Abstract

Background: Despite increased utilization of kidneys from deceased donors (DCD; Donation after Cardiocirculatory Death) for transplantation, delayed graft function (DGF) and primary non-function (PNF) incidence remain high for DCD kidneys while post-transplant graft survival is lower. An enhanced preservation technology will facilitate qualitative improvements in DCD kidneys and outcomes. Present study evaluated the ability of novel organ preservation solution Somah compared to University of Wisconsin (UW) solution for extended storage of DCD kidneys.

Methods: Porcine DCD kidneys procured 60 ± 10 minutes after cardiac death were flushed and preserved in storage solutions at 4°C for 72 hours. Gross morphology was evaluated and biopsies taken at 1, 6, 24 and 72 hours for histopathology (HP), tissue high energy phosphates (HEP; ATP+Creatine Phosphate) and western blotting for caveolin, endothelial-nitric oxide synthase (eNOS) von-Willebrands factor (vWF) and erythropoietin (EPO). Metabolic parameters of pH, pO₂, pCO₂, glucose and lactate levels in solutions were determined at similar time points.

Results: Somah-stored kidneys demonstrated normal gross morphology, greater metabolic activity and preservation of energy (HEP). In contrast, UW-kidneys showed mottled appearance, a greater degree of time-dependent increase in nuclear hyperchromacy of tubular epithelial cells, attenuation of metabolism, progressive decrease in HEP, and in the expression of eNOS, vWF and EPO, indicating tissue damage during storage.

Conclusion: Better preservation of renal tissue and higher energy status of Somah-preserved kidneys suggests that DCD kidneys stored in Somah are likely to perform better upon transplantation. Future studies including transplantation of DCD kidneys stored in Somah are required for further evaluation.

Keywords: Extracorporeal storage; Kidney transplantation; Preservation solution; Somah; Donation after cardiocirculatory death

Abbreviations: CP: Creatine Phosphate; eNOS: endothelial Nitric Oxide Synthase; EPO: Erythropoietin; HEP: High Energy Phosphates; vWF: von Willebrands Factor

Introduction

Eighty-six of every hundred patients requiring renal transplant last year, did not get one, adding to the ever mounting waitlist for transplant-requiring individuals [National Kidney Foundation; http://www.kidney.org]. This is despite tremendous progress in solid organ transplantations in last several decades and availability of renal replacement therapy (Hemodialysis and Peritoneal dialysis) which are not only associated with severely debilitating and/or life-threatening complications but hinder routine lifestyle of patients due to frequent requirements for hospital visits, imparting a huge financial burden on society.

While last year 14000 patients in US received renal transplant, 2500 patients are added to renal transplant waitlist each month, with around 100,000 patients awaiting kidney transplantation at the time of writing this paper; the average waiting time being three to five years [1]. Although, 65% of renal grafts are procured from deceased (DCD; Donation after Cardiac Death) donors, these grafts are twice as likely to develop delayed graft function (DGF) compared to standard-criteria donors [2], an increased incidence of primary non-function (PNF) [3], and halved overall graft survival. Whereas grafts exposure to warm and/or cold ischemia is an obligate inevitability of solid organ transplant, the resulting cellular/tissue damage during this period is consequential of the less than desired post-transplant operative renal function. The above data indicate an enormous potential to improve quality of DCD kidneys for transplantation by making advances in preservation technology; and thus more desirable long-term patient outcomes.

To this end, we formulated a novel organ storage solution, Somah, demonstrably superior in ex-vivo static preservation of porcine cardiac grafts, compared to current clinical norms such as Celsior and UW [4-7]. The rationally incorporated components of Somah, not only help attenuate reperfusion injury (by reactive oxygen species), the synthesis of high energy phosphates (HEP; ATP+Creatine Phosphate) is also enhanced significantly, thus appropriating metabolic requirements during storage and immediately upon reperfusion. The superiority of Somah has also been demonstrable in extracorporeal static storage of DCD Livers [8] and perfused storage of DCD Lungs [9].
Our studies on porcine hearts and livers demonstrate that preservation of energy metabolism during extracorporeal storage directly correlates to superior post-storage biochemical and functional outcomes. Present study was designed to evaluate the efficacy of Somah in extended extracorporeal static storage of DCD kidneys using insights gleaned from our experience of preservation of other organs in Somah and studies published in the field, as a standard for preservation of renal grafts cellular viability and energy requirements during storage.

Materials and Methods

Surgical procurement of kidneys

Female Yorkshire Swine weighing 40-50 Kg were used as per protocol approved by institutional animal studies committee. Swine were sedated with telazol 4-6 mg/kg i.m. and xylazine 2 mg/kg i.m., intubated and connected to ventilator. Anesthesia was maintained using i.v. propofol (10 mg/kg/hr) and remifentanil (40-60 µg/hr). Cis-atracurium (10-20 mg i.v.), a paralytic agent, was administered ten minutes prior to surgery. Upon midline sternotomy, animals were systemically heparinized (300 mg/kg) and aortic root cannulated. Ice cold cardioplegia (20 mM K+) was infused after aortic clamping to stop the heart which was then excised for other experiments as described [6,7]. Time of complete cessation of heart contraction was recorded as the beginning of warm ischemia of other body organs. Post median laparotomy, suprahepatic aorta was cannulated and abdominal organs flushed with 2 L of ice cold UW (CoStorSol; Preservation Solutions Inc., Elkhorn, WI) or Somah solution (Somahlution Inc., Jupiter, FL), at 100 mmHg pressure and a flow of 300 ml/min, till perfusate returning through inferior vena cava (IVC) was clear. Abdominal organ harvest was concluded first with hepatectomy for use in other experiments, and subsequent total bilateral nephrectomy after carefully dissecting the renal pedicles. Kidneys were immediately transferred to Somah or UW solution (Table 1) at 4°C and static stored for 72 hours. Kidney biopsies were obtained for histopathology, HEP and Western blot assays at time 0, and 6, 24 and 72 hour time-points. Time 0 corresponds to 1 hour in storage; time required to transport kidneys from animal research facility to lab before first biopsy.

Histopathology

Tissue was fixed in formalin and embedded in paraffin before cutting 10 µ thin sections that were melted onto glass slides for further processing. Tissue sections were dried in sequentially increasing ethanol and embedded in paraffin. Paraffin blocks were cut 10 µ thin sections that were stained with methylene blue and eosin staining after which slides were immersed in xylene clearing agent, covered with cover slip and examined under microscope. Images were acquired and analyzed using Olympus microscope and image analyzer system (BX51TRF; Olympus America Inc, USA) and assessed in blind fashion by independent observers.

ATP and creatine phosphate assay

ATP and creatine phosphate (CP) were measured in kidney tissue extracts as described [4-7,10]. In brief, 20 mg of renal tissue was suspended in 400 µl of 0.4 M ice-cold perchloric acid and homogenized twice for 30 secs. Homogenate was centrifuged at 1970 g for 10 mins at 0°C. An aliquot of supernatant was neutralized with equal volume of ice-cold 0.4 M KHCO₃ solution and centrifuged as above. Supernatant was stored at -80°C for ATP/CP measurements. The pellet was dissolved in equal volume of 0.1 M NaOH and centrifuged and used for protein assay. ATP/CP was measured using a bioluminescent assay kit (Sigma-Aldrich and GloMax Multi+Detection System, Promega) according to manufacturer’s protocol.

### Table 1: Composition of Somah and UW solutions

<table>
<thead>
<tr>
<th>Component</th>
<th>SOMAH (pH 7.5)</th>
<th>UW SOLUTION (pH 7.4)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potassium Phosphate (monobasic)</td>
<td>0.44 mmol/L</td>
<td>25.00 mmol/L</td>
</tr>
<tr>
<td>Potassium Chloride</td>
<td>7.00 mmol/L</td>
<td>100.00 mmol/L</td>
</tr>
<tr>
<td>Sodium Chloride</td>
<td>125.00 mmol/L</td>
<td>27.00 mmol/L</td>
</tr>
<tr>
<td>Sodium Bicarbonate</td>
<td>5.00 mmol/L</td>
<td>5.00 mmol/L</td>
</tr>
<tr>
<td>Calcium Chloride</td>
<td>1.30 mmol/L</td>
<td>3.00 mmol/L</td>
</tr>
<tr>
<td>Sodium Phosphate (dibasic; heptahydrate)</td>
<td>0.19 mmol/L</td>
<td>0.00 mmol/L</td>
</tr>
<tr>
<td>Magnesium Chloride (hexahydrate)</td>
<td>0.50 mmol/L</td>
<td>0.00 mmol/L</td>
</tr>
<tr>
<td>Magnesium Sulfate (heptahydrate)</td>
<td>0.50 mmol/L</td>
<td>0.00 mmol/L</td>
</tr>
<tr>
<td>D-Glucose</td>
<td>11.00 mmol/L</td>
<td>1.00 mmol/L</td>
</tr>
<tr>
<td>Glutathione (reduced)</td>
<td>1.50 mmol/L</td>
<td>0.50 mmol/L</td>
</tr>
<tr>
<td>Ascorbic Acid</td>
<td>1.00 mmol/L</td>
<td>0.00 mmol/L</td>
</tr>
<tr>
<td>L-Arginine</td>
<td>5.00 mmol/L</td>
<td>0.00 mmol/L</td>
</tr>
<tr>
<td>L-Citrulline Malate</td>
<td>1.00 mmol/L</td>
<td>0.00 mmol/L</td>
</tr>
<tr>
<td>Adenosine</td>
<td>2.00 mmol/L</td>
<td>0.00 mmol/L</td>
</tr>
<tr>
<td>Creatine Orotate</td>
<td>0.50 mmol/L</td>
<td>0.00 mmol/L</td>
</tr>
<tr>
<td>Creatine Monohydrate</td>
<td>2.00 mmol/L</td>
<td>0.00 mmol/L</td>
</tr>
<tr>
<td>L-Carnosine</td>
<td>10.00 mmol/L</td>
<td>0.00 mmol/L</td>
</tr>
<tr>
<td>L-Carnitine</td>
<td>10.00 mmol/L</td>
<td>0.00 mmol/L</td>
</tr>
<tr>
<td>Dichloroacetate</td>
<td>0.50 mmol/L</td>
<td>0.00 mmol/L</td>
</tr>
<tr>
<td>Insulin</td>
<td>10 mg/L</td>
<td>4.0 mg/L</td>
</tr>
</tbody>
</table>

### Western blotting

**Protein extraction:** 20 mg of kidney tissue was suspended in extraction buffer containing a protease inhibitor cocktail. Tissue was homogenized for 30 secs, centrifuged at 16,100 × g for 10 mins and supernatant was collected. Equal amounts of total protein (30 µg) from different samples were mixed with Laemmli sample buffer containing 5% β-Mercaptoethanol, and heated at 100°C for 5 mins.

**Eletrophoresis:** Proteins were resolved on 10% SDS-PAGE, and electro-blotted onto nitrocellulose membrane; proteins were identified using antibodies (anti-Caveolin, eNOS, vWF, and EPO) and chemiluminescence assays and band densities were normalized to beta-actin as described [4].

**Metabolic analysis**

Changes in pH and lactate, glucose metabolism, oxygen and carbon-dioxide concentrations (pO₂ and pCO₂) in Somah and UW were assessed at 0 (1 hour; vide supra), 6, 24 and 72 hour time-points during storage using VetScan iStat and VetScan VS2.

**Statistical analysis**

The measurements and data extraction were performed in blinded fashion. Comparison within two groups (UW vs. Somah, n=7 in each group) was conducted to evaluate effects of the two solutions. Quantified initial values of various assays were compared to subsequent time points within each group using one-way analysis-of-variance.
ANOV A, followed by Dunnet’s multiple comparison test, and t-test for comparative analysis between groups. Statistical significance was accepted at 95% confidence level (p<0.05). All values used were mean ± SEM unless otherwise indicated. All analyses were performed using GraphPad Prism 6 (v6.1). The authors had access to and take full responsibility for integrity of data. All authors have read and agree to the manuscript as written.

Results

Morphological and histological changes in explanted kidneys (Figure 1)

Gross morphology of kidneys: Kidneys were examined during 1-3 days of immersion static storage at 4°C. Kidneys stored in UW showed dusky and mottled appearance (Figure 1a), indicative of organ congestion. In contrast, kidneys stored in Somah appeared healthy, of uniform color and morphologically unaltered after 3 days storage (Figure 1d).

Histomorphology of kidneys: Irrespective of storage solution, there was no evident interstitial edema in all DCD kidneys at observed time-points, with well preserved overall structure of renal tissue (Figures 1b,1c,1e and 1f). The normal amorphous collection in tubular lumen was observed in proximal convoluted tubules (PCT) at all time-points and not increased with storage duration. Distal convoluted tubules (DCT) remained mostly clear of any debris at all time-points except at 72 hour where a minimal to moderate epithelial denudation was apparent in both UW and Somah-preserved kidneys (Figures 1c and 1f). Renal glomeruli exhibited normal architecture with normal appearing Bowman’s space and continuous parietal epithelium at all time-points, in both solutions (Figures 1b,1c,1e and 1f).

A gradual time-dependent loss of tubular epithelial nuclear heterogeneity with increased hyperchromaticity and occasional loss of cellular margins was observed in both UW and Somah-stored kidneys, indicative of tubular epithelial injury (Figures 1c and 1f). However, extent of these changes were significantly greater in UW-stored kidneys (p<0.05). On an average 3.5%, 24.4%, 39.7% and 37% in UW kidneys, and 4.4%, 6.2%, 10.9% and 11.6% in Somah kidneys, of the epithelial cell nuclei (of PCT/DCT’s) were severely hyperchromatic, at time 0, and 6, 24 and 72 hour storage time-points, respectively, suggesting that renal tubular epithelial cells in Somah-stored DCD kidneys were able to endure ischemia for a longer duration than UW-stored kidneys.

Metabolism in stored kidneys

DCD kidneys were evaluated for physiological/biochemical viability by assessing metabolic functions during extracorporeal storage; demonstrating differentially active metabolism in UW and Somah groups. While freshly reconstituted solutions started off with a more alkaline pH (more so in UW), a time-dependent fall in pH was apparent in Somah, but remained more acidic (6.8-7.2) compared to UW (7.5-7.4) (Figure 2A). Interestingly, while there was a temporal increase in glucose levels in UW solution, suggesting glycogenolysis, there was a comparative significant fall (p<0.05) in glucose levels in Somah, at and beyond 6 hour time-points, suggesting utilization of glucose present in Somah by kidney tissue/cells (Figure 2B). Inversely,
compared to Somah, there was a significantly greater (p<0.05) rise in lactate levels in UW at 72 hour time-point (Figure 2C).

Since in an open experimental system as ours, the solubility of atmospheric oxygen in solution is inversely related to temperature, and since oxygen consumption follows zero-order kinetics, and as Somah and UW both were supersaturated with oxygen at 4°C, with a pO\textsubscript{2} of 200 ± 13 mmHg at all times (Figure 2D), oxygen utilization by kidneys could not be clearly demonstrated. However, in contrast to UW, in which pCO\textsubscript{2} was significantly lower than in Somah throughout storage (7.28 ± 0.40 mmHg at 1-hour and 7.50 ± 0.48 mmHg at 72 hours), a significant initial increase in pCO\textsubscript{2} was observed in Somah (p<0.01) (from 5.8 ± 1.15 mmHg after initial immersion of kidney to 17.00 ± 0.45 mmHg) within 1 hour of DCD kidney storage, and remained consistently high during 72 hour period, indicating oxidative metabolic turnover right from the start of storage period (Figure 2E). It must be noted that pCO\textsubscript{2}, an indicator of dissolved CO\textsubscript{2}, does not change in Somah or UW solution in the absence of organs (not shown). Probability of bicarbonates present in Somah contributing to increase in pCO\textsubscript{2} was negated by the fact that HCO\textsubscript{3}– concentration remained mostly unaltered (4.86 mM/L) during 72 hour storage. Conversely, HCO\textsubscript{3}– concentration decreased from 6.30 to 4.33 mM/L in UW during 72 hour storage that may have contributed to the non-significant increase in pCO\textsubscript{2} (Figure 2E).

**High-energy phosphates in stored kidneys**

The UW kidney tissue ATP, Creatine Phosphate (CP) and total HEP concentrations decreased linearly and significantly (p<0.05) during hypothermic storage. HEP dropped by 20% within six hours (Figure 3), with a net decrease of 45% at the end of 72 hour storage. In contrast, ATP, CP and total HEP levels did not change appreciably in Somah kidneys, and any decrease in ATP was compensated with a parallel increase in CP concentration, thus maintaining superior total energy levels in renal tissue during storage.

**Markers of vascular endothelial function**

The expression of caveolin, eNOS, vWF and EPO was well preserved in DCD kidneys preserved in Somah, during the entire storage period (Figure 4). In contrast, while expression of caveolin protein was unaltered in UW-preserved kidneys as well, there was time-dependent decrease in expression of eNOS, vWF and EPO, indicative of possible renal tissue damage.

**Discussion**

Being one of the most resistant internal organs to ischemia, use of kidneys from deceased (DCD) donors for transplantation has been commonly practiced worldwide [11-13]. While DCD pool of kidneys is still highly underutilized, prognosis of DCD kidneys upon transplantation is associated with an observably increased incidence of DGF, PNF and lowered graft life expectancy in recipients. The results of our present study suggest that by maintaining energy levels in DCD kidneys during storage using Somah, subtle damage at cellular levels can be avoided; thus possible avenues to enhance DCD kidney storage and post-transplant prognosis could be uncovered.

Our results show that UW-preserved kidneys display patchy areas of discoloration within a few minutes of organ perfusion with UW, upon harvesting, and did not resolve during 72 hour storage. In contrast, Somah-preserved kidneys remained uniform in their external morphology at all time-points. While Somah has a viscosity close to that of normal saline, UW is characterized by a much higher viscosity due to hydroxyl-ethyl starch (HES; Table 1) [14,15]. While HES helps to prevent organ edema during storage, it increases solution density that can interfere with perfusion of all parts of the organ. Although gross morphology is not the best indicator of organ viability, it is probable that an uneven perfusion of kidneys during harvesting, despite copious use of solution for perfusion, could have resulted in patchy discoloration of kidneys, due to greater UW viscosity (Figure 1).

Remarkably, renal histopathology showed no gross ultrastructural changes in either cortical or medullary regions of UW or Somah-stored kidneys. However, higher magnifications revealed subtle changes in cellular nuclei, especially in tubular epithelial cells, characterized...
A significant loss of energy state (HEP) in explanted organs during storage leads to irreversible degenerative changes in the organ [21]. It is thus imperative for preservative solution to maintain organ in homeostasis and/or allow it to recover during extended storage by modulating organs metabolic pathways. Despite potential glycogenolysis-dependent increase in glucose concentration in UW during storage (Figure 2B), a significant depletion of renal HEP stores was apparent, in contrast to obvious preservation of energy state in Somah kidneys (Figure 3). This suggests that UW kidneys were highly catabolic, leading to loss of HEPs and tissue injury, (Figure 1). In contrast, greater oxidative phosphorylation of glucose in Somah-stored kidneys, facilitates greater HEP generation (for equivalent glucose molecules), than by anaerobic glycolysis alone [22] thus enhancing the organs energy state during storage. The low HEP levels observed in DCD UW-kidneys, could predictably result in delayed graft function (DGF) at the least, and even primary non-function (PNF). Therefore, despite the predominant use as preservation solution for explanted kidneys, UW may not provide optimal conditions for extended storage of DCD (or BHD) kidneys. In contrast, preservation in Somah may provide a viable alternative.

The blood vessels (capillaries) form the bulk of renal cortical tissue while tubular structures predominate in renal medulla. While glomerular tuft collapse within 6 hour storage in Histidine-Tryptophan-Ketoglutarate (HTK) solution has been reported [18], we did not find such drastic glomerular change in either Somah or UW stored kidneys, at any time-point (Figure 1). However, a steady decline in expression of eNOS (important in vasomotor function), von-Willebrand factor (vWF; marker for blood vessel endothelium) and erythropoietin (EPO; marker of specialized peritubular epithelial cells), during 72 hour DCD kidney storage in UW, suggests subtle damage to both vascular [23] and tubular structures [24] (Figure 4). This is consistent with our histological findings of increase in tubular nuclear hyperchromacity observed in UW-stored kidneys. In contrast, the expression of all investigated proteins including caveolin, eNOS, vWF and EPO were unaltered during the same periods of observation in Somah-preserved kidneys suggesting both cortical and tubular renal tissue preservation.

**Conclusion**

With the rising number of patients requiring renal transplantation, it has become extremely important to enhance availability and quality of DCD kidneys for transplant. While traditional preservation...
techniques are still associated with a major post-transplant inadequacy of DCD organ function, we provide preliminary evidence that use of Somah for static preservation of DCD kidneys may potentially decrease the incidence of DGF, PNF and improve graft life upon transplantation. Further studies involving different storage and perfusion conditions including renal transplantation are required for both BHD and DCD Somah preserved kidneys before our laboratory observations become a clinical reality.

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References


