Current perspectives in transplant medicine: hypothermic oxygenated perfusion

Sebastian G Michel1
Joren C Madsen2,3
1Department of Cardiac Surgery, Ludwig-Maximilians-University, Munich, Germany; 2Department of Surgery, Center for Transplantation Sciences, 3Division of Cardiac Surgery, Department of Surgery, Massachusetts General Hospital, Boston, MA, USA

Abstract: The purpose of this review was to provide an update on hypothermic oxygenated perfusion as a preservation technique for whole organ allografts. Clinical and experimental data in heart, lung, liver, kidney, and pancreas/islet transplantation are summarized with a special emphasis on marginal donors and donation after circulatory death. The rationale behind hypothermic machine perfusion as well as its advantages and disadvantages compared to conventional cold storage and the competing technique, normothermic machine perfusion, are reviewed.

Keywords: organ transplantation, organ preservation, hypothermic perfusion

Introduction

Organ transplantation remains the most effective way to treat organ failure. Unfortunately, the gap between available donor organs and potential recipients is wide and becoming wider (www.organdonor.gov). Therefore, various attempts have been made to increase the donor pool, for example, by accepting longer ischemic times or by using extended criteria donors (ECD) whose organs may be of inferior quality due to age or disease. Using donors after circulatory death (DCD) represents another way to increase the pool of transplantable organs but is, by definition, associated with a period of harmful warm ischemia time unlike organs retrieved from donors after brain death (DBD). Machine perfusion (MP) of donor organs (either hypothermic machine perfusion [HMP] or normothermic machine perfusion [NMP]) attempts to minimize and even compensate for harmful effects of warm and cold ischemia in order to improve organ quality and ultimately transplant outcomes while at the same time, increasing the number of potential donors.

Static cold storage (CS) in a suitable solution like University of Wisconsin (UW) or Celsior on ice, remains the standard method to preserve organs of good quality after DBD. However, the revival of MP in kidney transplantation has led to a 37% increased usage of this technique in the USA.1 Not only can MP mitigate some of the negative effects of ischemia on the donor organ, but it is also a helpful tool in evaluating the overall quality of the organ before it is transplanted. Moreover, in some cases, MP can provide a beneficial period of physiologic resuscitation which can convert a compromised organ into one that is suitable for transplantation.2 This review will provide an update on current clinical applications and latest experimental findings of MP technology in organ transplantation with a focus on “hypothermic oxygenated perfusion”.

Correspondence: Joren C Madsen
Center for Transplantation Sciences, Massachusetts General Hospital-East, Charlestown Navy Yard, Building 149, 13th Street, Charlestown, MA 02129-2020, USA
Tel +1 617 726 6506
Fax +1 617 726 4067
Email jcmadsen@mgh.harvard.edu

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HMP vs static CS

After procurement, the donor organ undergoes a period of ischemia because it is disconnected from the blood supply and therefore from oxygen and nutrients. Static CS at the recommended 4°C reduces the oxygen demand and the metabolism approximately 12-fold but not completely. This results in ongoing adenosine triphosphate (ATP)-depletion, a change from aerobic to anaerobic metabolism and the production of lactate depending on the length of the ischemic time.4 The result of the imbalance between supply and demand within the ischemic organ and its subsequent reperfusion lead to activation of the immune system and, ultimately, the result is cell death by apoptosis and necrosis, known as ischemia-reperfusion injury.5 Some organs are more sensitive toward ischemia (heart) than others (liver, kidney). Hypothermic oxygenated perfusion, that is, the ex vivo perfusion of donor organs on a machine at low temperatures prior to transplant (as it is defined by Taylor and Baicu6) is meant to prevent ATP depletion by providing oxygen and nutrients while eliminating toxic metabolic by-products such as lactate. In this way, ischemia-reperfusion injury is reduced and overall clinical results improved.4,7–9 Another, often overlooked advantage of HMP over CS is the fact that the temperature can be precisely regulated to remain in the desirable range of 4°C–8°C. In static CS on ice, temperatures often drop too low (0°C–2°C) resulting in tissue freezing10 and more severe tissue injury upon reperfusion.11

The importance of oxygenation

There is an ongoing debate on the optimum oxygen requirement of donor organs during hypothermia. On the one hand, we know that cellular metabolism is halved for every 10°C reduction in temperature (van’t Hoff’s rule). For example, when HMP is performed at 7°C cellular metabolism is only 12.5% compared to normothermia. However, if too much oxygen is delivered during hypothermia, free radicals are generated, which can lead to tissue damage.12 Conversely, there is clear evidence that ATP levels in the tissue can only be preserved when oxygen is supplied, even at low temperatures. Thus, “oxygenated” HMP can avoid oxidative stress for the donor organ.13,14

Interestingly, three different preclinical studies using a porcine kidney transplant model found a difference in the effects of oxygenation in MP depending on whether the organ was procured from a heart-beating donor or from a DCD donor. When heart-beating donors were used oxygenated MP was actually inferior to anoxic MP as evidenced in the lower creatinine clearance in the oxygenated group.15 However, in the DCD setting, associated with a greater period of warm ischemia, creatinine clearance was higher,16 peak creatinine levels were lower after kidney transplantation and there was less interstitial fibrosis in the 3-month follow-up when oxygenated MP was applied.17 This might be explained by the fact that oxygenated MP is more beneficial in organs undergoing a prolonged warm ischemia.2 It could also explain why in Watson’s multicenter randomized controlled trial of DCD-kidney transplantation, using “nonoxygenated” MP no statistically significant reduction in delayed graft function (DGF) could be shown when compared to static CS.9

HMP of the kidney

HMP of the kidney was originally used in the 1960s by Belzer’s group18 but the machine was so huge that the ease and low cost of static CS were superior, particularly when UW solution became available.19 A revival of nonoxygenated HMP has been seen recently as more and more marginal kidneys are accepted for transplantation to meet the demand of the fast growing numbers of recipients on the waiting list. These ECD include old donors with impaired kidney function as well as DCD donors. It has been shown that transplantation of such kidneys has a survival benefit for the patient compared to dialysis alone.20 There is solid evidence that the rate of DGF, defined as need for dialysis in the first week posttransplant, is lower after nonoxygenated HMP compared to conventional CS. This is true for standard criteria (good quality, brain dead) donors,21,22 ECD23 and DCD.8,22

In these studies, nonoxygenated HMP was applied. Testing the quality of the graft during MP by measuring flow and resistance could not have been shown to predict transplant outcome in some studies24–26 but elevated resistance was associated with DGF and graft failure at 1 year in the randomized Eurotransplant trial.27

The therapeutic approach of adding substrates to improve the graft quality during HMP is still under investigation. It could be shown in a porcine model that propofol, acting as an antioxidant, was able to reduce renal vascular resistance and improve renal function posttransplant.28

HMP of the liver

Clinical experience with liver HMP is limited. Three transplant centers (Columbia using nonoxygenated HMP, Groningen and Zurich using oxygenated HMP) currently use this technique29 as an end-ischemic perfusion after CS (i.e., not continuous starting after the organ procurement, but HMP after CS before implanting the liver). The initial report of 20 standard DBD human livers showed less enzyme release, less
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Early graft dysfunction, and a shorter hospital stay compared to control when nonoxygenated HMP was used. Guarrera et al also investigated marginal DBD donors and found less biliary complications when nonoxygenated HMP was applied compared to CS alone. The group from Zurich studied DCD livers using oxygenated HMP (liver assist, Figure 1) and showed less biliary complications and therefore better graft survival. Lastly, the Groningen group could show less biliary strictures in DCD livers using oxygenated HMP (liver assist, Figure 1). Randomized studies are underway to investigate this preliminary data further. There is still debate over whether perfusion should only be performed through the portal vein or through the portal vein and the hepatic artery.

HMP of the heart

Registry data of the International Society for Heart and Lung Transplantation show that ischemia of more than 200 minutes is associated with a higher mortality after heart transplantation. Therefore, it is critical to find ways to protect against longer ischemic times for the heart. Not only would extending the safe ischemic period time permit the transportation of hearts procured from more distant sites, but it could also enable human leukocyte antigen (HLA)-matching between donor and recipient. HLA-matching is not routine in heart transplantation at the moment because of the time constraints, however, it has been shown to improve survival in some reports. Currently, only “normothermic” perfusion is used clinically (see Normothermic machine perfusion (NMP) section). There is, however, preclinical experimental data showing that hypothermic oxygenated MP is beneficial for donor hearts.

Using a porcine model, our group showed that oxygenated HMP using the Paragonix Sherpa™ Cardiac Transport System (Figure 2) led to a better preserved cell structure of the myocytes after reperfusion of the donor heart after the currently accepted ischemic period of 4 hours and also after an extended ischemic period of 12 hours. Furthermore, we demonstrated that the perfused hearts had less depletion of tissue ATP stores and had preserved architecture of the mitochondria (Figure 3). There was also a nonsignificant trend toward lower endothelin-1 levels (marker for endothelial dysfunction) and less apoptosis in TUNEL-staining in perfused hearts after 12 hours compared to 12 hours conventional CS in Celsior. These benefits could potentially translate to better short-term (less primary graft failure) and long-term (less cardiac allograft vasculopathy) clinical outcomes after heart transplantation. We are currently working on finding the optimal perfusion pressure and flow to avoid edema formation on the one hand but to provide enough oxygen and nutrients on the other.

In a canine study, hearts transplanted after 24 hours of HMP were comparable to 4 hours statically stored hearts in terms of duration of cardiopulmonary bypass support, survival, inotropic support, and cardiac output. Another study in dogs showed superior systolic functional recovery on a Langendorff system after 24 hours of HMP compared to 8 hours of CS. A similar preclinical model found superior functional recovery after HMP but also reduced caspase-3 (marker for apoptosis), reduced malondialdehyde (marker for oxidative stress), and reduced endothelin-1-levels (marker for endothelial dysfunction) in the heart tissue as well as higher ATP levels compared to CS.
HMP of the lung

Hypothermic oxygenated MP of lungs has only been studied in preclinical experimental models. Using a DCD-lung transplant model in dogs, a positive effect of HMP was shown compared to conventional CS alone. Lungs showed less ischemia-reperfusion injury histologically, less proinflammatory cytokine levels, and higher tissue ATP levels after reperfusion using HMP with Steen solution at 10°C. The authors hypothesized that HMP is particularly helpful for the wash-out of microthrombi that are present in the donor lung before transplantation.

HMP of the pancreas

Until now, HMP has not been used in human pancreas transplantation. Although it has been shown to be feasible and safe in animal models, an advantage compared to CS could not be shown. HMP causes graft edema and congestion followed by venous thrombosis and graft failure despite attempts to adjust flow and pressure. However, there may be a theoretical advantage of edema formation in perfused pancreata because islets separate more easily from the extracellular matrix in perfused pancreata compared to CS and this can yield a greater number of the islets.

Normothermic machine perfusion (NMP)

NMP with blood undoubtedly comes closest to the physiological state of the donor organ. Aerobic metabolism can continue outside the body and there is no ATP depletion. The disadvantage of NMP is its complexity and its cost compared to CS or HMP. Oxygen and energy demands are much higher at 37°C, and therefore, technical failure leads to ischemic damage immediately. MP at hypothermia has much more room for error. In addition, normothermia has a higher risk of transmitting infection with the donor organ, as bacteria grow faster. Nevertheless, several reports have shown positive and promising results of this technique for different organs.

In clinical lung transplantation, normothermic perfusion using the Organ Care System™ (Transmedics, Andover, MA, USA) has been shown to be safe and effective in a pilot study of 12 patients. A randomized multicenter trial (INSPIRE) has been completed in 2014, and preliminary data presented at the International Society for Heart and Lung Transplantation Meeting in Montreal 2013 showed noninferiority compared to conventional CS in terms of the combined primary endpoint 30-day survival and absence of primary graft dysfunction grade 3 after 72 hours. The other established technique to preserve donor lungs is conventional CS followed by ex vivo lung perfusion (EVLP) to improve the quality of the donor lung. This technique is particularly helpful when DCD-lungs with inferior initial quality are being evaluated for transplant. It has been shown that transplantation of these high-risk donor lungs using EVLP led to similar results to those obtained with conventionally selected lungs without EVLP. EVLP is a clinical routine in some experienced lung transplant centers (e.g., Toronto and Lund), and is a helpful tool to evaluate the quality of marginal donor organs particularly for DCD.
System has been shown to be noninferior to conventional CS\(^6\) (PROCEED II trial, primary endpoint 30-day graft and patient survival). Importantly, the Organ Care System has also enabled the most successful cases of DCD heart transplantation in Australia\(^6\) and the UK. In kidney transplantation, Nicholson and Hosgood have described a very low DGF rate of only 5.6% after NMP, despite the fact that they used marginal donors.\(^{51}\) The rate of DGF in these 18 patients is significantly lower than the 20.8% seen in the Eurotransplant trial using HMP.\(^{22}\) The OCS\(^{\text{TM}}\) Liver PROTECT clinical trial is currently initiating and Peter Friend’s group from Oxford and Birmingham has just published the results of a Phase I study with 20 patients showing that NMP for liver transplantation is safe and feasible.\(^{52}\) Using the OrganOx metra\(^\text{®}\), they could even show significantly lower aspartate aminotransferase levels in the first 7 days in the NMP group compared to CS. Finally, there is one report on the successful use of NMP for quality assessment of discarded human pancreases (insulin secretion and amylase levels).\(^{53}\) Based on these data, normothermic perfusion appears to be highly efficient and may represent the future of organ preservation.

**Conclusion**

In conclusion, HMP is superior to the current standard of static CS of donor organs and can increase the donor pool by enabling the surgeon to more aggressively accept marginal donors and longer preservation times. For kidney transplantation, controlled randomized trials have shown that HMP improves outcomes by reducing DGF. The novel emerging technology of NMP seems to be promising in heart, lung, liver, and kidney transplantation and, although expensive, will probably be the future of organ preservation, as it is the most physiological way to preserve a donor organ.

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**Disclosure**

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