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**Anticoagulation and antiplatelets as prophylaxis for hepatic artery thrombosis after liver transplantation**

Algarni AA *et al*. Anticoagulation and antiplatelets for hepatic artery thrombosis

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**Abstract**

Hepatic artery thrombosis (HAT) is the most serious vascular complication after liver transplantation (LT). Multiple risk factors have been identified to impact its development. Changes in haemostasis associated with end stage liver disease and the disturbance of the coagulation and anticoagulation cascades play an important role in development of this lethal complication. Early recognition and therapeutic intervention is mandatory to avoid its consequences. Pharmacological prophylaxis, by the use of antiplatelet or anticoagulant agents, is an important tool to reduce its incidence and prevent graft loss. Only a few studies have shown a clear benefit of antiplatelet agents in reducing HAT occurrence, however, these studies are limited by being retrospective and by inhomogeneous populations. The use of anticoagulants such as heparin is associated with an improvement in the outcomes mainly when used for a high-risk patients like living related liver recipients. The major concern when using these agents is the tendency to increase bleeding complications in a setting of already unstable haemostasis. Hence, monitoring of their administration and careful selection of patients to be treated are of great importance. Well-designed clinical studies are still needed to further explore their effects and to formulate proper protocols that can be implemented safely.

**Key** **words**: Liver transplantation; Hepatic artery thrombosis; Haemostasis; Anticoagulation; Antiplatelets; Heparin

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**Core tip:** Hepatic artery thrombosis (HAT) is the most serious vascular complication after liver transplantation. Changes in haemostasis associated with liver disease play a role in its development. Pharmacological prophylaxis may reduce its incidence and prevent graft loss. Few studies have shown a clear benefit of antiplatelets in reducing HAT occurrence, however, these studies have several limitations. The use of anticoagulants showed an improvement in the outcomes when used for high-risk patients. Their major concern is the tendency to increase bleeding complication. Hence, monitoring of their administration and careful selection of patients to be treated are of great importance.

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**INTRODUCTION**

Vascular complications after liver transplantation (LT) are common despite the progressive improvement and innovations in anastomotic vascular techniques. They frequently result in hepatic failure and graft loss and thus the need for re-transplantation. Hepatic artery thrombosis (HAT) after LT remains the most serious vascular complication with an overall incidence varies from 2% to 9%[1]. It is usually associated with around 50% rate of mortality or re-transplantation[2].

The mechanism of HAT development is not fully understood. It is believed to be multifactorial including both operative and non-operative factors (Table 1) [3,4]. All these factors are involved in causing recipient hypercoagulability and to disturb the unstable rebalanced haemostasis that usually occurs in liver disease patients leading to a rise in the incidence of HAT[5,6].

An early diagnosis with immediate treatment is the key to proper management of HAT. Therapeutic options for managing HAT include either revascularization of the transplanted graft or re transplantation[7]. Revascularization can be achieved through arterial reconstruction, surgical thrombectomy, or radiologically guided thrombolysis. Shortage of organs and poor patients conditions may lead physicians to consider revascularization, which can achieve good success rates[8]. Re-transplantation is still the gold standard treatment for HAT but is confounded by both graft availability and the patient’s general condition.

Prophylaxis against HAT in the early postoperative period by usage of antiplatelet or anticoagulant agents is thought to be hazardous by many surgeons because of the risk of postoperative bleeding. However, postoperative pharmacological prophylaxis can reduce the risk of arterial anastomosis thrombosis when used in vascular surgical procedures[9]. The adoption of this in LT could help in reducing the incidence of HAT.

**HAEMOSTASIS IN END STAGE LIVER DISEASE**

In liver transplant recipient with cirrhosis, considerable changes in the haemostatic system are often found[10]. These changes include thrombocytopenia, platelet functional defects, decreased circulating levels of coagulative factors and decreased fibrinolytic proteins. Although routine diagnostic tests of haemostasis are commonly abnormal, interpretation of these tests is more difficult in patients with a complex haemostatic disorder as in cirrhotic patients. It is now well established that patients with cirrhosis and abnormal routine coagulation tests might not have an increased bleeding tendency and that thrombotic complications may still occur[11]. This happens through a state of a rebalanced haemostasis where the changes in the pro-haemostatic pathways are associated with changes in the anti-haemostatic pathways (Figure 1)[12]. The changes that promote bleeding include thrombocytopenia, platelet function defects, enhanced production of nitric oxide and prostacyclin, low levels of factors II, V, VII, IX, X, and XI, vitamin K deficiency, dysfibrinogenemia, low levels of a2-antiplasmin, factor XIII and thrombin-activatable fibrinolysis inhibitor and elevated tissue plasminogen activator level. Those alterations are encountered by others that counteract bleeding such as: elevated levels of von Willebrand factor (VWF), decreased levels of ADAMTS-13 (von Willebrand factor cleaving protease), elevated levels of factor VIII, decreased levels of protein C, protein S, antithrombin, a2-macroglobulin and heparin cofactor II and low levels of plasminogen[5,12].

The resultant effect of all these changes is a rebalanced haemostasis. Although it is a functional system, it is unstable compared to the haemostatic balance in healthy individuals, explaining the potential occurrence of both bleeding and thrombotic complications in patients with cirrhosis[12].

**CHANGES IN THE HAEMOSTATIC BALANCE AFTER LT**

In the first days after LT, multiple transplantation-related triggers initiate pro-coagulation, anticoagulation and pro-fibrinolytic cascades. Theses mechanisms might not be balanced properly leading to thrombosis in the area of anastomosis (Figure 1). The triggers include substantial surgical damage, stasis as a result of clamping of major vessels, release of activators from the donor liver and systemic inflammatory responses. Other factors are shown in Table 1[6].

Intraoperatively, the graft endothelium gets activated due to cold ischemia and reperfusion effects. As soon as the reperfusion happens, platelets adhere to the sinusoidal endothelium, which contributes to endothelial cell apoptosis causing more ischemia/reperfusion damage. This process will probably expose additional pro-coagulant triggers causing further platelet activation and initiation of coagulation[13]. As long as these pro-coagulant triggers go on, thrombosis can easily occur. However, the activated fibrinolytic system removes the clots immediately as they are formed. When this balance is disturbed, a state of hypercoagulability arises and a clinically evident thrombus might form in the anastomotic areas[14].

In addition, an acquired hypercoagulability may increase the risk for immediate graft thrombosis. It can be due to transplanting a liver from a donor with Factor V Leiden mutation or infections like cytomegalovirus (CMV). Perioperative haemostatic agent such as fresh frozen plasma, platelets, recombinant factor VIIa, and anti-fibrinolytics such as aprotinin and aminocaproic acid plays role as well (Table 2)[12].

Mechanisms of late thrombosis of the graft vascular anastomosis or even systemic thrombosis are less obvious. It could be because of pro-thrombotic side effect of immunosuppressant or as consequences of medical comorbidities like hypertension, hypercholesterolemia and diabetes mellitus. It could happen due to viral infection as well especially CMV[15].

Liver diseases are classically portrayed as having a substantial bleeding tendency. For this reasons, the postoperative use of anticoagulants or antiplatelets has been restricted, in order to limit bleeding complications. However, recent data showed that the bleeding diathesis of patients with liver disease is not only a result of poor haemostasis[16]. Other factors may play a significant role and augment bleeding tendency. Portal hypertension is an example of that. The minimal blood loss and the decreasing requirements of coagulation products in nowadays transplant support this concept. Moreover, improvements in surgical techniques and a restrictive fluid and transfusion protocols has led to a reduction in perioperative transfusion requirements[17]. Considering this fact, the limited use of anticoagulants or antiplatelets in the postoperative period is reconsidered.

**USE OF ANTIPLATELETS IN LT**

In the immediate periods after LT, the constant platelet activation and aggregation result in development of thromboxane leading to fibrinogen activation, which contribute to arterial thrombosis and graft failure[18]. Because of this pivotal role of platelets, antiplatelets therapy by inhibiting this platelet-activation may lead to reduction in the incidence of such complication. Aspirin, which is the most frequent used antiplatelet agents, interferes with platelet aggregation, which in turn leads to an endothelial cell–mediated inhibition of the coagulation cascade. This happens through irreversible inhibition of cyclooxygenase1 and hence, inhibition of thromboxane generation[19].

Very few studies focused on the efficacy and safety of antiplatelet therapy in patients after LT. In 1997, in a retrospective study, Wolf *et al*[20] found no significant difference in the incidence of early HAT (3.7%) *vs* (4.0%) between patients who had a prophylaxis aspirin (354 of 529 patients) and those who did not. Bleeding events occurred in 1.1% of patients treated with aspirin compared to 0.6% of the control group. The effect of aspirin prophylaxis on late HAT was not investigated[20].

Vivarelli *et al*[21] reported a single centre retrospective study where they looked to the effect of long-term aspirin administration (100 mg) on the incidence of late HAT in a large number of patients. In this study, they categorize the patients into high and low risk groups based on their finding on a previous study. One of two independent risk factors was used to categorize the patients into either group: grafts retrieved from donors who died from a cerebrovascular accident or the use of an arterial iliac conduit. Late HAT incidence was 3.6% (12 of 338) of the high-risk patients who were not given aspirin, whilst it was seen in only 0.6% (1 of 160) of the high-risk patients who received aspirin prophylaxis. The relative risk reduction was 82%. In the low-risk group, 1 of 330 patients developed late HAT and he was in the non-prophylaxis arm (254 patients). Out of 236 patients who received aspirin, there was no recorded episode of any bleeding complications throughout the follow up period (median of 1704 d). They could not assess the effect of antiplatelets on early HAT due to their inability to start patients immediately on aspirin with a known impaired coagulative function and a high risk of bleeding. This obvious benefit of aspirin on late HAT occurrence should be considered specifically in patients with an absence of bleeding complications.

Shay *et al*[22] showed that aspirin prophylaxis is safe and effective in decreasing early HAT in adult recipients. The incidence of overall HAT was found to be significantly higher at 4.9% in the control group *vs* 3.0% in the treated group. Early HAT incidence dropped from 3.9% in the control group to 1.8% with aspirin prophylaxis. Also the incidence of early HAT causing graft loss decreased significantly from 3.6% to 0% with the use of early aspirin prophylaxis. There was no difference in bleeding complications between the two groups. The main difference between these studies is the dosage and the time of initiation of therapy. Shay *et al*[22] used a higher dosage at 325 mg/d, initiated immediately after surgery with no evidence of significant bleeding.

There are several limitations to these studies that should be considered. First, not all of them include paediatric patients, where the incidence of HAT is higher and survival is better. Secondly, they were all retrospective studies with a higher risk of bias. Also the compared groups were mostly from different time periods where surgical techniques, donor selection, and postoperative management are different. However, these studies showed evidence of benefit from the use of antiplatelet agents as a prophylaxis for early and late HAT without increasing the risk of bleeding or other surgical complications. The current usage of antiplatelet agents in treatment of cardiovascular and peripheral vascular disease might support this conclusion. A further well-designed randomized study to explore this field would be appropriate.

**USE OF ANTICOAGULANTS IN LT**

The plasma concentration of coagulant and anticoagulant proteins is disturbed after LT. This imbalance leads to hypercoagulability condition that contributes to vascular thrombosis and possible loss of the graft[14]. The reducing level of antithrombin III and protein C are believed to be responsible for the development of this hypercoagulability state[23]. For this reason, supplementation of protein C and augmenting the effect of antithrombin III could help to overcome this haemostatic disturbance.

Fresh frozen plasma (FFP) is usually the source of protein C along with other clotting factors. Hashikura *et al*[24] reported that infusing FFP was helpful in reducing incidence of HAT and maintaining coagulation haemostasis. In contrast, Mazzaferro *et al*[25] found a higher incidence of HAT in paediatric liver recipient who received more FFP compared to those who did not receive FFP intraoperatively. Hatano *et al*[26] found a similar result where the occurrence of HAT was associated with a higher intraoperative infusion of FFP. Although the use of FFP might be associated with an increased risk of HAT, its usage is still warranted to maintain haemostasis and to reduce bleeding complication. Excessive use of FFP should be avoided as it might induce a state of hypercoagulability and could result in graft thrombosis.

Heparin appears to be the method of choice to anticoagulate liver transplant recipients. Most liver transplant centres have developed their own protocols for heparin infusions and the monitoring of its activity. It is important to note that the rationales of these protocols remain unclear, as they are designed based on empirical rules. The Shinshu group demonstrated that intensive anticoagulation should be established in living related liver transplantation (LDLT) in a paediatric Population[24]. Sugawara *et al*[27] showed a similar result in the setting of living donor LT. The reports on usage of heparin in cadaveric liver transplant are very limited to date. The anti-coagulatory effect of the unfractionated heparin (UFH) is known to be through its antithrombin III activity. The low molecular weight heparin (LMWH) selectively inhibit clotting factor Xa and to a lesser extent augment antithrombin III activity[28].

Bleeding complication can occur with the unmonitored use of heparin. Kaneko *et al*[28] reported that 9% of their living related liver recipients who used UFH developed haemorrhagic complications that required surgical treatment. In contrast to UFH, using LMWH is believed to be useful to reduce haemorrhagic complications due to its selective inhibition of coagulation factor Xa and because of its reduced ability to bind to endogenous plasma proteins such as platelet factor 4 and von Willebrand factor[29]. In addition, LMWH lessen liver damage in ischemia-reperfusion injury and hyperperfusion.[30] For these reasons, LMWH is more advantageous than UFH for intraoperative and postoperative anticoagulant therapy in LT. The major problem for the prophylactic use of LMWH is determining the optimal dose and monitoring serum factor Xa activity. The route of administration of LMWH depends on the indication. In patients with a high bleeding tendency, such as liver transplant recipients, an adjustable continuous infusion may be recommended to avoid increased plasma levels and to cope with continuous pathophysiological changes in the coagulation cascade[31].

Antithrombin is plasma glycoprotein synthesized in the liver and plays a major role in the coagulation after LT. The level of antithrombin III drops after transplant and remains low for up to two weeks. Replacing this drop might help to gain an adequate anticoagulation effect[32]. Kaneko *et al*[32] in his pilot study found that the combined use of AT and UFH might reduce fibrin degradation product D-dimer (FDP-DD) levels and prevent a postoperative drop in the platelet count. Taniai *et al*[33] concluded in his study that adding antithrombin to heparin could be the best approach. Further research should be done to prove its benefit and safety in LT[33.

Monitoring of anticoagulation after LT assures an adequate level without an increase in the bleeding complications. Activated clotting time (ACT) is the most commonly used monitoring method. Linkins *et al*[34] showed that UFH is more sensitive than LMWH in regards to ACT level. This indicates that the ACT cannot monitor the activity of factor Xa but is still a valuable tool in monitoring the anticoagulant effects of LMWH. The lower level of antithrombin activity in the LMWH compared to the anti- Xa activity could explain this lower sensitivity[29]. It has been shown that ACT can monitor the anticoaguatory effect of LMWH in coronary intervention procedures[35]. Uchikawa *et al*[31] showed that ACT measurement is a simple, reliable method for bedside monitoring of LMWH anticoagulant effects for LDLT. It is assumed that ACT level should be kept within the normal range in order to prevent haemorrhagic complication. Thus, the anticipated value of ACT is between 140 and 150 s, which is the upper limit of the normal range. However, Kaneko *et al*[32] tested the ACT level by measuring Plasminalpha2 plasmin inhibitor complex, thrombin-antithrombin III complex and fibrin degradation product pre and postoperatively with measurement of complete blood count, ACT, activated partial thromboplastin time and prothrombin time international normalized ratio for two weeks after surgery. They conclude that frequent monitoring of ACT is necessary to keep the ACT level in the target range in the first postoperative week. Because of hyper-fibrinolytic condition and the high rate of haemorrhagic complications after the first week, the dose of heparin should be adjusted to maintain lower ACT levels during this period.

In contrast to UFH, LMWH are cleared by renal route. Hence in patients with impaired renal functions, as commonly seen following LT, monitoring and adjustment of the dose according to the degree of renal injury are required. Observational studies have shown more bleeding complications in renal impaired patients[36]. Prophylactic LMWH doesn’t appear to increase the bleeding tendency and therefore might not need monitoring or adjustment[37]. Therapeutic LMWH bio accumulates and cause more bleeding if left unadjusted[38]. This bioaccumulation is more evident when creatinine clearance is less than 30 mL/min[38]. No clear guidelines are available for dose adjustment and further clinical and pharmacological studies are required for dosage guidance[39].

It is clear that a further study looking to the appropriate protocols of anticoagulation and the proper monitoring tools is needed. The rarity of studies and the empirical assumption of tailoring the protocols make drawing a conclusion difficult.

**CONCLUSION**

Pharmacological prophylaxis is probably beneficial in reducing the incidence of HAT. It is relatively safe if used carefully with a continuous monitoring and adjustment. Proper protocols need to be developed based on proper well-designed clinical studies.

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**Figure 1 Changes in haemostasis in normal situation, liver disease and after liver transplantation.**

**Table 1 Risk factors of hepatic artery thrombosis**

|  |  |
| --- | --- |
| Operative factors | Non operative factors |
| Surgical technique  Number of anastomosis,  Use of conduits  Vessels kinking  Small vessels size | Donor age of more than 60 yr  Long cold ischemia time  Preservation damage  Lack of blood group compatibility  Cytomegalovirus-positive donor status  Hypercoagulable recipient status  Recipient cigarette smoking  Rejection  Primary sclerosing cholangitis |

**Table 2 Factors involved in development of hypercoagulability after liver transplantation**

|  |  |
| --- | --- |
| Endogenous | Acquired |
| Substantial surgical damage  Stasis as a result of clamping of major vessels  Release of activators from the donor liver  Systemic inflammatory responses  The quality of the graft  The length of surgery  The technique of graft preservation  Cold ischemia/reperfusion effect | Donor with Factor V Leiden mutation  Infections  Viral infection like cytomegalovirus  Perioperative haemostatic agent:  Fresh frozen plasma  Platelets  Recombinant factor VIIa  Anti-fibrinolytics:  Aprotinin Aminocaproic acid |