Introduction

Colorectal cancer is the third most common malignancy and the third most common cause of cancer-related death in the United States. Most patients die from distant metastases, and 60% will develop liver metastases over the course of their disease (1,2). While curative-intent hepatectomy offers the best chance for long-term survival, fewer than 25% of patients are resectable (3). Over the last four decades, multimodal medical and surgical treatments have been developed in an effort to address historically modest response rates to systemic chemotherapy. It is in this setting that hepatic arterial infusion pump (HAIP) chemotherapy has emerged as an effective and unique form of regional chemotherapy.

Over the past five decades, the safety and efficacy of HAIP chemotherapy has been evaluated. Randomized control led trials conducted in the setting of unresectable liver metastasis have demonstrated improved response rates, quality of life, and overall survival (4-10). Studies examining the use of HAIP chemotherapy in the adjuvant setting have also shown benefit, with improved hepatic recurrence rates and survival (11,12).

Traditionally, HAIP placement has been performed via an open approach (13). In the context of unresectable metastatic disease, the morbidity of a laparotomy has been a limiting aspect of HAIP chemotherapy (14). The rising prevalence of laparoscopic and robotic approaches to colectomy and hepatectomy in colorectal cancer surgery represent yet another impulse to adopt minimally invasive (MIS) approaches to HAIP placement.

Herein, we describe the development of surgical approaches to HAIP placement, including the emergence of MIS HAIP surgery, relevant pre- and post-operative considerations, salient operative details in the MIS approach, and common complications related to HAIP placement and chemotherapy.
Historical development of hepatic arterial infusion chemotherapy

Anatomical and pathological research beginning in the 1950s established the dual blood supply of liver and the preferential arterial blood supply of intrahepatic tumors (15). Normal hepatocytes derive nutrients from portal venous blood flow, while liver tumors obtain nutrients almost exclusively from the hepatic artery.

In the 1960s, Sullivan and colleagues at the Lahey clinic made use of these insights to pioneer surgical techniques for inserting hepatic arterial catheters for regional delivery of chemotherapy. Though a small cohort with both heterogeneous tumor types and chemotherapeutic agents, they reported two important findings: (I) tumors showed favorable treatment response to regional chemotherapy, and (II) 5-fluoro-2’deoxyuridine (FUDR)—with its high first pass liver extraction and short half-life—maximized local efficacy while limiting systemic toxicity. Their work established the feasibility and general principles of surgical access for HAIP chemotherapy (16).

Initial HAIP chemotherapy protocols utilized an external pump system. This required either long-term hospitalization for continuous infusion, or the use of a bulky and cumbersome pump/harness with self-administration of chemotherapeutic drugs by patients (17). The development of a fully implantable pump system by Ensminger and colleagues by the 1980s transformed hepatic arterial chemotherapy into a truly outpatient treatment modality (18) (Figure 1).

Subsequent publications highlighted that although potentially beneficial, hepatic arterial chemotherapy was not without risk. In a period before stringent patient selection criteria, and with variable use of fully implantable pump systems, and procedural variation, mortality and major morbidity from surgery varied widely, with rates ranging from ~0–17% and 12–41%, respectively (20-25).

In the following decades, several developments led to improved safety and outcomes for HAIP surgery. These included formal criteria for patient selection, refinement in the management of variant hepatic arterial anatomy, and improved salvage of HAIP chemotherapy after complications (19,26-31). In modern series with good patient selection and confirmed in subsequent meta-analyses, mortality rates have approached 0%, with overall and pump-related complication rates reported between 10–20% (26,29).

One of the concerns regarding HAIP chemotherapy has been the morbidity associated with laparotomy in patients already debilitated with cancer, especially in the unresectable metastatic setting (32,33). With the emergence of laparoscopic surgery, several small series were published in the 1990s reporting the safety and feasibility of laparoscopic HAIP insertion as an alternative to laparotomy (34,35). To date, four series examining laparoscopic HAIP surgery and one examining robotic HAIP placement have been published (14,36-39). A single study has examined comparative outcomes in open vs. MIS approaches (39).

Overall rates of mortality for MIS HAIP surgeries are comparable to the open approach and range from 0–3%. As with open approaches, this rate is difficult to interpret given the frequency of concomitant colorectal and liver surgeries.
Table 1 Summary of studies examining MIS approaches to HAIP placement

<table>
<thead>
<tr>
<th>Author</th>
<th>Year</th>
<th>Study type</th>
<th>Approach</th>
<th>N</th>
<th>EBL (cc)</th>
<th>Concurrent surgical procedure</th>
<th>Convert to open</th>
<th>Median LOS (days)</th>
<th>Peri-op death</th>
<th>Reported complications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Qadan et al. (19)</td>
<td>2017</td>
<td>RCS</td>
<td>Open</td>
<td>53</td>
<td>369</td>
<td>Yes</td>
<td>NA</td>
<td>6</td>
<td>0</td>
<td>40: 27, grades 1–2; 13, grades 3–5</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Lap</td>
<td>21</td>
<td>160</td>
<td>Yes</td>
<td>14</td>
<td>5</td>
<td>0</td>
<td>5: grades 1–2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Robot</td>
<td>24</td>
<td>170</td>
<td>Yes</td>
<td>4</td>
<td>4</td>
<td>0</td>
<td>19: 15, grades 1–2; 4, grades 3–5</td>
</tr>
<tr>
<td>Dhir et al. (38)</td>
<td>2016</td>
<td>RCS</td>
<td>Robot</td>
<td>24</td>
<td>100 (IQR 20–200)</td>
<td>Yes</td>
<td>1</td>
<td>NR</td>
<td>8.4</td>
<td>1, multi-system organ failure; 1, bleeding; 3, catheter related; 1, catheter thrombosis; 1, partial catheter occlusion; 1, duodenal erosion</td>
</tr>
<tr>
<td>Franklin et al. (37)</td>
<td>2006</td>
<td>RCS</td>
<td>Lap</td>
<td>27</td>
<td>151 (range 20–300)</td>
<td>Yes</td>
<td>NR</td>
<td>8.4</td>
<td>2, ileus; 5, catheter related; 3, catheter thrombosis; 1, pulmonary embolus; 1, pump replacement</td>
<td></td>
</tr>
<tr>
<td>Cheng et al. (14)</td>
<td>2004</td>
<td>RCS</td>
<td>Lap</td>
<td>38</td>
<td>100 (range 25–1,200)</td>
<td>Yes</td>
<td>1</td>
<td>3</td>
<td>2, ileus; 5, catheter related; 3, catheter thrombosis; 1, pulmonary embolus; 1, pump replacement</td>
<td></td>
</tr>
<tr>
<td>Urbach et al. (36)</td>
<td>2001</td>
<td>RCS</td>
<td>Lap</td>
<td>8</td>
<td>50</td>
<td>Yes</td>
<td>1</td>
<td>4</td>
<td>0</td>
<td>1, pulmonary embolus</td>
</tr>
<tr>
<td>Allen et al. (26)</td>
<td>2005</td>
<td>RCS</td>
<td>Open</td>
<td>544</td>
<td>314 (SD 303)</td>
<td>yes</td>
<td>NA</td>
<td>5</td>
<td>120, non-catheter related; 49, catheter related; 33, arterial thrombosis; 16, extrahepatic perfusion</td>
<td></td>
</tr>
</tbody>
</table>

HAIP, hepatic arterial infusion pump; RCS, retrospective case series; EBL, estimated blood loss; Lap, laparoscopic; LOS, length of stay.

Overall complication rates are likewise similar, ranging from 12–21%. Operative duration, estimated blood loss (EBL), and complication rates are similar to open HAIP placement (Table 1).

Pre-operative evaluation

Pre-operative evaluation of all potential HAIP chemotherapy patients—regardless of operative approach—involves staging of disease; multidisciplinary discussion regarding the relative roles of systemic, regional, and local treatment modalities; and formal assessment of medical co-morbidities and surgical risk.

Clinical, endoscopic, and radiographic assessment [multidetector computed tomography (CT) scans without PET or MRI of the abdomen] are typically undertaken to assess the patient’s burden of disease and to rule out extrahepatic disease (EHD) (40). Provided they can adhere to scheduled refilling and necessary follow-up, individuals with good performance status, no obvious sites of EHD, and preserved liver function can be considered for HAIP chemotherapy.

Relative contraindications for placement of HAIP can be broadly classified into: (I) patient, (II) tumor, and (III) anatomic factors (Table 2).

In patients deemed appropriate candidates, pre-operative evaluation of the patient’s hepatic arterial anatomy is paramount. One of the main aims of pre-operative
assessment of hepatic arterial anatomy is to identify potential variant arterial anatomy. Such findings necessarily shape the operative plan and may alter the point of arterial cannulation for insertion of the HAIP. At present, CT angiography represents the gold standard for evaluation of hepatic arterial anatomy, providing high-resolution arterial anatomy without artifacts that is technically adequate for pre-operative planning.

### The approach to variant hepatic arterial anatomy

Variant hepatic arterial anatomy has been a well-known phenomenon since the 1960s with the publication of Nicholas Michels’s systematic study of variant anatomy based on 200 cadaveric subjects. Normal hepatic arterial anatomy is defined by: (I) common hepatic arterial origin off the celiac axis, and (II) when the gastroduodenal artery arises from the common hepatic artery (CHA) proximal to the bifurcation of left and right hepatic arteries (Figure 2). This is present only 50–60% of the time. The most common variants involve the presence of “replaced” or “accessory” right (15%) or left (11%) hepatic arterial anatomy (42).

Early approaches to variant hepatic arterial vasculature in HAIP surgery were highly variable. Techniques ranged from ligation of accessory vessels, technically complex anastomoses between gastroduodenal artery (GDA) and replaced or accessory vessels, or retrograde catheterization via the splenic artery (20,30,43). The use of dual lumen pump systems was also utilized; however, extended operative time and uneven hepatic perfusion led to its disuse (44).

The surgical management of variant anatomy was greatly simplified with the concept of ligation of all accessory and replaced vessels. Rayner and colleagues demonstrated the safety of such an approach in 15 patients, with complete bi-lobar hepatic perfusion documented in 87% of patients (27). While early experience with a simplified approach demonstrated higher rates of technical complications in patients with variant anatomy (23), modern series have shown that outcomes are equivalent (44).

In the largest review of variant anatomy, Allen and colleagues examined 265 consecutive HAIP placements over a five-year period to determine whether patients with variant anatomy experienced increased rates of catheter-related complications and to determine optimal technique. They reported an overall morbidity rate of 20% and a 12% rate of pump-related complications. Patients with variant anatomy were more likely to experience pump-related complications if a vessel other than the GDA was cannulated (28% vs. 4%, P<0.001) and if they had multiple variant vessels versus only a single variant vessel (23% vs. 6%, P<0.05) (30). A follow-up study examining technical complications in 544 patients undergoing HAIP insertion reported similar overall mortality and technical

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**Table 2** Relative contraindications to HAIP chemotherapy. Adapted from (19) with permission

<table>
<thead>
<tr>
<th>Patient</th>
<th>Tumor</th>
<th>Anatomic factors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Poor performance status (Karnofsky &lt;60%)</td>
<td>Extrahepatic disease</td>
<td>Portal vein thrombosis</td>
</tr>
<tr>
<td>Liver insufficiency (total bilirubin &gt;1.5 mg/dL)</td>
<td>Extensive hepatic metastasis &gt;60%</td>
<td></td>
</tr>
<tr>
<td>Irreversible coagulopathy</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current infection</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inability to follow up for routine follow-up</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

HAIP, hepatic arterial infusion pump.
complication rates. Of note, on multivariable analysis, only non-GDA insertion of the catheter and surgeon experience (<25 cases) were associated with complications (26).

These results helped standardize the management of variant hepatic arterial anatomy by establishing the safety and efficacy of ligating all variant vessels and underscored the importance of GDA insertion of HAIP catheters whenever possible. The techniques standardized in open surgery serve as the foundation for MIS HAIP surgery.

The procedure

Once a patient is determined to potentially benefit from HAIP chemotherapy, is found to be a reasonable surgical candidate, and found to have suitable hepatic arterial anatomy on pre-operative imaging, the patient is taken to the operating room.

The patient is placed in supine position with a footboard, general anesthesia is induced, and the patient is prepped and draped in sterile fashion. Selection of the pump site is performed early in the operation and marked with a 7–8 cm transverse line—typically on the left side—2–3 finger breadths below the costal margin and above the anterior superior iliac spine. When patients are morbidly obese, the pocket is created above the costal margin allowing the pump to rest on the left chest wall to minimize pump migration and flipping. Marking of the pump pocket at this stage of the operation avoids inadvertent placement of trocars through the eventual pump pocket (Figure 3).

After obtaining intraperitoneal access and performing diagnostic laparoscopy to rule out EHD, the patient is placed into steep reverse Trendelenburg. Cholecystectomy is then performed to prevent chemotherapy induced cholecystitis. In cases where hepatic arterial anatomy is normal (the most common scenario), the common hepatic (CHA), GDA, proper hepatic artery (PHA), right hepatic artery (RHA), and left hepatic artery (LHA) are identified. When hepatic arterial anatomy is variant, the accessory or replaced artery is ligated (Figure 4). Sharp dissection with hook cautery is initiated 2 cm proximal to the takeoff of the GDA. The distal CHA, GDA, and proper hepatic artery (PHA) are circumferentially mobilized, and the right gastric artery is ligated. All accessory and collateral branches should be ligated to prevent extrahepatic perfusion. The LHA should be cleared up to first order branches, and the RHA dissected and cleared as far as is safe and feasible, as systematic review of cases of extrahepatic perfusion revealed that collaterals off the RHA account for the overwhelming majority of cases (45).

In nearly all instances, the pump catheter can be inserted in the GDA, which is the preferred vessel for pump catheter insertion. Doing so maximizes bi-lobar perfusion, minimizes turbulent arterial flow, and has been associated with long-term patency of the pump (30).

Pneumoperitoneum is released and a subcutaneous pump pocket is created at the site marked earlier in the case. Subcutaneous flaps are raised as needed to accommodate the pump, and the pocket should remain superficial to the fascia to facilitate needle access at subsequent oncology visits. Only after the GDA is ready for cannulation and the pump pocket created is the pump device brought into the surgical field. It is first filled with heparinized saline and the catheter copiously flushed with heparinized saline. A small rent in the fascia is made in the center of the pocket and

Figure 3 Port site placement for laparoscopic (A) and robotic (B) approaches to HAIP placement. Reproduced from (39) with permission. HAIP, hepatic arterial infusion pump.
the catheter is fed into the abdominal cavity. The pump is placed atop the remaining extraperitoneal length of catheter and, after ensuring the catheter has a straight unobstructed path into the peritoneal cavity, the pump is secured to the fascia with stay sutures.

Pneumoperitoneum is re-established and attention is then turned to cannulating and securing the catheter into the GDA. The CHA, GDA, and PHA are coated with papaverine injected through the abdominal wall with a spinal needle. Control of the distal GDA is obtained by ligation with non-absorbable suture and a surgical clip. Proximal control is achieved with small, straight bulldog clamps inserted through a 12 mm port, which are then placed on the CHA and PHA. Alternatively, a single curved bulldog clamp can be placed on the CHA/PHA to exclude flow to the GDA and allow for placement of the catheter with the tip at the junction of the GDA with the CHA/PHA. A non-absorbable suture is placed posterior to the GDA and will be used to secure the catheter once introduced.

An arteriotomy is made with a No. 11 blade in the anterior wall of the GDA and a plastic introducer is used to facilitate introduction of the catheter into the GDA to avoid dissection. The previously placed suture is tied to secure the catheter in place. An additional two ties are then placed around the GDA to avoid catheter migration (Figure 4).

Once the catheter is in place, adequacy of hepatic perfusion is assessed by injecting methylene blue into the pump. The liver is grossly assessed for uniform color change immediately post injection and after several minutes, making note of the presence of any extrahepatic perfusion.

After ensuring the absence of kinking in the catheter and confirming easy flushing of the pump with heparinized saline, ports are removed under direct visualization and pneumoperitoneum is released. All port sites larger than 8 mm are closed at the level of the fascia. Finally, all port sites closed with absorbable suture.

### Confirming adequacy of hepatic perfusion

Prior to the first dose of HAIP chemotherapy, appropriate hepatic perfusion via pump is evaluated by means of a radionuclide pump-flow study. Radiolabeled sulfur-colloid (SC) is injected intravenously and technetium-labeled macroaggregated albumin (MAA) via the pump. Perfusion scans are obtained and the two images overlaid: SC representing intrahepatic perfusion and MAA the region perfused by the pump. If the fused study is normal, HAI chemotherapy is initiated, usually 1–4 weeks post-operatively (46).

Incomplete hepatic perfusion occurs in 2% of cases and typically results from failure to ligate an aberrant vessel or failure of cross lobar collateralization. These patients are followed with repeat perfusion scans in 2–4 weeks, and nearly all have resolution of incomplete perfusion (30). In instances where this is not the case, angiography with embolization of any remaining aberrant vessels can be performed.

Extrahepatic perfusion may be detected on the post-operative perfusion scan or based on clinical presentation during chemotherapy. The MAA-labeled scan will demonstrate an area of perfusion beyond the territory delineated in the SC scan, indicating the potential for perfusion of the duodenum, pancreas, or stomach. In cases not detected prior to initiation of HAIP chemotherapy,
severe epigastric pain or diarrhea with infusion is typical. The underlying etiology of such symptoms is pancreatitis or ulcers. Up to 80% of pumps with extrahepatic perfusion can be salvaged with endovascular embolization procedures (47).

**Post-operative/technical complications**

Large single-institution case series as well as several meta-analyses have been published characterizing the complication profile of HAIP surgery in detail. Pump-related complication rates in these modern analyses range from 10–20%. The most common complications are arterial thrombosis (6–7%), perfusion abnormalities (5%), catheter dislodgment (3–5%), and pump pocket infection (1–3%) (26,29,48,49). In the most comprehensive series evaluating technical complications in 544 consecutive HAIP placements (open and MIS approaches), the overall salvage rate for complications was 45% and the complication-related HAIP failure rate was 12% (26).

Complications from MIS HAIP insertion are similar to the open technique (Table 1). Where published, rates of catheter-related complications in laparoscopic HAIP range from 11–13% (14,37,38). Overall, the majority of complications are mild, with 60-100% of complications reported as grade 1–2, with few grade 3–5 complications reported (36,38,39). To date, there have been no reported differences in rates of complications.

Of relevance in evaluating MIS approaches to HAIP insertion is the conversion rate to open. Rates of conversion range from 3–67% (14,36-39) in laparoscopic cases. Conversion during robotic operations is similar, with reported rates of 4–17% (38,39). Of note, in the only study systematically comparing open, laparoscopic, and robotic approaches to HAIP insertion, conversion rates of robotic approaches were significantly lower than laparoscopic (17% vs. 67%, P<0.01), with no differences in overall complication rate or length of stay.

**Chemotherapy/drug-related toxicity**

The adverse event profile of HAIP chemotherapy is unique in that patients experience both technical- and chemotherapy-related toxicities. FUDR and 5FU are the most commonly used regional chemotherapeutic agents in HAIP chemotherapy. With almost entire first pass liver extraction, systemic toxicity from FUDR is limited and adverse events are local. Chemical hepatitis is the most common adverse event in FUDR-based regimens (34%), and typically presents as liver function test abnormalities (29).

By contrast, 5FU extraction by the liver is modest, ranging from 20–40%, and common adverse events relate to systemic toxicities. The most commonly reported adverse events are gastrointestinal, with nausea, vomiting, and diarrhea reported in 40% of patients (29). For these patients, the dosing schedule is modified and treatment resumes with normalization of lab work, as outlined in published dosing guidelines (50).

Biliary toxicity is a related but distinct complication of HAIP chemotherapy and occurs in 5–30% of patients (29,51). Unlike the portal venous-based perfusion of hepatocytes, the biliary tree depends on arterial perfusion. HAIP chemotherapy, therefore, places patients at significant potential risk for biliary toxicity and injury. The use of dexamethasone with FUDR has been shown to mitigate the potential biliary toxicity of FUDR HAIP chemotherapy and is a standard part of current FUDR-based regimens (52).

The most significant and serious manifestation of biliary toxicity is biliary sclerosis. Published incident rates of biliary sclerosis in the setting of HAIP chemotherapy range from 1–26% and are more prevalent in the adjuvant setting, with FUDR, and with co-administration of the anti-VEGF drug bevacizumab (53). Based on published results, the etiology is thought to be related to ischemic and drug toxicity.

Early recognition of biliary toxicity and prompt intervention in cases of biliary sclerosis are critical in preventing long-term harm. Should dose modification or cessation (temporary or definitive) fail to resolve laboratory abnormalities, radiographic studies (CT or MRI) should be obtained to discover potential strictures, which may be treated with endoscopic retrograde cholangiopancreatographic stent/dilation. In patients with isolated strictures from biliary sclerosis who can be stented or dilated, survival is not compromised (51).

**Summary**

Since its initial development, HAIP chemotherapy has emerged as an important therapeutic modality in the treatment of colorectal cancer liver metastasis. With standardized patient selection, operative technique, and familiarity with post-operative pump-related complications, published outcomes have improved and are more consistent. Laparoscopic HAIP surgery has comparable safety and efficacy to standard open approaches.

The role of laparoscopic HAIP in an era of robotic surgery remains an outstanding question. Two published
series report comparable outcomes for robotic HAIP placement (38,39). Rates of conversion, however, do appear to be lower in robotic approaches. It does appear to be a procedure that is ideal for the robotic platform, given the small operative field that requires very fine vascular dissection. As an operative platform, the articulating, wristed movement of robotic instruments is particularly well-suited for cannulating and securing the arterial catheter. At our own institution, we no longer routinely perform laparoscopic HAIP placement, favoring the robotic approach (Figure 4). On the other hand, laparoscopic HAIP surgery remains an important MIS surgical technique in the setting of concomitant laparoscopic colorectal or liver surgery. Regardless of approach, surgeon and institutional experience with HAIP chemotherapy remain critical factors for ensuring optimal outcomes and rescuing patients from pump-related complications.

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Footnote

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