Outcome of liver transplantation with Histidine-Tryptophan-Ketoglutarate vs. University of Wisconsin preservation solution: A retrospective observational double-center trial

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Abstract

Background: The question whether the choice of preservation solution affects outcome after liver transplantation is still not satisfactorily answered.

Methods: A double-center retrospective study of short- and long-term results of 3134 consecutive liver transplantations with follow-up periods up to 23 years was performed using multivariate, risk-adjusted analysis with a subset for living-donor transplants, pediatric transplants and cases with prolonged cold ischemic times. An additional focus was put on biliary complications.

Results: Although long-term graft survival appears to be increased by Histidine-Tryptophan-Ketoglutarate-use (p=0.018), this effect could not be confirmed in risk-adjusted analysis (p=0.641). Multivariate regression analysis revealed that 3-month mortality (p=0.120), 3-month graft survival (p=0.103) and long-term patient survival (p=0.235) were not influenced by the choice of preservation solution. There was no difference in the occurrence of common complications after liver transplantation. This was confirmed for living donor and pediatric transplantation and cases with prolonged cold ischemic time. Analysis of the preservation solutions' impact on length of hospital (p=0.113) and intensive care unit stay (p=0.481) revealed no significant difference. Histidine-Tryptophan-Ketoglutarate solution is easier to handle and less expensive.

Conclusions: University of Wisconsin and Histidine-Tryptophan-Ketoglutarate solutions are clinically equivalent. Histidine-Tryptophan-Ketoglutarate solution could have an economically superior profile. The notion that the choice of preservation solution can have an impact on the onset of biliary complications after liver transplantation remains a matter of controversy.
Key Words: organ procurement, risk-adjusted analysis, prolonged cold ischemic time, biliary complications, economic evaluation
Background

University of Wisconsin (UW) and Histidine-Tryptophan-Ketoglutarate (HTK) solutions are both used for organ preservation in liver transplantation (LTX). Both have been applied for decades largely depending on local preferences and several differences are known. UW is characterized by low osmolarity, the containment of colloid and several radical scavengers, such as adenosine and glutathione [1]. It revolutionized organ preservation in 1987 by facilitating "semi-elective" LTX [2]. HTK was developed by Bretschneider and colleagues in 1975, originally for cardioplegia [1]. In 2002, the US Food and Drug Administration approved HTK for preservation of donor livers. It is characterized by a lower osmolarity than UW, a strong histidine buffer system which increases the osmotic effect of mannitol and lower electrolyte concentrations [1, 3].

The effects of the choice of preservation solution (PS) on outcome after organ transplantation are still a matter of controversy. A systematic review of reports published until 2007 with a total of 1200 liver transplant recipients states that both are equally effective [4]. More recent data from the UNOS database points to preferable characteristics of UW, especially in means of graft survival [5]. Further, the impact of PS on the onset of biliary complications still remains unclear, since there are reports suggesting equivalent [6, 7] as well as differing risk profiles [8-10]. However, most of the available literature is limited by its single-center study character, small sample sizes and only short- to medium-term follow-up schemes.

We performed a double-center, data-base driven statistical analysis of the short- and long-term results of more than 3100 liver transplant recipients. Narrowly scoped subset-analyses were performed additionally for living-donor transplants, pediatric
transplants, as well as cases with prolonged cold ischemic time (CIT). Additionally, a focused investigation of biliary complications is part of this analysis.

**Patients and Methods**

This is a double-center, retrospective, observational study from two German liver transplant centers within the Eurotransplant community. Hannover Medical School provided data of 2554 consecutive liver transplants (1376 males [53.9%]; 1177 females [46.1%]) with a median recipient age of 43 years at transplant (range 0-74 years) performed between the 01.06.1987 and the 31.12.2012. Mean follow-up in this cohort is 5.7 years (median 3.9 years; range 0-22 years). The University Hospital Kiel provided data of 580 consecutive transplants (336 males [57.9%]; 244 females [42.1%]) with a median recipient age of 52 years at transplant (range 0-75 years) performed between the 01.11.1988 and the 09.02.2013. Mean follow-up in this cohort is 3.1 years (median 3.2; range 0-23.3 years). Only patients with either HTK or UW usage were included. The primary study endpoints were short- and long-term patient survival as well as death-censored graft survival. Short-term patient and graft survival was defined as 90 days post-transplant. The secondary study endpoints were the occurrence of post-transplant complications (hepatic artery thrombosis (HAT), portal vein thrombosis (PVT), caval vein thrombosis, post-operative hemorrhage, biliary complications), the necessity of operative revisions, the length of hospital stay, and the length of intensive care unit (ICU) stay.

**Statistical methods**

This is an analysis of prospectively stored and retrospectively compiled data. Age at transplant, recipient sex, allocation era (I: early phase center allocation 01/01/1987-
31/12/1992; II: late phase center allocation 01/01/1993-30/06/2000; III: Eurotransplant liver allocation system 01/07/2000-16/12/2006; IV: Model of End-Stage Liver Disease-based allocation 17/12/2006-09/02/2013), living-donor transplant, experience of the operating surgeon (defined as previously performed LTX as leading surgeon), CIT, pediatric patient (defined as younger than 18 years), operative duration, combined transplant, number of post-operative revisions, complications (as listed above), as well as the cohort's five most common indications leading to transplant (see Table 1) were analyzed as possible risk factors with influence on the primary and secondary study endpoints. Their relevance was identified with univariate binary regression and Cox regression analysis respectively. To avoid over-fitting all variables with an alpha-level <0.1 were considered for risk-adjusted multivariate regression analyses, which were performed to examine variations in risk according to usage of either UW or HTK solution. Kaplan-Meier analysis with Log Rank Tests, Pearson's Chi²-test or ANOVA-analysis were applied where appropriate. For all statistical tests a p-value <0.05 was defined as significant. The IBM SPSS statistics-software version 21.0 was used to perform statistical analysis.

Additional subset analysis

An additional detailed subset investigation was performed for the Hannover cohort's living donor related transplants (n=125; 4.9%), pediatric patients (n=575; 22.5%), and patients with prolonged CIT (n=346; 13.5%). Prolonged CIT was defined as the mean CIT (622 min) plus the standard deviation (239 min) which results in a CIT>861 min (the CIT was normally distributed, Kolmogorov-Smirnov-Test: p<0.001).
Amount of preservation solution

The Kiel data-base stored data on the perfused amounts of UW and HTK solution for 507 patients (87.4%). This data was investigated as a continuous variable in risk-adjusted regression modeling to assess its influence on the primary and secondary study endpoints. We aimed to reveal a potential dose-dependent effect of the preservation solution on outcome with this analysis. Further, addition of the Kiel cohort served to confirm the results from the Hannover analysis and aimed to reduce possible center bias.

Clinical handling of UW and HTK

Before application several additives, such prostaglandin E1 and/or dexamethason have to be added to UW solution. Once these components have been mixed, the solution must be used within 24h. In contrast, HTK solution can be used right away and no further additives are needed. From a clinical point of view, HTK can be quicker perfused without any need for pressurized perfusion. This is due to HTK’s comparatively low viscosity, which equals the one of water (2.0 cP). The viscosity of UW is higher (6.2 cP) because of the presence of colloids [1, 11].

Results

Descriptive statistics of the two cohorts and the distribution of UW vs. HTK solution are summarized in Table 2. In Hannover, UW was used in 1091 transplants (42.7%) and HTK in 1463 transplants (57.3%). In Kiel, livers were perfused 223 times with UW (38.4%) and 375 times with HTK (61.6%). The distribution and development of solution usage above time is depicted in Figure 1.
The Hannover cohort

Patient survival

In Kaplan-Meier analysis the type of PS has no statistically significant impact on long-term patient survival (p=0.235) (Fig. 2A). Three-month mortality was not significantly affected by the type of preservation solution (p=0.120).

Graft survival

Univariate Kaplan-Meier analysis demonstrated that HTK is beneficial for long-term graft survival (p=0.018; Log-Rank) (Fig. 2B). This result could not be confirmed in multivariate Cox regression (p=0.641), which was adjusted for necessity of operative revisions (p<0.001; HR:1.943; 95%-CI:1.711-2.206), occurrence of HAT (p<0.001; HR:4.306; 95%-CI:3.447-5.379), PVT (p<0.001; HR:1.973; 95%-CI:1.414-2.784), post-operative hemorrhage (p<0.001; HR:1.540; 95%-CI:1.346-1.763), biliary complications (p=0.003; HR:1.260; 95%-CI:1.086-1.461), age at transplant (p<0.001; HR:1.007; 95%-CI:1.004-1.011), living-donor LTX (p<0.001; HR:0.539; 95%-CI:0.370-0.785), CIT (p<0.001; HR:1.001; 95%-CI:1.000-1.001), pediatric LTX (p=0.001; HR:0.775; 95%-CI:0.662-0.908), era in which the transplant was performed (p<0.001; HR:0.811; 95%-CI:0.760-0.866), the indication hepatocellular carcinoma (HCC) (yes/no) (p=0.022; HR:1.247; 95%-CI:1.038-1.498), the indication primary sclerosing cholangitis (PSC) (yes/no) (p=0.001; HR:0.651; 95%-CI:0.501-0.847) and the indication biliary atresia (yes/no) (p=0.007; HR:0.702; 95%-CI:0.535-0.921).

Three-month graft survival was not affected by the type of PS (p=0.103).

Complications after LTX

The choice of PS did neither influence the occurrence of HAT (p=0.850), nor of PVT (p=0.761), caval vein thrombosis (p=0.053), post-operative hemorrhage (p=0.273),
biliary complications (p=0.178) and necessity of operative revisions (p=0.120) after LTX.

Biliary complications (n=488; 19.1%) have been further analyzed on a more detailed level, because of previously reported associations of these with the PS [11-13]. They were categorized into biliary leaks (n=218; 44.7%), stenosis (n=77; 15.8%), cholangitis (n=80; 16.4%) and not further specified biliary complications (n=129; 26.4%). In univariate regression analysis, beneficial effects on biliary leakage (p=0.035; OR:0.735; 95%-CI: 0.531-0.982) and biliary stenosis (p=0.011; OR:0.532; 95%-CI: 0.323-0.879) were associated with HTK-preservation. This result for biliary leakage could not be verified in a multivariate binary regression model (p=0.572), which was adjusted for the occurrence of HAT (p=0.001; OR:2.566; 95%-CI:1.525-4.317), caval thrombosis (p=0.043; OR:2.227; 95%-CI:1.025-4.835), post-operative hemorrhage (p=0.001; OR:1.690; 95%-CI:1.256-2.274), duration of LTX-procedure (p=0.011; OR:0.998; 95%-CI:0.997-1.000), living-donor LTX (p<0.001; OR:2.826; 95%-CI:1.779-4.489), CIT (p<0.001; OR:0.999; 95%-CI:0.998-0.999), the use of hepaticojejunostomy (p<0.001; OR:2.146; 95%-CI:1.618-2.848), PVT (p=0.012; OR:0.439; 95%-CI:0.213-0.905), pediatric LTX (p=0.005; OR:1.571; 95%-CI:1.156-2.136), era in which the transplant was performed (p<0.001; OR:1.584; 95%-CI:1.360-1.894), the indications viral hepatitis (yes/no) (p=0.001; OR:0.410; 95%-CI:0.231-0.727), acute liver failure (yes/no) (p=0.021; OR:0.524; 95%-CI:0.288-0.953) and biliary atresia (yes/no) (p<0.001; OR:2.251; 95%-CI:1.489-3.405).

Further, the beneficial influence of HTK on the rate of biliary stenosis could not be confirmed in a multivariate regression model (p=0.122), which was risk-adjusted for the occurrence of HAT (p<0.001; OR:5.398; 95%-CI:2.865-10.171) and the era in which the transplant was performed (p=0.001; OR:1.486; 95%-CI:1.162-1.899).
Analysis of cases with post-transplant cholangitis ($p=0.353$) as well as not specified biliary complications ($p=0.254$) remained insignificant.

**Length of hospital and ICU stay**

Neither the length of hospital stay ($p=0.113$) nor the length of ICU stay ($p=0.481$) after LTX were associated with the applied PS.

**Pediatric and living donor transplant**

22.5% of the Hannover cohort were pediatric patients ($n=575$). HTK was used more often in this subgroup ($n=302$; 52.5%) than UW ($n=273$; 47.5%) ($p<0.001$; Chi$^2$). None of the solutions influenced long-term patient ($p=0.104$) and graft survival ($p=0.059$) or three-month mortality ($p=0.056$) and three-month graft survival ($p=0.126$). There was no association of one of the investigated PS with biliary complications in the pediatric subgroup ($p=0.059$).

125 patients (4.9%) received a living-donor related transplant during follow-up. In 114 cases (91.2%) a HTK-preserved organ was transplanted; in 11 cases (8.8%) UW-solution was used ($p<0.001$; Chi$^2$). None of the solutions influenced long-term patient ($p=0.125$) and graft survival ($p=0.085$) or three-month mortality ($p=0.258$) and three-month graft survival ($p=0.181$). 33 patients (26.4%) developed biliary complications after living-donor related LTX. All of these transplanted grafts had been conserved with HTK, thus there is a statistically significant association between HTK-use and biliary complications in this sub-cohort ($p=0.036$).

**Prolonged cold ischemic time**

Definition of prolonged CIT is outlined in the Patients and Methods section. 346 (13.5%) patients received an organ that was exposed to a prolonged CIT. 116 of
these (33.5%) donor organs have been preserved with HTK and 230 (66.5%) using UW (p<0.001; Chi²). In this sub-cohort, the choice of PS had no statistically significant influence on long-term patient (p=0.989) and graft (p=0.796) survival as well as three-month mortality (p=0.550) and three-month graft survival (p=0.724). Further, there was no statistically significant difference between the rate of biliary complications in the two PS cohorts (p=0.097).

The Kiel cohort

In Kiel, 223 patients (38.4%) received a UW-preserved organ and 375 patients (61.6%) a HTK-preserved donor liver (see Figure 1B). Part of this cohort were 80 living donor related LTX (13.8%) and 87 pediatric recipients (15%). The mean CIT was 602 minutes (SD: 152 minutes).

Patient (p=0.798) and graft survival (p=0.454) were not influenced by the PS in multivariate analysis (adjusted for living donor transplantation, number of operative revisions, recipient age at transplant and era; all p<0.001). Investigation of short-term outcome also revealed no significant results in risk-adjusted regression analysis (three-month mortality: p=0.576; three-month graft survival: p=0.396). The PS had no influence on the occurrence of biliary complications (p=0.728) or the necessity of operative revisions (p=0.256).

An average of 4,869 ml UW solution was used for donor organ preservation (median 5,000 ml; range 1,500-10,000 ml). HTK preservation needed higher amounts of averaged 8,303 ml (median 8,000; range 1,000-17,000). Neither the amount of UW nor of HTK was associated with three-month mortality (UW: p=0.518; HTK: p=0.417) or graft survival (UW: p=0.789; HTK: p=0.053), biliary complications (UW: p=0.518; HTK: p=0.955) as well as long-term patient (UW: p=0.395; HTK: p=0.021, HR:1.000) or graft survival (UW: p=0.454; HTK: p=0.026, HR:1.000).
Economical evaluation

The costs for HTK per liter range between 50 and 56 Euro based on the amount of ordered units in Germany. There is currently no governmental approval for UW, thus no UW solution available on the German market. We therefore retrieved the costs of UW per liter from the literature, where costs of 181 Dollar per liter (≈136 Euro) are stated for UW on the US market [1]. Based on this information and taking the mean amount of PS into account, preservation of an average liver graft with HTK costs 415 Euro, whereas an average UW-perfused graft costs 662 Euro. Further, the costs for the said additives for preparation of UW solution have to be taken into account.

Discussion

Both HTK and UW are widely used for graft preservation in solid organ transplantation [4, 14, 15]. There are several studies available on the influence of the PS on outcome defined as survival, onset of complications, especially in the biliary tract, and rejection rates [4, 5, 14, 15]. The results of these studies range from equality of the solutions [4] to favoring one or the other [5, 10], thus the evidence-based choice of preservation solution remained a matter of controversy. Our local clinical situation is actually sharpened by the concerning fact that UW has currently no valid license for organ preservation and is thus not available in Germany. This has become problematic, since there are organs donated in other Eurotransplant partner nations, e.g. Belgium, where UW is still approved and used, leaving German transplant physicians without the possibility to re-perfuse an organ flown in from these countries. We recently experienced such an unfortunate and evitable event. A clinical, evidence-based standard to support the choice of PS use is thus indispensable.
We therefore performed an analysis with the data of two liver transplant centers (Hannover and Kiel, Germany), which are both within the Eurotransplant community and applied widely established statistical methods rigorously to compare the preservation solutions' influence on short- and long-term patient and graft survival, complications after LTX as well as the length of hospital and ICU stay. Competing risks were identified as possible confounders and systematically taken into account in multivariate analysis. An advantage of this study is the large number of included cases (total n=3134), which represents, at least to our knowledge, the largest non-registry-based investigation of this kind available at the moment. Further, UW and HTK were both applied in nearly fifty-fifty distribution, thus an analysis with sufficient comparable sample sizes was warranted (see Figure 1A/B).

Short- and long-term patient survival was not influenced by the choice of PS. Although long-term graft survival seemed to be increased by HTK-use (see Figure 2A), this effect could not be confirmed in risk-adjusted analysis. Furthermore, 3-month graft survival did not differ in the two PS cohorts. Since no association between the PS and common complications could be shown, we assume that HTK and UW are clinically equivalent choices and no PS can be preferred in terms of influence on the short- or long-term outcome after LTX. This can be confirmed for living donor as well as pediatric LTX. The data further revealed that it does not matter how much solution is actually used for preservation, as long the amount seems clinically sufficient (the investigated amounts of PS range from 1,500-10,000 ml UW and 1,000-17,000 ml HTK).

To answer the question whether one of the solutions is preferable in situations with extensively prolonged CIT, this subgroup was defined, identified and subsequently analyzed. Also in this subset analysis the choice of PS made no statistically
significant difference regarding short- and long-term patient and graft survival as well as the onset of post-transplant complications.

The notion that the choice of PS can have an impact on the onset of biliary complications after LTX is still a matter of controversy [6-10]. This aspect was therefore analyzed in depth resulting in no statistically significant difference between UW and HTK in means of onset of either biliary leakage (p=0.572), stenosis (p=0.122), post-transplant cholangitis (p=0.353) and not further specified biliary complications (p=0.254). The result that there might be more biliary complications after living-donor LTX is probably biased by the overall larger number of HTK-perfused grafts in this subgroup (HTK-use in 91.2%). Another possible explanation for this observation may be caused by differing practices in perfusion of the segment IV artery and the segment IV bile duct with PS. Further, it is known that partial liver grafts are associated with increased bile duct leaks from the liver resection surface.

Since our data suggests that there is no clinical difference between UW and HTK in LTX, economic considerations can play an important role for the choice of PS. As one important economical factor we analyzed the PS's impact on length of hospital and ICU stay, which was both not affected by the PS. HTK is clinically easier to handle, because it is sold in ready-to-use units while UW must be mixed with additives prior to use. although this might not be done anymore in some centers without deteriorating outcome. Furthermore, the quicker perfusion of the graft with HTK compared to UW can be seen as cost-benefit advantage ("time is money"). However, this effect is reduced by the fact that in average twice the amount of HTK is flushed. The routine use of expensive medications, including the PS, has been identified as one driving factor of cost-increase in transplantation [16]. The costs per liter vary drastically between the two investigated PS leading to the conclusion that HTK is the solution with the economically superior profile. A study aiming to find a
cost prediction model for LTX from 2008 [17] nevertheless reported that the median total cost of LTX significantly differs regarding two cohorts with either UW or HTK usage in favor of UW, although this result was not validated by a multivariate analysis. Also the number of included patients was smaller (n=139) than our study-population. Further investigations of these important economic aspects are needed to fully understand these interesting findings.

In conclusion, it is hard to really determine the economical impact of the PS, especially in the setting of this study. Due to center volume and individual negotiations there are different prices even among centers in the same region or country.

This study is limited due to the fact that non heart-beating donation is legally prohibited in Germany and thus no such cases could be included. This aspect might be partly responsible for the discrepancy between our results and the UNOS-registry study by Stewart et al. from 2009 [5], in which grafts donated after cardiac death were included.

It is somehow surprising that the rates of HAT and PVT are currently increasing as compared to the previous eras (see Table 2A), which might be due to the improved diagnostic capacities and thus higher detection rates rather than worsening results. Another important aspect in this context is the notion that with the introduction of MELD-based liver allocation and the “sickest-first-principle” in Germany, patients accepted for a transplant tend to be sicker nowadays than in earlier eras [18, 19]. This might be also reflected in the increasing lengths of hospital and ICU stay (see Table 2A).

Recently published data on a biochemically improved HTK solution may provide a promising option for the future [20]. Liu and colleagues reported that HTK-N solution, which is modified with N-acetylhistidin, amino-acids and iron chelators, protects liver
grafts with microvesicular steatosis better than common HTK in a rat model [20]. This result is especially interesting given the expanding use of extended donor criteria in LTX. Celsior solution might also be able to show some advantages, which was underlined in a recent prospective multicenter trial from France [21]. Further studies are needed, preferably with higher levels of evidence such as prospective multicenter trials or randomized registry trials to investigate the preliminary results of our study. In times of chronically diminishing donation rates in Germany [19], it is the transplant society's responsibility to ensure the best possible preservation of the scarce donor organs. Thus, further investigations regarding this topic are warranted.

Conclusions

UW and HTK are clinically equivalent. HTK could have an economically superior profile. However, it is hard to really determine the economical impact of the PS, especially in the setting of this study. The notion that the choice of preservation solution can have an impact on the onset of biliary complications after liver transplantation remains a matter of controversy.
List of abbreviations (in alphabetical order)

CIT     cold ischemic time
HAT     hepatic artery thrombosis
HCC     hepatocellular carcinoma
HR      hazard ratio
HTK     Histidine-Tryptophane-Ketoglutarate solution
ICU     intensive care unit
LOS     length of hospital stay
LTX     liver transplantation
OR      odds ratio
PS      preservation solution
PSC     primary sclerosing cholangitis
PVT     portal vein thrombosis
SD      standard deviation
UW      University of Wisconsin solution
95%-CI  95%-confidence interval

Competing interests: The authors declare that they have no competing interests.

Author contributions: AK, HS and BR participated in the design of the study and performed the statistical analysis. The data was collected by AK, FL, JK, HS and BR. The manuscript was drafted by AK, JG, HS and BR and critically revised by AK, JG, VA, CK, FL, FB, TB, JK, HS and BR. All authors read and approved the final manuscript.
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Figure Legends

**Figure 1:** Distribution of preservation solution in the Hannover (A) and Kiel (B) cohort as well as usage of solutions over the follow-up time in the Hannover cohort (C).

**Figure 2:** Results of univariate Kaplan Meier analysis of patient (A) and graft (B) survival distributed by the preservation solution in the Hannover cohort.
**Figure 1:** Distribution of preservation solution in the Hannover (A) and Kiel (B) cohort as well as usage of solutions over the follow-up time in the Hannover cohort (C).
Figure 2: Results of univariate Kaplan Meier analysis of patient (A) and graft (B) survival distributed by the preservation solution in the Hannover cohort.

A

B
**Table 1:** The five most common indications for liver transplantation in the Hannover and Kiel cohort.

<table>
<thead>
<tr>
<th>Hannover cohort</th>
<th>Number of patients (% of cohort)</th>
<th>Kiel cohort</th>
<th>Number of patients (% of cohort)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Indication</strong></td>
<td><strong>1 viral hepatitis 334 (13.0%)</strong></td>
<td><strong>alcoholic cirrhosis 119 (20.5%)</strong></td>
<td></td>
</tr>
<tr>
<td><strong>2 hepatocellular carcinoma</strong></td>
<td><strong>281 (11.0%)</strong></td>
<td><strong>viral hepatitis 86 (14.8%)</strong></td>
<td></td>
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<tr>
<td><strong>3 acute liver failure</strong></td>
<td><strong>246 (9.6%)</strong></td>
<td><strong>hepatocellular carcinoma 43 (7.4%)</strong></td>
<td></td>
</tr>
<tr>
<td><strong>4 primary sclerosing cholangitis</strong></td>
<td><strong>196 (7.7%)</strong></td>
<td><strong>biliary atresia 41 (7.1%)</strong></td>
<td></td>
</tr>
<tr>
<td><strong>5 biliary atresia</strong></td>
<td><strong>192 (7.5%)</strong></td>
<td><strong>acute liver failure 36 (6.2%)</strong></td>
<td></td>
</tr>
</tbody>
</table>
Table 2 A: Distribution of primary and secondary study endpoints as well as preservation solutions to the different eras, 2554 patients from the Hannover liver transplant program.

<table>
<thead>
<tr>
<th>Era</th>
<th>I</th>
<th>II</th>
<th>III</th>
<th>IV</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>n (%)</td>
<td>346 (13.5%)</td>
<td>773 (30.3%)</td>
<td>811 (31.8%)</td>
<td>624 (24.4%)</td>
<td>n.a.</td>
</tr>
<tr>
<td>HTK</td>
<td>32 (9.2%)</td>
<td>470 (60.8%)</td>
<td>439 (54.1%)</td>
<td>522 (83.7%)</td>
<td>&lt;0.001¹</td>
</tr>
<tr>
<td>UW</td>
<td>314 (90.8%)</td>
<td>303 (39.2%)</td>
<td>372 (45.9%)</td>
<td>102 (42.3%)</td>
<td>&lt;0.001¹</td>
</tr>
<tr>
<td>3 months mortality</td>
<td>82 (23.7%)</td>
<td>142 (18.4%)</td>
<td>112 (13.8%)</td>
<td>98 (15.7%)</td>
<td>&lt;0.001¹</td>
</tr>
<tr>
<td>graft loss</td>
<td>224 (64.7%)</td>
<td>374 (48.4%)</td>
<td>281 (34.6%)</td>
<td>171 (27.4%)</td>
<td>&lt;0.001¹</td>
</tr>
<tr>
<td>3 month graft survival</td>
<td>238 (68.8%)</td>
<td>582 (75.3%)</td>
<td>645 (79.5%)</td>
<td>482 (77.4%)</td>
<td>0.001¹</td>
</tr>
<tr>
<td>HAT</td>
<td>14 (4.0%)</td>
<td>19 (2.5%)</td>
<td>30 (3.8%)</td>
<td>37 (6.0%)</td>
<td>0.010¹</td>
</tr>
<tr>
<td>Portal vein thrombosis</td>
<td>6 (1.9%)</td>
<td>7 (0.9%)</td>
<td>24 (3.1%)</td>
<td>27 (4.4%)</td>
<td>0.001¹</td>
</tr>
<tr>
<td>Caval vein thrombosis</td>
<td>1 (0.3%)</td>
<td>4 (0.5%)</td>
<td>10 (1.3%)</td>
<td>31 (5.0%)</td>
<td>&lt;0.001¹</td>
</tr>
<tr>
<td>Post-operative hemorrhage</td>
<td>73 (22.7%)</td>
<td>197 (25.9%)</td>
<td>162 (20.8%)</td>
<td>169 (27.3%)</td>
<td>0.020¹</td>
</tr>
<tr>
<td>Biliary complications</td>
<td>34 (16.9%)</td>
<td>127 (16.7%)</td>
<td>169 (21.7%)</td>
<td>138 (22.3%)</td>
<td>0.015¹</td>
</tr>
<tr>
<td>Mean LOS in days (SD)</td>
<td>44 (34)</td>
<td>40 (30)</td>
<td>41 (35)</td>
<td>51 (43)</td>
<td>&lt;0.001²</td>
</tr>
</tbody>
</table>

HTK=Histidine-Tryptophane-Ketoglutarate solution; UW=University of Wisconsin solution; SD=Standard Deviation; HAT=hepatic artery thrombosis; LOS=length of hospital stay

¹Pearson's Chi² Test

²ANOVA
**Table 2 B**: Distribution of primary and secondary study endpoints as well as preservation solutions to the different eras, 580 patients from the Kiel liver transplant program.

<table>
<thead>
<tr>
<th>Era</th>
<th>I</th>
<th>II</th>
<th>III</th>
<th>IV</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>n (%)</td>
<td>9 (1.6%)</td>
<td>96 (16.6%)</td>
<td>152 (26.2%)</td>
<td>323 (55.7%)</td>
<td>n.a.</td>
</tr>
<tr>
<td>HTK</td>
<td>0 (0%)</td>
<td>8 (8.3%)</td>
<td>68 (44.7%)</td>
<td>281 (87.0%)</td>
<td>&lt;0.001&lt;sup&gt;1&lt;/sup&gt;</td>
</tr>
<tr>
<td>UW</td>
<td>9 (100.0%)</td>
<td>88 (91.7%)</td>
<td>84 (55.3%)</td>
<td>42 (13%)</td>
<td>&lt;0.001&lt;sup&gt;1&lt;/sup&gt;</td>
</tr>
<tr>
<td>3 months mortality</td>
<td>3 (33.3%)</td>
<td>22 (22.9%)</td>
<td>35 (23.0%)</td>
<td>32 (9.9%)</td>
<td>&lt;0.001&lt;sup&gt;1&lt;/sup&gt;</td>
</tr>
<tr>
<td>Graft loss</td>
<td>8 (88.9%)</td>
<td>59 (61.5%)</td>
<td>75 (49.7%)</td>
<td>85 (26.5%)</td>
<td>&lt;0.001&lt;sup&gt;1&lt;/sup&gt;</td>
</tr>
<tr>
<td>3 month graft survival</td>
<td>6 (66.7%)</td>
<td>68 (70.8%)</td>
<td>106 (69.7%)</td>
<td>262 (81.1%)</td>
<td>0.021&lt;sup&gt;1&lt;/sup&gt;</td>
</tr>
<tr>
<td>Biliary complications</td>
<td>1 (11.1%)</td>
<td>33 (34.4%)</td>
<td>47 (30.9%)</td>
<td>18 (19.4%)</td>
<td>0.099&lt;sup&gt;1&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

HTK=Histidine-Tryptophane-Ketoglutarate solution; UW=University of Wisconsin solution; SD=Standard Deviation

<sup>1</sup>Pearson's Chi² Test