Primary graft dysfunction of the liver: definitions, diagnostic criteria and risk factors

Disfunção primária do enxerto hepático: definições, critérios diagnósticos e fatores de risco

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ABSTRACT

Primary graft dysfunction is a multifactorial syndrome with great impact on liver transplantation outcomes. This review article was based on studies published between January 1980 and June 2015 and retrieved from PubMed database using the following search terms: “primary graft dysfunction”, “early allograft dysfunction”, “primary non-function” and “liver transplantation”. Graft dysfunction describes different grades of graft ischemia-reperfusion injury and can manifest as early allograft dysfunction or primary graft non-function, its most severe form. Donor-, surgery- and recipient-related factors have been associated with this syndrome. Primary graft dysfunction definition, diagnostic criteria and risk factors differ between studies.

Keywords: Transplantation; Primary graft dysfunction/diagnosis; Liver transplantation; Donor selection; Tissue and organ procurement; Risk factors

INTRODUCTION

Liver transplantation is the only choice for patients suffering from end-stage liver disease. Primary graft dysfunction (PGD), one of the major complications of liver transplantation, is associated with higher morbidity in the early post-transplant period and may compromise graft survival.(1-3)

Primary graft dysfunction is related to ischemia-reperfusion injury(1,2) to the transplanted organ and may be subdivided into early allograft dysfunction (EAD) and primary nonfunction (PNF), its more severe manifestation.(3,4) Primary graft dysfunction occurs in 38.7% of liver transplants,(3) with PNF incidence ranging from 0.9% to 8.5%.(1,4)

Several studies attempted to establish a definition of PGD and determine related diagnostic criteria and risk factors. Along with surgery-related factors, clinical and epidemiological characteristics of donors and recipients, which differ between European, North and Latin American countries, were associated with PGD.(1)

The lack of universally accepted PGD definition and diagnostic criteria and the wide variation in risk factors described in literature preclude the adoption of standardized procedures across organ transplant centers. This study set out to compile definitions, diagnostic criteria and risk factors for this significant liver transplant complication in its different manifestations (i.e., EAD and PNF).

This review was based on articles published from January 1980 to June 2015; articles were retrieved from PubMed database using the following search terms: “primary graft dysfunction”, “early allograft dysfunction”,

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“primary non-function”, and “liver transplantation”. Original and review articles from Europe, United States, Canada and Asia were included. Latin American studies with sufficient statistical power were lacking. Studies involving pediatric populations, live donors, double transplants or experimental models were excluded.

DEFINITION

Early allograft dysfunction and PNF describe different degrees of graft functional impairment arising during the intraoperative period, and therefore reflect different stages of the same syndrome. However, clear definitions of EAD and PNF are lacking in medical literature. While EAD refers to a set of transient clinical and laboratory changes reflecting post-transplant graft malfunction, PNF is a more severe, catastrophic event characterized by liver necrosis, rapid increase in serum transaminase, coagulopathy, increased lactate levels, hemodynamic instability, hypoglycemia and respiratory and renal failure. Different from EAD, a transient condition with potential graft function recovery, PNF leads to graft failure requiring emergency retransplantation.

EARLY ALLOGRAFT DYSFUNCTION

Variables and cut-off points employed in EAD definition and diagnosis differ between studies. Laboratory changes reflecting liver metabolism and function in the first few days after transplantation, such as serum alanine aminotransferase (ALT) and aspartate aminotransferase (AST) levels, prothrombin time (PT), international normalized ratio (INR), and serum lactate and ammonia levels, are the most commonly used parameters. Early allograft dysfunction is known to be associated with lower graft and patient survival rates, longer stay at intensive care unit (ICU) and increased postoperative morbidity and mortality.

Previous studies taking time to dysfunction diagnosis as the driving factor for implementation of patient support therapy established the third postoperative day as the cut-off time for EAD. Nanashima et al. used serum AST or ALT levels over 1,500IU/L in two consecutive tests, within 72 hours of surgery, to define EAD. According to Dhillon et al., EAD diagnosis should be based on transaminase scores [(AST+ALT)/2] on the second postoperative day; values <285IU/L and between 285 and 986IU/L would reflect good and intermediate graft function, respectively, while values above 986IU/L would define EAD.

Graft function scores were also developed. González et al. described early graft function using a point scoring system based on serum ALT levels, bile production and prothrombin time within the first 72 hours post-transplant; laboratory test results scored 3-4, 5-6 and 7-9 would reflect good graft function, moderate graft function and EAD, respectively. Heise et al. created a survival-based classification system using serum AST and ALT levels, bile production and prothrombin time within 1, 3, 7 and 14 days of surgery. Postoperative laboratory values are known to change over time; therefore, in this system, patients initially diagnosed with EAD can be reallocated to different subgroups after the first 72 hours of surgery based on laboratory test result fluctuations. Hence, graft function is graded Berlin A (4-5 points; good graft function), Berlin B (6 points; intermediate graft function) or Berlin C (7-8 points; EAD).

The widely used concept proposed by Ploeg et al. is based on hepatocyte injury and liver synthesis capacity assessment (serum transaminase levels and PT, respectively). In this unicenter study, EAD was defined as serum AST levels >2,000IU/L, PT over 16 seconds and serum ammonia levels >50μmol/L between second and 17th day post-surgery.

In the first definition of EAD derived from a multicenter study, post-transplantation liver function, donor- and graft-related factors, and recipient pre-transplantation status were described as risk factors. Serum bilirubin levels, PT and the occurrence of hepatic encephalopathy between post-transplant days 2 and 7 were taken into account. Increased PT and serum bilirubin levels prior to transplantation, recipient hospitalization due to clinical condition deterioration, donors aged 50 years or over or hospitalized for more than 3 days, and ischemic time over 15 hours were also associated with EAD.

A definition of EAD based on graft function variables (injury, coagulopathy and cholestasis) up to postoperative day 7 was proposed by Olthoff and a liver transplant expert group. Rather than deriving novel variable cut-offs, the study was aimed at validation of previous definitions; objective variables that could be easily obtained over the course of patient follow-up were selected. In that study, EAD diagnosis was established when one or more of the following variables were present: serum bilirubin levels ≥10mg/dL; INR ≥1.6 on postoperative day 7; and serum AST or ALT levels >2,000IU/L within the first 7 days of surgery.

Friedman et al. hypothesized that serum inflammatory protein expression in response to surgery might reflect
EAD;\(^{18}\) hence, inflammatory cytokine profiles could be used to diagnose the condition. Multivariate analysis revealed that patients suffering from EAD had higher interleukin (IL) 6 and IL-2R levels prior to than after surgery; in contrast, MCP-1 (CCL2), IL-8 (CXCL8) and RANTES (CCL5) levels were higher immediately after than before surgery. These findings suggest potential association of several biomarkers with EAD.

Croome et al.\(^{19}\) employed MELD (Model for End-Stage Liver Disease) score derived donor data to predict EAD. The value of the MELD score in predicting EAD within the first week, or mortality within 90 days post-transplant, was emphasized by Wagener et al.\(^{20}\)

A study involving 66 transplanted patients\(^{21}\) suggested peak serum AST investigation should be performed within the 5-11-hour post-reperfusion time window, given AST levels are known to drop by half within the first postoperative day.\(^{21}\) This finding is highly significant, bearing in mind the lack of a consensus among organ transplant centers as to the best time point for the first serum AST level determination. Direct liver function assessment tests, such as indocyanine green clearance, can be employed for early graft dysfunction diagnosis.\(^{22}\) The application of the liver function test LiMAx (maximal liver function capacity)\(^{22,24}\) 24 hours following transplant was reported by Lock et al.\(^{22}\)

This test reflects real-time liver function and was shown to be successful in graft function assessment immediately after surgery.\(^{22}\) The main definitions of EAD are given on chart 1.

### Chart 1. Main definitions of early allograft dysfunction

<table>
<thead>
<tr>
<th>Author</th>
<th>Definition</th>
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<tbody>
<tr>
<td>Ottroff et al.(^{23})</td>
<td>At least one of the following parameters: bilirubin ≥10mg/dL on day 7, INR ≥1.6 on day 7, ALT or AST &gt;2,000IU/L up to day 7</td>
</tr>
<tr>
<td>Ploeg et al.(^{26})</td>
<td>AST &gt;2,000IU/L, TP &gt;16 seconds and ammonia &gt;50μmol/L from day 2 to day 7</td>
</tr>
<tr>
<td>Nanashima et al.(^{27})</td>
<td>AST or ALT &gt;1,500IU/L in 2 consecutive tests within the first 72 hours</td>
</tr>
<tr>
<td>Dhillon et al.(^{21})</td>
<td>[(ALT + AST)/2] &gt;986IU/L on day 2</td>
</tr>
</tbody>
</table>

INR: international normal ratio; ALT: alanine aminotransferase; AST: aspartate aminotransferase; TP: prothrombin time.

**PRIMARY NONFUNCTION**

Primary nonfunction (PNF) is the most severe manifestation of graft dysfunction.\(^{1,4}\) Just as with EAD, a clear, universally accepted definition of PNF is lacking. However, there is one common denominator across studies: the need for early retransplantation or progression to patient death.\(^{4,7-9}\)

Makowka et al. defined PNF as patient death or need for retransplantation within the first 72 hours of surgery, other causes of liver failure excluded.\(^{25}\) Ploeg et al.\(^{8}\) described PNF as liver function inconsistent with recipient survival, either progressing to patient death or requiring retransplantation within 7 days of surgery. A similar PNF definition was adopted by Broering et al.,\(^{12}\) albeit with a 10-day interval from transplantation to diagnosis. On the other hand, Uemura et al.\(^{1}\) defined PNF as poor early graft condition requiring retransplantation or progressing to patient death within the first 7 days of surgery and emphasized the absence of other causes of liver failure, particularly vascular conditions, such as hepatic artery and portal vein thrombosis.\(^{1,26}\)

Other authors tried to define PNF based on objective laboratory cut-off values. The following criteria were proposed by Kremers et al.\(^{27}\) serum ALT levels >2,500IU/L, glucose levels <60mg/dL, INR >2.5 or bile flow <50mL/day. Dhillon et al.\(^{11}\) associated their own EAD laboratory criteria with poor recipient progression to define PNF.\(^{11}\) PNF diagnosis would correspond to [(ALT + AST)/2] >986IU/L, on the second post-transplantation day, with need for retransplantation or progression to patient death up to day seven. Máthé et al. also based their PNF definition on laboratory criteria and clinical progression; in that study, PNF was described as all clinical cases with serum AST or ALT levels >1,500IU/L, in two consecutive tests, within the first 72 hours post-transplantation and requiring retransplantation or progressing to recipient death.\(^{28}\)

According to the strict criteria established by the United Network for Organ Sharing (UNOS; American transplantation regulatory body), PNF is defined as serum AST levels ≥3,000 associated with at least one of the following: NR ≥2.5, acidosis corresponding to arterial pH ≤7.30 or venous pH ≤7.25 and/or serum lactate levels ≥4mMol/L. Researchers supporting definitions based on objective criteria argue this would simplify PNF diagnosis and promote fast reinclusion in organ transplant waiting lists, the only alternative to prevent patient death.\(^{4}\)

From the clinical perspective, PNF may be defined as hyperpotassemia, increased serum lactate levels, hemodynamic instability and oliguric renal failure up to the second week post-transplant.\(^{26}\) Other clinical manifestations include persistent encephalopathy and metabolic acidosis, marked hypoglycemia, coagulopathy and reduced or absent bile production associated with progressive increase in serum AST levels.\(^{29}\) The main PNF definitions are given on chart 2.
RISK FACTORS FOR EARLY ALLOGRAFT DYSFUNCTION AND PRIMARY NONFUNCTION

Early allograft dysfunction and PNF are multifactorial syndromes. Surgery, donor- and recipient-related factors have been described with these outcomes.\(^{11,4,7,28}\)

Donor age is often emphasized in the literature; however, a clear age cut-off has not been established to date. Increased risk of EAD has been associated with liver transplantation from donors aged over 49,\(^{8}\) 65\(^{30}\) or 45\(^{7}\) years. In the latter Olthoff et al.,\(^{7}\) univariate analysis revealed a trend towards increased incidence of EAD in transplants from donors aged over 49.6 years when age (years) was treated as a continuous variable.

Hepatic steatosis is also frequently associated with EAD in the literature. Steatosis type (i.e., macro or microvesicular) or severity (mild, moderate or severe; <30%, 30 to 59%, and >60%, respectively) are known to be associated with the occurrence of EAD and PNF.\(^{31,32}\) Primary nonfunction incidences of 1.8% and 5.1% were reported following transplantation of nonsteatotic grafts and grafts with macrovesicular steatosis greater than 30%, respectively.\(^{33}\) In contrast, grafts with up to 30% steatosis had similar PNF indices compared to nonsteatotic grafts.\(^{42}\) Grafts with moderate steatosis (30-59%) can be used, provided they are well allocated to recipients (i.e., recipients free from other risk factors).\(^{34,35}\) Transplantation of grafts with severe (over 60%) steatosis is limited to a few centers; most groups choose not to use such grafts out of increased EAD and PNF risk concerns.\(^{35,36}\)

Given the limited availability of grafts compared to organ demands, several transplant centers use marginal donors. Such donors are defined as expanded criteria donors according to the following: age ≥60 years, body mass index >27 to 30kg/m\(^2\), macrovesicular steatosis ≥30%, length of stay at ICU >4 to 5 days, use of vasoactive drugs (dopamine at doses >10μg/kilogram minute or any other catecholamine, regardless of dosage), long-standing hypotension (over 1 hour), serum sodium levels >150-155mmol/L, cold ischemia time >8 hours, hot ischemia time >40-45 minutes, controlled sepsis, serum creatinine levels >1.2mg/dL, history of alcoholism, serum bilirubin, ALT and AST levels >2.0mg/dL, >170IU/L and >140IU/L, respectively.\(^{37,39}\) The use of expanded criteria donors has been largely associated with EAD.\(^{37,39}\) Donors presenting with three or more of the above described parameters were expanded criteria donors by most authors.\(^{4,37,38}\)

The Donor Risk Index (DRI) concept was introduced by Feng et al.,\(^{40}\) in 2006, to predict the risk of graft failure. The DRI corresponds to a score system based on the following donor characteristics: donor age, race and height, cause of death, donation after cardiac death, use of split/partial grafts, geographic location and cold ischemia time. While not contraindicating donor use, the DRI provides supporting data for improved graft allocation.\(^{41}\) Different from other studies addressing risk factors in a categorical manner, the index developed by Feng et al.\(^{40}\) is a quantitative graft dysfunction risk index. However, difficulties in donor racial classification and, more importantly, the use of deceased donor organs from heart-beating donors, limit DRI application in Brazil. The combined use of DRI and recipient MELD scores to predict graft survival was reported by Avolio et al.\(^{42}\)

Cold and hot ischemia times were associated with EAD and PNF in several articles.\(^{14,7,8,43}\) However, cold ischemia time and hot ischemia time cut-offs were not clearly determined.\(^{4}\) Cold ischemia times in excess of 10 hours were shown to be associated with higher incidence of EAD, PNF and biliary complications in the long term.\(^{8,43}\) Associations between prolonged hot ischemia time and greater hepatocyte damage were reported and may explain higher EAD incidences.\(^{4}\) According to Cameron et al.,\(^{39}\) Hot ischemia time in excess of 40 minutes increased the risk of EAD and PNF. Hot ischemia times in excess of 45 minutes were also incriminated as independent risk factors for the same outcomes by Sirivatanauksorn et al.\(^{44}\)

Recipient related factors are also known to be associated with EAD.\(^{7,43,44}\) Recipient age and race have been arguably recognized as risk factors.\(^{5,39}\) Younger recipient age has been associated with lower risks of EAD and PNF.\(^{39}\) However, given the superior overall clinical condition of younger compared to older patients, younger recipients tend to receive marginal grafts; this fact may explain the higher EAD and PNF risks documented in this age group in some studies.\(^{6,43}\) Donor, surgery and recipient related risk factors for EAD are listed on chart 3.

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**Chart 2. Main definitions of primary nonfunction**

<table>
<thead>
<tr>
<th>Author</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ploeg et al.(^{26})</td>
<td>Liver function inconsistent with life; need for retransplantation or progression to death within seven days of surgery</td>
</tr>
<tr>
<td>Nanushima et al.(^{16})</td>
<td>AST or ALT &gt;1,500IU/L in 2 consecutive tests within the first 72 hours, requiring retransplantation or progressing to recipient death</td>
</tr>
<tr>
<td>Dhillon et al.(^{17})</td>
<td>[ALT + AST] &gt;96IU/L on day 2, requiring retransplantation or progressing to recipient death up to day 7</td>
</tr>
<tr>
<td>Broering et al.(^{18})</td>
<td>Need for retransplantation up to day 10 or death due to graft nonfunction</td>
</tr>
<tr>
<td>Máthé et al.(^{28})</td>
<td>AST or ALT &gt;1,500IU/L in 2 consecutive tests within the first 72 hours, leading to retransplantation or recipient death</td>
</tr>
<tr>
<td>Kremers et al.(^{29})</td>
<td>ALT &gt;2,500IU/L; blood glucose levels &lt;60mg/dL; INR &gt;2.5 or bile flow &lt;50mL/d</td>
</tr>
</tbody>
</table>

AST: aspartate aminotransferase; ALT: alanine aminotransferase; INR: international normal ratio.
CONCLUSION

Primary graft dysfunction can be divided into early allograft dysfunction and primary nonfunction. Early allograft dysfunction is characterized by laboratory changes within the first postoperative week and full graft recovery; in contrast, primary nonfunction is associated with clinical and laboratory catastrophic deterioration. Primary nonfunction includes primary dysfunction criteria but requires retransplantation or progresses to recipient death within the first seven days of surgery, after ruling out acute vascular complications.

Variables related to donor (age and graft steatosis) and to surgery (Cold ischemia times and Hot ischemia times) are the most commonly reported risk factors for primary graft dysfunction. The relation between recipient-related variables and graft dysfunction is less clear. The lack of a uniform, widely accepted graft dysfunction definition precludes the establishment of reproducible cut-offs across organ transplant centers.

Therefore, efforts must be directed towards the establishment of uniform early allograft dysfunction diagnostic criteria to determine the need for rapid implementation of support therapy at organ transplant centers. The use of a single test for standardized graft dysfunction diagnosis is recommended and may eliminate potential confounding factors derived from the adoption of different criteria in risk factor assessment studies. Indocyanine green clearance is thought to be one such promising tests.

REFERENCES


