of all hepatic veins (V), or portal bifurcation (P), contiguous extrahepatic tumor (E), multifocal tumor (F), and spontaneous rupture (R), tumor size and pathological pattern. Tumor size and VPEFRM were judged by contrast abdominal computed tomography (CT), abdominal ultrasound scan and intraoperative conditions. In general, P1 and P2 were defined as P+, V1, V2, and V3 were defined as V+ (16). None of the patients underwent liver transplantation.

Chemotherapy

According to the COG risk stratification system, the chemotherapy regimens of each group were different. Chemotherapy was not performed in the very low-risk group. The CSV(cisplatin, 5-FU, vincristine) chemotherapy regimen was used in the low-risk group for a total of four to six courses. Forty-five patients were defined as intermediate risk, of whom 30 patients were treated with the CSV(cisplatin, 5-FU, vincristine)/PLADO(cisplatin, doxorubicin) chemotherapy regimen, eight patients were treated with the PLADO(cisplatin, doxorubicin) chemotherapy regimen, and seven patients were treated with the C5V(cisplatin, 5-FU, vincristine, doxorubicin) chemotherapy regimen.
regimen for a total of six to eight courses. The C5VD (cisplatin, 5-FU, vincristine, doxorubicin)/IIV (ifosfamide, irinotecan, vincristine) chemotherapy regimen was used in the high-risk group for a total of eight courses (Tables S2, S3). Alpha-fetoprotein (AFP) was assessed before each course of chemotherapy and tumor efficacy (17) was evaluated every two courses (CT or MR assessment).

Follow-Up
After the end of treatment, patients were followed-up with a comprehensive review included alpha-fetoprotein (AFP), complete blood routine, biochemical routine, echocardiogram, chest and CT abdomen and so on. In the first year, patients were followed every 3 months based on assessment of their alpha-fetoprotein (AFP) level, complete blood and biochemical (liver and kidney function and electrolytes) routine tests, abdominal ultrasound, chest X-ray, and electrocardiogram. Abdominal CT and echocardiogram were reviewed every 6 months. Patients were re-examined every 6 months from the second year and every year from the third year. When a metastasis was suspected, a chest CT scan, bone scintigraphy, positron emission tomography (PET), and biopsy (if necessary) were performed to confirm the metastasis and/or recurrence. The last follow-up date for patients who were still alive was August 2019.

Causes of death and sites of recurrence were determined from death certificates, medical interviews, and radiological findings. The study endpoints were overall survival (OS), defined as the time period from the date of initial diagnosis until death or last follow-up, and disease-free survival (DFS), defined as the time period from the date of initial diagnosis until occurrence of an event (recurrence, progressive disease, death, or diagnosis of a second malignant neoplasm) or last contact, whichever occurred first.

Statistical Analyses
The statistical analyses except C-index analyses were performed using the SPSS software, version 19.0 (SPSS Company, Chicago, Illinois, USA). The OS and DFS were calculated by the Kaplan–Meier method and compared using the log-rank test. The prognostic varieties in predicting OS and DFS were assessed by multivariate Cox proportional hazards regression analyses. All covariates that had clinical significance were included in a multivariate Cox proportional hazards model. Results were given as mean ± S.D. All statistical tests were two-sided, and a significant difference was considered for P < 0.05. The concordance index method (c-index) was used to rank the different risk-stratified staging systems according to their capacity of discriminating patients according to the outcome. The c-index was calculated using R with the rms package (18) of Dr. Frank Harrell.

RESULTS
Baseline Characteristics
A total of 100 patients who were diagnosed as HB were identified during that period. Based on our inclusion criteria, 14 patients were excluded from this study, of whom two died during the perioperative period (2%, 2/100), three died while on chemotherapy (3%, 3/100), and nine who were lost in follow-up (9%, 9/100). The patients’ baseline characteristics are summarized in Table 1. A total of 86 patients were included in the study, comprising of 54 (62.8%) male and 32 (37.2%) female, with a median age at diagnosis of 26.94 months (range: 0.9–139.6 months). AFP was elevated (>100 ng/ml or age-specific upper limit) in 84 (97.7%) patients. Based on the results of the imaging studies, patients were classified using the PRETEXT staging system and judged whether they had distant metastases and VPEFR. 22 (25.6%) patients had distant metastasis and 44 (51.2%) were positive for VPEFR at diagnosis. Fetal histology (n=29) was the most common pathological subtype in our patients on postoperative pathological results. All patients were classified using the COG risk group and CHIC risk group. Using the COG risk stratification system, 3 (3.5%) patients were considered as having very low-risk, 5 (5.8%) as low-risk, 45 (52.3%) as intermediate-risk, and 33 (38.4%) as high-risk. Using the CHIC risk stratification system, five (5.8%) patients were considered as having very low risk, 13 (15.1%) as low risk, 38 (44.2%) as intermediate risk, and 33 (38.4) as high-risk.

| Table 1 | Characteristics of patients included in the study. |
| --- | --- |
| N | % |
| ALL | 86 | 100 |
| Gender | | |
| Male | 54 | 62.8 |
| Female | 32 | 37.2 |
| Age(months) | | |
| <36 | 51 | 59.3 |
| 36-96 | 31 | 36.0 |
| ≥96 | 4 | 4.7 |
|AFP at diagnosis (ng/ml) | | |
| ≤100 | 2 | 2.3 |
| 101–1000 | 7 | 8.1 |
| >1000 | 77 | 89.5 |
| PRETEXT staging | | |
| I | 9 | 10.5 |
| II | 38 | 44.2 |
| III | 14 | 16.3 |
| IV | 25 | 29.1 |
| Metastasis at diagnosis | | |
| No | 64 | 74.4 |
| Yes | 22 | 25.6 |
| VPEFR | | |
| VPEFR (+) | 44 | 51.2 |
| VPEFR (-) | 42 | 48.8 |
| Histological subtype | | |
| Epithelial | 4 | 4.7 |
| Fetal(Pure Fetal excluded) | 29 | 33.7 |
| Pure Fetal | 2 | 2.3 |
| Embyronal | 5 | 5.8 |
| Mixed embyronal-fetal | 7 | 8.1 |
| Mixed epithelial-mesenchymal | 22 | 25.6 |
| HB (unknown) | 17 | 19.8 |
| COG risk group | | |
| Very low risk | 3 | 3.5 |
| Low risk | 5 | 5.8 |
| Intermediate risk | 45 | 52.3 |
| High risk | 33 | 38.4 |
| CHIC risk group | | |
| Very low risk | 5 | 5.8 |
| Low risk | 13 | 15.1 |
| Intermediate risk | 36 | 44.2 |
| High risk | 30 | 34.9 |
**Univariate Analysis of Prognostic Factors**

All factors listed in Table 2 were included in univariate analysis (Table 2), of which, age at diagnosis (OS $P=0.029$, DFS $P=0.027$), PRETEXT staging (OS $P<0.001$, DFS $P<0.001$), and metastasis at diagnosis (OS $P<0.001$, DFS $P=0.007$) were identified as significant prognostic factors for OS and DFS. For OS, the 3-year OS of patients younger than 36 months, aged from 36 months to 96 months and older than 96 months were 83.4%, 67.6%, and 50.0%, respectively. The 3-year OS of the PRETEXT stage I, II, III, and IV were 87.5%, 91.4%, 71.4%, and 52.0%. The 3-year OS of M (+) and M (-) were 44.4% and 87.4%. For DFS, the 3-year DFS of patients from different age groups were 64.7%, 35.9%, and 50.0%, respectively. The 3-year DFS of the PRETEXT stage I, II, III, and IV were 87.5%, 71.6%, 42.9%, and 20.0%. The 3-year DFS of M (+) and M (-) were 29.2% and 63.0%.

**Multivariate Analysis of Prognostic Factors**

Multivariate analysis showed that metastasis at diagnosis (HR 3.628, 95% CI 1.404-9.378, $P=0.008$), PRETEXT staging (HR 2.176, 95% CI 1.230-3.849, $P=0.008$) and age at diagnosis (HR 2.268, 95% CI 1.033-4.982, $P=0.041$) were independent prognostic factors for OS (Table 3). However, AFP at diagnosis, VPEFR and histological subtype were not (P>0.05) for OS. For DFS, multivariate analysis showed that PRETEXT staging (HR 2.241, 95% CI 1.533-3.277, $P<0.001$) and age at diagnosis (HR 1.792, 95% CI 1.018-3.154, $P=0.043$) were significant prognostic factors. However, metastasis at diagnosis, AFP at diagnosis, VPEFR and histological subtype were not (P>0.05) for DFS (Table 3).

**Survival**

The median follow-up period was 33.9 months (range 2.4-57.0 months). The 1-, 3-year OS of the 86 investigated patients were 86.0%, 76.3%, respectively (Figure 1A). The median OS was 33.9 months. The 1-, 3-year DFS were 74.4%, 54.0%, respectively (Figure 1B). The median DFS was 26.6 months.

There was a statistically significant difference between patients with very low-risk, low-risk, intermediate-risk and high-risk based on the COG risk stratification system for OS ($P<0.001$, Figure 1C). But statistically significant differences were not observed between patients using the COG risk stratification system for DFS ($P = 0.123$, Figure 1D). There was a statistically significant difference between patients with very low-risk, low-risk, intermediate-risk and high-risk using the CHIC risk stratification system for both OS ($P<0.001$, Figure 1E) and DFS ($P = 0.001$, Figure 1F).

The prognostic value of the COG and CHIC risk stratification system was compared by means of Harrell’s concordance index. The CHIC risk stratification system had a higher c-index (OS 0.740, DFS 0.660) compared to the COG risk stratification system (OS 0.730, DFS 0.600).

**DISCUSSION**

Based on the proposal of CHIC risk stratification system (7), we compared the accuracy of the COG risk stratification system to the CHIC risk stratification system for predicting the prognosis of Chinese children with HB. First of all, our results showed that the 1-, 3-year OS of children with HB were 86.0%, 76.3%, and 74.4%, 54.0% in DFS, respectively. What’s more, our study indicated that age at diagnosis played a significant role in prognosis for both OS and DFS, along with the PRETEXT staging and metastasis at diagnosis. Above all, our results revealed that compared to the COG risk stratification system we used, the CHIC risk stratification system was better to predict the survival in children with HB, especially for DFS.

With the use of adjuvant and neo-adjuvant chemotherapy, the 1-, 3-year OS and DFS were 86.0%, 76.3%, and 74.4%, 54.0% in our cohort, respectively. The literature on the treatment outcome of children with HB is inadequate. Chinese Children’s
Cancer Group (19) (CCCG) reported 6-year OS of standard risk and high risk groups were 90.3% ± 2.9% and 67.6 ± 7.0%. Liu et al. (20) revealed 5-year OS and DFS of children with HB in Hong Kong were 77.6% ± 5.5% and 69.2% ± 6.1%, respectively. Qiao et al. (21) showed the 5-year OS and EFS rates were 63.4% and 57.1%. Recent clinical trials initiated by four major organizations generally reported that the 3-year OS was above 80% (22). The SIOPEL-4 (23) results suggested that 3-year OS of children with high-risk HB was 83% and EFS was 76%. The JPLT-2 (24, 25) study of 279 patients indicated the 3-, 5-year OS were 82.4%, 80.9%, and 69.9%, 62.4% in DFS, respectively. The GPOH (26) Study HB99 showed the 5-year OS was 58% for high risk patients and 94% for standard risk patients.

Results of the GPOH (26) Study HB99 showed the 5-year OS was 58% for high risk patients and 94% for standard risk patients. Similarly, our study also showed unsatisfactory results that the 3-year OS and DFS were 54.0%, 41.0% for the COG high-risk group, and 52.1%, 33.5% for the CHIC high-risk group. The survival of patients in high risk group was gloomy, especially in DFS. First of all, due to economic and medical limitations, all high-risk patients had not undergone liver transplantation. Some data suggested that the overall long-term outcome of primary liver transplantation for HB was excellent (27). Secondly, the chemotherapy regimen-C5VD/IIV for high-risk patients in this
study was derived from a clinical trial, which was terminated in advance due to poor efficacy. In other words, our treatment of high-risk patients had failed. In addition, the lack of molecular variables at first diagnosis affected our judgment of prognosis. Therefore, it’s necessary to seek more effective treatment to improve the survival of patients in high-risk groups.

Compared to the COG risk stratification system, the CHIC risk stratification system excluded pathologic subtypes and included age at diagnosis to form stratification trees based on four clinically familiar PRETEXT groups. Over the past 10 years, four major organizations have been devoted to exploring effective prognostic factors in children with HB (23, 24, 26–29). In 2013, based on a database involving 1605 HB cases treated between 1988 and 2008 from eight completed HB trials, CHIC identified novel prognostic factors for HB (15). Advanced PRETEXT group, macrovascular venous or portal involvement, contiguous extrahepatic disease, primary tumor multifocality, and tumor rupture significantly increased risk for EFS-event. Higher age (≥8 years), low AFP (<100 ng/ml) and metastatic disease significantly increased risk for OS-event. Recently, CHIC stressed that risk of an event for children with HB increased with increasing age at diagnosis (30). In China, several results about the prognostic factors of children with HB have appeared recently. Qiao et al. (21, 31) presented AFP <100 or >1000 (ng/ml), multifocality, vascular invasion, metastases, and PRETEXT stage were associated with poor prognosis. Another multicentric study demonstrated that female, age over 5 years, PRETEXT III, PRETEXT IV, incomplete surgical resection, and with metastasis had poorer prognosis (19). Our findings confirmed that age at diagnosis played a significant role in prognosis for both OS and DFS, along with the PRETEXT staging and metastasis at diagnosis. Although AFP at diagnosis, VPEFR, and histological subtype did not show statistical differences in univariate analysis and multivariate analysis, but we cannot deny that these were acknowledged factors associated with prognosis. In our cohort, only 2 (2.3%) patients had low AFP (≤100 ng/ml) at diagnosis and no one was confirmed to be SCU histologic subtype. The absence of the above cases may be the reason for the non-statistical difference in AFP at diagnosis and pathological type.

Our results showed that both COG and CHIC risk stratification systems could effectively predict the prognosis of children with HB for OS. For DFS, the CHIC risk stratification system was effective, but the COG risk stratification system was not. In addition, the CHIC risk stratification system had a higher c-index (OS 0.743, DFS 0.730), compared to the COG risk stratification system (OS 0.726, DFS 0.594). Yoon et al. (32) provided supporting evidence of utility of the CHIC risk stratification system by providing good prognostic value of its variables. The study from Hong Kong also demonstrated the CHIC risk stratification system was effective to classify different risk groups for predicting the prognosis (20). However, the results clearly indicated that both the COG and CHIC risk stratification system did not had good efficacy to distinguish between the prognosis of very low-risk group and low-risk group. Therefore, the risk stratification system more suitable for children with HB in China needs further development.

Our results confirmed good prognostic value of age at diagnosis, the PRETEXT staging and metastasis at diagnosis. More importantly, our study validated the feasibility of the risk stratification system proposed by CHIC and its application to Chinese children with HB. Although retrospective, the data we have presented were based on a relatively long duration of follow-ups (median, 33.9 months) from the children who were diagnosed as HB from January 2014 to December 2017. We also used sensitive and specific imaging technologies and histological confirmation in order for us to provide a valid base to evaluate the prognostic value of the risk stratifications. However, there were some limitations in the present study that should be commented. Firstly, it is a single-institution study of a fairly homogenous population. Due to the rarity of HB, the inadequate number of cases may be difficult to define several prognostic factors, such as AFP at diagnosis. Secondly, patients with intermediate-risk, were offered different chemotherapy regimens which might have contributed to the differences in patients’ prognosis. Thirdly, our study did not analyze the effect of molecular variables which would be potential prognostic factors alternative to clinical factors in the future. Several prognostic markers have been reported in association with HB (22, 33). Sumazin et al. demonstrated high NFE2L2 activity and high expression of LIN28B, HMG A2, SALL4 were associated with poor prognosis and marked high-risk group (34). Another research indicated that the four novel tumor suppressor candidates including GPR180, MST1R, OCIAD2, and PARP6 were potentially useful molecular markers for predicting poor prognosis in HB patients (35). In the future studies, we should try to include genetic mutation in the prognostic correlation analysis.

CONCLUSION

In this cohort of Chinese children with HB, age at diagnosis showed a significant association with patient outcome, along with the PRETEXT staging and metastasis at diagnosis. Compared to the COG risk stratification system, the CHIC risk stratification system was superior in predicting the survival of Chinese children with HB.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: http://www.researchdata.org.cn (RDD number RDDA2020001362).

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by the institutional review board (IRB) of Sun Yat-sen University Cancer Center (SYSUCC; Guangzhou, China). Written informed consent to participate in this study was
provided by the participants’ legal guardian/next of kin. Written informed consent was obtained from the minor(s)’ legal guardian/next of kin for the publication of any potentially identifiable images or data included in this article.

**AUTHOR CONTRIBUTIONS**

JH, YX, ZJ, and ZZ contributed conception and design of the study. HJ, YX, and SL organized the database. FS, JZ, and JW performed the statistical analysis. JH and YW wrote the first draft of the manuscript. HJ, YZ, JL, and ZZ wrote sections of the manuscript. All authors contributed to the article and approved the submitted version.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fonc.2020.552079/full#supplementary-material
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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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