Partial ALPPS with a longer wait between procedures is safe and yields adequate future liver remnant hypertrophy

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Backgrounds/Aims: Associating Liver Partition and Portal Vein Ligation for Staged Hepatectomy (ALPPS) has generated controversy due to high morbidity and mortality. We present our series of patients with 30-40% parenchymal transection and minimal hilar dissection.

Methods: Patients who had partial ALPPS between April 2015 and April 2016 were included. Patients with colorectal liver metastases (CRLM) had their future liver remnants (FLR) cleared with metastasectomies. The liver was divided along the future line of transection to 30-40%, right portal vein was stapled and divided without extensive hilar dissection, with minimal handling of right liver, which was not mobilised. We preserved the middle hepatic vein. Data were collected prospectively for hypertrophy of the FLR, morbidity and mortality.

Results: Among the 8 patients (age 25-68) investigated, one patient with cholangiocarcinoma had portal vein embolization prior to partial ALPPS. All patients completed two stages with adequate FLR hypertrophy at a median of 28 days. No mortality was found. The median length of stay after stages 1 and 2 was 9 and 9.6 days, respectively. The median increase in FLR was 38%.

Conclusions: A limited transection of 30-40%, minimal hilar dissection and longer wait between stages yielded adequate FLR hypertrophy with low morbidity and no mortality. (Ann Hepatobiliary Pancreat Surg 2019;23:13-19)

Key Words: Liver resection; Future liver remnant; ALPPS; Hypertrophy; Colorectal liver metastases

INTRODUCTION

Liver resection offers the best chance for long-term survival of patients with colorectal liver metastases (CRLM) and hilar cholangiocarcinoma. A combination of better chemotherapeutic regimens and surgical innovations has increased the resection rates in patients with CRLM. Several technical limiting factors such as location of the tumor and distribution preclude liver resection. A future liver remnant (FLR) of 20-30% is deemed safe in patients with normal background liver and at least 40% FLR in patients with compromised liver (steatohepatitis, fibrosis, cirrhosis).¹ The techniques that have proved successful in increasing resection rates include neoadjuvant chemotherapy in patients with CRLM,² ³ portal vein embolization (PVE),⁴ portal vein ligation (PVL),⁵ two-stage liver resections,⁶ Associating Liver Partition and Portal vein ligation in Staged hepatectomy (ALPPS)⁸ and Associating Portal Embolization and Artery Ligation (APEAL).⁹

The ALPPS procedure created tremendous interest among liver surgeons because of the pace of liver regeneration that it allowed, enabling completion of the two stages of liver resection within 9 to 11 days. The procedure has yet to be widely adopted mainly due to concerns with increasing morbidity, mortality⁸ and potentially higher recurrence rates.¹¹ ¹²

We performed the first ALPPS procedure in April 2015 using only partial transection of liver and other technical modifications involving minimal dissection. This approach produced significant hypertrophy of the FLR and we have performed 8 procedures with no mortality and acceptable morbidity. We proceeded to the second stage in all pa-
tients after a longer interval than the classical ALPPS. The aim of this report is to show the feasibility of safe two-stage liver resection via transection of only 30-40% of liver (partial ALPPS). A longer interval between the two procedures may lead to less liver failure.

PATIENTS AND METHODS

In order to describe the modified version of the ALPPS procedure, the first operation is briefly explained below.

The patient was a 68-year-old lady with obstructing carcinoma of descending colon and synchronous liver metastases. She underwent left hemicolectomy with end stoma in January 2014 for a T4bN0M1 tumor. The liver metastases involved all segments except segment 1 and 5 (Fig. 1). The patient had palliative chemotherapy with 4 cycles of XELOX (capecitabine plus oxaliplatin) following progression with FOLFIRI (leucovorin calcium, 5-fluorouracil, and irinotecan) regimen as second-line treatment. Aflibercept was added to augment the response, and was completed in February 2015. An Fludeoxyglucose 18F (FDG)-positron emission tomography (PET) scan revealed liver only metastases and a pair of foci in the left lateral abdominal wall suggestive of local recurrence. However, review of cross-sectional images in this area revealed no masses and after discussion with the hepatobiliary multidisciplinary team (MDT) meeting it was decided to explore the possibility of two-stage liver resection.

Using laparotomy, the tumors in the segments 2, 3 and 4b were removed as metastasectomies (Fig. 2) with macroscopically clear margins. The FLR was inadequate. Cholecystectomy was carried out and the right portal vein (RPV) was ligated with minimal dissection by incising the peritoneum on the right and posterior aspects of the hilum. The RPV was stapled and divided (Fig. 3). Metastasectomy of the tumor in the segment 4b led to a large defect in the liver with only a small sliver of tissue between this defect and the gall bladder fossa. This was divided down to the hilum resulting in transection of nearly a third of liver. The right liver was not mobilized. A silastic drain

![Fig. 1. Metastases detected in the hepatic segment 6 (A), segments 2, and 7 (B), segments 3, 4, and 8 (C).](image1)

![Fig. 2. Intraoperative photograph showing metastasectomies in segment 4.](image2)

![Fig. 3. Right portal vein stapling (arrow) with minimal hilar dissection.](image3)
was placed along the cut surface of the liver. The patient recovered adequately and was discharged on postoperative day 6 with the drain in situ. Histological examination revealed complete resection with a margin of at least 1 mm. A computed tomography (CT) scan was performed to measure the volume of the FLR 12 days after the first stage procedure and volumes were assessed using standard procedure described below.\(^{13}\)

The CT assessment showed an increase in FLR from 408 ml to 630 ml, or an increase of 54%. She then underwent a completely extended right hemihepatectomy 21 days after the first stage procedure, using and completing the previous transection plane. Histologically, the tumor in the segment 7 close to the inferior vena cava extended to within 1 mm of the resection margin. The patient recovered without any complication and was discharged 7 days after the operation.

The above experience indicated that FLR hypertrophy was feasible with partial transection of the liver. We performed similar procedures in seven other patients reported here. We defined 30-40% transection for an extended right hepatectomy as resection of the liver along the falciform ligament and removing the segment 4b pedicle. The cut was extended up to the segment 4a pedicle which was left in situ. The middle hepatic vein was not divided in the first operation.

**Liver volume assessment**

Liver volume was assessed using commercial volumetry software, Volume Viewer advanced imaging platform (GE Healthcare Version 2.0, AW Server) and calculated by manually defining the contours of the liver parenchyma on representative sequential axial 0.625 mm slices of the CT acquisition on a liquid crystal display (1920×1200 resolution, 24-inch wide screen display, LA2405, HP Compaq).

The volume rendering tool automatically generates a 3D volume and calculates the total liver volume (TLV) from the selected slices. This process was repeated for all definable liver lesions to yield the estimated total lesion volume and the two volumes subtracted to determine the estimated Total Functional Liver Volume (TFLV).

The selected liver volume was used to define the predicted Functional Residual Liver Volume (FRLV) by segmenting and excluding the hepatic parenchyma planned for resection as directed by the liver surgeon. The ratio of predicted functional residual liver volume (%FRLV) was then calculated.

The actual FRLV was subsequently measured on the post-ALPPS axial CT performed on day 14 to determine the extent of hypertrophy and feasibility of complete hepatectomy.

**Hepatobiliary (HIDA) assessment**

We performed a HIDA scan as part of the evaluation following the first stage for differential evaluation of the function of FLR. Hepatobiliary scintigraphy was performed by injecting 70 MBq of \(^{99m}\)Tc-mebrofenin intravenously after a 4-h fast. Dynamic anterior and posterior images were acquired for 60 min, with the patient lying supine on a large field-of-view dual-headed gamma camera (Millennium Hawkeye, GE) equipped with low-energy high-resolution collimators. Images were grouped into twelve 5-minute frames. The images were assessed, both to calculate the function of the future liver remnant, and also to exclude biliary leak.

Quantitation of the FLR was performed by drawing regions of interest around the left and right liver on both the anterior and posterior 0-5-min grouped image, as defined by the prior hepatic cut. The % contribution of each half of the liver was calculated by geometric mean.

In summary, all patients with CRLM had a CT chest, abdomen and pelvis, magnetic resonance imaging of the liver and a FDG PET scan before surgery. The patients were discharged after the first stage and exposed to a CT scan of abdomen and a HIDA scan 2 weeks from the procedure. The second stage procedure was scheduled 3 weeks from first stage. Chemotherapy was completed at least 6 weeks before the first stage procedure in all patients with CRLM.

We recorded the complications and classified them according to the Clavien-Dindo system.\(^{14}\) Post-hepatectomy liver failure (PHLF) was classified according to the grades proposed by the International Study Group of Liver Surgery.\(^{15}\)

We followed up patients at 4 weeks after surgery and at 3 monthly intervals. The CT scan of chest abdomen and pelvis was carried out at 3 months, 6 months and a year from the second stage procedure. Recurrence data and mortality were censored at December 2016.
The procedure was approved by the Clinical Effectiveness Committee at the University Hospital of Wales (UK).

RESULTS

A total of 8 patients, with a median age of 61 years (25-68), 6 with CRLM and 2 with hilar cholangiocarcinoma were studied. All patients proceeded to a second stage operation with adequate FLR hypertrophy at a median interval of 32 (21-36) days between the procedures. No mortality was detected. The pre and post mini-ALPPS FLR, morbidity and mortality data are presented in Table 1.

The last patient had a pre-operative PVE and a trial dissection for a hilar cholangiocarcinoma. The liver volume was inadequate at the time of surgery despite a reasonable response to PVE and a partial parenchymal transection was performed. This patient developed post-operative bile leak requiring surgical washout. Following recovery, the second-stage resection was performed on day 36. Otherwise, following first-stage resection, 1 patient developed a bile leak and another patient presented with hospital-acquired pneumonia. Following second-stage resection, 1 patient developed transient liver failure and 2 patients developed bile leaks (1 requiring endoscopic retrograde cholangiopancreatography). All of the patients underwent CT scan at a median of 14 days (11-35) of first-stage surgery to assess the volume of the FLR. A hypertrophy of 10% to 78% was achieved. Second-stage operations included right trisectionectomies. Patients were followed up for a median of 390 days (233-472).

DISCUSSION

We have shown that FLR can hypertrophy enough to allow second-stage resection by only dividing the liver partially. Besides, other techniques listed below were used to reduce morbidity and mortality. The right liver which was scheduled for resection in the second stage was not mobilized at all. The hilar dissection only involved an incision of the peritoneal layer on the right and posterior aspects of the free edge of the lesser omentum to allow access to the right portal vein. This was divided using a stapler. No bag was used to isolate the liver for subsequent resection, thus reducing the risk of infection. A silastic drain was left where the liver was divided to en-

| Table 1. Demographics, pre and post partial ALPPS FLR, morbidity and blood transfusion in patients undergoing partial ALPPS |
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| **Age** | **Pathology** | **Preop FLR (ml)** | **Post 1st stage FLR (ml)** | **S1 blood transfusion** | **Time to second stage (days)** | **Morbidity stage 1** | **S2 blood transfusion** | **Morbidity stage 2** |
| --- | --- | --- | --- | --- | --- | --- | --- | --- |
| 68 | CRLM | 408 | 630 | 1167 | 54 | None | 21 | None |
| 25 | CRLM | 285 | 400 | 1026 | 66 | None | 52% | 21 | None |
| 62 | CRLM | 555 | 749 | 1548 | 36 | 28% | 28 | 50 | None |
| 68 | CRLM | 351 | 476 | 1470 | 36 | 40% | 35 | None | 42% |
| 63 | CRLM | 342 | 476 | 1470 | 36 | 40% | 35 | None | 42% |
| 65 | Hilar cholangiocarcinoma | 419 | 586 | 1563 | 40 | 42% | 35 | Wound infection | 21 |
| 43 | CRLM | 285 | 400 | 1026 | 66 | None | 52% | 21 | None |
| 68 | CRLM | 351 | 476 | 1470 | 36 | 40% | 35 | None | 42% |
| 69 | Hilar cholangiocarcinoma | 419 | 586 | 1563 | 40 | 42% | 35 | Wound infection | 21 |
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CRLM, colorectal liver metastases; FLR, future liver remnant; SLV, standard liver volume; S1, stage 1; S2, stage 2
able easy access to the second procedure. We left a long polydioxanone (PDS) suture around the hilum as a loop for easy identification of the hilum to facilitate Pringle’s maneuver during the second operation.

de Santibañes et al.\textsuperscript{16} have described a similar procedure in four patients. However, they did not perform any hilar dissection during the first stage. Instead, they performed intraoperative PVE using the inferior mesenteric vein and designated the procedure as Mini-ALPPS. They claim that the lack of hilar dissection is beneficial for the second stage. We perform minimal hilar dissection and only the right and posterior peritoneum on the hilum is dissected to allow access to the right portal vein, which is stapled. We have not encountered any difficulties related to this during the second stage. We started to perform the procedure before the Santibanes paper was published, and hence our approach of minimal hilar dissection. This approach also avoids any potential logistic difficulties in performing PVE in the theatre. Li et al.\textsuperscript{17} described similar operation of PVE 2 days after the first procedure in patients with tumor involving the hilum.

The advantages of ALPPS include the higher completion rates of the two stages compared with two-stage hepatectomies (TSH). However, critics have shown a poorer long-term outcome in patients undergoing ALPPS compared with TSH.\textsuperscript{18} Our study is an observational study and hence prone to many biases. A randomized controlled trial is the only way to compare the long-term outcomes of ALPPS vs. TSH. Although no randomized controlled trial compared ALPPS and TSH, a systematic review and meta-analysis of all the comparative studies showed that the overall survival was not different between the two approaches. ALPPS was associated with higher morbidity and mortality.\textsuperscript{19} In addition, higher rates of liver failure followed ALPPS despite volumetric increase in the remnant liver. Matsuo et al. have shown that the hepatocytes that regenerate after ALPPS were morphologically immature compared with PVE.\textsuperscript{20} Only one patient developed ISGLS grade B liver failure in our series, with a 35-day interval between the procedures. The median interval between the two procedures was 21 days and 5 out of the 8 patients underwent the second procedure within 30 days of the first intervention. This longer interval compared with the 9-11 days described in most series may have reduced the liver failure, which requires confirmation in a larger series or a randomized controlled trial. The last two patients in the series did not manifest significant hypertrophy (15% and 10% respectively), which is most likely due to portoportal collaterals found on the post-partial ALPPS CT scan. The existence or development of portoportal collaterals is a significant factor impeding hypertrophy following PVE.\textsuperscript{21}

The median increase in hypertrophy in our patients was 38% (10-78%), which was lower than in classical ALPPS, where the range varied from 58% to 110%.\textsuperscript{22} and attributed to unidentified portoportal collaterals. Indeed the mechanism for significant hypertrophy following ALPPS is not fully understood. Our technique yielded enough hypertrophy such that all patients proceeded to the second stage operation.

It could be argued that the initial liver volumes were adequate to perform a single-stage procedure. However, we preferred two stages based on the quality of the liver at the time of the first stage along with the volume. Rosok et al.\textsuperscript{23} reported the Scandinavian experience involving a small number of patients, who were converted to ALPPS during a planned single-stage operation. The factors underlying the decision included detection of additional lesions or suspected poor quality of liver, which was critiqued by Belghiti et al.\textsuperscript{24} in their editorial in the same issue of the journal, suggesting that many of these patients could have been operated with a single-stage procedure. Liver failure after resection carries high mortality. Currently, except for the volume, no predictors are available for the development of liver failure. However, Cieslak et al.\textsuperscript{25} showed that $^{99m}$Tc-mebrofenin scintigraphy with a cut-off value of 2.7/min/m\textsuperscript{2} may be used to assess adequacy of FLR function. Further, no strict correlation with increase in volume was observed with CT volumetry. The HIDA scan was not used in the same way. The differential volume of the liver was calculated and in the presence of adequate FLR (i.e. >25%) we proceeded to the second stage of ALPPS.

We have shown that partial ALPPS is a safe alternative to the ALPPS originally described. It facilitates safe and rapid two-stage procedure for tumor clearance in patients with CRLM. Many previous studies report a near 100% success in proceeding to the second stage. However, subsequent to this report we found a 79-year-old patient who failed to proceed to second-stage ALPPS due to a heart
block and was thus contraindicated for the procedure. The patient lacked adequate hypertrophy of the FLR and underwent PVE due to abnormal anatomy that allowed perfusion of the right liver. The long-term outcomes in patients undergoing ALPPS for CRLM are questionable as well.

There are several limitations to our study. This is a small study of only 8 patients, and the technique used to measure volumes by HIDA scan was not validated. It is possible that the recorded volumes may have been adequate for a single-stage procedure. We feel that the safety of the procedure facilitates liver surgeons to consider this option that is intermediate between traditional ALPPS and TSH. A longer interval between the two procedures along with the use of $^{99m}$Tc-mebrofenin scintigraphy to assess FLR function may prevent postoperative liver failure.

This preliminary experience may allow prospective consideration of interventions in patients requiring FLR clearance that is inadequate without an additional procedure such as PVE or PVL. Modifications to the original ALPPS described facilitate safe two-stage hepatectomy, mainly in patients with CRLM.

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