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# Metabolic differences between cold stored and machine perfused porcine kidneys:

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### **Accepted Manuscript**

Metabolic differences between cold stored and machine perfused porcine kidneys: A <sup>1</sup>H NMR based study

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1	Metabolic differences between cold stored and machine perfused porcine
2	kidneys: A <sup>1</sup> H-NMR based study.
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24	Key words: hypothermic machine perfusion, kidney, transplantation,
25	metabolism, NMR, organ preservation.

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32	Abstract
33	Hypothermic machine perfusion (HMP) and static cold storage (SCS) are the
34	two methods used to preserve deceased donor kidneys prior to transplant
35	This study seeks to characterise the metabolic profile of HMP and SCS
36	porcine kidneys in a cardiac death donor model.
37	
38	Twenty kidneys were cold flushed and stored for two hours following retrieval
39	Paired kidneys then underwent 24 hours of HMP or SCS or served as time
40	zero controls. Metabolite quantification in both storage fluid and kidney tissue
41	was performed using one dimensional <sup>1</sup> H-NMR spectroscopy. For each
42	metabolite, the net gain for each storage modality was determined by
43	comparing the total amount in each closed system (i.e. total amount in
44	storage fluid and kidney combined) compared with controls. 26 metabolites
45	were included for analysis.
46	
47	Total system metabolite quantities following HMP or SCS were greater for 14
48	compared with controls (all p<0.05). In addition to metabolic differences with
49	control kidneys, the net metabolic gain during HMP was greater than SCS for
50	8 metabolites (all p<0.05). These included metabolites related to central
51	metabolism (lactate, glutamate, aspartate, fumarate and acetate).
52	
53	The metabolic environments of both perfusion fluid and the kidney tissue are
54	strikingly different between SCS and HMP systems in this animal model. The
55	total amount of central metabolites such as lactate and glutamate observed in
56	the HMP kidney system suggests a greater degree of de novo metabolic

57	activity than	in the SCS	system.	Maintenance	of	central	metabolic	pathways
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may contribute to the clinical benefits of HMP.

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61	Introduction
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Hypothermic Machine Perfusion (HMP) and Static Cold Storage (SCS) are the two methods of kidney preservation that are used widely in clinical practice during the time period between organ retrieval and implantation [16]. A key concept for both preservation modalities is that cellular metabolism, and therefore cellular metabolic requirements, are minimised in these hypothermic conditions and the rate of metabolism reported to be about 5-8% at temperatures below 4°C [29] with a similar decrease in oxygen requirement [1].

The superiority of HMP over SCS is well documented [4,17,22,23,27] but the mechanisms by which this occurs are not clear. Improvement in flow dynamics, with fall in the intra-renal resistance is likely to be one factor but the

additional metabolic support derived from the circulation of nutrient-containing

perfusion fluid may also help preserve organ function and have a beneficial

effect [7,30].

Metabolomic analyses of preservation fluid during HMP using 1D-<sup>1</sup>H-NMR (One-dimensional proton nuclear magnetic resonance) spectroscopy, by groups including our own, have demonstrated this to be reproducible and highly specific for metabolite identification and quantification [2,10,24]. However, surprisingly, to our knowledge there are no studies that have sought to compare the metabolomic profiles, or metabolome, of HMP and SCS kidneys.

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Porcine kidneys are widely used in transplantation studies owing to their similar physiological and anatomical properties to human organs [9,11]. In addition, the metabolic profiles during periods of HMP for porcine and human kidneys are comparable [24], with a correlation between metabolite profiles during storage and post transplant outcome [2]. For HMP preserved human kidneys, the metabolic profile from perfusates of immediate graft function kidneys differs from those with delayed function [10] and reinforces the concept that significant metabolism occurs during HMP and that metabolism reflects functional outcome.

The aims of this study were twofold. Firstly, to determine the distribution of metabolites between the two different compartments (fluid and tissue) during the organ preservation period. Secondly, to determine the total amount of each metabolite within HMP and SCS kidneys systems after 24 hours of organ storage and through comparison with control kidneys, the metabolic changes that occur.

1	03	Methods	
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### Animal Research

Abattoir/slaughterhouse pig kidneys (F.A. Gill, Wolverhampton, UK) were used and no animals were sacrificed solely for the purposes of this study, negating any need for ethical board approval. Experiments were performed on 22-26 week old male 'bacon weight' pigs, weighing 80-85kg. All experiments were performed following the principles of laboratory animal care according to NIH standards. Animals were sacrificed by electrical stunning and exsanguination. Initial organ preservation was performed following organ retrieval and occurred within 14 minutes of death, replicating deceased cardiac death (DCD) donor conditions. Kidneys were cold flushed (4°C) with 1L SPS-1 (UW) solution at a pressure of 100mmHg. Organs were then stored at 4°C in SPS-1 for 2 hours to replicate the clinical period of organ transportation.

### Experimental groups

Paired kidneys were randomly allocated to receive either HMP or SCS for 24 hours. HMP kidneys were perfused with 1L of KPS-1 using the LifePort Kidney Transporter 1.0 (Organ Recovery Systems, Chicago, IL). (Perfusion pressure 30mmHg). SCS Kidneys were submerged in 1L of fresh chilled SPS-1 solution with a surrounding ice bath. Preservation fluid was sampled for each kidney at baseline and 2, 4, 8, 12, 18, and 24 hours. After 24 hours,

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128	organs were rapidly dissected and tissue samples (1cm <sup>3</sup> sections) flash frozen
129	and stored (-80°C). All experiments were performed in a cold room (4°C) to
130	ensure consistency.
131	
132	
133	Control kidneys
134	
135	To ascertain metabolism during SCS or HMP storage conditions, baseline
136	values prior to storage conditions were needed (time 0). Large volume tissue
137	sampling precludes effective organ perfusion and therefore 'Control kidneys'
138	were used to establish baseline metabolite levels. These were (n=6) flushed
139	and cold transported in identical fashion to experimental kidneys and tissue
140	samples obtained as described above (i.e. not subjected to 24hrs of SCS or
141	HMP).
142	
143	Sample processing and metabolite quantification
144	
145	NMR samples were prepared from storage fluid by mixing 150 $\mu$ L of 400 mM
146	(pH 7.0) phosphate buffer containing 2 mM DSS (4,4-dimethyl-4-silapentane-
147	1-sulfonic acid) and 8mM imidazole with 390 $\mu L$ of each fluid sample and 60
148	$\mu L$ of deuterium oxide (D2O) to reach a final phosphate buffer concentration
149	of 100 mM and a final DSS concentration of 500 μM. After mixing, the 600 μL
150	samples were pipetted into 5mm NMR tubes, sonicated and centrifuged.
151	Technical replicates of samples (x3) were prepared for each timepoint.

For cell extract studies, 500mg of renal cortex was manually cryohomogenised in liquid nitrogen. 5.1ml of both methanol (-80°C) and chloroform was added to the powdered tissue and samples diluted with 4.65ml of  $dH_20$  at 4°C. Samples were centrifuged to separate into polar and non-polar layers and 1.5ml of the upper polar layer was dispensed into a cryovial and dried. Three technical replicates were performed for each tissue sample. Dried polar residue was then dissolved in 390 $\mu$ L of  $dH_20$  and 210  $\mu$ L of buffer solution as described above.

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The protocol used for <sup>1</sup>H-NMR analysis has been described previously [10,24]. Briefly, this entailed processing on a Bruker AVII 500 MHz spectrometer, acquisition of one dimensional spectra and then metabolite identification and quantification using Matlab based 'Metabolab' software [18] and Chenomx 8.1 (ChenomxInc) software respectively. Metabolites were deemed to be present if they exhibited non-ambiguous spectral patterns or their presence deemed biologically plausible and confirmed on ultra performance liquid chromatography mass spectrometry. Any metabolites that were present in different concentrations in the SCS and HMP fluid (e.g. glucose, gluconate, mannitol, adenine, adenosine etc.) were excluded from comparative analysis. Metabolite quantifications were corrected to allow for sample dilution with sample buffer. When determining concentrations of metabolites using Chenomx, the researchers were blind to the storage group. Quantification of the total amount of metabolite in the storage fluid, tissue and total system was calculated based upon the weight of the kidney at time of sample acquisition and final volume of storage fluid.

Statistical analysis

For each timepoint, three results were obtained (technical replicates) and the median value used. For comparison of SCS and HMP conditions, analysis was performed using Wilcoxon paired signed rank test (two tailed) as one kidney from each pair was subjected to each condition and normality was not consistent on prior analysis. When comparing SCS or HMP with control kidneys, Mann-Whitney u test (two tailed) was used, as these were non-paired samples. Data were reported as median concentrations and interquartile (IQ) range. All analysis was performed using GraphPad Prism version 6.00 for Mac OS X, GraphPad Software, La Jolla California USA, with p<0.05 deemed to be indicative of statistical significance.

190	Results
191	
192	Metabolic optimisation of cadaveric kidneys is a potential target to improve the
193	function of kidneys for transplantation. This study seeks to establish the
194	degree of metabolism, if any, that occurs in the two widely used methods of
195	kidney organ storage prior to transplantation (HMP and SCS).
196	
197	The total quantity of each metabolite after 24 hours of either HMP or SCS was
198	calculated using <sup>1</sup> H-NMR methods and compared with control organs to
199	determine the net metabolic change during each storage method.
200	
201	We found evidence of metabolite production for both storage modalities with
202	14 metabolites present in significantly greater quantities in the HMP or SCS
203	system compared with controls (all p<0.05) (table 1)(Fig 1, Fig 1(Suppl)).
204	There were significantly more metabolites with a net increase in the HMP
205	system (13/14) compared with the SCS system (7/14) (p=0.033).
206	
207	Table 1. Total amount of metabolite present in each of the storage modalities
208	at time zero (controls) or after 24 hours of preservation (SCS or HMP). Data
209	reported as Median (Interquartile Range), unless stated otherwise. Statistical
210	test: $^{\Psi}$ Mann-Whitney u test (two tailed) $^{\#}$ Wilcoxon paired signed rank test
211	(two tailed). *Significant at p<0.05.
212	
213	Fig. 1. Metabolites significantly elevated in the HMP system compared with
214	both SCS and control kidneys. Metabolite levels represent total amounts

215	(mmol) in the storage fluid, kidney tissue and entire system for porcine
216	kidneys after 24hrs of HMP or SCS or time zero controls. Highly significant
217	(**p<0.01) and significant (*p<0.05) differences between HMP system versus
218	both controls and SCS kidneys.
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219	
220	
221	Eight of the metabolites were significantly elevated in the HMP system
222	compared with both the control and SCS systems (all p<0.05), indicating a
223	greater degree of metabolite production. These included lactate, glutamate,
224	aspartate, fumarate, acetate, myo-inisitol, niacinamide and formate (Fig 1).
225	
226	Despite the additional 24 hours of organ preservation, albeit in static
227	conditions, the amount of lactate in the SCS system was comparable to
228	controls (1.37 vs 1.11mmol p=0.138). However the amount in the HMP
229	system (2.13mmol) was almost twice the amount of either controls or SCS
230	systems (p=0.002 and p=0.031). However, despite greater amounts overall,
231	the amount present in the HMP tissue (0.76mmol) was actually lower than
232	SCS tissue (1.14mmol) or control tissue (1.11mmol) (p= 0.031 and p=0.002
233	respectively), reflective of lower intracellular concentrations for HMP kidneys.
234	
235	The distribution of metabolites between the extracellular storage fluid and
236	tissue samples for both storage conditions are detailed in table 2. As
237	expected, there were greater quantities of metabolites in the circulating HMP
238	fluid compared with the static conditions of SCS at most time-points. After 24
239	hours, the total amount of metabolite in the perfusate for the HMP kidneys

240	was significantly greater than the SCS group for $(21/26 = 80.8\%)$ of
241	metabolites. Whilst concentrations rose most rapidly in the first 2 hours of
242	perfusion and therefore may be in part due a metabolite washout
243	phenomenon, there was an increase in most metabolites over sequential
244	timepoints as would be expected with on-going production (fig 2a-c).
245	
246	Table 2. Metabolites present in tissue and storage fluid in HMP or SCS kidney
247	systems at 24 hours. Data reported as Median (Interquartile Range), unless
248	stated otherwise. Statistical test: #Wilcoxon paired signed rank test (two
249	tailed). *Significant at p<0.05.
250	Fig. 2. Concentration of metabolites in the storage fluid of SCS and HMP
251	kidneys over 24 hour time period for four example metabolites. Values plotted
252	as median (interquartile range).
253	
254	
255	Reduced glutathione is a constituent of both KPS-1 (used in HMP) and SPS-1
256	(used in SCS) fluids at equal concentrations. Whilst this remained at stable in
257	the SCS environment, the glutathione was clearly consumed by the HMP
258	group and after 8 hours concentrations were 17.6 fold higher in the SCS fluid
259	(1.60mM vs. 0.091mM, p=0.001) (fig 2d). Despite apparent organ uptake of
260	reduced glutathione, there was no evidence of this in the tissue samples from
261	either group.
262	



265	Discussion
266	
267	The aim of this study was to determine any metabolic differences between the
268	two clinically used methods of organ storage in this animal model.
269	
270	Whilst the calculation of the total amount of metabolite within the system does
271	rely on several assumptions (complete metabolite extraction from tissue and
272	metabolite homogeneity within tissue), we felt this was imperative to draw
273	meaningful comparison between groups and enables the calculation of net
274	metabolite production/consumption in these two closed systems (HMP and
275	SCS).
276	
277	Although the storage fluid used in each experimental group differs (most
278	notably absence of glucose in the SCS fluid) and therefore caution should be
279	exercised in attributing any differences merely to the parameters of storage
280	(i.e. HMP or SCS), this study was designed to assess metabolism during the
281	two clinically used organ preservation techniques, not merely the storage
282	modality in isolation.
283	
284	This study clearly demonstrates the presence of major central metabolites
285	such as lactate, glutamate, fumarate, aspartate and acetate at greater levels
286	in the HMP system compared with both controls and SCS (fig. 2). This
287	strongly suggests that these metabolites are being produced during HMP.
288	Furthermore, the accumulation of these metabolites into the circulating

289	perfusion fluid demonstrates effective homeostatic mechanisms are active to
290	prevent over accumulation within the local cellular environment.

The list of metabolites reported in this study is not exhaustive and is a limitation of this study. Some interesting substrates (eg glucose) were excluded as this is only present in one of the storage fluids (KPS-1). For others the 1D <sup>1</sup>H NMR spectral pattern is either ambiguous or can be hidden under more domineering peaks from other compounds.

The increased total lactate in the HMP system is likely to reflect increased glycolysis in the HMP model. Although new glycolytic activity of the glucose within the HMP fluid is one likely contributor, this is unlikely to the singular cause. This is supported by evidence that the HMP fluid glucose concentrations did not decrease during the study period and replicates findings from previous human studies [10]. However conversion of a proportion of perfusion fluid glucose into lactate through glycolytic pathways has been corroborated by work demonstrating activity of these pathways using <sup>13</sup>C labelled glucose tracers[25].

The net gain of glutamate, fumarate, aspartate and acetate during HMP is also intriguing. Whilst identification of responsible metabolic pathways is difficult to ascribe solely with <sup>1</sup>H NMR studies, one explanation could be increased oxygen dependent tricarboxylic acid (TCA) cycle activity. Although uniform upregulation of TCA intermediates would support this theory, as discussed, many are not easily identifiable using <sup>1</sup>H NMR methods[6] and are

rarely found in equipoise even *in vivo* [14]. Several (<sup>13</sup>C) NMR studies have reported glutamate as a valid marker of TCA activity [3,5,20].

For some metabolites, the total system amounts for HMP and SCS kidneys were comparable to the controls, suggesting that either *de novo* production does not occur during the 24 hour preservation or that consumption mirrors production (table 1 *supplementary*). However, for metabolites with similar total amounts, the compartment in which they were located varied per metabolite. Some metabolites were entirely contained within the HMP kidney tissue (e.g. ADP, AMP, NAD+) and presumably in the intracellular compartment. Other metabolites were evident in both the tissue and storage fluid but at higher concentrations in the HMP fluid. These discrepancies in metabolite location further highlight that cellular transport processes are active in this environment but that movement of metabolites into the extracellular fluid is not indiscriminate.

Reduced glutathione is a constituent of the preservation fluid KPS-1 and is thought to play a role in the removal of Reactive Oxygen Species (ROS) generated during metabolism [19] In contrast to the SCS kidney, there is a rapid decrease in the concentration of glutathione in the preservation fluid of HMP stored kidneys and is about 5% of the SCS values after 8 hours (fig 1c.). The rate of glutathione depletion observed in this study is similar to a previously reported animal model [28] and is likely to reflect cellular uptake of this protective antioxidant. Interestingly, glutathione concentration remained relatively constant in the SCS kidney group. This further reinforces the

concept that HMP exerts its beneficial effects, at least in part, by providing access to the cellular components of the kidney during perfusion. Absence of reduced glutathione in tissue demonstrates that not only is this protective antioxidant readily absorbed by the kidney during perfusion but that even after a few hours it is not longer available in the reduced state.

Although the number of organs in each experimental group is small (n=7), it is comparable to other porcine kidney transplant reports [8,12,15,21,26,30]. To improve validity, samples were processed in triplicate at each timepoint and over 250 NMR spectra were analysed. One strength of this study is that the kidneys stored by HMP or SCS were paired, i.e. from the same pig, thus minimising any metabolic differences arising from polymorphism in cellular mediators of porcine metabolism. Although this approach does not provide functional outcome information for the preserved organ, previous studies have demonstrated good function for otherwise healthy porcine organs stored by either SCS or HMP for similar time periods[2,8,13,15,21,26].

This study demonstrates that in a porcine model, the distribution and amounts of metabolites vary significantly with the storage method (HMP or SCS). The net gain of many central metabolites during HMP conditions further supports the notion that significant metabolism occurs during HMP and this may contribute to the beneficial role of machine perfusion.

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470	Fig. 1. (suppl) Metabolites with comparable total amounts between SCS and
471	HMP systems but significantly elevated compared with controls. Metabolite
472	levels represent total amounts (mmol) in the storage fluid, kidney tissue and
473	entire system for porcine kidneys after 24hrs of HMP or SCS or time zero
474	controls. Highly significant (**p<0.01) and significant (*p<0.05) differences
475	between HMP and SCS systems versus controls.
476	Fig. 2. (suppl) Chemical shift used for metabolic quantification. Localised
477	spectral plots for metabolites of interest with shaded figures illustrating
478	metabolite quantification via best-fit analysis using Chenomx metabolite
479	database.
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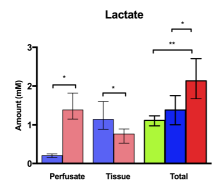
	Storage Modality			p-Values		
	Control System	SCS System	HMP System	Control vs	Control vs	SCS vs
	(mmol)	(mmol)	(mmol)	$scs^{\Psi}$	нмр Ψ	HMP <sup>#</sup>
Glutamate	1.54 (1.12- 1.84)	1.38 (1.11- 1.66)	3.97 (3.69- 4.71)	0.731	0.002*	0.031*
Myoinositol	1.18 (1.16- 1.19)	1.29 (1.01- 1.52)	2.16 (1.85- 2.41)	0.731	0.002*	0.031*
Lactate	1.11 (0.976- 1.23)	1.38 (1- 1.75)	2.13 (1.67- 2.71)	0.138	0.002*	0.031*
Hypoxanthine	0.454 (0.356-	0.710 (0.641- 0.762)	1.05 (0.909- 1.17)	0.001*	0.002*	0.156
	0.515)					
Formate	0.442 (0.274-	0.643 (0.589- 0.779)	0.842 (0.750- 0.943)	0.101	0.004*	0.031*
	0.638)					
Acetate	0.210 (0.206-	0.296 (0.253-0.301)	0.552 (0.494-0.654)	0.234	0.041*	0.031*
	0.212)					
Alanine	0.302 (0.243-	0.486 (0.339- 0.499)	0.501 (0.368- 0.558)	0.035*	0.015*	0.313
	0.360)					
Succinate	0.283 (0.267-	0.462 (0.312- 0.52)	0.434 (0.307- 0.541)	0.001*	0.015*	0.844
	0.297)					
Inosine	0.588 (0.561-	1.08 (0.885- 1.12)	0.185 (0.146- 0.233)	0.001*	0.002*	0.031*
	0.628)					
Aspartate	0.114 (0.104-	0.107 (0.0879- 0.11)	0.165 (0.140- 0.215)	0.234	0.041*	0.031*
	0.118)					
Leucine	0.0476 (0.0441-	0.0667 (0.0513-	0.0693 (0.0495-	0.014*	0.026*	0.688
	0.0517)	0.0820)	0.0773)			
Niacinamide	0.0196 (0.0181-	0.0289 (0.0243-	0.0490 (0.0420-	0.001*	0.002*	0.031*
	0.0207)	0.0319)	0.0557)			
Tyrosine	0.0262 (0.0217-	0.0434 (0.0339-	0.0387 (0.0332-	0.001*	0.013*	0.438
	0.0302)	0.0462)	0.0431)			

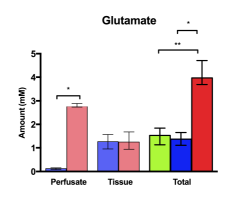
Fumarat	te	0.00412 (0.00339-	0.00308 (0.00145-	0.0133 (0.0077-	0.064	0.002*	0.031*
		0.00418)	0.00348)	0.0212)			

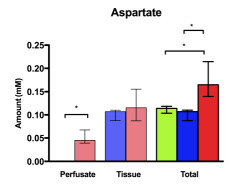


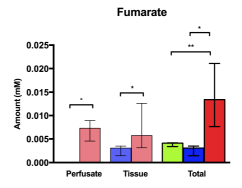
	Storage	Total storage fluid amount	p-value <sup>#</sup>	Total kidney tissue amount	p-Value <sup>#</sup>
		(mmol)		(mmol)	
Glutamate	SCS	0.0812 (0.125- 0.155)	0.0312*	0.952 (1.26- 1.58)	0.6875
	НМР	2.72 (2.75- 2.89)		0.94 (1.24- 1.68)	
Myoinositol	SCS	0.316 (0.399- 0.879)	0.0625	0.596 (0.676- 0.853)	0.5625
	НМР	1.05 (1.25- 1.38)		0.653 (0.816- 1.3)	<b>Y</b>
Lactate	SCS	0.153 (0.205- 0.245)	0.0312*	0.89 (1.14- 1.59)	0.0312*
	НМР	1.15 (1.38- 1.82)		0.521 (0.755- 0.895)	
Hypoxanthine	SCS	0.294 (0.328- 0.404)	0.0312*	0.289 (0.407- 0.424)	0.0625
	НМР	0.705 (0.781- 0.867)		0.189 (0.258- 0.31)	
Formate	SCS	0.132 (0.136- 0.186)	0.4375	0.434 (0.486- 0.545)	0.0312*
	НМР	0.151 (0.16- 0.169)		0.688 (0.599- 0.774)	
Acetate	SCS	0.073 (0.0808- 0.0912)	0.0312*	0.167 (0.201- 0.229)	0.0312*
	НМР	0.239 (0.257- 0.331)	Y	0.252 (0.289- 0.344)	
Alanine	SCS	0.0465 (0.0643- 0.0815)	0.0312*	0.303 (0.415- 0.435)	0.0312*
	НМР	0.253 (0.306- 0.358)		0.116 (0.187- 0.207)	
Succinate	SCS	0.0104 (0.0155- 0.0184)	0.0312*	0.298 (0.446- 0.498)	0.0312*
	НМР	0.104 (0.131- 0.208)		0.203 (0.294- 0.347)	
Inosine	SCS	0.703 (0.852- 0.961)	0.0312*	0.145 (0.182- 0.201)	0.0312*
	НМР	0.0877 (0.108- 0.128)		0.058 (0.0723- 0.109)	
Aspartate	SCS	-	0.0312*	0.0879 (0.107- 0.11)	0.3125
	НМР	0.039 (0.0452- 0.0682)		0.0874 (0.115- 0.155)	
Leucine	SCS	0.00442 (0.00506- 0.00761)	0.0312*	0.0486 (0.0591- 0.0775)	0.0312*
	НМР	0.0285 (0.038- 0.0468)		0.0222 (0.0304- 0.0318)	
Niacinamide	SCS	-	0.0312*	0.0243 (0.0289- 0.0319)	0.0938
	НМР	0.0221 (0.028- 0.0282)		0.0194 (0.0221- 0.0278)	

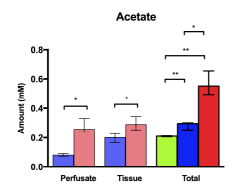
Tyrosine	SCS	0.00336 (0.0071- 0.00843)	0.0312*	0.0306 (0.0371- 0.0391)	0.0312*
	НМР	0.0197 (0.0228- 0.0276)		0.0112 (0.0143- 0.0171)	
Fumarate	SCS	-	0.0312*	0.00145 (0.00308- 0.00348)	0.0312*
	НМР	0.00456 (0.00737- 0.00895)		0.00314 (0.00574- 0.0126)	

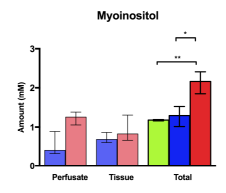


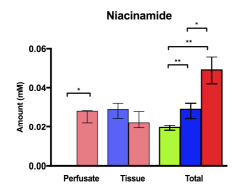


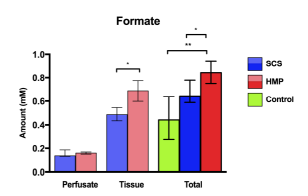


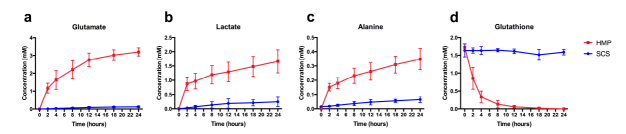












Metabolic differences between cold stored and machine perfused porcine kidneys: A <sup>1</sup>H-NMR based study.

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