INTRODUCTION

With advances in imaging modalities, perioperative care, surgical techniques, and indications for liver resection have been expanded. Both right and extended right hepatic lobectomy are occasionally associated with postoperative liver failure (1) because it involves a resection of a large portion of the liver. Therefore, choosing which procedure to perform must be done with care for each individual. Removing more than 70% of liver parenchyma has been known to increase the risk of postoperative mortality (1). Previous clinical and experimental reports have shown that when portal veins are occluded or stenotic, it leads to atrophy of the affected lobe or segment along with compensatory hypertrophy of the unaffected hepatic lobe (2, 3). Based on this result, preoperative portal vein embolization (PVE) has become a well-established means to upsize the future remnant liver as well as to prevent postoperative liver insufficiency in patients considered for extensive liver resection (4-6). Additional hepatic artery embolization combined with PVE has been reported to be beneficial for compensatory hypertrophy compared with PVE alone (7, 8).

Our study has been performed to compare the volume change and the regenerative capacity between portal vein ligation (PVL) and heterochronous PVL with hepatic artery ligation (HAL) in a rodent model.
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**MATERIALS AND METHODS**

Experimental Design and Surgical Procedures

This study was approved by Dankook University’s "Animal Care and Use Committee" (South Korea). All procedures were conducted in accordance with the ‘Guide for Care and Use of Laboratory Animals’ published by the National Institutes of Health and the ethical guidance of the International Association for the Study of Pain.

Male Sprague-Dawley rats (Samtako Co, Seoul, Korea) aged 13 weeks and weighing between 350 and 450 g were used for all of the experiments. All the animals were kept in identical housing units with an alternating 12 h light-dark cycle. Food and water were available ad libitum.

After an acclimation period of at least 7 days, the animals were separated into three groups as follows: group I, performed only ligation of the left lateral and median portal vein branches (which supplies approximately 70% of total liver volume); group II, first completed ligation of the portal vein, followed by ligation of the same branches of the hepatic artery after 48 hours; control group, conducted laparotomy without ligation.

Prior to the operations, all of the animals were anesthetized with an intraperitoneal injection of zolazepam mixed with xylazine and normal saline at room temperature, using a non-sterile, clean technique. In group I, via midline incision, the branch of the portal vein supplying the median and the left lateral lobes of the liver was ligated using 6-0 silk. The abdominal wall and skin were closed with 4-0 silk. In group II, the second operation was performed after 48 hours, by performing ligation of the same branch of the hepatic artery. Abdominal cavity irrigation using a mixture of normal saline and antibiotics was performed as the last step of each operation.

Postoperatively, the animals had free access to water and food. Rats from each group were sacrificed using the same anesthesia method described above on 1, 3, 5, and 7 days after the operation. At least five animals per group were employed for every study time point. Blood was collected by the heart puncture method just before sacrifice. The liver was removed, and the weight of the body and non-ligated lobes were weighed.

**Immunohistochemistry**

Whole liver was dissected and postfixed in 10% neutral-buffered formalin for 24 hours. Primary antibodies used for immunostaining were MIB-5 (a novel antibody reactive with the equivalent Ki-67 protein, 1 : 800, LabVision, Cheshire, UK), and c-kit (1 : 400, DAKO, Glostrup, Denmark). MIB-5 is used for the detection of cells in the active phases of the cell cycle and c-kit is used for the detection of stem cell expression. The liver was diluted in normal goat serum buffer. We used the Zymed non-biotin amplification system (Zymed Laboratories Inc., South San Francisco, CA, USA). Only nuclear-stained cells in Ki-67 and cytoplasmic-stained cells in c-kit were considered meaningful. The number of Ki-67-positive cells and c-kit-positive cells were counted using × 400 magnification in 10 randomly selected visual fields, using samples from five animals per group from each of the three experimental groups. Two pathologists evaluated the staining independently, and any discrepancy was resolved by a consensus review.

**Assessment of Liver Proliferation and Function**

The ratio of the weight of the nonligated lobe to the body weight was assessed using the following equation: R = weight of nonligated lobes/body weight (nLW/BW). Plasma was separated by centrifuge using blood samples collected by a direct heart puncture. The collected plasma was used for the standard liver function tests. The alanine aminotransferase (ALT), aspartate aminotransferase (AST), total bilirubin, alkaline phosphatase, gamma glutamyl transpeptidase, albumin, prothrombin time, and the international normalized ratio (INR) of the blood samples were determined. All of the parameters were measured using the standard laboratory methods.

**Statistical Analysis**

The values for the results were expressed as means ± standard error of the mean. Student’s t-test and Kruskal-Wallis analyses were used for the statistical analysis. The probability values lower than 0.05 were considered significant.

**RESULTS**

Two animals were lost in group II following the operation. During the initial stage of the experiments, rats that had under-
There was an overall increase in the number of c-kit-positive hepatocytes in group I detected on day 1; yet, the peak level was significantly higher in group II than in group I through days 3 and 7 after surgery. The nLW/BW was higher in group II than in group I at the operation site, and eventually died. Thereafter, all rats were separated and remained alive until the end of the study.

### Results of nonligated liver weight/body weight

There was an overall increase in the number of c-kit-positive hepatocytes in group I detected on day 1; yet, the peak level was significantly higher in group II than in group I through days 5 and 7. These results are summarized in Table 1.

![Table 1. Regenerated Liver Weight and Serologic Profiles of Liver Injury and Function](chart)

**Note.** - *The p value estimated with the t-test between PVL and PVL + HAL.

*The p value estimated with the Kruskal–Wallis analysis among PVL, PVL + HAL, and the control group.

ALP = alkaline phosphatase; ALT = alanine aminotransferase; AST = aspartate aminotransferase; HAL = hepatic artery ligation; nLW/BW = the ratio of the weight of the nonligated lobes to body weight at sacrificing; POD = postoperative day; PVL = portal vein ligation.
in group II was not detected until 3 days after surgery, as shown in Figs. 2, 3.

Expression of MIB-5

There was an overall increase in the nuclear expression of MIB-5 in group I compared to group II. The number of MIB-5-positive hepatocytes was significantly higher in group I than in group II from day 1 to 5 (p < 0.05). Unlike c-kit, the peak level of MIB-5-positive hepatocytes was noted on day 3 in both groups I and II. There were little differences in the expression of MIB-5 between groups I and II on day 7 (p = 0.80), as presented in Figs. 4, 5.

**DISCUSSION**

Liver regeneration following PVL has been studied in depth...
for several decades (9). Many studies have reported that PVE is an effective method for liver regeneration prior to liver resection (4-6). However, other studies have reported that adding hepatic artery embolization yields better results (6, 7, 10, 11). Kong et al. (10) reported that compensatory hypertrophy of the unaffected lobes was evident. Data from their study revealed that when PVL and HAL are performed simultaneously, the ligated liver tissues undergo massive necrosis (10). Nagino et al. (7) reported that additional hepatic artery embolization was effective in inducing compensatory hypertrophy; however, they also reported adverse events, such as necrosis and abscess formation. Like the results of Nagino et al., in our study, necrosis of the ligated lobe was grossly present in group II in which both PVL and HAL were performed at an interval of 48 h. As the evaluation of the ligated lobe was not the major point of our study, we did not

Fig. 4. Results of MIB-5 expression. Between group I and II, there are statistically significant differences in the expression of MIB-5 from postoperative days 1 to 5 \( (p < 0.05) \).

Note.—HAL = hepatic artery ligation, HPF = high power field, POD = postoperative day, PVL = portal vein ligation

Fig. 5. Microscopic images of nonligated lobes with MIB-5 staining \( (x = 200) \). Stained cells are more apparent in group I \( (A) \) than II \( (B) \) on day 3. On day 7, there are no significant differences between group I \( (C) \) and II \( (D) \).
Kong et al. (10) also reported that the ratio of the weight of the nonligated lobe to the body weight was significantly higher at 48 h in a group that underwent PVL plus the HAL heterochronous procedure than in a PVL-only group and a simultaneous PVL plus the HAL group. The time of the peak level of hypertrophy (168 h) was similar to our results. At first, hypertrophy of the nonligated lobe seemed more prominent in group II (PVL + HAL); however, there were no significant differences on day 7 in the two groups due to the delayed progressive hypertrophy in group I. Also, group II showed a slightly decreased volume ratio on day 7. This is in common with the reports by Veteläinen et al. (12) who noted that the abrupt increase in the volume followed by a subsequent decrease in the volume can be explained by a massive, progressive apoptosis. According to the report by Kong et al. (10), massive necrosis of the ligated lobe found in group II may have interfered with the regeneration of the nonligated lobe. However, further investigation regarding this was not performed in this study since apoptosis could not be proven due to the relatively short follow-up time of 7 days.

Rozga et al. (9) reported almost total resorption of the necroses in the portal vein of the ligated lobes, which was seen in 4 days after ligation. However, in our study, grossly necrotic tissue was found in all rats in group II at all time points. In our opinion, this may have been due to the fact that there was not enough time for the healing process from PVL because of the subsequent HAL.

Liver function tests revealed significantly higher levels of AST/ALT and total bilirubin in group II. Severe liver injury was also confirmed in group II with additional HAL. Albumin levels were also lower in group II, and INR was increased in group II. These results imply a decreased synthetic function of the liver in group II. Nakao et al. (13) also reported similar results.

In 1975, it was established that portal venous blood flow promoted hepatic cell regeneration (14). For the evaluation of liver regeneration, immunohistochemical staining for MIB-5 and c-kit were used in this study. MIB-5 is a novel antibody reactive with the rat equivalent to Ki-67 protein. The Ki-67 reacts exclusively with the nuclei of the proliferating cells and is expressed during all active parts of the cell division cycle (G1, S, G2, and M); however, it is absent in the resting cells (G0) (15). C-kit protein binds to the stem cell factor, a substance that causes certain types of cells to grow. Signaling through c-kit protein plays a role in cell survival, proliferation, and differentiation (16). In contrast to the results of Kong et al. (10), the MIB-5-labeling index showed higher levels in group I in our study. The levels of c-kit-positive hepatocytes were also higher in group I. Considering the increase in liver enzyme levels and bilirubin, acute damage to the hepatocytes in group II may have interfered with liver regeneration of the nonligated lobe. These results are similar to a previous report by Rozga et al. (9), who reported the highest values of DNA-synthetic activity in rats in a 70% ligation group at 24 h. In these rats, the increased DNA-synthetic activity persisted at least until 2 weeks after the procedure. The most vigorous mitotic activity in PVL in rats was noted at 48 h and persisted for 2 weeks (12). In group I of our study, the expression of c-kit and Ki-67 cells was highest on day 1 and day 3, respectively. On day 7, the levels of both markers were very low, possibly due to the markedly decreased regeneration activity. As a result, it is difficult to determine whether significant volume hypertrophy occurred after 7 days or at the time interval of the two markers has any correlation with liver regeneration.

There are several limitations to our study. First, portal vein ligation and portal vein embolization is not the same procedure. Broering et al. (17) and Iida et al. (18) reported that PVE was more efficient than PVL for the induction of hypertrophy of the contralateral lobe. They explained that one of the reasons for the larger liver volumes is the portal-portal collateral vessels. However, as in other several studies, fortnight normalization of increased portal blood flow induced by portal vein occlusion in humans and the early peak of hepatocytes proliferation after portal occlusion in rodents suggests that liver hypertrophy after portal occlusion is induced earlier than the formation of portal-portal collateral formation (19-21). Hence, this may have little impact on liver hypertrophy. Therefore we considered that our results can be applicable to PVE. Second, this study was conducted over a short period of time. Third, we did not perform a microscopic evaluation of the ligated lobe. Without this, the relationship between ligated and nonligated lobes was not revealed from our results. Fourth, we had not confined the c-kit-positive expression cells to the stem cells of the liver; therefore, the relevance of the stem cell expression cannot be concluded.

In conclusion, PVL alone was found to be safe and effective to induce compensatory liver regeneration. Performing both PVL
and HAL does not confer additional benefits.

REFERENCES

백서에서 단독 간문맥 결찰과 간문맥과 간동맥의 순차적 결찰 후 간재생의 비교 연구

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목적: 백서에서 간문맥 단독결찰과 간문맥과 간동맥의 순차적 결찰 후 반대측 간의 재생에 대하여 비교하고자 한다.
대상과 방법: 동물들은 세 그룹으로 분류하였다. 1) 좌측 간문맥 결찰만 시행한 그룹, 2) 좌측 간문맥 결찰 후 48시간 이 후에 동일한 간동맥 결찰을 시행한 그룹, 3) 혈관 결찰 없이 개복술만 시행한 경우. 술 후 1, 3, 5, 7일째 되는 날 각 그룹 에서 최소 5마리씩을 희생시켜 간 부피 변화와 간 기능 검사 결과, 간 재생에 관한 면역염색화학 결과를 얻었다.
결과: 결찰하지 않은 반대측 간의 부피 변화는 5일과 7일째 첫 번째와 두 번째 그룹에서 유의한 차이는 없었다. Alanine aminotransferase와 총 빌리루빈 값은 두 번째 그룹에서 유의하게 높았으며, 일부만값은 첫 번째 그룹에서 유의하게 높았 다. 간 재생의 과정에 있는 세포 확인을 위한 c-kit와 MB-5 면역세포학적 검사 양성세포는 술 후 1, 3, 5일에 첫 번째 그룹에서 통계적으로 유의하게 많은 분포를 보였다. 수술의 직접적인 결과로 사망한 경우는 없었다.
결론: 단독 간문맥 결찰은 대상성 간재생에 비교적 안전하고 유의한 방법이다. 본 연구에서는 추가적인 간동맥 결찰은 부 가적인 이득이 없을 것으로 보인다.

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