Validation of Messenger Ribonucleic Acid Markers Differentiating Among Human Acute Respiratory Distress Syndrome Subgroups in an Ovine Model of Acute Respiratory Distress Syndrome Phenotypes

Karin Wildi1,2,3*, Kieran Hyslop1†, Jonathan Millar1,2,4, Samantha Livingstone1,2, Margaret R. Passmore1,2, Mahé Bouquet1,2, Emily Wilson1,2, Gianluigi LiBassi1,2, John F. Fraser1,2 and Jacky Y. Suen1,2

1 Critical Care Research Group, Brisbane, QLD, Australia, 2 Faculty of Medicine, The University of Queensland, Brisbane, QLD, Australia, 3 Cardiovascular Research Group, Basel, Switzerland, 4 Roslin Institute, University of Edinburgh, Edinburgh, United Kingdom

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Edited by:
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*Correspondence:
Karin Wildi
wildik@hotmail.com
† These authors have contributed equally to this work and share first authorship

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Background: The discovery of biological subphenotypes in acute respiratory distress syndrome (ARDS) might offer a new approach to ARDS in general and possibly targeted treatment, but little is known about the underlying biology yet. To validate our recently described ovine ARDS phenotypes model, we compared a subset of messenger ribonucleic acid (mRNA) markers in leukocytes as reported before to display differential expression between human ARDS subphenotypes to the expression in lung tissue in our ovine ARDS phenotypes model (phenotype 1 (Ph1): hypoinflammatory; phenotype 2 (Ph2): hyperinflammatory).

Methods: We studied 23 anesthetized sheep on mechanical ventilation with observation times between 6 and 24 h. They were randomly allocated to the two phenotypes (n = 14 to Ph1 and n = 9 to Ph2). At study end, lung tissue was harvested and preserved in RNAlater. After tissue homogenization in TRIzol, total RNA was extracted and custom capture and reporter probes designed by NanoString Technologies were used to measure the expression of 14 genes of interest and the 6 housekeeping genes on a nCounter SPRINT profiler.

Results: Among the 14 mRNA markers, in all animals over all time points, 13 markers showed the same trend in ovine Ph2/Ph1 as previously reported in the MARS cohort: matrix metalloproteinase 8, olfactomedin 4, resistin, G protein-coupled receptor 84, lipocalin 2, ankyrin repeat domain 22, CD177 molecule, and transcobalamin 1 expression was higher in Ph2 and membrane metalloendopeptidase, adhesion G protein-coupled receptor E3, transforming growth factor beta induced, histidine ammonia-lyase, and sulfatase 2 expression was higher in Ph1. These expression
INTRODUCTION

More than five decades after its first description (1), acute respiratory distress syndrome (ARDS) still has an unacceptably high morbidity and mortality (2). After years of little success in ARDS research focusing on improving patient outcomes (3), analysis of large ARDS cohorts from randomized controlled trials identified two distinct biological subphenotypes amongst the heterogenous population of ARDS patients (4, 5): a discovery with potential for future targeted treatment in ARDS. In brief, a hypoinflammatory (or uninflamed; named P1) and a hyperinflammatory (or reactive; named P2) subphenotype, defined by specific functional and biological parameters, have been proposed and corroborated with existing retrospective clinical data (4–6). P2 was clearly associated with a more severe shock state and metabolic acidosis as well as worse clinical outcomes (4, 7). Importantly, the response of patients to specific medical measures and pharmacological treatment appears to be highly dependent on the respective subphenotype (4, 5, 8). Even though the concept of subphenotypes in ARDS is now widely accepted, there are still major gaps in our current knowledge regarding the in-depth comprehension of underlying biological driving factors and mechanisms.

Animal models have always played an important role in biological discovery and therapeutic translation in ARDS (9). We reported previously about ovine ARDS phenotype models that mimics some key features of human ARDS subphenotypes (10): phenotype 1 (Ph1: hypoinflammatory) and phenotype 2 (Ph2: hyperinflammatory). Additionally, a recent analysis revealed matching gene expression among animal models induced by lipopolysaccharides (LPS) as in our Ph2 model and the human P2 subphenotype (11).

To validate our ovine model on a molecular level, we compared a subset of messenger ribonucleic acid (mRNA) patterns could be found when different sources of mRNA – such as blood leukocytes and lung tissue – were compared.

Conclusion: In human and ovine ARDS subgroups, similar activated pathways might be involved (e.g., oxidative phosphorylation, NF-κB pathway) that result in specific phenotypes.

Keywords: acute respiratory distress syndrome, phenotypes, mRNA expression, up-and downregulation, precision medicine, predictive and prognostic enrichment, biomarker

MATERIALS AND METHODS

Animal Model

This study is a secondary analysis of pooled control animals of ovine Ph1 and Ph2 ARDS from two blinded, randomized, controlled preclinical trials. A more detailed explanation of the two studies is provided in the Supplementary Method.

Animal ethics was approved by The Queensland University of Technology Office of Research Ethics and Integrity (No 1600001108 and No 1800000606). All experiments were performed in accordance with the Australian Code of Practice for the Care and Use of Animals for Scientific Purposes and the Animal care and Protection Act 2001 (QLD).

A total of 23 female non-pregnant Merino-Dorset crossbred ewes, aged 1–3 years, mean weight 49 kg (47–52), participated in this analysis and were randomly allocated to one of the two groups: Ph1 (n = 14) and Ph2 (n = 9); (10). Eight animals were observed up to 6 h (Ph1 n = 5, Ph2 n = 3), 7 animals up to 12 h (Ph1 n = 4, Ph2 n = 3) and 8 animals up to 24 h (Ph1 n = 5, Ph1 n = 3; Figure 1).

In short, after induction of general anesthesia and intubation, the animal was mechanically ventilated in a lung-protective fashion (13). After completion of instrumentation, the animal was turned in prone position and ARDS was induced: (1) animals randomized to Ph1 received oleic acid (OA; O1008; Sigma-Aldrich, Australia) sequentially in 0.03 ml/kg doses intravenously (IV) until a PaO2/FiO2 ratio (PF) of <150 mm Hg was reached (2) animals assigned to Ph2 received Ph2 received aforementioned oleic acid IV followed by 0.5 μg/kg of lipopolysaccharide (LPS: E. coli O55:B5, Sigma-Aldrich, Australia) infused over 1 h. Intra-experimental monitoring, management and data collection has been reported in detail previously (10, 14).

Sample Collection and Processing

At study end, lung tissue was harvested and samples from the upper, middle and lower lobe of the right lung were stored in RNAlater (In Vitrogen, United States) at 4°C overnight and then at –80°C until extraction.

For histopathological assessment, lung tissue was preserved in 10% neutral buffered formalin for 24 h for embedding in paraffin afterward, then tissue was sectioned to 5 μm thickness.
and stained with hematoxylin & eosin. Slides were assessed by an independent and blinded veterinary pathologist using the lung injury score (LIS) as recommended by the ATS for experimental ARDS in animal models (15). In short, the LIS scores neutrophils in the alveolar space (A), neutrophils in the interstitial space (B), hyaline membranes (C), proteinaceous debris filling the airspace (D), and alveolar septal thickening (E). Every item is given a score between 0 and 2. The LIS is then calculated by: 

\[ \frac{(20 \times A) + (14 \times B) + (7 \times C) + (7 \times D) + (2 \times E)}{\text{number of fields}} \times 100, \]

leading to a score between zero (no injury) and one (severe lung injury). Twenty random high-power fields (400× total magnification) were scored per section, and the LIS was calculated per animal (mean ± SD).

**Selected Messenger Ribonucleic Acid Markers**

The 16 most upregulated mRNA markers in blood leukocytes in P2 and P1 subphenotypes were chosen as reported previously by Bos et al. (12). These were mainly markers of oxidative phosphorylation, cholesterol metabolism, and the Nuclear Factor Kappa-light-chain-enhancer of Activated B Cells pathway (NF-κB) as well as markers related to neutrophil and macrophage function in P2: matrix metalloproteinase 8 (MMP8), olfactomedin 4 (OLF4M), resistin (RETN), G protein-coupled receptor 84 (GPR84), CEA cell adhesion molecule 1 (CEACAM1), lipocalin 2 (LCN2), zinc finger DHHC-type palmitoyltransferase 18 (ZDHHC18), ankyrin repeat domain 22 (ANKRD22), CD177 molecule (CD177), and transcobalamin 1 (TCN1). For P1, mRNA markers involved in regulation of vital cell functions like proliferation and differentiation, motility and survival were among the most upregulated ones: membrane metalloendopeptidase (MME), retinol binding protein 7 (RBP7), adhesion G protein-coupled receptor E3 (ADGRE3), transforming growth factor beta induced (TGFBI), histidine ammonia-lyase (HAL), and sulfatase 2 (SULF2) were selected.

**Extraction of Messenger Ribonucleic Acid and Quality Control**

Tissue from the upper, middle and lower regions of the right lung were independently homogenized in Trizol (Ambion, United States) using a Polytron PT 2100 (Kinematica AG, Switzerland). Following homogenization, a PureLink™ RNA Mini Kit (Invitrogen, United States) was used to purify total RNA and a DNase treatment was performed using a DNA-free™ Kit (Invitrogen, United States). Sample concentration and purity was determined using a NanoDrop 1000 Spectrophotometer (Thermo Scientific, United States) and RNA integrity number (RIN) was assessed on an Agilent Bioanalyzer 2100 (Agilent Technologies, United States; Supplementary Table 1). Samples with a RIN ≥ 6 were standardized to a concentration of 100 ng/µL in nuclease-free water. Final concentration was determined on a Qubit 2.0 Fluorometer (Invitrogen, United States), before regional samples were pooled.

**Nanostring Gene Expression Analysis**

A custom NanoString nCounter assay (NanoString Technologies, United States) was used to assess the expression of the 16 genes...
of interest and the six housekeeping genes (MAU2, POLR2A, PGK1, RPL19, TBP, and YWHAZ) used for data normalization. The assay consisted of a nCounter Elements TagSets (reporter and capture tags) and oligonucleotide probe pairs (reporter/probe A and capture/probe B) designed by NanoString Technologies. Oligonucleotides were supplied by Integrated DNA Technologies (Iowa, United States) as pools (one containing every Probe A and the other every probe B) at a final concentration of 5 nM per oligo and 25 nM per oligo, respectively. Probe pools were used to prepare the working pools described in subsequent steps, as per the manufacturer’s instructions. Prior to assay, pooled RNA samples were diluted in nuclease-free water to 20 ng/µL and concentration was measured using a Qubit 2.0 Fluorometer (Invitrogen, United States). 140 ng of pooled RNA was combined with a master mix containing, nCounter Elements TagSets and oligonucleotide probe working pools (Integrated DNA Technologies, United States) in 0.2 mL PCR tubes, as per the manufacturer’s instructions (Supplementary Table 2). Samples were hybridized with reporter and capture probes at 67°C in a T100™ thermal cycler (Bio-Rad Laboratories, United States) for 18 h. To minimize potential evaporation, the thermal cycler lid was maintained 5°C higher than the block for the duration of the hybridization. Following hybridization, 15 µL of RNase-free water was added to the mixture. A total of 30 µL was loaded onto each lane of the NanoString microfluidic SPRINT™ cartridge (Seattle, United States) for automated processing on a nCounter SPRINT™ profiler (NanoString Technologies, United States).
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FIGURE 3 | Heatmap of relative changes in means in log2 normalized count in mRNA markers, according to time point of tissue harvesting. Figure legend: 8 animals available for analysis at 6 h (Ph1 n = 5, Ph2 n = 3), 7 at 12 h (Ph1 n = 4, Ph2 n = 3) and 8 animals at 24 h (Ph1 n = 5, Ph2 n = 3). Relative changes shown on the right y-axis were calculated as log2-normalized count of OA-IV-LPS divided by log2-normalized count of OA for each time point. Abbreviations: OA, oleic acid; OA-IV-LPS, oleic acid and lipopolysaccharide intravenously; MMP8, matrix metalloproteinase 8; OLFM4, olfactomedin; RETN, resistin; GPR84, protein-coupled receptor 84; CEACAM1, CEA cell adhesion molecule 1; LCN2, lipocalin 2; ANKRD22, ankyrin repeat domain 22; CD177, CD177 molecule; TCN1, transcobalamin 1; MME, membrane metalloendopeptidase; RBP7, retinol binding protein 7; ADGRE3, adhesion G protein-coupled receptor E3; TGFBI, transforming growth factor beta induced; HAL, histidine ammonia-lyase; and SULF2, sulfatase 2.

Data Analysis
The raw count data quality control assessment was conducted in nSolver™ Analysis Software 4.0 (NanoString Technologies, United States). Assay background correction was performed by subtracting the mean of the negative controls plus two standard deviations. Corrected counts were normalized to the geometric mean of the top 3 housekeeping genes (POLR2A, RPL19 and TBP), as determined by the geNorm algorithm. Normalization was performed using the nCounter Advanced Analysis 2.0 plugin.

Normalized mRNA counts are log2 transformed and levels were compared using a Mann–Whitney U test. Data in figures is expressed as mean and standard deviation (SD). All hypothesis testing was two-tailed and a p-value of less than 0.05 was considered statistically significant. All statistical analysis was performed using SPSS 27.0 (SPSS Inc, Chicago, United States).

RESULTS

Studied Population
Baseline characteristics of experimental animals did not differ among Ph1 and Ph2 and are provided in Supplementary Table 3. Clinical parameters during study observation period are presented at 6, 12, and 24 h of all animals alive at this time point (Supplementary Table 4). Ph2 animals displayed a higher heart rate and need for vasoactive drugs, indicating a more altered hemodynamic situation. While the PF ratio recovered to a certain degree in both injury models, Ph2 showed higher lactate levels combined with a more negative base excess, as well as lower platelet and higher neutrophil counts, pointing toward more disturbances in metabolic situation and more inflammation activation.

Levels of pro-inflammatory (IL-6 and -8) and anti-inflammatory cytokines (IL-10) were higher in Ph2 than Ph1 throughout the study observation time after induction of ARDS (Supplementary Figure 1). The LIS was 0.28 (±0.06) in Ph1 and 0.35 (±0.06) in Ph2 (p 0.06 for comparison; Supplementary Figure 2). In addition, there were more alveolar thrombi present and more lung tissue was affected by necrosis in Ph2 animals.

Levels of Messenger Ribonucleic Acid Markers Among Ovine Acute Respiratory Distress Syndrome Phenotypes All Time Points Pooled
Zinc finger DHHC-type palmitoyltransferase 18 and RBP7 were excluded because the counts were below the detection limit of the assay. Among the remaining 14 mRNA markers, in all animals over all time points, 13 markers showed similar trends in the ovine phenotypes as previously reported (12) in the MARS cohort (Figure 2): MMP8 (p 0.14), OLFM4 (p 0.15), RETN (p 0.37), GPR84 (p 0.009), LCN2 (p 0.89), ANKRD22 (p 0.31), CD177 (p 0.12), TNC1 (p 0.44) were more upregulated in Ph2 whereas MME (p 0.06), ADGRE3 (p 0.05), TGFBI (p 0.12), HAL (p 0.31), and SULF2 (p 0.02) expression was higher in Ph1. Only one mRNA marker, CEACAM1 (p 0.007), reported as upregulated in...
are that mRNA expression is likely to change over time in the animals assessed at 24 h. Potential explanations for this finding compared to human ARDS subphenotypes was mostly seen in described ARDS populations but not to other ARDS cohorts. It needs to be considered that because of the additional LPS injury, our ovine injury model caused by OA and additional LPS IV. It also needs to be considered: the same breed of sheep but a different flock was used for experiments in study 2, this could explain parts of the reported heterogeneity. Additionally, while the induction of ARDS phenotypes and the intra-experimental handling was the same, a potential influence of sample age between between study 1 and 2 on the expression results at 24 h cannot be excluded.

Some important limitations of our study have to be considered: First, mRNA expression has previously been reported in peripheral leukocytes (12) rather than lung tissue. We can safely assume that mRNA expression in circulation differs from the alveolar space but to what extent cannot be determined in this study as we did not have mRNA in plasma available for analysis. Second, the number of biological replicates per time point is limited and consequently so is the translatability of results. Third, as the observed changes in gene expression may not reflect translation into proteins, any conclusion about underlying biological mechanisms is limited. Fourth, for a more homogeneous view, we combined lung tissue from upper, middle and lower lobes of the right lung. However, due to the heterogeneous affection of different parts of the lungs in ARDS, localized differences in mRNA expression should also be investigated. Fifth, we did not sample for blood cultures, therefore we cannot exclude the development of an infection over the course of the study, potentially introducing a bias in the later results. Sixth, since all studied animals were female, we cannot determine whether the factor gender might contribute to the findings.

### DISCUSSION

In two different ovine ARDS phenotypes, mimicking some key features of the presumably inflammatory subgroups in human ARDS (10), we aimed to validate the most differentially expressed mRNA markers in human ARDS subphenotypes (12). We report two major findings: First, 13 of the 14 genetic markers tested in ovine lung tissue have shown similar trends in ovine Ph2/Ph1 phenotypes as reported in human blood leukocytes in the MARS cohort (12). However, only 3 markers reached statistical significance, most likely due to small samples sizes per assessed group. Second, mRNA expression among tissue harvesting time points showed some differences, indicating that the gene expression in ARDS phenotypes is time-dependent over the course of the disease.

Our results indicate that similar pathways might be involved (e.g., oxidative phosphorylation, NF-κB pathway) in human and ovine ARDS subgroups. This assumption has been underlined in a recent analysis, showing important similarities in gene expression patterns in human tracheal aspirate in P2 with animal models of ARDS induced by LPS (11). In the MARS cohort, mRNA markers of oxidative phosphorylation (MMP8) and neutrophil activation and function (MMP8, OLFM4, LCN2, and CD177) were shown to be a hallmark of the hyperinflammatory (or “reactive”) subphenotype (12). This finding adds evidence to the hypothesis that differentiation among presumably inflammatory ARDS subphenotypes is likely based on inflammatory pathways.

NF-κB is an important pathway in clinical ARDS (16) and LPS is known to cause NF-κB induction (17). Therefore, the higher expression in Ph2 animals of OLFM4, LCN2, TCN1, ANKRD22, RETN, and ANKRD22 can at least partially be explained by the injury model caused by OA and additional LPS IV. It also needs to be considered that because of the additional LPS injury, our ovine Ph2 model might reflect more a sepsis-related ARDS model, potentially better comparable to the MARS cohort than other described ARDS populations but not to other ARDS cohorts.

Deviation from matching trends in ovine mRNA levels as compared to human ARDS subphenotypes was mostly seen in animals assessed at 24 h. Potential explanations for this finding are that mRNA expression is likely to change over time in the dynamic biological process of ARDS. However, this assumption could not be assessed in the MARS cohort with early sampling within 24 h of ICU admission. As the animals assessed at 24 h were all part of the second study, completed 2 years after the first one, the time gap between the completion of the two studies has to be considered: the same breed of sheep but a different flock was used for experiments in study 2, this could explain parts of the reported heterogeneity. Additionally, while the induction of ARDS phenotypes and the intra-experimental handling was the same, a potential influence of sample age between between study 1 and 2 on the expression results at 24 h cannot be excluded.

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### CONCLUSION

Thirteen of 14 mRNA markers, overexpressed in human P2 and P1 ARDS subphenotypes, respectively, revealed similar expression patterns in our ovine model of ARDS phenotypes.

### DATA AVAILABILITY STATEMENT

The datasets presented in this study are publicly available in the GEO repository, accession number: GSE206888.

### ETHICS STATEMENT

The animal study was reviewed and approved by Queensland University of Technology, Brisbane, Australia: Office of Research Ethics and Integrity.

### AUTHOR CONTRIBUTIONS

KW, KH, JM, JF, and JS: conception and design of work, analysis and interpretation of data. KW, KH, JM, SL, MP, MB, EW, GL, JF; and JS: acquisition of data, drafting the work or revising it critically for important intellectual content, final approval

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of the version submitted for publication, and agreement to be accountable for all aspects of the submitted work. All authors contributed to the article and approved the submitted version.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmed.2022.961336/full#supplementary-material

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