Preclinical development of the AMC-HepaRG-Bioartificial Liver

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CHAPTER 6

HepaRG-progenitor cell derived hepatocytes cultured in Bioartificial Livers are protected from healthy- and acute liver failure-plasma induced toxicity

Submitted

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ABSTRACT

Background and Aims: For applicability of cell-based therapies aimed at the treatment of liver failure, such as bioartificial livers (BALs) and hepatocyte transplantation, it is essential that the applied hepatocytes tolerate exposure to the patient plasma. However, plasma from both healthy donors and acute liver failure (ALF) patients is detrimental to hepatocytes and hepatic cell lines, such as HepaRG. We aimed to elucidate the underlying mechanisms of plasma-induced toxicity against HepaRG cells in order to ultimately develop methods to reduce this toxicity and render HepaRG-BAL treatment more effective.

Methods: Differentiated HepaRG cells cultured in monolayers and laboratory-scale BALs were exposed to culture medium, healthy human plasma, healthy porcine plasma and ALF porcine plasma. Healthy human plasma was fractionated based on size and polarity, albumin depleted and heat treated to characterize the toxic fraction. The cells were assessed for viability by total protein content and trypan blue staining. Their hepatic differentiation was assessed on transcript level through qRT-PCR and microarray analysis, and on functional level for Cytochrome P450 3A4 activity and ammonia elimination. Mitochondrial damage was assessed by JC-1 staining and mitochondrial gene transcription.

Results: Sixteen hours of healthy human plasma exposure did not affect viability, however, hepatic gene-transcript levels decreased dramatically within four hours of exposure. These changes were associated with early NF-κB signaling and a shift from mitochondrial energy metabolism towards glycolysis. Healthy human plasma-toxicity was associated with the dose-dependent presence of heat-resistant, albumin-bound and (partly) hydrophobic toxic compound(s).

HepaRG cells cultured in BALs were partially protected from plasma-toxicity, which was mainly attributable to medium perfusion and/or 3D configuration applied during BAL culturing. The detrimental human plasma effects were reversible in BAL-cultured cells. Porcine ALF-plasma elicited mitotoxicity additional to the basal detrimental effect of porcine healthy plasma, which were only partially reversible.

Conclusion: A specific fraction of human plasma reduces hepatic differentiation of HepaRG cultures, in association with early NF-κB activation. In addition, ALF-plasma elicits mitotoxic effects. These findings allow for a targeted approach in preventing plasma-induced cell damage.
INTRODUCTION

Acute liver failure (ALF) and acute-on-chronic liver failure are syndromes associated with substantial mortality, for which there is an unmet need of therapeutic options [1], as orthotopic liver transplantation is limited by donor organ shortage. Two types of cell-based treatment modalities are under development: Bioartificial Liver (BAL) therapy [2-4] and hepatocyte transplantation [5, 6]. The performance of the applied cells in the presence of human plasma is of paramount importance, especially in BALs, that are designed as extracorporeal hepatocyte-bioreactors perfused with patient plasma through plasmapheresis. Healthy-donor human plasma (hplasma) is known to induce intracellular lipid accumulation, stress and a decrease in hepatic functionality of primary hepatocytes and hepatic cell lines, through unclarified mechanisms [7, 8]. In addition, plasma from liver failure patients contains not only detrimental compounds which are normally detoxified by the liver, such as ammonia, bile acids, and lactate, but also compounds associated with inflammation and infection, such as endogenous damage-associated molecular patterns released from necrotic cells, cytokines and chemokines, as well as lipopolysaccharides and other stimulators of innate immune response due to bacterial translocation [9-11].

The AMC-BAL is a bioreactor that holds liver cells in 3D-configuration, in a spirally wound, non-woven matrix, interwoven with capillaries which supply oxygen-enriched gas [12, 13]. Liver cells come in direct contact with patient plasma during treatment. Currently, the device is loaded with the human liver progenitor cell line HepaRG [14, 15] and was proven efficacious in prolonging survival time of rats with ischemic ALF [4]. Previously we described that healthy- and ALF-rat plasma induced toxicity in HepaRG cells cultured in monolayers and in BALs [16, 17].

In this study we show that hplasma has a rapid detrimental effect on hepatic differentiation and functionality of HepaRG cells. In order to develop protective strategies, we studied the underlying mechanism of plasma-induced toxicity and the toxic fraction of hplasma. We further studied whether culture procedures or different culture platforms could inhibit or reverse the plasma-induced toxicity. Finally we assessed whether ALF plasma induced additional toxicity compared to plasma of healthy subjects, using pig plasma.
MATERIALS AND METHODS

Cell culture and plasma exposure procedures
HepaRG cells (Biopredic International) were cultured in supplemented Williams’ E medium without dimethyl sulfoxide (DMSO), with 10% fetal bovine serum, as described [14, 18]. Differentiated monolayers were acquired after seeding 1:5 in 12-well plates and culturing for 28 days under control conditions, i.e. statically under an atmosphere of 5% CO₂, 75%N₂ and 20% of O₂. Monolayers cultured under hyperoxia were kept under normoxia until reaching confluence at day 14 and then transferred to 5% CO₂, 55%N₂ and 40% of O₂ until day 28.

3D ‘Bal-in-a-dish’ (BALIAD) cultures were seeded into 28,2 mm² pieces of non-woven matrix and kept in medium in 12-well plates under control conditions until day fourteen, after which they were transferred to an orbital-shaker incubator under normoxia and shaken at 60 RPM until day 28. Laboratory-scale BALs (9 mL volume) were loaded with 0.6g cryopreserved HepaRG cells and cultured for 21 days as described previously [19] before commencing plasma-exposure-experiments.

To test the effect of plasma exposure, culture medium was replaced by undiluted plasma, except for the experiment with serial dilutions, for which plasma was diluted with fresh culture medium as indicated. As a negative control, cells were kept in fresh culture medium. Plasma exposure times varied between experiments, as indicated.

To test whether HepaRG-BAL cultures could recover from plasma-toxicity, plasma was removed from the BALs after 16h exposure by a single-pass flush with fresh culture-medium, after which the system was filled entirely with fresh culture medium and allowed to recover for 24 hours. Matrix samples for RNA isolation were obtained before plasma exposure (control) and immediately after plasma exposure, as described previously [20].

Plasma
Healthy human plasma (hplasma) derived from mixed-gender pooled donors (Sera Laboratories International Ltd.). Healthy porcine plasma (pplasma) derived from landrace pigs. For porcine ALF plasma (pALFplasma), ALF was induced as described [21] in a 31 kg female landrace pig by intravenous administration of paracetamol 1.5 g/kg body weight over 12 hours. After 21 hours the animal went into cardiovascular failure, and subsequently plasma was acquired by exsanguination. Liver failure was confirmed by biochemical analysis (Table 1). All procedures involving animals were conducted in agreement with the Animals (Scientific Procedures) Act 1986 under (UK Home Office) Project License 60/4557, and after approval by the Roslin Institute’s Animal Welfare and Ethical Review Board.
All plasmas were anticoagulated with Lithium-Heparin, filter-sterilized and supplemented with penicillin (100 U/mL) and streptomycin (100 µg/mL) before exposure to HepaRG cells.

**Plasma fractionation or treatment**
To characterize the toxic component in plasma, we fractioned or treated this plasma with different techniques. Fractionation by size was performed by filtering with 100kDa Molecular weight cut-off filters (Amicon Ultra-4, Merck), which are specified to retain molecules of over 100kDa, including albumin, which we confirmed.

Fractionation by polarity was performed by Bligh and Dyer extraction [22], after which the hydrophobic fraction was dried under a stream of nitrogen, resuspended and sonicated in DMSO, which was then diluted in culture medium to equal volume as the original plasma sample (final concentration of DMSO was 1%). Vehicle control treatment was performed with culture medium that was treated the same way (n=6, 2 independent experiments).

Albumin depletion was done by concentrating 3mL of plasma with 100kDa Molecular weight cut-off filters until 40 uL retentate, which was then resuspended in a 20 mM sodium phosphate binding-buffer (pH 7.0), as recommended by manufacturer, and exposed to 7 mL albumin-binding Blue Sepharose 6 Fast Flow beads (GE Healthcare) for 3 hours at 4°C and 30 min at room temperature on an orbital shaker at 100 rpm. Subsequently, the mix was concentrated over 100kDa Molecular weight cut-off filters and the retentate was diluted in culture medium to a final volume of 3 mL. Control plasma was treated with the same protocol, except for the exposure to Blue Sepharose beads (n=6, 2 independent experiments). Albumin was under the level of detection in the albumin-depleted plasma, as confirmed by the clinical chemistry laboratory (data not shown).

Heat treatment of 100% plasma consisted of a 45-minutes exposure to 58°C in a block heater (n=6, 2 independent experiments).

### Table 1. Biochemical profile of plasma.

<table>
<thead>
<tr>
<th></th>
<th>Baseline</th>
<th>Time of death</th>
</tr>
</thead>
<tbody>
<tr>
<td>AST (U/L)</td>
<td>37</td>
<td>243</td>
</tr>
<tr>
<td>Ammonia (µM)</td>
<td>58</td>
<td>316</td>
</tr>
<tr>
<td>Mir-122 (Fold change vs baseline)</td>
<td>1</td>
<td>160</td>
</tr>
<tr>
<td>FV/FVIII (ratio)</td>
<td>1.07</td>
<td>0.13</td>
</tr>
</tbody>
</table>

Baseline value is before paracetamol administration. AST=Aspartate Aminotransferase; Mir-122 = MicroRNA-122; FV= clotting factor five activity; FVIII= clotting factor eight activity.
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Cell viability assessment
HepaRG monolayer cultures were exposed to plasma or culture medium for 16 hours, washed 2x with PBS and then assessed for toxicity by two methods (n=4, 2 independent experiments). Firstly the cultures were incubated with trypan blue 0.4% solution (Sigma) diluted 1:4, after which the cultures were immediately assessed for viability by microscopy. Secondly, the total protein content/well was determined as described [18].

Function tests
Hepatic function tests were performed as previously described[4]. Briefly, test medium was prepared from HepaRG medium supplemented with NH₄Cl (1.5 mM), L-lactate (2.5 mM), D-galactose (2.27 mM) and testosterone (125 μM). Monolayers and BALs were exposed to 1.5 and 110 mL of test medium respectively and samples taken after 0, 6 and 24 hours for monolayers and 0, 30, 60, 120, 240, 480 and 1440 min for BALs. Concentrations of ammonia were quantified using the Megazyme Ammonia Assay kit (Megazyme International). Cytochrome P450 3A4 (CYP3A4) activity was measured in BALs by the quantification of 6b-hydroxytestosterone as described [4], and in monolayers using the P450-Glo™ CYP3A4 with Luciferin-IPA assay according to the manufacturer’s instruction (Promega). All metabolic rates were normalized to total protein as described [18]. Synthetic properties could not be assessed adequately, due to some retention of plasma proteins after plasma exposure.

RNA isolation, qRT-PCR and microarray analysis
Cells in monolayers or representative samples of BAL-matrix were lysed in 600 μL RLT buffer (RNeasy minikit, Qiagen) from which RNA was isolated according to manufacturer’s instructions. Transcript levels were determined by qRT-PCR using gene-specific reverse transcriptase (RT)-primers and touchdown qPCR protocol and normalized to 18S ribosomal RNA, as previously described [23]. Where expressed as % of control, the data were normalized to the average of non-treated control cells within each independent experiment. For the microarray experiment RNA was isolated after 0, 1, 2, 4 or 8 hours exposure to plasma (n=3 independent ~1 cm² monolayer cultures per group) RNA was biotinylated with the cRNA labeling kit (Ambion) and hybridized to Illumina HumanHT-12 v4 arrays (Illumina) after randomization of the samples. Scanning was performed on the Illumina iScan (Illumina). Image analysis and extraction of raw expression data was performed with Illumina GenomeStudio v2011.1 Gene Expression software with default settings (no background subtraction and no normalization).

The microarray data were analyzed with Bioconductor packages (version 3.0) using the statistical software environment R (version 3.1.3). Raw data normalization was performed on the Illumina sample and control probe profiles by normexp-by-control background correction, quantile normalization, and log2 transformation using the limma package (version 3.22.7).
The arrayQualityMetrics package (version 3.22.1) was used to assess that the microarray data was of good quality. Probes with a detection P-value > 0.05 (non-expressed) on all arrays (16,560 of 47,323 probes) were filtered out. Differential expression between time points was assessed using a moderated t-test using the linear model framework from the limma package. Resulting P-values were corrected for multiple testing using the Benjamini-Hochberg false discovery rate. Corrected P-values ≤ 0.05 were considered statistically significant. Probes were reannotated using the IlluminaHumanv4.db package (version 1.24.0). Gene sets from the hallmark collection and two liver-specific gene sets from the C2 collection, HSIAO_LIVER_SPECIFIC_GENES and SU_LIVER [24, 25], were retrieved from the Molecular Signatures Database (MSigDB) v5.1 (Entrez Gene ID version). Gene set enrichment analysis was performed using CAMERA (limma package) and gene set variation analysis (GSVA) was performed using the GSVA package (version 1.14.1). Sample-specific gene set enrichment scores calculated by GSVA were clustered using Pearson correlation as distance measure and complete linkage as agglomeration method (function hclust). Top-10 upregulated genes were cross-referenced against the Boston University Gilmore Lab NF-κB target gene set (http://www.bu.edu/nf-kb/gene-resources/target-genes/).

**Mitochondrial membrane potential and mitochondrial abundance**

Mitochondrial membrane potential was determined using JC-1 staining. This cationic dye emits green fluorescence in the cytosol of the cells (monomeric form) and red fluorescence when aggregates are formed (dimeric form) in active mitochondrial membrane [26]. BAL matrix samples (6mmx6mm) were incubated with 0.5 mL 4 µM JC-1 (Invitrogen) for 30 min at 37°C. Simultaneously, the cells were incubated with 1µM verapamil (Sigma) to inhibit the efflux of JC-1 through the activity of ATP binding cassette subfamily B member 1. Red to green fluorescence (Fλ585/Fλ510) was quantified on a NOVOstar microplate reader (BMG Labtech).

The ratio of mitochondrial DNA (mtDNA) to nuclear DNA (nucDNA) was assessed as a measure of mitochondrial abundance expressed as the weighted mean. Total DNA was isolated from AMC-BAL matrix samples (6 mmx6 mm) using the QIAamp DNA kit (QIAGEN) according to the manufacturer’s specifications. 2 nuclear encoded genes, *i.e.* CCAAT/enhancer binding protein alpha (*CEBPa*) and N-acetyl transferase (*NAT*), and 2 mitochondrial-encoded genes, *i.e.* mitochondrial-NADH dehydrogenase subunit1 (*MT-ND1*) and mitochondrial cytochrome c oxidase subunit 3 (*MT-CO3*) were quantified by qPCR.

**Statistical analyses**

Data were analyzed and processed for graphical representation in Prism 7.01 (GraphPad). Values are expressed as mean ± standard deviation (SD). Student’s t-tests were used when comparing two groups and 1-way ANOVA when comparing multiple groups. All results were corrected for multiple testing according to the Holm-Sidak method. Significance was reached...
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if $P<0.05$. The data produced in BALs were $n=3$, 3 independent experiments, compared to monolayer cultures $n=8$, 3 independent experiments. Plasma-treatment experiments were performed as indicated in text.

RESULTS

Healthy-donor human plasma has a rapid, dose-dependent detrimental effect on the hepatic differentiation of HepaRG-derived hepatocytes

We determined the effects of plasma exposure on HepaRG monolayer cultures on total protein content, morphology, ammonia elimination rate, CYP3A4 activity and the transcript levels of the hepatic genes CYP3A4, hepatic nuclear factor 4 Alpha ($HNF4\alpha$) and arginase 1 ($ARG1$), which were previously established to be highly responsive to plasma exposure [16, 17]. HepaRG monolayers showed a rapid decrease in transcript levels of these hepatic genes when exposed to 100% plasma ($\text{ARG1}$). After 4 hours exposure, $HNF4\alpha$ transcript level was decreased to 17±2% of the level in control cultures, while CYP3A4 and $\text{ARG1}$ transcript levels decreased more gradually to 5±2% and 7±2% respectively after 24 hours. Decrease in transcript levels of these genes was accompanied by profound morphological changes. The clusters of hepatocyte-like polygonal cells in HepaRG cultures disappeared (Fig. 1B) and cell-cell contact was lost after 16 hours exposure to plasma. There was no significant cell death as determined by total protein content (Fig. 1C) or trypan blue staining (Fig S1). However, rates of ammonia elimination and CYP3A4 activity had decreased to 48±21% and 29±7% of levels in control cultures respectively (Fig. 1D).

A 16 hours exposure of monolayers to plasma in escalating concentrations revealed a plasma-concentration-dependent decrease of hepatic gene transcript levels of CYP3A4, $HNF4\alpha$ and $\text{ARG1}$ (Fig. 1E). The transcript levels of the control (0% plasma-exposure) group were significantly higher ($P<0.05$) compared to all other groups, with the exception of $\text{ARG1}$ versus 25% plasma. The gene transcript levels were also significantly lower after exposure to 100% compared to 25% plasma for all three genes ($P<0.05$). Together, these data indicate that plasma has a rapid and dose-dependent toxic effect on the differentiation of HepaRG monolayers, although their viability is still unaffected.

Detrimental plasma fraction is hydrophobic, albumin-bound, and heat-stable

Next, we determined whether the detrimental effect of plasma was due to a lack of indispensable culture medium components or to the presence of a toxic fraction, by exposing monolayers to different fractions of plasma for 16h and testing the effect on transcript levels of $\text{ARG1}$, $HNF4\alpha$, and CYP3A4.
Figure 1. Effect of plasma on differentiated HepaRG cells. A) Transcript levels of the hepatic genes ARG1, CYP3A4 and HNF4A in HepaRG cells after 0.5, 1, 2, 4, 8, 12 and 24 hours of exposure to plasma, expressed as a % of the transcript levels in non-exposed cells. B) Morphology of HepaRG cells before and after 16 hours exposure to plasma, bar= 50µm. C) Total protein content per well after 16 hours exposure to plasma or culture medium. D) Hepatic functions ammonia elimination and CYP3A4 activity before and after 16 hours exposure to plasma. E) Hepatic gene transcript levels after 16 hours exposure to plasma in different concentrations. *= P<0.05 versus control.
Plasma filtered through 100kDa molecular weight cut-off filters (which also depleted albumin, Mw 67kDa), did not negatively affect transcript levels of ARG1 or HNF4A, while CYP3A4 was even significantly upregulated compared to control cultures (Fig. 2A). This supported the presence of a toxic fraction rather than depletion of essential culture medium components by 100% plasma incubation. Plasma that was albumin-depleted induced a significantly smaller decrease in transcript levels of ARG1 ($P<0.01$) and HNF4A ($P<0.001$) and an increase in CYP3A4 transcript levels ($P<0.001$) compared to control-treated plasma (Fig. 2B). Heat treated plasma (58°C 45 min) yielded similar effects as untreated plasma (Fig. 2C) and finally, the hydrophobic plasma fraction also reduced transcript levels of ARG1 and HNF4A, but not of CYP3A4 (Fig. 2D).

These data show that the cytotoxic effect of plasma is due to an albumin-bound, heat-stable fraction, that is, at least partly, hydrophobic, and not due to a lack of essential culture medium factors.

**Figure 2. Toxicity of plasma fractions.** Transcript levels of the hepatic genes ARG1, CYP3A4 and HNF4A in differentiated HepaRG monolayers exposed to different fractions of plasma for 16 hours, expressed as a % of the transcript levels in non-exposed cells. A.) Plasma filtered through a molecular weight cut-off filter compared to untreated plasma. B.) Albumin depleted plasma compared to control-treated plasma. C.) Heat treated plasma compared to untreated plasma. D.) The hydrophobic plasma fraction compared to treatment vehicle control. *= $P<0.05$ versus monolayers exposed to untreated plasma. &= $P<0.05$ versus monolayers exposed to control-treated plasma. $=$ $P<0.05$ versus monolayers exposed to vehicle-treated control.
Plasma-induced damage is associated with rapid activation of NF-κB target genes

To investigate the pathways involved in plasma-induced toxicity we compared the whole-genome transcriptomes of HepaRG monolayers exposed to plasma for different durations by microarray analysis. Since loss of hepatic gene transcription commenced within eight hours (Fig. 1A), we compared monolayers after 1, 2, 4 and 8 hours of plasma exposure with control monolayers. There were profound and rapid changes in transcriptomes; after 1, 2, 4 and 8 hours there were 134, 1154, 5548 and 6584 genes differentially expressed, respectively, compared to control cultures (Fig. 3A).

![Figure 3. Differentially expressed genes and gene sets.](image)

A.) Venn diagram of differentially expressed (adjusted \( P < 0.05 \)) genes of the monolayer cultures exposed to plasma for 1 to 8 hours compared to unexposed cultures. B.) Heatmap of the sample-specific gene set enrichment scores calculated by gene set variation analysis (GSVA) on the hallmark gene sets and two previously described human liver-specific gene sets. A high detail full-resolution version is included in Fig. S1, the results of the CAMERA analysis are summarized in Table S1. EMT= Epithelial to mesenchymal transition, Ox. phosphorylation = Oxidative phosphorylation.
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Of the top-10 upregulated genes at the different time points versus control cultures (Table 2), 11 out of 26 unique genes were NF-κB targets (depicted in bold).

Table 2. Top-10 upregulated genes after different durations of plasma exposure compared to non-exposed controls

<table>
<thead>
<tr>
<th>Gene</th>
<th>1h exposure</th>
<th>2h exposure</th>
<th>4h exposure</th>
<th>8h exposure</th>
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<td>CYR61</td>
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<td>EDN2</td>
<td>8.6</td>
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<td></td>
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<td>EGR1</td>
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Values represent fold change versus control cultures. Genes written in bold are recognized targets of NF-κB according to the Boston university Gilmore Lab NF-κB target gene set.
Gene set enrichment analysis on the hallmark collection of the Molecular Signatures Database (MSigDB) v5.1 showed that genes regulated by NF-κB in response to TNF were the most upregulated gene set after two hours exposure (Fig. 3B, Fig. S2). Energy metabolism changed drastically, as oxidative phosphorylation was strongly downregulated at later time points. In line with the qRT-PCR data (Fig. 1A), enrichment analysis of two previously described gene sets of liver-specific genes (1: [25] and 2:[24]) showed a profound decrease of hepatic gene transcript levels at 4 and 8 hours after exposure (Fig. 3B, Fig. S2).

**BAL-culturing protects against plasma-induced deterioration, which may be explained by 3D configuration and medium perfusion**

HepaRG cultures in BALs were less sensitive to plasma-induced damage compared to monolayers. After 16h of exposure, ammonia elimination was significantly decreased in monolayer cultures, but not in BAL cultures (Fig. 4A), while CYP3A4 activity was significantly decreased in both cultures, although significantly less in BAL cultures than in monolayers (2.2-fold versus 3.4-fold reduction) (Fig 4B). After exposure, transcript levels of ARG1, CYP3A4 and HNF4A were declined 6.9-, 5.4- and 4.0-fold, respectively, in monolayer cultures, while in BAL cultures CYP3A4 transcript levels were unchanged after plasma exposure and ARG1 and HNF4A transcript levels were less reduced, (2.2- and 1.8-fold) (Fig. 4C-E).

Three main differences between the monolayer and BAL culture systems are the level of oxygenation (20% vs 40% O2, respectively), the configuration (2D vs 3D, respectively) and medium perfusion (absent or present, respectively). To assess which of these factors contribute to the relative resistance to plasma toxicity in BAL cultures versus monolayer cultures, we exposed medium-perfused 3D cultures under ambient normoxia (BALIAD cultures) and static monolayer cultures under 40% of ambient oxygen to plasma and compared the decrease in transcript levels of ARG1, CYP3A4 and HNF4A between standard monolayer and BAL cultures (Fig. 4C-E). The effect of plasma on transcript levels did not differ between normoxic or hyperoxic monolayers. In BALIAD cultures, the transcript levels of ARG1 and CYP3A4, but not of HNF4A, were less decreased by plasma exposure compared to monolayer cultures. In addition, the transcript levels of all three tested genes were not significantly different after plasma exposure in BAL- and BALIAD-cultures, indicating that medium perfusion and 3D culture may be important contributing factors to the protective effect of BAL-culturing.

**Loss of hepatic gene transcription is reversible in BALs after exposure to healthy human plasma**

The previous experiments were performed with healthy human plasma, because of the scarcity of human ALF plasma. However, when applied clinically, the BAL-system will be exposed to plasma from patients suffering from ALF, which has a different composition.
Figure 4. Effect of culture platform on plasma toxicity. A-B.) Functionality of HepaRG monolayer or BALs cultures in culture medium (control) or after exposure to plasma for 16h, expressed as a % of unexposed cultures in the same culture-platform: A.) Ammonia elimination, B.) CYP3A4 activity. C-E.) Transcript levels of hepatic genes in HepaRG cells cultured in monolayers under normoxia or hyperoxia, in BALIADs or in BALs, exposed to plasma for 16h, expressed as a % of unexposed cultures in the same culture-platform: C) ARG1, D) CYP3A4, E) HNF4A. *= P < 0.05 versus plasma exposed monolayer. #= P < 0.05 versus plasma exposed monolayer. $= P < 0.05 versus unexposed cultures in the same culture platform.
from healthy plasma. The effects of the ALF contribution on plasma toxicity were therefore analyzed by investigating hepatic transcript levels of BALs exposed to plasmas from: healthy pigs (pplasma) and pigs with acetaminophen-induced ALF (pALFplasma). As a control for species differences, pplasma-exposed BALs were also included. After plasma exposure, all BALs were allowed to recover in culture medium for 24 hours and transcript levels were again measured to assess the reversibility of the plasma-induced toxicity.

HepaRG BALs exposed to pplasma, pplasma and pALFplasma all showed a significant decrease in transcript levels of ARG1 and HNF4A (Fig. 5), while only the latter two groups showed a decrease in CYP3A4 transcript levels. The level of recovery from the plasma exposure varied with the origin of the plasma. After recovery from pplasma exposure, transcript levels of all three genes were higher compared to post-exposure levels, up to or exceeding control levels. After recovery from pplasma exposure, only CYP3A4 transcript levels increased significantly.

Figure 5. Effect of pplasma, pplasma and pALFplasma on BAL-cultured HepaRG cells and recovery procedure. Transcript levels of hepatic genes ARG1, CYP3A4 and HNF4A in BAL-cultured HepaRG cells before exposure (control), directly after 16 hours of plasma-exposure and after 24 hours recovery in culture medium, expressed as a % of the transcript levels in non-exposed BAL-cultured HepaRG cells. Three types of plasma were applied: A) pplasma, B) pplasma, C) pALFplasma. *=P<0.05 versus control. #= P<0.05 versus post-plasma.
although HNF4A transcript levels no longer differed significantly from the pre-exposure levels. After recovery from pALF plasma exposure, there was no significant upregulation of any of the three genes compared to post-exposure levels, although there was no significant difference anymore to pre-exposure level for HNF4A transcripts. Thus we conclude that the decrease of gene transcript levels after 16 hours plasma exposure is reversible, although the degree of reversibility varies with the origin of plasma.

Porcine ALF plasma induces additional mitochondrial damage compared to healthy plasma

Since hepatocytes are dependent on mitochondrial activity for their high energy demanding functions, and microarray analysis indicated a shift in energy metabolism from oxidative phosphorylation towards glycolysis upon plasma exposure (Fig. 3B), we assessed

Figure 6. Effect of plasma and pALF plasma and recovery procedure on mitochondria of BAL-cultured HepaRG cells. A-B,) Transcript levels of mitochondrially encoded genes MT-CYTB and MT-ND5 in BAL-cultured HepaRG cells before exposure, directly after 16 hours of plasma-exposure and after 24 hours recovery in culture medium, expressed as a % of the transcript levels in non-exposed BAL-cultured HepaRG cells. C,) Membrane potential as determined by JC-1 staining after exposure to plasma, or pALF plasma (geometric mean ± geometric SD). D,) Mitochondrial abundance as reflected by the ratio between DNA encoding the mitochondrial genes MT-CO or MT-ND1 and DNA encoding the nuclear gene CEBPA (geometric mean ± geometric SD). ^=P<0.05 versus control. ^=P<0.05 versus post-plasma.
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mitochondrial abundance, membrane potential and mitochondrial gene transcript levels on the \(_p\) plasma and \(_{pALF}\) plasma exposed BALs.

After exposure to \(_{pALF}\) plasma, transcript levels of the mitochondrially encoded genes \(MT-CYTB\) and \(MT-ND5\) decreased by 3.2 and 3.6 fold respectively, after recovery they increased 1.9 and 1.7 fold respectively compared to control BALs, that were not exposed to any plasma (Fig. 6A-B). BALs exposed to \(_p\) plasma showed no change in transcript levels, suggesting an additional mitotoxic effect of the ALF plasma. This was confirmed by JC1-staining on samples of BAL matrix after recovery (Fig. 6C), which indicated a significant loss of mitochondrial membrane potential for \(_{pALF}\) plasma exposed BALs, but not \(_p\) plasma exposed BALs. However, the ratios between DNA of the mitochondrial genes \(MT-CO\) and \(MT-ND1\) and the nuclear gene \(CEBPA\), were unaffected in both groups (Fig. 6D), indicating that the abundance of mitochondria had not changed.

These results indicate that \(_{pALF}\) plasma, but not \(_p\) plasma impairs mitochondrial gene transcription, which is reversible after 16h plasma exposure, and also that mitochondrial functionality is negatively affected by \(_{pALF}\) plasma.

DISCUSSION

It is of paramount importance for BALs and other cell-based therapies that the applied cell source is resistant to human blood plasma. In order to improve this resistance it is vital to unravel the underlying mechanism. In this study we showed that \(_p\) plasma did not affect viability after 16 hours of exposure, but had a dose-dependent detrimental effect on the hepatic functionality and differentiation grade of HepaRG-derived hepatocytes, in close association with NF-κB signaling. Cells were partly protected from this basic plasma-induced damage when cultured in BALs and partial restoration of hepatic gene transcription could be achieved by a 24-hour regeneration period on standard culture medium after plasma exposure. In addition we found that toxicity can be attributed to an albumin-bound, and heat-resistant plasma fraction that is, at least partly, hydrophobic. In \(_p\) plasma that was filtered or albumin depleted, \(CYP3A4\) transcript levels were found to be increased, which is likely to reflect preservation of the detoxification system’s capability to be induced by exposure to ligands.

The 3D configuration and/or medium perfusion contribute in a large part to the increased plasma-resistance observed in BALs compared to monolayer cultures. This is in accordance with a previous report that primary rat hepatocytes are protected from plasma-induced toxicity by 3D-culturing [27]. The applied 3D culture platform provides a flexible culture
surface, which may be essential for maintaining cell-cell contact. Loss of adhesion is one of the first events in plasma-induced toxicity, and anchorage proteins, such as E-cadherin and the beta1-integrin receptor are known to protect against hepatocyte dedifferentiation and apoptosis [28-31]. In addition, matrix rigidity is known to limit hepatocyte differentiation, in part through transcription factor HNF4A [32].

Plasma was more toxic compared to hplasma, and pALF plasma exerted additional mitochondrial toxicity, underlining that ALF plasma is a hostile environment for both the patient and cell-based medicinal therapies. Due to the scarcity of human ALF plasma there was no opportunity to assess the effect of plasma from ALF patients. In this study we used a model of paracetamol-induced liver failure which is commonly associated with mitotoxicity and disruption of cell tight junctions [33], although also ALF-plasma of other origins has been reported to impair mitochondrial activity to varying degrees [31, 34]. However, the detrimental effects of ALF-plasma are likely to vary between etiologies, patients and clinical status, implicating the necessity of close monitoring of biocomponent functionality during therapy. This way, the BAL can be replaced after reaching critical plasma-induced toxicity of its biocomponent. Yet, it would be advantageous to inhibit at least the basic plasma-induced toxicity caused by toxic components already present in healthy plasma, which likely varies to lesser extent between individuals.

Hepatic gene transcript levels started to decline as early as four hours after hplasma exposure, preceded by upregulation of pro-inflammatory genes within one hour, suggesting a rapid mechanism, probably mediated through the activity of the pro-inflammatory NF-κB signaling-pathway. NF-κB -associated pro-inflammatory cytokines, such as IL-6, are known to have a dedifferentiating effect on primary hepatocytes [35, 36]. Especially for drug-detoxification enzymes, it is well documented that pro-inflammatory cytokines, and particularly IL-6, downregulate functionality through inhibition of transcription [37, 38]. These mechanisms were confirmed in HepaRG cells by others: pro-inflammatory cytokines suppressed transcription of phase 1 and phase 2 detoxification enzymes and drug transporters, as well as CYP450-enzyme activities [39].

There are several isoforms of NF-κB, with partially overlapping signaling cascades, exerting a multitude of effects. Pan-inhibition of NF-κB is known to induce and exacerbate hepatocyte apoptosis [40, 41], therefore this is not an attractive strategy to counter plasma toxicity. It has been described that the RELA/p65 isoform leads to inflammatory cytokine production, posing a more specific target. However, genetic disruption of this pathway is also known to sensitize hepatocytes to apoptosis [42, 43].
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Ideally, plasma toxicity should be inhibited by targeting more upstream processes. We have established that the detrimental compound(s) reside(s) in a plasma fraction that is albumin bound, non-polar and heat resistant. NF-κB can be activated through extracellular receptor signaling, such as Toll-Like Receptors, which are classically stimulated by bacterial and viral particles, mitochondrial reactive oxygen species (ROS)-production and hypoxia [44, 45]. We found that antioxidant treatment of HepaRG cells with N-acetyl cysteine did not decrease the detrimental plasma effects (data not shown), indicating that ROS is unlikely to be a main contributor to plasma toxicity [45]. NF-κB activation through hypoxia-induced pathways might be an option, as our microarray data showed that oxidative phosphorylation is severely affected. This does not necessarily imply a direct association with hypoxia; culturing the HepaRG cells and exposing them to plasma under hyperoxic conditions did not yield any protection against plasma toxicity.

In case of direct receptor stimulation, there is the possibility to inhibit plasma toxicity by molecular interference. Plasmapheresis filters with high affinity for Toll-like receptor 2 ligands ameliorated clinical symptoms of ALF in pigs with paracetamol overdose [46]. However, for specific targeting, further studies into the nature of the toxic component in plasma are required. Specific detoxification modules could easily be integrated into the plasmapheresis set-up, potentially protecting both the biocomponent and the native liver. A hybrid system, combining BAL-treatment with albumin dialysis may reduce damage of plasma to the biocomponent. One study, performed with human ALF plasma, addressed this option and showed, however, that albumin dialysis of the plasma did not reduce its pro-apoptotic effect on hepatocytes [31]. This may indicate that the toxic fraction has a very high affinity for albumin, since dialysis only removes the unbound fraction, and compounds with high binding affinities are therefore not removed efficiently.

Alternatively, the basic plasma-induced toxicity can be prevented by filtration for molecular size; plasma passed through a molecular weight cut-off filter of 100 kDa was no longer toxic. A disadvantage of small-pore plasma filters is that the mass-transfer of compounds during BAL therapy may be limited, which will decrease its efficacy. Others have previously studied the optimal plasmapheresis filter pore size, and concluded that a high-convection 400 kDa cut-off membrane offers the optimal balance between protection of the biocomponent and mass-transfer of toxins [47].

As an alternative strategy, the negative effects of basic plasma on hepatocytes can also be reversed by limiting the exposure-time to plasma so that damage to the biocomponent is still reversible. Our data show that gene transcription of HepaRG-BAL cultures can be restored after 16 hours of plasma exposure by recirculating culture medium through the
device, indicating that alternated treatment- and restoration runs are also a viable strategy to increase the life-span of BALs, as proposed by others previously [7].

In conclusion, plasma has a detrimental effect on differentiated HepaRG cultures, which is associated with early NF-κB activation. Strategies to extend functional time in BAL set-up include intermitted exposure- and recovery runs, the use of small-pore size and/or cytokine scavenging plasmapheresis filters and direct inhibition of involved pathways such as NF-κB signaling. ALF plasma may elicit additional toxic effects, which may vary between patients. Therefore, close monitoring of the functionality will be required for optimal BAL therapy of ALF patients.
REFERENCES


SUPPLEMENTAL INFORMATION

Figure S1. Cell viability evaluated by trypan blue exclusion test (stained cells have lost membrane integrity and are not viable), of differentiated HepaRG monolayers exposed for 16 hours to: A.) culture medium control, or B.) plasma. Red arrows indicate trypan blue positive cells.
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Figure S2. Heatmap of sample-specific gene set enrichment scores. A high detail full-resolution version of Fig. 3B.