The Effects of 6-Chromanol SUL-138 during Hypothermic Machine Perfusion on Porcine Deceased Donor Kidneys

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Abstract: Diminishing ischemia-reperfusion injury (IRI) by improving kidney preservation techniques offers great beneficial value for kidney transplant recipients. Mitochondria play an important role in the pathogenesis of IRI and are therefore interesting targets for pharmacological interventions. Hypothermic machine perfusion (HMP), as a preservation strategy, offers the possibility to provide mitochondrial-targeted therapies. This study focuses on the addition of a mitochondrial protective agent SUL—138 during HMP and assesses its effect on kidney function and injury during normothermic reperfusion. In this case, 30 min of warm ischemia was applied to porcine slaughterhouse kidneys before 24 h of non-oxygenated HMP with or without the addition of SUL—138. Functional assessment was performed by 4 h normothermic autologous blood reperfusion. No differences in renal function or perfusion parameters were found between both groups. ATP levels were lower after 30 min of warm ischemia in the SUL–138 group (n.s, \( p = 0.067 \)) but restored significantly during 24 h of HMP in combination with SUL—138. Aspartate aminotransferase (ASAT) levels were significantly lower for the SUL—138 group. SUL—138 does not influence renal function in this model. Restoration of ATP levels during 24 h of HMP with the addition of SUL in combination with lower ASAT levels could be an indication of improved mitochondrial function.

Keywords: SUL—138; machine perfusion; ischemia-reperfusion injury; kidney; DCD; mitochondria

1. Introduction

The global observatory on donation and transplantation calculated that less than 10% of the global need for donor organs is met [1]. This shortage has led to the use of sub-optimal quality organs donated from expanded criteria donors (ECD) and donation after circulatory death (DCD) donors [2]. Especially, kidneys donated from DCD donors are more susceptible to ischemia-reperfusion injury (IRI) [3] than those donated from brain death (DBD) donors, and this is reflected by higher incidences of delayed graft function (DGF) [4–6]. DGF has far-reaching consequences for the recipients of these kidneys since it requires a compulsory return of the patients to undergo haemodialysis until recovery of kidney function. Furthermore, the chance of acute cellular rejection and poorer long-term outcomes increases [7,8].

Preventing or diminishing IRI during the donation—and transplantation setting would be of great beneficial value by decreasing DGF rates and increasing kidney quality. Hypothermic machine perfusion (HMP), instead of static cold storage (SCS), as a preservation technique has already proven to be superior to SCS in terms of better—preserved
kidneys concerning reduced duration and incidence of DGF [4,9–11], and is therefore one of the several strategies that could be applied to reduce IRI.

Mitochondria play a pivotal role in the pathogenesis of IRI [12]. Mitochondrial reactive oxygen species (ROS) production is a fundamental early driver of IRI and is a nonspecific effect of the interaction of oxygen present during reperfusion with dysfunctional mitochondrial respiratory chains [13]. Since the proximal tubule compartments of kidneys contain large numbers of mitochondria, they are especially vulnerable to hypoxia [14,15]. Considering their crucial role, the prevention of mitochondrial injury during kidney preservation seems a logical approach.

In this study, we combined HMP with SUL—138, a compound in the group of 6—chromanols, which are shown to have a mitochondrial protective effect on cells during hypothermia [16]. SUL—138 is the (S)—enantiomer of SUL—109, an agent that has shown to maintain mitochondrial function of cells during hypothermia and rewarming through the activation of complexes I and IV of the electron transport chain [17,18]. In cold stored porcine kidneys, the addition of SUL—121 resulted in α1—adrenoceptor mediated vasodilation upon warm reperfusion which could be attributed to the (R)—enantiomers of SUL—121 (SUL—150) [19].

To exclude any flow—disturbance we choose the (S)—enantiomer of SUL—109 to study the potential of this 6—chromanol to decrease ischemia induced renal injury during dynamic kidney preservation. To resemble potential clinical application, we explored the effect of SUL—138 during 24 h HMP of porcine DCD kidneys followed by assessment of kidney function in a normothermic reperfusion model.

2. Materials and Methods

2.1. Animal Model

Porcine (female Dutch landrace pigs, 5–6 months, 100 kg on average) kidneys (349 ±42, 56) grams on average) were retrieved from two local abattoirs after the pigs were killed according to the standardised procedure of a sedative electrical shock followed by exsanguination. Blood was immediately collected in a container containing 25,000 IU of heparin (LEO Pharma A/S, Ballerup, Denmark). No animal ethics committee approval was necessary since slaughterhouse waste material was used for these experiments.

2.2. Experimental Design

A total duration of 30 minutes warm ischemia (WI) was chosen to induce ischemic injury as a model of DCD donation. Subsequently, the kidneys were preserved for 24 h by hypothermic machine perfusion with (n = 6) or without SUL—138 (n = 6). After the preservation period the kidneys were reperfused in an ex vivo normothermic machine perfusion (NMP) setup for four hours. A total of 12 kidneys were randomised into either the SUL or the vehicle group.

2.3. Hypothermic Machine Perfusion

During the 30 min of WI the kidneys were prepared for HMP by removing excess fat surrounding the kidney, ureter and renal artery. After 30 min kidneys were flushed with 180 mL of cold (4 °C) saline (Baxter BV, Utrecht, The Netherlands). Saline was supplemented with SUL—138 dissolved in dimethylsulfoxide (DMSO) with a final concentration of 1 × 10^{-4} mol/L in the SUL—138 group and with equimolar DMSO only in the vehicle group. After flushing a needle biopsy (23 mm) was taken (In vivo, Best, The Netherlands) from the cortex of the kidney and stored in sonification solution (SONOP containing 0.372 g EDTA in 500 mL 71% ethanol (v:v) and NaOH (pH 10.9)) and 4% buffered formaldehyde for further analysis. The renal artery was cannulated to enable connection to the HMP machine (Kidney Assist Transport, Organ Assist, Groningen, The Netherlands). A total volume of 500 mL University of Wisconsin machine perfusion solution (Belzers MP, Bridge to life Ltd., London, UK) supplemented with SUL—138 (end concentration: 1 × 10^{-4} mol/L) or vehicle (equimolar concentration of DMSO) was used as preservation solution in the
SUL—138 and vehicle group, respectively. HMP was initiated after the kidney was placed in the machine for a total duration of 24 h at a temperature of 4 °C. The perfusion was pressure-controlled with a mean arterial pressure of 25 mmHg. Samples of the perfusion solution were taken after 15, 60 min and 24 h. Furthermore, pressure, temperature and flow rates were continuously monitored.

2.4. Ex Vivo Normothermic Machine Perfusion to Assess Renal Function

Renal function was assessed in an isolated ex vivo normothermic machine perfusion (NMP) setup after the preservation time of 24 h according the model described earlier [20].

In brief: The renal artery and ureter were cannulated with a 12 and 8 French cannula, respectively. Just prior to attachment of the kidney to the NMP the remaining preservation solution was flushed away with 50 mL cold saline. Kidneys were weighed, and another biopsy was taken and stored as described above. To assess function, kidneys were attached to a specially designed organ chamber and were pressure-controlled perfused with a mean arterial pressure of 75 mmHg (Kidney Assist Transport, Organ Assist, Groningen, The Netherlands) at a temperature of 37 °C for 4 h. An oxygen mixture of 95% O₂ and 5% CO₂ was supplied to the oxygenator (Hilite LT 1000, Medos Medizin technik AG, Stolberg, Germany) at a fixed rate of 500 mL/min. A blood-based perfusion solution was used during NMP which consisted of 500 mL whole blood depleted of leukocytes by using a leukocyte filter (Bio R O2 plus, Fresenius Kabi, Zeist, The Netherlands) and 300 mL lactated Ringer’s (Baxter BV, Utrecht, The Netherlands), supplemented with 6 mg Mannitol (Sigma–Aldrich, St Louis, USA), 6 mg Dexamethasone (Centrafarm, Etten–Leur, The Netherlands), 10 mL 8,4% sodium bicarbonate (B Braun Melsungen AG, Melsungen, Germany), 90 mg creatinine (Sigma–Aldrich, St Louis, MO, USA), 1000 mg/200 mg Amoxicilline/Clavulanic acid (Sandoz BV, Almere, The Netherlands), 100 µL 20 mg/mL sodium nitroprusside (Sigma–Aldrich, St Louis, USA), 10 mL glucose 5% (Baxter BV, Utrecht, The Netherlands) and during NMP a constant infusion of an amino acid mixture (10% Aminoplasmal, Braun Melsungen AG, Melsungen, Germany), 2.5 mL 8,4% sodium bicarbonate and 17 IU Novorapid, (Novo Nordisk, Bagsvaerd, Denmark) was given at 20 mL/h. When arterial glucose levels dropped below 5 mmol/L, levels were correct with 5% glucose (Baxter BV, Utrecht, The Netherlands). At the end of NMP, biopsies were taken and stored as described above for further analysis.

2.5. Renal Function Testing

During NMP, renal flow rate and urine flow were monitored every 15 min. Perfusate and urine samples were taken after 15, 60, 120, 180 and 240 min for storage and blood gas measurements (ABL90 FLEX, Radiometer, Zoetermeer, The Netherlands).

Vascular resistance during HMP and NMP was calculated. The following formula was used:

\[
\text{Vascular resistance} = \frac{\text{Arterial pressure} \times \left( \frac{\text{min}}{\text{mL}} \right)}{\text{Arterial flow}}
\]

Creatinine clearance and fractional sodium excretion levels were calculated with concentrations of plasma and urine creatinine and sodium that were measured using routine procedures at the clinical chemistry lab of the University Medical Center Groningen (UMCG). Furthermore, the level of proteins in the urine was determined by the University Medical Center Groningen Department of Laboratory Medicine using standardised protocols on a modular analyser (Roche, Almere, The Netherlands).
2.6. Mitochondrial Function, Integrity and Damage

Renal oxygen consumption (QO$_2$) was calculated as an indication of the metabolic activity of the kidney. Venous and arterial pO$_2$ and saturation were measured for this purpose. The following formula was used:

$$\text{Oxygen consumption (mlO}_2\text{min}^{-1}\text{100gr)} = \left(\left(\text{Hb} \times 2.4794 + (\text{pO}_2\text{arterial} \times K)\right) - \left((0.024794 \times \text{Hb} \times \text{SO}_2\text{venous} ) + (\text{pO}_2\text{venous} \times K)\right) \times Q \right) / g \times 100$$

(2)

where Hb is the perfusate’s hemoglobin content in mmol/L, pO$_2$ is the venous or arterial partial oxygen pressure in kPa, K is the solubility constant of oxygen in water at 37 °C and equals 0.0225 (mL O$_2$ per kPa), SO$_2$ is the saturation in %, Q is the renal blood flow in L/min and g is the kidney weight in grams.

Total sodium reabsorption (T sodium) was calculated with the following formula:

$$T_{\text{Sodium}} (\text{mmol Sodium mmol O}_2\text{100gr)} = \left(\left(\text{CrCl} \times \text{P}_{\text{Na}}\right) - \left(\text{U}_{\text{Na}} \times \text{U}\right)\right) / g \times 100$$

(3)

where CrCl represents creatinine clearance (mL/min), P$_{\text{Na}}$ perfusate sodium concentration (mmol/L), U$_{\text{Na}}$ urine sodium concentration (mmol/L), U urine production (mL/min) and g is kidney weight.

Adenosine triphosphate (ATP) was measured in all biopsies that were taken during the experiment (after WI, preservation and NMP) with methods described previously [21]. ATP concentrations were expressed as µmol/g protein.

2.7. Kidney Injury Markers

Lactate dehydrogenase (LDH) was determined at the clinical chemistry lab of the UMCG according to standard procedures. Urinary N—acetyl—beta—D—glucosaminidase (uNAG) was determined following a protocol described previously by our lab [21,22]. ASAT levels were measured as indicator of mitochondrial damage. A standardised protocol by the clinical chemistry lab of the UMCG was used.

2.8. Oxidative Stress Due to Active Oxygenation

Thiobarbituric acid—reactive substances (TBARS) were analysed in preservation fluid perfusate (TBARS$\text{per fusate}$) and urine samples (TBARS$\text{urine}$), at all sample moments. The protocol for this analysis has been described in detail before [21]. In brief, the TBARS assay measures the level of products of lipid peroxidation present in the sample. In plasma, these products will consist mainly of malondialdehyde (MDA). TBARS concentrations are expressed in µmol/L. Total TBARS production was calculated with the following formula:

$$\text{TBARS production (IU)} = (\text{TBARS}_{\text{urine}} \times U) + (\text{TBARS}_{\text{per fusate}} \times (P + I))$$

(4)

U represents urine production (mL/min), P priming volume of the NMP setup (L) and I is the volume of infusion during NMP (L).

2.9. Statistics

Results are reported as means with standard deviation. Statistical analysis was performed with Graphpad Prism 7.02 (San Diego, CA, USA). The area under the curve (AUC) was calculated for the renal function parameters flow, creatinine clearance, fractional sodium excretion, proteins in urine, oxygen consumption, total sodium reabsorption. The AUC were also calculated for renal injury markers, such as ASAT and total TBARS production. For LDH and uNAG was chosen to show the increase over time. For LDH, the total duration of 4 h was chosen and for uNAG the period between 120 and 240 min of NMP was calculated since an increase in this tubular injury marker was seen during this
period. All values were tested for significance using a Mann-Whitney U test. $p < 0.05$ was considered to indicate statistical significance.

3. Results

3.1. Hypothermic and Normothermic Perfusion Parameters

During HMP no differences in flow rates were observed between the vehicle and SUL—138 group. Flow started with a steep increase during the first 20 min and slowly increased thereafter until the end of the 24 h preservation period (Figure 1A). In addition, no differences in vascular resistance were observed between the groups. The vascular resistance decreases during the first hour of HMP and then remains stable (Figure S1A in Supplementary Materials).

Renal flow during NMP was comparable between the vehicle and SUL—138 group (Figure 1B). During the first hour flow rates increased to approximately 80 mL/min/100 gr and then slowly decreased during 4 h of NMP, whereas the vascular resistance decreases during the first hour and then slowly increases during the remaining NMP time (Figure S1B in Supplementary Materials).

3.2. Renal Functionality during Normothermic Perfusion

No significant difference was seen between the vehicle and SUL—138 group in terms of glomerular function represented by creatinine clearance. Both groups show a similar trend over time (Figure 2A).
Immediately after start of NMP fractional sodium excretion is 100%, suggesting a total absence of tubular function. Over time both groups restored tubular reabsorption, decreasing FEsodium to approximately 30%. No differences are seen between the vehicle and SUL—138 group (Figure 2B).

Urine production during NMP was also comparable between the two groups. Urine production was the highest the first 15 min and decreased over time (Figure 2C).

Protein levels in the urine were comparable between the vehicle and SUL—138 group (Figure 2D).

Figure 2. Kidney functionality parameters during normothermic machine perfusion. Porcine kidneys were tested for kidney function for 4 h with NMP after 24 h HMP with (SUL—138 ●) and without (Vehicle ⊗) SUL—138. (A) Creatinine clearance rates, (B) Fractional sodium excretion, (C) Urine production, (D) Protein content in urine. NMP, normothermic machine perfusion; HMP, hypothermic machine perfusion. The data are shown as mean ± SD.

3.3. Mitochondrial Function (Integrity) and Metabolic Activity

Comparable oxygen consumption rates were found for both groups. These rates slowly increased over the four-hour reperfusion time from 1 mL O₂/min/100 gr to 2 mL O₂/min/100 gr (Figure 3A). Total sodium reabsorption was equal between both groups during NMP (Figure 3B).

Arterial (Figure 3C) lactate levels were comparable in both groups during the reperfusion.

ATP levels measured after warm ischemia (timepoint 0.5) were not significantly higher in the vehicle group compared to the SUL group ($p = 0.067$). ATP levels significantly increased during 24 h of preservation in the SUL group. No significant differences between groups were found in ATP values after 24 h preservation (timepoint 24.0) and after NMP (timepoint 28.0) (Figure 3D).
Figure 3. Parameters dependent on mitochondrial function. Parameters were assessed in porcine kidneys that were preserved with 24 h of non-oxygenated hypothermic machine perfusion with (SUL—138 *) and without (Vehicle ⊗) the addition of SUL—138, (A) Oxygen consumption rates during NMP, (B) Total sodium reabsorption rates during NMP, (C) Arterial lactate levels during NMP, (D) ATP levels were measured during the experiment after 30 min warm ischemia (timepoint 0.5), after 24 h HMP (timepoint 24) and at the end of 4 h NMP (timepoint 28), * p < 0.05 significant increase in ATP values during 24 h HMP with the addition of SUL—138. NMP, normothermic machine perfusion; ATP, adenosine triphosphate; WIT, warm ischemic times; ASAT, Aspartate Aminotransferase; AUC, Area under the curve. The data are shown as mean ± SD.

3.4. Kidney Injury and Oxidative Stress Markers

The total increase of LDH as a sign of leakage of cell content was comparable between the groups. The tubular injury marker, urinary N—acetyl—beta—D—glucosaminidase (uNAG), was not significantly lower for the SUL—138 group (p = 0.063) (Figure 4A,B).

The AUC for ASAT was significantly lower in the SUL—138 group compared to the vehicle. No differences were seen regarding oxidative stress represented by similar total TBARS production levels (Figure 4C,D).
mic machine perfusion on kidney function. We found that the addition of SUL—138 during hypothermic preservation did not result in improved renal function during 4 h of normothermic reperfusion. We did find a significant increase in ATP levels during HMP in the SUL—138 group and a lower release of ASAT during reperfusion.

In this study we used oxygen consumption, total sodium reabsorption and ATP levels as indirect markers for mitochondrial function. In this experiment, we were not able to detect a clear effect of SUL—138 on ATP levels. There was a tendency of lower ATP levels after 30 min of warm ischemia in the SUL—138 group. Since SUL—138 was already present in the first flush out it could be that the observed ATP depletion can be attributed to the addition of the 6—chromanol. Since we did not measure ATP levels directly after death, we cannot be sure about this effect of SUL. However, we compared these ATP levels to a historical cohort of kidneys in which we measured ATP at the same moment. The addition of the 6—chromanol. Since we did not measure ATP levels directly after death, it could be that the observed ATP depletion can be attributed to the

**4. Discussion**

The aim of this study was to evaluate the effect of SUL—138 during 24 h of hypothermic machine perfusion on kidney function. We found that the addition of SUL—138 during hypothermic preservation did not result in improved renal function during 4 h of normothermic reperfusion. We did find a significant increase in ATP levels during HMP in the SUL—138 group and a lower release of ASAT during reperfusion.

Warm and cold ischemic periods are unavoidable during a donation—and transplantation setting and this results in ischemia—reperfusion injury (IRI). ROS production by mitochondria is a well-known early effect of IRI [13] and preventing this would be a valuable target for diminishing IRI. SUL compounds have shown to be protective against cold—induced ischemia and mitochondrial dysfunction [18,19]. Both SUL compounds 109 and 121 have shown to increase ATP levels after hypothermic preservation followed by a period of rewarming of adipose—derived stem cells and in rat kidneys [17,18]. Furthermore, both compounds reduced the production of ROS in various hypothermia/rewarming and disease models that are characterised by mitochondrial dysfunction [17,23,24]. SUL—138, the compound used in this study, has already shown to maintain cell growth and morphology during hypothermic storage of various cell lines in vitro (3T3—LI, HUVEC, HEK293 and NRK—52E) [16] but has no published data on its effect on mitochondrial integrity.

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we cannot be sure about this effect of SUL. However, we compared these ATP levels to a historical cohort of kidneys in which we measured ATP at the same moment. The kidneys flushed with SUL have a significant lower ATP content compared to this historical cohort \((n = 47, p = 0.0003)\).

After 24 h of HMP the relative increase ATP levels was larger in the SUL—138 treated kidneys indicating a mitochondrial protective action. This is in line with earlier findings in cells and during hypothermia induced renal injury. We have reported earlier using the same model as presented here, that the presence of oxygen during HMP leads to significant higher ATP levels in kidneys [20]. In addition, other studies have demonstrated that the addition of oxygen during HMP of both kidneys and livers is beneficial to support cellular respiration and subsequent ATP production [25–30]. The lack of oxygen during HMP in this study may have lead to the underestimation of the potential of SUL—138 as additive during HMP. For the current experiment we chose not to include oxygen during HMP since non—oxygenated HMP is the current clinical standard [11]. In other studies where the mitochondrial protective capacity of SUL compounds have been shown was oxygen present [18]. One could therefore assume that the lack of efficacy of the SUL compound may be due to the absence of oxygen in our model. Which is likely as SUL activates complex IV of the mitochondrial membrane transporter, which utilises oxygen as final electron acceptor. The significant restoration of ATP values in the SUL-138 group during 24 h HMP, without the addition of oxygen could therefore be a sign of improved mitochondrial integrity.

Another indication of mitochondrial protection by SUL—138 is presented by the significant lower ASAT levels during NMP in the SUL-group compared to the vehicle group. This observation is in line with the earlier study in hypothermic rats where a decrease in ASAT levels were seen in combination with increased cortical ATP levels in SUL—121 and SUL—109 treated rats [18].

Despite the potential protection of mitochondrial integrity by SUL compounds found in this study and in other studies, [17,18] we were not able to show functional differences. This is also in line with the rat study where creatinine levels equally increased after hypothermia induced renal injury. This does however not imply that the use of 6—chromanol compounds are futile in renal transplantation. Recently a direct correlation between mitochondrial membrane potential and delayed graft function following kidney transplantation was found. [31] Our model only allows a short term follow up and is not suitable to study DGF since all kidneys show immediate function.

In conclusion, we were not able to show direct functional improvement of the addition of SUL—138 during non—oxygenated HMP in terms of renal function during short-term normothermic reperfusion. The ATP levels in combination with significant lower ASAT levels found in this study could be an indication that mitochondria are better preserved in the SUL—group. However, more in—depth assessment on mitochondrial function markers are necessary to support this finding. In addition, the use of SUL compounds in oxygenated HMP could be considered especially given the recent study in which DCD kidneys were better preserved. [32] Furthermore, it could be of interest to study whether SUL would be able to shorten the minimum (oxygenated) HMP duration for its maximum restorative effect.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/transplantology2030031/s1, Figure S1: Vascular resistance during preservation and testing. Porcine kidneys were preserved with HMP with SUL-138 (•) and without the addition of SUL-138 (Vehicle ⊗) for 24 h. (A) Vascular resistance during HMP, (B) Vascular resistance during normothermic functionality testing. HMP, hypothermic machine perfusion. The data are shown as mean ± SD.

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**Informed Consent Statement:** Not applicable.

**Data Availability Statement:** All data is presented in this study.

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**Conflicts of Interest:** The authors declare no conflict of interest. Pieter C Vogelaar is research scientist and Guido Krenning is the chief scientific officer at Sulfateq (Groningen, The Netherlands), a company that owns patents on SUL-138 and produces and markets these and similar compounds.

**References**


