Combination Therapy of Metformin and Statin May Decrease Hepatocellular Carcinoma Among Diabetic Patients in Asia

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Abstract: Previous studies have shown that metformin or statins may decrease hepatocellular carcinoma (HCC) in diabetic patients. Accordingly, this article evaluates whether combination therapy may further reduce HCC.

Newly diagnosed type 2 diabetes mellitus (DM) patients, excluding those with history of malignancy prior to the date of DM diagnosis, were recruited to a DM cohort. DM patients developed HCC as the cancer cohort and the date for HCC diagnosis as index date. Non-cancer cohort was frequency matched with 4:1 according to age, sex, DM-year, and index date as case group from DM cohort.

Patients who were treated with statins showed a 63% decreased risk of HCC (odds ratio [OR] = 0.37; 95% confidence interval [CI] = 0.27–0.49). Patients who consumed simvastatin, atorvastatin, or rosuvastatin significantly decreased risk for HCC (OR = 0.32, 0.31, and 0.22; 95% CI = 0.18–0.58, 0.19–0.52, and 0.08–0.61, respectively). Metformin combinations with simvastatin, atorvastatin, or rosuvastatin may decrease HCC (OR = 0.30, 0.30, and 0.24; 95% CI = 0.15–0.59, 0.16–0.54, and 0.08–0.70, respectively). The comorbidities for HCC were decreased by consuming simvastatin and atorvastatin (OR = 0.31 and 0.29; 95% CI = 0.14–0.67 and 0.15–0.57, respectively). Only combination therapy of metformin and simvastatin may significantly decreased HCC comorbidities (OR = 0.26; 95% CI = 0.11–0.60) in our study.

In Asia, not all metformin combinations with statins may reduce the incidence of HCC and not all of this kind of combination therapy may decrease the HCC comorbidities.

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Abbreviations: CI = confidence interval, DM = diabetes mellitus, HBV = hepatitis B virus, HCC = hepatocellular carcinoma, HCV = hepatitis C virus, ICD-9-CM = International Classification of Diseases, Ninth Revision, Clinical Modification, NAFLD = non-alcoholic fatty liver disease, NHIRD = National Health Insurance Research Database, OR = odds ratio.

INTRODUCTION

The 4 most critical diseases, as emphasized by the World Health Organization in 2011, are diabetes mellitus (DM), cancer, heart disease, and lung disease. However, DM can increase the comorbidities and mortality rates of all these diseases.

Since 1959, studies have reported an association between DM and certain cancers. The mortality rate among diabetic patients with cancer is more than 1.28 times that of diabetic patients who lack cancer. Previous studies have also shown that DM can increase the risk of hepatocellular carcinoma (HCC) independently of the hepatitis B virus (HBV), hepatitis C virus (HCV), cirrhosis, heavy alcohol consumption, and non-alcoholic fatty liver disease (NAFLD). More than 90% of patients with HCC in Taiwan test positive for the HBV surface antigen or HCV antibodies, and these carriers generate the severe public health problems as DM. Most guidelines recommend administration of metformin and statins as first-line medications for the treat of DM and dyslipidemia, ensuring primary or secondary prevention. Abundant literature has discussed the additional benefits provided by metformin or statins such as anti-cancer and anti-inflammatory activity. This article examines whether metformin combined with statins may further reduce HCC.

METHODS

For this nested-case–control study, we referenced the Longitudinal Health Insurance Database 2000 (LHID2000), which was established by the Taiwan Bureau of National Health...
Insurance. The Taiwan National Health Insurances is a single-payer program that covers over 99% of the population in Taiwan. The LHID2000 contains data of 1 million insurants. Beneficiaries of this program were randomly selected from the 2000 registry. This database contains all medical records of each insured people from 1996 to 2010. The insurant identities re-coded before the data samples were released to researchers. This research was approved by the institutional review board of the China Medical Hospital, Taiwan. Details of the LHID2000 database are presented on the National Health Research Institute’s website (http://w3.nhri.org.tw/nhird/date_01.html).

Patients newly diagnosed with type 2 DM (International Classification of Diseases, 9th Revision, Clinical Modification (ICD-9-CM): 250.X0 and 250.X2) between 2000 and 2010, excluding those with a history of malignancy (ICD-9-CM: 140–208) before the DM diagnosis date, were recruited to forma DM cohort (Figure 1). DM patients who were diagnosed with HCC (ICD-9-CM: 155) were classified as the case group and the date of HCC diagnosis was defined as the index date. Controls were randomly assigned after being selected among DM patients who had not developed HCC, and matched at a 4:1 frequency according to age (stratum 5 years), sex, the year of DM diagnosis, and index date.

The risk factors in this investigation comprised comorbidities and medication use. The comorbidities included HBV (ICD-9CM: 070.2, 0710.3, and V02.61), HCV (ICD-9-CM: 070.41, 070.44, 070.51, and V02.62), NAFLD (ICD-9-CM: 571.8), and alcoholic liver damage (ALD; ICD-9-CM: 571.0, 571.1, and 571.3). Because there was a higher relationship between HCC and cirrhosis duration, the duration between the date for cirrhosis diagnosed (ICD-9-CM: 571.2, 571.5, and 571.6) and the index date was as a cofounder. All comorbidities were defined before the index date. Regarding medication use, we evaluated statins and metformin. Metformin use was defined before the index date, and statin use was defined between January 1, 2000 and the index date. We considered the 6 common statins simvastatin, atorvastatin, pravastatin, fluvastatin, lovastatin, and rosuvastatin. Medication users were further divided into 2 groups: “usually” and “occasionally” based on their duration rate of medication use during the study period. A duration rate of <50% was defined as occasionally, and a rate of ≥50% was defined as usually.

Logistic regression was used to assess the odds ratio (OR) and 95% confidence interval (CI) for patients with HCC in statin users compared with non-users. The association between HCC and the duration of each statin exposure was assessed. The relevant variables in Table 1 were included in the adjusted logistic regression analysis. The trend test results were also assessed using adjusted logistic regression. The statistical significance level was set as a 2-tailed \( P \leq 0.05 \). Furthermore, all analyses were performed using SAS software, Version 9.3 (SAS Institute, Carry, NC).

**RESULTS**

The DM cohort in this study comprised 340 patients with HCC and 1360 controls. The mean patient age was 62.6 years (standard deviation = 11.3). The proportion of men exceeded

| TABLE 1. Distribution of Risk Factor and Odds Ratio for HCC |
|-----------------|-----------------|-----------------|-----------------|-----------------|
|                  | **HCC (N = 340)** |                  | **Non-HCC (N = 1360)** |                  |
|                  | **n** | **%** |                  | **n** | **%** | **OR** | **95% CI** |
| **Age, y**       | 62.6 ± 11.3 | 62.4 ± 11.3 | 1.00 | 0.99–1.01 |
| **Men**          | 265   | 77.9  | 1060   | 77.9  | 1.00  | 0.75–1.33 |
| **Risk factor**  |                  |                  |                  |                  |
| **NAFLD**        | 9     | 2.65  | 30     | 2.21  | 1.21  | 0.57–2.56 |
| **ALD**          | 14    | 4.12  | 19     | 1.40  | 3.03  | 1.50–6.11 |
| **HCV**          | 107   | 31.5  | 45     | 3.31  | 13.4  | 9.22–19.5** |
| **HBV**          | 142   | 41.8  | 80     | 5.88  | 11.5  | 8.40–15.7** |
| **Statin**       | 63    | 18.5  | 522    | 38.4  | 0.37  | 0.27–0.49** |
| **Metformin**    | 230   | 67.7  | 972    | 71.5  | 0.84  | 0.65–1.08 |
| **Cirrhosis duration, y** | 1.77 ± 2.91 | 0.11 ± 0.85 | 1.81 | 1.62–2.02** |

**ALD** = alcoholic liver damage, CI = confidence interval, **HBV** = hepatitis B virus, **HCC** = hepatocellular carcinoma, **HCV** = hepatitis C virus, **NAFLD** = non-alcoholic fatty liver disease, OR = odds ratio.

\(^{1}\) Between the date for cirrhosis diagnosed and the index date.

\(^{2}\) \( P < 0.01\).

\(^{**}\) \( P < 0.0001\).
that of women (77.9% vs 22.1%) in the HCC group (Table 1).

The logistic regression analysis results indicated that patients with cirrhosis duration (per year), ALD, HCV, or HBV demonstrated an increased risk of HCC (OR = 1.81, 3.03, 13.4, and 11.5, respectively). However, patients who received statins showed a 63% decreased risk of HCC (OR = 0.37; 95% CI = 0.27–0.49).

Compared with patients who did not receive metformin and statins, no significant difference was observed for various duration rates of statin or metformin use after controlling for cirrhosis duration, ALD, HCV, and HBV (Table 2). However, a significant decrease was observed as the duration rates of statin use increased among patients who received no or occasional metformin treatment (trend test P = 0.01 and 0.0008).

Table 3 shows the association between HCC and various statins. Compared with patients who did not receive statins, those who consumed simvastatin, atorvastatin, or rosuvastatin exhibited a significantly reduced risk of HCC (OR = 0.32, 0.31, and 0.22; 95% CI = 0.18–0.58, 0.19–0.52, and 0.08–0.61, respectively). The same trend was observed in patients who received metformin (OR = 0.30, 0.30, and 0.24; 95% CI = 0.15–0.59, 0.16–0.54, and 0.08–0.70). The risk of HCC decreased with the exposure duration (per month) of simvastatin, atorvastatin, and rosuvastatin which was increased compared without statin treatment (OR = 0.82, 0.95, and 0.79; 95% CI = 0.73–0.94, 0.92–0.99, and 0.63–0.99). In patients with metformin treatment, the risk reduced with the duration of simvastatin and atorvastatin (OR = 0.81 and 0.95; 95% CI = 0.70–0.93 and 0.92–0.99, respectively).

Table 4 shows the association between HCC and various statins in patients with any comorbidities included NAFLD, ALD, HCV, HBV, and cirrhosis. Overall, the risk of HCC was decreased among patients treated with simvastatin and atorvastatin compared with patients who did not receive statin (OR = 0.31 and 0.29; 95% CI = 0.14–0.67 and 0.15–0.57, respectively). Compared to non-statin use, longer duration of simvastatin and atorvastatin use could reduce HCC risk (OR = 0.81 and 0.96; 95% CI = 0.68–0.97 and 0.92–0.99, respectively). The patients treated with metformin showed similar results.

**DISCUSSION**

**DM and HCC**

The relationship between DM and HCC was identified in 1986. One systematic review showed that DM patients are 3.64 times more likely to develop HCC compared with non-DM patients. Other studies have also suggested that DM is a potential risk factor for HCC. The possible reasons for this inference are explained as follows: First, liver cirrhosis may cause glucose intolerance and up to 30% of cases may become diabetic. Terminal or end-stage hepatic disease can further increase glucose intolerance, which eventually progresses into diabetic. Terminal or end-stage hepatic disease can further cause glucose intolerance and up to 30% of cases may become diabetic.

Second, non-alcoholic steatohepatitis (NASH) is among the most severe manifestations of NAFLD, which is a risk factor for HCC. NASH can cause a wide range of liver disorders, including steatosis and steatohepatitis, demonstrating pathological features similar to those of alcohol-related liver injury, fibrosis, and occasionally cirrhosis. Unfortunately, DM is a risk factor for NAFLD. Third, both HCV and hemochromatosis can cause liver disease, and have been associated with an increased risk of DM. Fourth, insulin is produced and secreted by β-cells located in the pancreas.

**TABLE 2. Adjusted Odds Ratio for HCC Among Metformin and Statin Use Duration Rate**

| Metformin | Statin | None | Sometimes | Usually | HCC Case/N | Model 1 | Model 2 | P for Trend in Model 2 |
|-----------|--------|------|-----------|---------|------------|---------|---------|------------------------|
| None      | 99/383 | 1.00 | 0.52      | 0.24    | 3.51       | 0.007   | 0.06    | 0.07                   |
| Sometimes | 155/639| 1.00 | 0.52      | 0.24    | 3.51       | 0.007   | 0.06    | 0.07                   |
| Usually   | 31/251 | 0.49 | 0.29      | 0.14    | 9/133      | <0.0001 | 0.0008 | 0.09                   |

Statins: None, Sometimes, Usually. Model 1: adjusted for cirrhosis duration and ALD. Model 2: adjusted for cirrhosis duration, ALD, HCV, and HBV.
High concentrations of endogenous insulin pass directly into portal circulation in the liver, potentially explaining the association between DM and HCC. Fifth, other factors including obesity and inflammation may exert influence; these factors have been reported for DM and cancer. Furthermore, chronic inflammation can stimulate, promote, and advance cancer cell development. In this research, the risk factors for HCC were cirrhosis, HBV, HCV, and ALD, rather than NAFLD and metformin use (Table 1). However, additional studies may be required to clarify the effects on Asian DM patients with HCC.

### TABLE 3. Association Between HCC and Different Statin Use and Exposure Duration

| OR (95% CI) | HCC Case/N | | |
| --- | --- | --- | --- |
| All | | | |
| Non-statin | 277/1115 | 1.00 | 1.00 |
| Statin | | | |
| Simvastatin | 19/211 | 0.32 (0.18–0.58)** | 0.82 (0.73–0.94)** |
| Atorvastatin | 24/309 | 0.31 (0.19–0.52)** | 0.95 (0.92–0.99)** |
| Pravastatin | 16/74 | 1.35 (0.70–2.59) | 1.04 (0.99–1.09) |
| Fluvastatin | 11/110 | 0.54 (0.27–1.08) | 0.94 (0.88–1.01) |
| Lovastatin | 16/138 | 0.54 (0.29–1.02) | 0.93 (0.86–1.00) |
| Rosuvastatin | 5/96 | 0.22 (0.08–0.61)** | 0.79 (0.63–0.99)** |
| Without Metformin use | | | |
| Non-statin | 99/383 | 1.00 | 1.00 |
| Statin | | | |
| Simvastatin | 4/35 | 0.43 (0.11–1.69) | 0.91 (0.73–1.12) |
| Atorvastatin | 5/65 | 0.39 (0.14–1.08) | 0.96 (0.89–1.03) |
| Pravastatin | 3/15 | 1.79 (0.46–6.96) | 1.05 (0.98–1.13) |
| Fluvastatin | 0/22 | — | — |
| Lovastatin | 1/26 | 0.14 (0.02–1.29) | 0.92 (0.82–1.04) |
| Rosuvastatin | 0/11 | — | — |
| With Metformin use | | | |
| Non-statin | 177/732 | 1.00 | 1.00 |
| Statin | | | |
| Simvastatin | 15/176 | 0.30 (0.15–0.59)** | 0.81 (0.70–0.93)** |
| Atorvastatin | 19/244 | 0.30 (0.16–0.54)** | 0.95 (0.92–0.99)** |
| Pravastatin | 13/59 | 1.29 (0.61–2.76) | 1.04 (0.98–1.11) |
| Fluvastatin | 11/88 | 0.81 (0.40–1.66) | 0.98 (0.91–1.04) |
| Lovastatin | 15/112 | 0.65 (0.33–1.29) | 0.93 (0.84–1.03) |
| Rosuvastatin | 5/85 | 0.24 (0.08–0.70)** | 0.80 (0.64–1.02) |

Model 1: the risk for HCC TBL in each statin users compared with non-statin users adjusted for cirrhosis duration, ALD, HCV, and HBV. Model 2: the risk for HCC in patients with each statin exposure duration (month) compared with people without statin exposure adjusted for cirrhosis duration, ALD, HCV, and HBV. ALD = alcoholic liver damage, CI = confidence interval, HBV = hepatitis B virus, HCC = hepatocellular carcinoma, HCV = hepatitis C virus, OR = odds ratio.

**P < 0.05.
***P < 0.01.
****P < 0.0001.

### Statins and HCC

Previous studies have reported that statins can not only reduce low-density lipoprotein (LDL) levels but also decrease HCC, infection, or inflammation. Statins reduce HCC cells by inducing cell apoptosis. Pravastatin can be prescribed in chemotherapy treatments for HCC to improve the survival rates of patients with advanced HCC. The enzyme 3-hydroxy-3-methylglutaryl coenzyme-A reductase (HMG-CoAR), which is a rate-limiting step in the mevalonate pathway for LDL reduction, can induce the production of isoprenoids, which contribute to the activation of Ras. HMG-CoAR is a rational molecular target for innovative anti-neoplastic treatments of HCC. Chemotherapy treatment that employs statin therapy can decrease resistance to cytotoxic drugs by activating the Ras/Raf/MEK/ERK signal transduction cascade and increasing cholesterol levels in cancer cells. Other statins such as fluvastatin and cerivastatin have been reported to inhibit hepatic tumor cell growth in rats and to induce tumor-specific apoptosis. One study showed that pitavastatin and atorvastatin can effectively and safely reduce elevated hepatic enzyme levels in patients with NAFLD. The results of our analyses supported the findings of previous studies. Specific statins, such as simvastatin, atorvastatin, and rosuvastatin, may reduce the risk of HCC (Tables 1 and 3). These findings indicate that various statins demonstrate distinct potential anti-cancer effects, and that the 2 statins investigated herein can yield benefits for Asian patients with HCC. The limitation of our patient numbers in different statin groups can also offer the same results as the previous studies for different statins.
Metformin and HCC

A systematic review and meta-analysis showed that metformin can reduce the mortality rates and incidence of all cancers compared with other treatments for DM. Metformin significantly reduced the risk of colorectal cancer, liver tumors, and lung cancer. One article stated that DM patients treated with metformin were associated with an estimated 70% reduction in HCC risk. Another systematic review showed that metformin yields more benefits for treating pancreatic and hepatic cancers compared with colon, breast, and prostate cancers. Several possible mechanisms for the anti-cancer effects of metformin have been discussed. Indirect pathways include the avoidance of weight gain, as compared with other anti-diabetic medications, and the amelioration of insulin resistance by reducing hyperinsulinemia. However, both pathways may promote carcinogenesis. The direct pathways of metformin can stimulate AMP-activated protein kinase (AMPK) through LKB-1 (serine–threonine liver kinase B1), which is a tumor-suppressing protein kinase. AMPK inhibits protein synthesis and hepatic glucose output during cellular stress and inhibits the mammalian target of rapamycin (mTOR), which is a downstream effector of growth factor signaling commonly present in malignant cells. Regarding breast cancer, metformin can inhibit mTOR activity, thereby reducing HER-2 protein expression. Metformin can also induce cell-cycle arrest and apoptosis, reducing growth factor signaling with hyperinsulinemia in diabetic or non-diabetic patients, thereby decreasing the risk of colorectal cancer. These articles have suggested other possible mechanisms by which metformin can decrease HCC risk, including reducing inflammation and endogenous reactive oxygen species. Studies have reported varying results regarding the anti-cancer effects of metformin, including no apparent benefit for reducing the risk of colorectal cancer. A recent nested case–control study conducted in Asia determined no significant relationship between metformin use and the incidence of cancer (such as liver cancer and female breast cancer). According to these articles, it remains ambiguous whether metformin exerts anti-neoplasm effects in DM patients. Our analysis showed that metformin cannot reduce HCC (Table 1). The possible reasons for this finding are explained as follows: First, diabetes is a progressive disease and inhibits the mammalian target of rapamycin (mTOR), which is a downstream effector of growth factor signaling commonly present in malignant cells. Another systematic review showed that metformin yields more benefits for treating pancreatic and hepatic cancers compared with colon, breast, and prostate cancers. Several possible mechanisms for the anti-cancer effects of metformin have been discussed. Indirect pathways include the avoidance of weight gain, as compared with other anti-diabetic medications, and the amelioration of insulin resistance by reducing hyperinsulinemia. However, both pathways may promote carcinogenesis. The direct pathways of metformin can stimulate AMP-activated protein kinase (AMPK) through LKB-1 (serine–threonine liver kinase B1), which is a tumor-suppressing protein kinase. AMPK inhibits protein synthesis and hepatic glucose output during cellular stress and inhibits the mammalian target of rapamycin (mTOR), which is a downstream effector of growth factor signaling commonly present in malignant cells. Another systematic review showed that metformin yields more benefits for treating pancreatic and hepatic cancers compared with colon, breast, and prostate cancers. Several possible mechanisms for the anti-cancer effects of metformin have been discussed. Indirect pathways include the avoidance of weight gain, as compared with other anti-diabetic medications, and the amelioration of insulin resistance by reducing hyperinsulinemia. However, both pathways may promote carcinogenesis. The direct pathways of metformin can stimulate AMP-activated protein kinase (AMPK) through LKB-1 (serine–threonine liver kinase B1), which is a tumor-suppressing protein kinase. AMPK inhibits protein synthesis and hepatic glucose output during cellular stress

### TABLE 4. Association Between HCC and Different Statin Use in Patients With Comorbidity

| Statin | HCC Case/N | Model 1 | Model 2 |
|--------|------------|---------|---------|
| Non-statin | 223/334 | 1.00 | 1.00 |
| Statin | | | |
| Simvastatin | 14/36 | 0.31 (0.14–0.67)** | 0.81 (0.68–0.97)** |
| Atorvastatin | 16/47 | 0.29 (0.15–0.57)** | 0.96 (0.92–0.99)** |
| Pravastatin | 9/13 | 1.42 (0.39–5.17) | 0.98 (0.89–1.09) |
| Fluvastatin | 7/17 | 0.48 (0.17–1.32) | 0.94 (0.85–1.03) |
| Lovastatin | 11/22 | 0.57 (0.23–1.43) | 0.94 (0.86–1.02) |
| Rosuvastatin | 4/11 | 0.32 (0.09–1.16) | 0.84 (0.66–1.06) |
| Without Metformin use | | | |
| Non-statin | 79/116 | 1.00 | 1.00 |
| Statin | | | |
| Simvastatin | 3/5 | 0.99 (0.11–8.95) | 0.91 (0.70–1.20) |
| Atorvastatin | 3/8 | 0.47 (0.10–2.30) | 0.95 (0.88–1.03) |
| Pravastatin | 1/2 | 4.16 (0.13–139) | 0.99 (0.83–1.18) |
| Fluvastatin | 0/3 | — | — |
| Lovastatin | 1/3 | 0.24 (0.02–2.82) | 0.94 (0.86–1.03) |
| Rosuvastatin | 0/1 | — | — |
| With Metformin use | | | |
| Non-statin | 144/218 | 1.00 | 1.00 |
| Statin | | | |
| Simvastatin | 11/31 | 0.26 (0.11–0.60)** | 0.80 (0.66–0.97)** |
| Atorvastatin | 13/39 | 0.26 (0.12–0.55)** | 0.96 (0.92–1.00) |
| Pravastatin | 8/11 | 1.25 (0.30–5.11) | 0.98 (0.86–1.12) |
| Fluvastatin | 7/14 | 0.78 (0.25–2.42) | 0.97 (0.87–1.08) |
| Lovastatin | 10/19 | 0.64 (0.23–1.76) | 0.94 (0.81–1.08) |
| Rosuvastatin | 4/10 | 0.35 (0.09–1.37) | 0.87 (0.67–1.12) |

Model 1: the risk for HCC in each statin users compared with non-statin users adjusted for cirrhosis duration, ALD, HCV, and HBV. Model 2: the risk for HCC in patients with each statin exposure duration (month) compared with people without statin exposure adjusted for cirrhosis duration, ALD, HCV, and HBV. ALD = alcoholic liver damage, CI = confidence interval, HBV = hepatitis B virus, HCC = hepatocellular carcinoma, HCV = hepatitis C virus, OR = odds ratio.

**P < 0.05.
**P < 0.01.
***P < 0.001.
accompanied by continuous chronic inflammation that results from hyperglycemia or hyperinsulinemia, which play key roles in cancer cell activity, including its initiation, promotion, and progression. Metformin can decrease insulin resistance but cannot directly reduce abnormal insulin secretion. Second, high concentrations of insulin secreted by β-cells pass directly into liver tissues and increase the risk of HCC. Metformin has no effect on this pathway. Third, DM results from chronic inflammation and can cause additional oxidative stress. Compared with metformin, statins more effectively reduce inflammation. Forth, the sub-analysis results (Table 2) showed that the duration of metformin use may be another issue worth exploration in relation to its anti-cancer effects. Certain studies have highlighted that low concentrations of metformin (<0.5 mM) selectively inhibit CD133+ cell proliferation and reduce cancer stem cell activity. The cancer stem cell hypothesis suggests that cancer stem cells play a key role in tumor genesis, recurrence, and the resistance to adjuvant cancer therapies. Another study stated that low doses of metformin can induce p53-dependent senescence in hepatoma cells. One study conducted in Asia showed that low doses of metformin (<1000 mg/day) may be associated with a reduced risk of cancer in diabetic patients, and another article reported a trend of dose–response relations to cancers (treatment that employed less than 250 mg/day for 1 month were discussed). The results of the current study indicated a significant trend of decreased HCC risk in DM patients who received occasional metformin treatment (Table 2, trend test P = .03). This may imply that low doses or short durations of metformin use are sufficient to yield anti-cancer effects in Asian DM patients with HCC. Fifth, according to the results in Table 1, NAFLD and metformin do not contribute to the risk of HCC. Racial or genetic differences may be a possible explanation.

**Metformin Combined With Statins and HCC**

Our research did not involve observing all the positive effects of combining metformin and statins regarding reduced HCC risk. However, specific statins, such as simvastatin and atorvastatin alone or combined with metformin, may lower the risk of HCC (Table 3). We infer that this may be caused by the interaction of metformin with simvastatin or atorvastatin. DM patients treated with metformin combined with simvastatin exhibited a relatively low risk of comorbidities when they presented with HCC (Table 4). This finding indicates that administering this type of combination therapy to DM patients with HCC is fairly beneficial in Asia.

**SUMMARY**

Our study showed no benefit of metformin to HCC. Treatment with pravastatin alone or with metformin also showed no benefit to HCC. Further basic research is required for clarification.

**STUDY LIMITATIONS**

Two limitations to our studies should be considered. First, the LHID2000 cannot offer detailed information such as alcohol consumption, which may be risk factor for HCC. Second, all patient data in the LHID2000 are anonymous. Relevant clinical clues, such as child score for liver cirrhosis, imaging results, or pathologic findings, cannot be followed in our study.

**CONCLUSION**

According to the research results, not all metformin combined with statins may reduce the incidence of HCC in Asia and not all of this combination therapy may decrease the development of comorbidities in diabetic patients with HCC.

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