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Systems Biology behind immunoprotection of both Sheep and Goats after Sungri/96 PPRV vaccination

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16 Abstract

17 Immune response is a highly coordinated cascade involving all the subsets of PBMCs. In this study, RNA-Seq analysis of PBMC subsets - CD4+, CD8+, CD14+, 18 19 CD21+ and CD335+ cells from day 0 and day 5 of Sungri/96 Peste des Petits 20 Ruminants vaccinated sheep and goats was done to delineate the systems biology 21 behind immune - protection of the vaccine in sheep and goats. Assessment of the immune response processes enriched by the differentially expressed genes in all the 22 23 subsets suggested a strong dysregulation towards development of early inflammatory microenvironment, which is very much required for differentiation of monocytes to 24 25 macrophages, and for activation and migration of dendritic cells into the draining lymph 26 nodes. The protein - protein interaction networks among the antiviral molecules (IFIT3, ISG15, MX1, MX2, RSAD2, ISG20, IFIT5 and IFIT1) and common DEGs across 27 PBMCs subsets in both the species identified ISG15 to be an ubiquitous hub, that helps 28 29 in orchestrating antiviral host response against PPRV. IRF7 was found to be the key master regulator activated in most of the subsets in sheep and goats. Most of the 30 31 pathways were found to be inactivated in B - lymphocytes of both the species indicating 32 that 5 dpv is too early a time point for the B - lymphocytes to react. The cell mediated immune response and humoral immune response pathways were found more enriched 33 34 in goats than in sheep. Though, animals from both the species survived the challenge, a 35 contrast in pathway activation was observed in CD335+ cells.

36 Importance

Peste des petits ruminants (PPR) by PPRV is an OIE listed acute, contagious transboundary viral disease of small ruminants. Attenuated Sungri/96 PPRV vaccine used all over India against this PPR, provides long-lasting robust innate and adaptive immune response. The early antiviral response was found mediated through type I 41 interferon independent ISGs expression. However, systems biology behind this immune 42 response is unknown. In this study, in vivo transcriptome profiling of PBMC subsets 43 (CD4+, CD8+, CD14+, CD21+ and CD335+) in vaccinated goats and sheep (at 5 days 44 of post vaccination) was done to understand this systems biology. Though there are a 45 few differences in the systems biology across cells (specially the NK cells) between sheep and goats, the co-ordinated response that is inclusive of all the cell subsets was 46 47 found to be towards induction of strong innate immune response, which is needed for 48 an appropriate adaptive immune response.

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50 **1. Introduction**

51 Peste des petits ruminants (PPR) is an OIE listed acute, highly contagious 52 transboundary viral disease of small ruminants, caused by PPR virus of genus 53 Morbillivirus and family Paramyxoviridae [1]. Morbidity and mortality can be as high as 54 100% and 90%, respectively [2]. The disease manifests as fever, discharge from the eyes and nose, stomatitis, pneumonia and enteritis [3]. PPRV vaccine developed by 55 56 continuous passage (N=59) of Sungri/96 strain in Vero cells is widely used throughout 57 India [4]. The vaccine provides long-lasting robust innate and adaptive - humoral and a 58 strong cell-mediated immunity [2], which, however, warrants further investigation [5], [6], 59 [7]. PPRV is lymphotropic and epitheliotropic [8], [9], [10]. The primary receptors for 60 PPRV include the signalling lymphocyte activation molecule (SLAM) on activated T cells, B cells, and dendritic cells and Nectin-4 receptor on epithelial cells [8], [9], [11]. 61

62 Immune response is complex within a host and different cell types respond differently to infection as different classes of receptors receive cues and produce distinct 63 effector molecules [12]. It is a highly coordinated effort of distinctly programmed 64 hematopoietic cell types, and a product of various direct and indirect effects and 65 66 interactions between similar or different cell types [13]. Moreover, the tissue 67 microenvironment also affects the elicited immune response. In case of viruses, the 68 complexity of the host response depends on variations in genetic makeup, cell tropism 69 and replication kinetics [12], [13], [14], [15]. PBMCs include T helper cells (CD4+), T 70 cytotoxic cells (CD8+), B-lymphocytes (CD21+), monocytes(CD21+), natural killer 71 cells(CD335+) and dendritic cells (CD320+), which play an important role in virus 72 recognition and induce immune response for host defence. While analysing whole blood 73 or PBMCs, the response of under-represented cell populations can be masked [13].

74 Despite advances in our understanding of vaccines, the mechanisms by which 75 protective immune responses are orchestrated among the cell subsets are little known. Molecular patterns and gene signatures detected in blood post vaccination represent a 76 77 strategy to prospectively determine vaccine efficacy [16]. The conventional 78 immunological methods like ELISA, ELISpot etc. are of utmost importance in this regard 79 and may continue to remain so in future [17]. However, these approaches are inept at 80 predicting the systems biology behind immune - protection. Delineating the systems 81 biology would help understanding the molecular mechanisms of vaccine induced 82 immune responses. RNA sequencing is a widely used quantitative transcriptome 83 profiling system for deciphering the systems biology comprehensively [18]. Previously, 84 RNA sequencing was used to unravel transcription factors, which modulate immune 85 response to PPRV Sungri/96 live attenuated vaccine strain in vitro in PBMCs [6]. Also, a

predicted immune signalling pathway of PPRV Sungri/96 vaccine induced immune response with predominant role of IRFs, TRIMs and ISGs in creation of robust antiviral state *in vitro* in PBMCs has been proposed [7].

Till now there are no *in vivo* reports of transcriptome profiling of PBMC subsets in PPRV vaccinated goats and sheep. Herein, transcriptional profiling of circulating CD4+, CD8+, CD14+, CD21+ and CD335+ cells of PPRV vaccinated sheep and goats at 0 day (control i.e. just before vaccination) and before the development of antibody response (5 days post vaccination i.e. 5dpv) to decipher the vaccine induced immune response was done.

95 **2. Materials and methods**

96 **2.1 Animal experiment, ethics statement and virus**

Live attenuated PPR vaccine virus (Sungri/96) was used as vaccine virus.
Permission for studies on animal subjects was obtained and protocols approved from
IVRI – Institutional Animal Ethics Committee (IAEC) under CPCSEA, India vide letter
no. 387/CPCSEA. The vaccine potency testing experiment was carried out as per the
guidelines of Indian Pharmacopeia – 2014 (page.no: 3626).

In this study, healthy sheep and goats (n=5, age=12 months) confirmed negative for PPRV antibodies (c-ELISA and SNT) and PPRV antigen (s-ELISA) [19], [20] were used. On c-ELISA, the samples with a percent inhibition (PI) value of >40% were considered positive. The animals were acclimatised for 14 days, followed by vaccination on day 0 with a 10^3 TCID50 field dose of Sungri/96 strain through sub-cutaneous route, as mentioned in our previous report [21]. All the animals vaccinated survived the challenge from the virulent PPRV in the vaccine potency testing experiment.

109 2.2 Isolation of T helper cells, T cytotoxic cells, B lymphocytes, monocytes and 110 natural killer cells by magnetic assisted cell sorting technology (MACS)

111 Blood was collected from the animals (n=5) in heparin coated vacutainer vials at 0 112 day (just before vaccination) and 5 days post vaccination (5 dpv). PBMCs were isolated by using Ficol Histopaque gradient method. PBMCs were strained through cell strainer 113 114 of 0.40-micron size. The PBMCs cell subsets were enriched by positive selection using 115 indirect magnetic assisted cell sorting technology (Miltenyi Biotech). Cell sorting was done as per the manufacturers protocol. Initially, the cell-specific surface marker FITC-116 117 conjugated primary antibodies, anti CD4+ (T helper cells, #MCA2213F), anti CD8+ (T cytotoxic cells, #MCA2216F), anti CD14+ (Monocytes, #MCA1568F), and anti CD21+ 118 119 (B lymphocytes, #MCA1195F), were used. For CD335 (NK cell) anti CD335+ (#MCA5933GA as primary antibody) and FITC labelled secondary antibody (#F9137) 120 121 were used. Subsequently, the cells were magnetically labeled with anti - FITC 122 MicroBeads. Then the cell suspension was loaded on a miniMACS® column which was 123 placed in the magnetic field of a MACS Separator. The magnetically labeled cells were 124 retained in the column while the unlabeled cells run through. After removal of the 125 column from the magnetic field, the magnetically retained cells were eluted as positively 126 selected cell fraction. The purity of the cells was further checked by flow cytometer. The 127 cells were stored in RNA later for further use at -80°C. Cells were kept on ice and cold buffers were employed to minimize alterations in gene expression during labelling andsorting.

130 **2.4 RNA-sequencing of the samples**

131 Total RNA from each of the PBMC subsets was isolated using the RNeasy Mini kit 132 (Qiagen GmbH, Germany) as per the manufacturer's protocol. The integrity and quantity of isolated RNA were assessed on a Bioanalyzer (Agilent Technologies, Inc). The 133 134 library was prepared using NEBNext Ultra RNA Library Prep Kit for Illumina 135 (NewEngland Biolabs Inc.) following the manufacturer's protocol. Approximately, 100ng 136 of RNA from each sample was used for RNA library preparation. The quality of the 137 libraries was assessed on Bioanalyzer. Libraries were quantified using a Qubit 2.0 Fluorometer (Life technologies) and by gPCR. Library (1.3ml, 1.8pM) was denatured, 138 139 diluted and loaded onto a flow cell for sequencing. cDNA library preparation and 140 Illumina Sequencing was performed at Bioserve Pvt. (Hyderabad, India). RNA-Seq data 141 was generated in FASTQ format.

142 **2.5 Raw data processing**

Raw sequence data from each sample was subjected to quality control checks using FastQC (Babraham Bioinformatics). Moreover, low quality reads with a mean phred score less than or equal to 25 and reads shorter in length than 50 nt were removed using prinseq-lite software [22] before downstream analysis. The data was submitted to the GEO database with accession number GSE155504.

148 2.6 Differential expression and identification of differentially expressed genes149 (DEGs)

Figure 1 summarizes the steps used in the analysis. Quality filtered reads from control and vaccinated samples (0 day and 5 dpv) were mapped to the *Capra hircus* or *Ovis aries* reference genome for the respective subsets. The gene counts were obtained using Bowtie2.0 in RSEM [23]. The counts were used for calculating differentially expressed genes (DEGs) by use of R packages - EBSeq, DESeq2, edgeR. The common DEGs from the three packages were used for downstream analysis while fold changes for the corresponding genes were taken from DESeq2 [24].

157 **2.7 Gene Ontology Analysis**

Initially, DEGs of each subset (CD4+, CD8+, CD14+, CD21+ and CD335+) in sheep and goats were functionally annotated in g:Profiler to identify the significant immune system KEGG pathways. The expression of common DEGs of each subset between sheep and goats that are involved in immunological KEGG pathways is represented in a heatmap. Finally, to understand the co-ordinated response across all the subsets, genes expressed in all cell subsets were functionally annotated in g:Profiler (a gene is considered expressed if it is expressed in one subset).

165 **2.8 Comparison analysis using Ingenuity Pathway Analysis (IPA) analysis.**

Ingenuity Pathway Analysis (IPA) IPA is an all-in-one, web-based software application that enables analysis, integration, and understanding of data from gene expression, miRNA, and SNP microarrays, as well as metabolomics, proteomics, and RNAseq experiments.The DEGs from all the subsets in both species were overlaid in 170 IPA against its Ingenuity Knowledge Base (IKB) to perform a comparison analysis.

171 Canonical pathways activated (Z score > 2) or inactivated (Z score < -2) across all the 172 subsets were identified.

173 **2.9 Protein-protein interaction networks**

Using knowledge based approach [25], [26], antiviral genes - IFIT3, ISG15, MX1, MX2, RSAD2, ISG20, IFIT5 and IFIT1 were selected based on their expression in at least one subset. Protein - protein interactions between these antiviral molecules and the common genes among all the subsets for each species were extracted using STRING [27] and customized scripts. The degree or connectivity was calculated using igraph package [28]. The complete interaction networks were visualized in Cytoscape 3.8.0 [29].

181 **2.10** Validation of DE genes by quantitative real time PCR (qRT-PCR)

182 gRT-PCR was performed using Applied Biosystems 7500 Fast system to validate the expression of key genes using GAPDH as an endogenous control by TagMan 183 184 chemistry in PBMC subsets. GAPDH was employed as the internal control as it was found to be suitable endogenous control in earlier studies in PPR [30]. Key genes used 185 186 in the study for validation by q-RT-PCR and their TaqMan probe IDs are - DDX58, Ch04684385_m1; IFIT3, AIAA1E0; IRF7, AI89L87; MX1, Oa04659431_m1; ISG15, 187 188 AI70N2Z; and GAPDH, AIFAT31. All the samples were run in triplicates. The relative expression of each sample was calculated using the $2^{-\Delta\Delta CT}$ method with control as 189 190 calibrator [31].

191 Results

192 In the present study, CD4⁺, CD8⁺, CD14⁺, CD21⁺ and CD335⁺ cells were enriched 193 (Supplementary Figure 1) from the blood collected (5 goats and 5 sheep) at 0 day and 194 5 dpv (5 days post vaccination). RNA was isolated from these subsets to profile the 195 transcriptome with an aim to delineate the systems biology behind the Sungri/96 196 vaccine induced immuno-protection at 5 dpv in sheep and goats. The number of DEGs in CD4⁺, CD8⁺, CD14⁺, CD21⁺ and CD335⁺ cells were 1834, 1641, 2343, 3910 and 197 3607, respectively, in goats and 1464, 1586, 1847, 721 and 4019, respectively, in sheep 198 199 (Figure 2A and 2B). Venn diagram was generated to examine the common and unique 200 DEGs among cells. On comparison, 618 and 139 DEGs were found common among CD4⁺, CD8⁺, CD14⁺, CD21⁺ and CD335⁺ cells, in goats and sheep, respectively. The 201 number of unique DEGs was highest in CD21⁺ cells of goats and CD335⁺ cells of sheep 202 203 (Figure 2C and 2D).

204 Gene Ontology Analysis

Initially to evaluate the changes within a subset, functional annotation for genes expressed in each subset was done using g:profiler. The immune system KEGG pathways enriched in each subset were assessed. In all the cells an innate immune response leading to cell mediated adaptive immune response was observed (Figure 3 and Figure 4).

210 CD4+ cells of Sheep and Goats

211 On comparing CD4+ cells in sheep and goats, Fc gamma R-mediated phagocytosis,

Toll-like receptor signaling pathway, NOD-like receptor signaling pathway, Th1 and Th2

cell differentiation, T cell receptor signaling pathway and Th17 cell differentiation were

found significantly enriched in both the species. Besides these, in goats, Fc epsilon RI

signaling pathway, C-type lectin receptor signaling pathway, TNF signaling pathway,

Chemokine signaling pathway and NF-kappa B signaling pathway were found enriched

in CD4+ cells.

218 CD8+ cells of Sheep and Goats

In CD8+ cells,Th17 cell differentiation, C-type lectin receptor signaling pathway, NOD like receptor signaling pathway, Antigen processing and presentation,Th1 and Th2 cell
 differentiation, T cell receptor signaling pathway, Fc gamma R-mediated phagocytosis,
 Chemokine signaling pathway, NF-kappa B signaling pathway were found enriched in
 both sheep and goats. Besides these, three more pathways - Rap1 signaling pathway,
 Fc epsilon RI signaling pathway and MAPK signaling pathway were enriched in sheep.

225 CD14+ cells of Sheep and Goats

In CD14+ cells of both the species, Th17 cell differentiation, C-type lectin receptor signaling pathway, NOD-like receptor signaling pathway, TNF signaling pathway, Fc gamma R-mediated phagocytosis, Chemokine signaling pathway and NF-kappa B signaling pathway were found enriched. Additionally, in goats, Toll-like receptor signaling pathway, Rap1 signaling pathway and Phagosome Th1 and Th2 cell differentiation were enriched.

232 CD335+ cells of Sheep and Goats

In CD335+ cells, Th17 cell differentiation, Toll-like receptor signaling pathway, C-type lectin receptor signaling pathway, Necroptosis, MAPK signaling pathway, NOD-like receptor signaling pathway, TNF signaling pathway, Fc gamma R-mediated phagocytosis, Chemokine signaling pathway and NF-kappa B signaling pathway were enriched in sheep and goats (**Figure 3 and Figure 4**). Th1 and Th2 cell differentiation and FoxO signaling pathway were found enriched additionally in goats CD335+ cells.

239 Common DEGs in each subset between Sheep and Goats

240 The common genes in sheep and goats that are involved in immunological processes in 241 each subset are represented in a heatmap (**Supplementary Figure 2**). In CD4+, CD8+, 242 CD14+, CD21+ and CD335+ cells the numbers of common genes involved in 243 immunological processes were found to be 67, 91, 122, 17 and 179, respectively. Most 244 of the common DEGs in CD4+, CD8+ and CD!4+ were found upregulated in both the 245 species. However, in CD21+ and CD335+, a contrast in expression of these genes was 246 observed between sheep and goats. Most of the DEGs were upregulated in goats but 247 downregulated in sheep.

248 Coordinated response

To understand the co-ordinated response across all the subsets, genes expressed in all cell subsets were functionally annotated. A total of 5512 and 5297 genes were found expressed across all subsets **(Supplementary File 2)** in goats and sheep, respectively. Among these, in goats and sheep, 689 and 703 genes, respectively, were found associated with innate immune response biological processes (**Supplementary Figure 3**). A subset of 544 immune response genes were found to be common between sheep and goats with 144 and 158 genes being unique, respectively. This shows that in both sheep and goats the coordinated vaccine response at 5dpv across all the subsets is towards triggering a strong innate immune response as evident from the upregulation of innate immune genes.

Comparison analysis across subsets using Ingenuity Pathway Analysis (IPA) analysis.

261 Ingenuity pathway analysis (IPA) evaluates the DEGs and predicts activation or 262 inactivation of pathways. A comparative analysis was done to evaluate the canonical pathways that are activated/inactivated across all subsets in both species using IPA. 263 264 Pattern Recognition Receptors (PRR) are the first line of defense against any pathogen. The role of RIG-I-like receptors (RLRs) - RIG-1, LGP2 and MDA-5 that sense viral 265 infection, [32] was found to be predominant in CD4⁺, CD8⁺ and CD14⁺ cell subsets at 5 266 dpv in goats and, in CD4⁺ and CD14⁺ cell subsets of sheep (Figure 5). This RIG-I 267 recognition of viral RNA induces anti-viral state in cells by phosphorylating the IRFs [33] 268 269 and regulating NF-kB activity through binding to Nf-kb1 3'-UTR mRNA [34]. This 270 activation of IRFs by cytosolic pattern recognition receptors was found to be significant 271 in CD4⁺ cells of goats and was triggered, though not significant in CD4⁺ cells of sheep, 272 and in CD8⁺ and CD14⁺ cells of both the species (Figure 5). IRF3 was upregulated in CD4+, CD14+, CD21+ and CD335+ of goats, and in CD335+ of sheep; IRF7 was 273 274 upregulated in CD4+, CD8+, CD14+ and CD335+ of sheep, and in all cell subsets of goats (Supplementary File 1). IRF7 was also identified to be the most prominent 275 276 upstream regulator across subsets in both the species. RNA viruses are also recognized by TLR3 (dsRNA) and/or by TLR7/8 (sRNA) [35]. At 5 dpv, role of PRRs in 277 recognition of viruses was found activated in CD14⁺ cells of both sheep and goats, and 278 279 in CD8⁺ cells of goats. TLR2 and TLR4 were upregulated in CD14⁺ cells of both sheep and goats and in CD8⁺ cells of goats. This TLR signaling results in activation of 280 281 NF-kB and induction of IFN-inducible genes and co-stimulatory molecules [36].

The NF-κB activation by viruses was found activated in CD4⁺. CD8⁺ and CD14⁺ 282 283 cells of sheep and in CD8⁺ and CD14⁺ cells of goats but was inactivated in CD21⁺ cells 284 of both sheep and goats (**Figure 5**). The genes involved in this NF-κB pathway - CD4, 285 LCK, IKK, ERK 1/2, PKR and RIP were upregulated in CD4+ cells of sheep; LCK, RAS, 286 MEKK1, C-RAF, ERK1/2, IkB and CCR5 were upregulated, and CD21 and CXCR5 287 were downregulated in CD8+ cells of sheep; RAS, PKC, ERK 1/2, IkB, NFkB & PKR 288 were upregulated in CD14+ cells of sheep; CD4, LCK, RAS, PKR, ERK 1/2 and IkB were upregulated and CXCR5 was downregulated in CD8+ cells of goats; RIP, PKR, 289 290 AKT, IKK, ERK 1/2, IkB & c-RAF were upregulated and CXCR5, CD4 & LCK were 291 downregulated in CD14+ cells of goats(Supplementary File 1). NF-KB acts as a 292 mediator of pro-inflammatory and anti - inflammatory gene induction and plays a role in 293 regulating T-cell differentiation and effector function [37]. Several interleukin and chemokine signaling pathways were found activated in CD4⁺, CD8⁺ and CD14⁺ cells of 294 295 both the species i.e. IL-1 signalling in CD8+ cells of goats and CD14+ cells of sheep; IL-296 15 signalling in CD8+ cells of sheep and goats; IL-2 signalling in CD8+ cells of sheep 297 and goats and CD14+ cells of sheep; IL-22 signalling in CD4+ cells of sheep and CD8+

cells of sheep and goats; IL-6 signalling in CD8+ and CD14+ cells of sheep and goats;
IL-8 signalling in CD4+ cells of sheep, and CD8+ cells and CD14+ cells of sheep and
goats, and ; chemokine signalling in CD8+ cells and CD14+ cells of sheep and goats
(Figure 5).

302 Dendritic cell (DC) maturation was found significantly activated in CD21+ and 303 CD8+ cells of both the species. DCs are known to present antigenic peptides 304 complexed with MHC class I molecules to CD8-expressing T cells in order to generate 305 cytotoxic cells [38]. The Interferon signalling pathway that is essential for increased 306 cellular resistance to viral infection was found activated at 5 dpv in CD4⁺, CD8⁺ and 307 CD14⁺ cells of both the species. Interestingly, IFN alpha and beta were not 308 dysregulated in any of the subsets in both sheep and goats. The IFN receptors IFNAR1 309 and IFNAR2 were downregulated in most of the subsets. The absence of expression of type-I interferons in our study suggested IFN-independent ISG stimulation as reported 310 311 previously for PPR [7]. However, IFNgamma receptors were found to be activated in 312 most of the subsets. Further, most of the canonical pathways were identified to be 313 inactivated in the CD21+ cells. This indicated that the CD21+ cells are activated later for 314 the production of antibodies as significant increase in antibody production against PPRV 315 vaccination was observed 14 dpv [39]. The enrichment (-log p value) of genes in cell 316 mediated immune response and humoral immune response biofunctions was significantly higher in goats than in sheep in CD4⁺, CD14⁺, CD21⁺ and CD335⁺(Figure 317 318 **6**).

319 **Protein-protein interaction networks**

320 The protein-protein interaction network includes hubs connected with interacting 321 genes. The hubs in a network reflect the functional and structural importance of the 322 network. A total number of 618 and 139 DEGs were found to be commonly expressed in 323 Goats and sheep, respectively in all the subsets (Figure 2). On deciphering the 324 interactions between these DEGs and the 8 antiviral molecules (IFIT3, ISG15, MX1, 325 MX2, RSAD2, ISG20, IFIT5 and IFIT1) considered under the knowledge based 326 approach, most of the antiviral molecules formed the hubs in the network. ISG15 in both 327 species was found be the major hub with a connectivity of 75 and 16 in goats and 328 sheep, respectively (Figure 7A & Figure 7C). Heatmap of the genes involved in the 329 networks revealed that most of these antiviral genes in both the species are upregulated 330 (Figure 7B & Figure 7D).

331 Realtime PCR

The key genes identified from RNA-seq data - *DDX58, IFIT3, IRF7, ISG15* and *MX1* were validated by qRT-PCR. The expression of all the validated genes was in concordance with RNA sequencing results **(Table 1).**

335 **Discussion**

Vaccines protect against an infectious agent by inducing cells or molecules capable of rapidly controlling their replication or by inactivating their toxins. Primarily, vaccines trigger an inflammatory reaction, mediated by cells of the innate immune system - dendritic cells, monocytes and neutrophils. These cells recognize PAMPs through PRRs to get activated to produce cytokines and chemokines [40], [41], [42], 341 [43]. This inflammatory microenvironment is essential for differentiation of monocytes to 342 macrophages, and for activation and migration of dendritic cells into the draining lymph 343 nodes [44]. In the absence of this inflammatory response the dendritic cells remain 344 immature and the naive T cells in the lymph nodes do not differentiate into CD4+ T 345 cells. It is evident that PPRV - Sungri/96 live attenuated vaccine triggers activation of 346 the innate immune system after it is phagocytosed by monocytes/dendritic cells at the 347 site of administration [45]. This RNA virus may be then recognized by TLR3/7 on the 348 endosome or by the RIG-1 or MDA5 in the cytosol to induce an inflammatory response. 349 This induction of inflammatory response is evident in both sheep and goats with the 350 triggering of several pathways viz. role of RIG1-like receptors in antiviral innate 351 immunity, role of pattern recognition receptors in recognition of viruses, production of 352 nitric oxide and reactive oxygen species in macrophages, NF-KB activation by viruses, 353 and several IL signaling pathways in CD14+ cells. This triggering in inflammatory 354 response is much needed for the activation of dendritic cells and monocytes, and for further draining of these cells to the nearest lymph node where naive T cells are 355 356 activated [44]. This activation of T cells is clearly seen by the activation of pathways in 357 CD8+ (T- cytotoxic) and CD4+ (T-helper) cells.

358 Out of the several pathways activated in both CD4+ and CD8+ cells, NOD-like 359 receptor signaling pathway, Th1 and Th2 cell differentiation, T cell receptor signaling pathway and Th17 cell differentiation were found significantly enriched in both the 360 species. The differentiation of T cells to Th1 and Th2 is crucial for inducing the immune 361 362 response. Th1 cells stimulate cellular immune response, participate in the inhibition of 363 macrophage activation and stimulate B cells to produce IgM and IgG1 [46]. Th2 stimulates humoral immune response, promotes B cell proliferation and induces 364 365 antibody production [46]. The distinct subsets of helper T cells - TH1, TH2 and TH17, are effective at protecting against pathogens [47]. Additionally, activation of C-type 366 367 lectin receptors (CLRs) signaling in CD8+ cells of Sungri/96 vaccinated sheep and 368 goats and in CD4+ cells of goats indicates induction of adaptive immune response. C-369 type lectin receptors (CLRs) are important pattern recognition receptors involved in 370 recognition and induction of adaptive immunity to viruses [48]. However, in CD21+ cells 371 most of the pathways were found inactivated/not activated as 5 dpv may be too early a 372 time point to detect activation in the CD21+ cells.

373 NK cells (CD335+) are known to mediate both innate immune and adaptive 374 immune responses by modulating both CD8+ and antibody production [49]. In this study 375 most of the pathways - interferon signaling, crosstalk between dendritic cells and natural 376 Killer cells, chemokine signaling, inflammasome pathway, iNOS signalling and 377 complement system in NK cells were found activated in vaccinated goats than in sheep. 378 Upregulation of RIG-1 and MDA5 in NK cell of goats reflects setting off of the innate 379 immune response [50]. Also, activation of interferon signaling pathway in infected NK 380 cells of goats suggests evoking of both the innate and adaptive immune responses [51]. 381 The activation of iNOS signaling invokes immune response in virus infected cells 382 [52]. The activation of complement system in NK cells aids in antibody production by 383 bridging both innate and adaptive immune response [53]. Upregulation of CD69, 384 NKp30, FAS & TNFR2 and activation of crosstalk between dendritic cells and natural 385 killer cells pathway, in goats must be embarking innate immune response, followed by adaptive response on antigen presentation after vaccination. The activation and 386

triggering of several pathways in NK cells of goats at this early time point may be
 because of Sungri/96 vaccine strain being of goats origin and that the activation of
 these pathways at a later time point in sheep cannot be ruled out.

390 The network of antiviral molecules (IFIT3, ISG15, MX1, MX2, RSAD2, ISG20, 391 IFIT5 and IFIT1) with the DEGs commonly expressed in the subsets in both sheep and 392 goats, reflected ISG15 as a major hub. The network was found to be dense in goats in 393 comparison to sheep. ISG15 is one of the most highly induced ISGs in viral infections 394 [25], [54] and was also found to be directly induced by IRF3/IRF7, independent of IFNs 395 [55], [56], [57]. It is an ubiquitin-like protein that covalently attaches to target proteins in 396 a process known as ISGylation [25], [58]. HERC5 is considered as the major ligating 397 enzyme in ISGylation. This ISGylation of viral proteins was reported to have an 398 inhibitory effect on the viral infection [59], whereas ISGylation of host proteins leads to 399 either activation [59] or increase in stability [60]. HERC6 instead of HERC5 is 400 considered as the major ligating enzyme in mice [61]. In our study, HERC5 and HERC6 401 were found upregulated in goats. Further, the antiviral gatekeeper MX1 acts prior to 402 genome replication at an early post entry step of the virus life cycle. Similarly, MX2 403 specifically targets viral capsid and affects nuclear entry of the HIV-1 [25], [62], [63], 404 [64]. IFIT family (IFN-induced protein with tetratricopeptide repeats) are a group of ISGs 405 that inhibit virus replication by binding and regulating the functions of cellular and viral 406 proteins and RNAs [65]. IFITs were also characterized to play a critical role in protecting hosts from viral pathogenesis. RSAD2, also known as Viperin, is the another most 407 highly induced antiviral effector found in ER and ER-derived lipid droplets [66]. RSAD2 408 409 was characterized to have various modes of antiviral action to inhibit enveloped viruses 410 [67]. It can also affect virus life cycle at an early stage by inhibiting RNA replication [68]. 411 All these genes - MX1,MX2,IFIT1, RSAD2, IFIT3 and IFIT5 were found upregulated in 412 both sheep and goats suggesting a strong antiviral response in both the species.

413 It is important to note the in our study both sheep and goats survived PPRV 414 virulent virus challenge post vaccination, indicating an adequate immune response to 415 counter the virus. In an independent study it was reported that Sungri/96 vaccine is 416 equally potent in both sheep and goats [69]. In our study, though there are a few 417 differences in the systems biology across cells (specially the NK cells) between sheep 418 and goats, the co-ordinated response that is inclusive of all the cell subsets was found 419 to be towards induction of strong innate immune response, which is needed for an 420 appropriate adaptive immune response.

421 **Conflict of interest**

- 422 None of the authors have a conflict of interest to declare.
- 423 Author contributions

RKS, BPM, and RKG conceived and designed the research. KKR and DM
performed the vaccine testing experiment. SAW and ARS conducted the wet lab work.
SAW, MRP, RINK and RKG analyzed the data. SAW, MRP, RKG, APS, and BM helped
in manuscript drafting and editing. RKS, BPM, and RKG proofread the manuscript.

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- 639
- 640
- 641 Legends
- 642 **Figure 1:** Workflow for RNA sequencing data analysis
- 643 **Figure 2:** Number of dysregulated DEGs in PBMC subsets (A) Goats and (B) Sheep.
- 644 Green colour represents downregulation and red colour represents upregulation.
- 645 (C)Venn diagrams representing unique/common DEGs among cells in goats (D) Venn
- 646 diagrams representing unique/common DEGs among cells in sheep
- **Figure 3:** Functional annotation of DEGs involved in immunological processes for each subset of PBMCs: (A) CD4+, (B) CD8+, (C) CD14+, (D) CD21+ and (E) CD335+ using q:Profiler in Goats

Figure 4: Functional annotation of DEGs involved in immunological processes for each subset of PBMCs: (A) CD4+, (B) CD8+, (C) CD14+, (D) CD21+ and (E) CD335+ using g:Profiler in Sheep

Figure 5: Comparison analysis of canonical pathways related to immunological processes among the subsets of PBMCs in both sheep and goats using IPA. Blue colour represents Z score < 0 and red colour represents Z score > 0. Z score ≥ 2 means activation of canonical pathways and Z score ≤ -2 means inactivation of canonical pathways

Figure 6: Comparison of significant enrichment (-log p value) of genes in cell mediated immune response and humoral immune response bio functions in (A) CD4+, (B) CD8+, (C) CD14+, (D) CD21+ and (E) NK cells (CD335+) between vaccinated goats and sheep.

Figure 7: Protein-protein interaction network of antiviral genes - IFIT3, ISG15, MX1,
MX2, RSAD2, ISG20, IFIT5 and IFIT1 with the common DEGs across all subsets of
each species (A) Goats and (B) Sheep. Size of circle indicates the degree of interaction.
(C) Heatmap for fold change (Log₂FC) values of DEGs involved in the network among
the subsets of PBMCs of goats. (D) Heatmap for fold change (Log₂FC) values of DEGs
involved in the network among the subsets of PBMCs of sheep. Green colour indicates
downregulation and red colour indicates upregulation.

Supplementary Figure 1: Purity of PBMCs subsets - T helper cells (CD4+), T cytotoxic cells (CD8+), monocytes (CD14+), B lymphocytes (CD21+), and natural killer cells (CD335+) by flow cytometry. 'PBMCs' means unstained cells. 'PBMC' with Ab' means before magnetic beads cell separation. Enriched cells mean post magnetic beads cell separation.

Supplementary Figure 2: Heatmap for fold change (Log₂FC) value of common immune
DEGs between goats and sheep for each subset of PBMCs: (A) CD4+, (B) CD8+, (C)
CD14+, (D) CD21+ and (E) CD335+. Green colour indicates downregulation and red
colour indicates upregulation.

- 678 **Supplementary Figure 3:** Functional annotation related to immunological processes of 679 genes expressed in all the subsets or in one subset of PBMCs associated with innate 680 immune response (A) Goats (B) Sheep
- 681 **Supplementary File 1:** Individual list of DEGs expressed in the subsets of goats and 682 sheep
- 683 **Supplementary File 2:** List of 5512 and 5297 DEGs expressed across the subsets in goats and sheep, respectively

685



Validation of RNA Sequencing data by qRT-PCR using TaqMan assays

DEGs in Goat





(C)

(A)

DEGs in Sheep



(D)



(B)











Canonical Pathways	CD4+(Goat)	CD4+(Sheep)	CD8+(Goat)	CD8+(Sheep)	CD14+(Goat)	CD14+(Sheep)	CD21+(Goat)	CD21+(Sheep)	CD335+(Goat)	CD335+(Sheep)
Activation of IRF by Cytosolic Pattern Recognition Receptors	2.524	1.213	1.941	0.775	1.414	1.147	0.378	-3.162	1.347	-1.347
Acute Phase Response Signaling	1.826	1.091	4.426	2.683	4.323	3.43	-1.571	-0.302	1.387	-2.661
B Cell Activating Factor Signaling	0	1.633	N/A	1.633	-1.265	0	-2.4	N/A	-2	-2.837
B Cell Receptor Signaling	-1.896	-0.378	1.029	0.714	0.516	1.109	-4.088	-1.789	-2.278	-3.71
CCR5 Signaling in Macrophages	1	2.121	1.414	2.111	0.333	1.414	-1	-2	1.155	-1.387
CD28 Signaling in T Helper Cells	0.169	2.043	1.4	3.333	-0.926	0.928	-2.887	-2.673	-0.687	-4.621
CD40 Signaling	-0.258	1.291	0.905	0.832	0.471	0.5	-3.157	-2.646	-1.671	-3.182
Chemokine Signaling	1.387	1.604	3.638	2.828	2.2	2.236	0	-1.633	2.746	-0.2
CNTF Signaling	-0.577	0.832	2,138	3 357	1.886	1.604	-3,138	-2 449	-0.408	-3.157
Complement System	2.236	N/A	2,449	N/A	1.89	1.633	0.816	N/A	2.646	N/A
Crosstalk between Dendritic Cells and Natural Killer Cells	2 84	1.414	2 673	2.828	1.698	2 333	0	-1.342	2 828	-1.5
CXCR4 Signaling	-0.186	1 732	3.087	1 732	2 534	2 556	-1 786	-2 887	0.42	-2 722
bioRxiv preprint@i/hm/my/my/To_110/2000 broversteen eligitesteele Amontargis TerfcoTygoropelle (Dellespreprint	1 414	0 447	1 89	1.89	-1.342	1	1.342	N/A	1 89	-2 236
(which was not certified by peer review) is the author/runder, who has granted bioRxiv a ficense to display the preprint in perpetuity. It is made Dendrific Cell Mailable under a CC-BY 4.0 International license.	1 761	2 4 4 9	3 272	1.80	2.832	2 967	-1 511	-2.324	1 414	-3 727
Ec Epsilon RI Signaling	-0.6	1 213	1.606	0 784	1 414	2.6	-2 654	-2 121	-0.309	-2.832
GM-CSE Signaling	-0.688	0.535	2 828	2 982	3 128	2.0	-2 921	-1 89	0.000	-3 182
Granzyme B Signaling	-1	0.000 N/Δ	Ν/Δ	Ν/Δ	-0 447	-0 447	-2 236	Ν/Δ	-1 633	-1.89
HMGB1 Signaling	0.209	1	3 138	3 1/1	2 705	3 651	-2.200	-2 71/	0 781	-7.00
iCOS-iCOSL Signaling in T Helper Cells	0.203	2	1 201	2 268	_1 005	0	-1.230	-2.714		-4.450
II -1 Signaling	1 265	1 807	2 236	1 667	1 604	2 309	-2.402	-2.121 _1	-0. 4 03	-2 132
IL-15 Signaling	-0.218	1.037	2.230	3	0.816	2.303	-3.087	-1	U -0 686	-3.43
IL 2 Signaling	0.210	1.231	2.105	3 578	1 270	2 128	2.8	-5	0.365	3 024
IL-2 Signaling	0.471	2.226	2.0	2.226	1.279	2.130	-3.0	-2.040	-0.303	-3.024
IL-22 Signaling	0.6	2.230	2 71	2.230	2.902	2.656	-1.155	-2	0.302	-2.111
IL-0 Signaling	0.0	0.943		3.13	0	0.729	-2.037	-0.032	1.09	-3.200
IL-7 Signaling Falliway	-0.210	1.213	0.032	2.651	0	0.720	-3.413	-2.333	-1.09	-4.042
IL-0 Signaling	0.622	2.191	4.11	3.001	4.341	4.422	-2.073	-3.030	1,609	-3.437
IL-9 Signaling	0.577	1 242	0.370	1.007	0	0.707	-3	-2.040	-1.090	-4
	1.414	-1.342	2.449	N/A	2.230	1.007	-0.447	N/A	2.490	-1.033
INOS Signaling	2.309	2.111	3.102	2.03	2.490	2.073	-0.218	-1.134	2.005	-2.524
Interferon Signaling	2	2.309	3.5	3.000	3	3.742	1.342	-0.707	3	0
MIF Regulation of Innate Immunity	1.897	2.449	3	1.342	1.007	2.309	1.732	-0.447	3.357	-0.277
Natural Killer Cell Signaling	0.603	1.183	2.611	4.082	1.633	2.53	-4.276	-2.183	0.361	-4.028
$NF-\kappa B$ Activation by Viruses	0.426	2.524	2.837	2.236	2.191	2.828	-3.452	-2.714	0	-4.217
NF-κB Signaling	-0.324	0	1.257	0.18	1.82	1.982	-5.333	-2.324	-1.756	-4.677
p38 MAPK Signaling	0.577	2.138	2.84	2.5	1.706	2.4	-1.095	-0.378	0.539	-3.43
PI3K Signaling in B Lymphocytes	-0.905	1.4	2.137	1.406	1.414	1.761	-2.286	-3.207	-0.64	-2.994
PKCe Signaling in T Lymphocytes	0.667	2.887	2.887	2.967	-0.295	1.095	-2.889	-2.53	-0.662	-4.811
Production of Nitric Oxide and Reactive Oxygen Species in Macrophages	1.622	1.1//	3.43	2.556	2.771	3.086	-1.886	-1.964	0.254	-2.853
Role of JAK1, JAK2 and TYK2 in Interferon Signaling	N/A	0.447	2.236	N/A	1.342	1.342	-0.707	N/A	1.342	-1.633
Role of NFAT in Regulation of the Immune Response	0.152	3.053	3.087	2.771	0.412	1.061	-2.213	-3.5	-0.372	-5.08
Role of Pattern Recognition Receptors in Recognition of Viruses	1.4	0.5	2.449	0.535	2.785	3.266	-1.915	-1.633	1.298	-3.272
Role of PI3K/AKT Signaling in the Pathogenesis of Influenza	-0.775	0.378	0.707	1.667	0.832	-0.302	-2.183	-2.828	-0.655	-3.153
Role of PKR in Interferon Induction and Antiviral Response	2.041	0.2	2.117	0.655	1.257	1.768	-2.343	-1.291	0.469	-2.722
Role of RIG1-like Receptors in Antiviral Innate Immunity	1.667	0.707	1.342	0	2.121	1.633	-0.905	-2	0	-2.496
Th1 Pathway	0	1.342	2.746	1.512	0.343	0.816	-1.234	-2.138	0.152	-3.893
Th17 Activation Pathway	0.277	-0.277	-0.707	0.728	-0.943	0.535	-3.4	-0.378	-2.353	-3.771
Th2 Pathway	0.557	1.043	1.46	2.117	0.354	2.065	-1.947	-2.121	0.762	-2.874
TNFR1 Signaling	1.941	1.667	1.414	2.121	2.111	2.138	-2	-1	0	-2.236
Toll-like Receptor Signaling	1.265	0.577	2.111	-0.632	1.807	2.324	-2.236	1.342	0.688	-2.065
TREM1 Signaling	0.447	0	1.5	0.426	2.711	2.837	-3	0	1.095	-3.78







(C) _{Genes}

	CD4+(Goat)	CD8+(Goat)	CD14+(Goat)	CD21+(Goat)	CD335+(Goat)	Genes	CD4+(Goat)	CD8+(Goat)	CD14+(Goat)	CD21+(Goat)	CD335+(Goat)
	1.985919432	1.71876284	1.15136785	2.596312933	2.793894552	PSMB10	0.733634947	0.811814649	0.471405892	1.797708577	1.614186152
	-0.846500334	-0.759261474	-0.730885501	-2.133744367	-1.666365376	PSMA6	0.510611356	0.655455073	0.693812493	1.205230743	1.007605142
6	1.583485414	1.768262174	1.030203217	2.24159954	2.218249521	PSMA3	0.557769618	0.784140139	0.520217636	1.66137058	0.997966857
	1.474048446	0.69800351	1.013086907	0.640549219	1.665751527	PLSCR1	1.873187503	1.996412272	1.564759696	2.403630449	3.507333561
	0.603191954	0.762936426	0.399184392	1.731154544	1.227057625	PLAC8	0.271393827	1.041879029	0.788244067	1.796782612	0.331973468
2A	0.834909533	1.376929861	3.502396279	1.288194232	3.002356344	PARP9	0.451327594	0.677068178	0.298457762	0.527184722	0.498137374
	0.867704673	0.905132405	0.727333005	0.39565109	1.124896732	OAS1	1.742622357	2.312926759	1.455953723	1.248023064	1.921085233
	-1.249225733	-0.627167681	0.425388161	-0.657745261	-0.422975202	NUP153	-0.437629202	-0.524970959	-0.488578097	-2.23713924	-1.419258684
	-2.93621845	-2.676344862	-1.850399723	-1.406230815	-2.863219945	NLRC5	0.4550748	0.253996148	0.337083634	-1.107697252	-0.213593162
	0.481424937	0.709629829	0.375608441	-1.103549185	0.588575974	MX2	2.229995482	2.455105126	1.265294526	2.269962652	2.669472068
	0.686534554	0.979474724	0.412903447	0.347633714	0.761319	MX1	1.436277669	1.928714159	1.406726256	1.872160903	1.669321064
Л1	0.352276035	0.299236491	0.298744598	0.48397953	0.469815216	MOV10	0.565233775	0.343280646	0.30756072	0.587073914	0.257462
CA4	-0.578276522	-0.510995083	-0.570504563	-0.578081186	-1.04716526	LY6E	1.629232915	1.347876258	-1.449384456	1.977711152	1.549066722
G	1.198313102	1.453634	1.021035919	2.063245421	1.688438395	ITSN2	-0.55278938	-0.352319861	-0.424744518	-1.376095373	-1.319074279
2	1.683033244	1.544952511	1.323374254	1.044086576	2.573485598	ISG20	2.690769097	2.210485397	2.522078248	3.469713783	4.847516641
	0.777564021	0.90868795	0.556943188	2.068525254	1.252645748	IRF7	1.975621957	2.041883882	1.39959564	2.008508899	2.423957195
	0.405565069	0.428737023	0.127851061	1.70468363	0.781272276	IRF1	0.690274379	1.066038277	0.641875829	0.947798974	1.726629791
	0.620685866	0.660895494	0.157868984	1.733896081	0.991905629	IFITM3	1.147254384	1.492668765	1.632635537	1.93728427	2.719419248
	0.493112145	0.658913473	0.322262188	1.58/516454	0.934397007	IFIT5	1.576467091	1.876552182	1.48437405	1.294382276	2.55568389
A	0.622541955	0.632672895	0.290766873	1.767862173	1.056662945	IFIT3	2.088394134	2.055511315	1.260277191	2.566300219	3.351404589
	0.774925566	0.94882869	0.634872728	1.9/294/9/8	1.227759202	IFIT2	2.089832269	1.439416054	0.84672498	1.251797719	2.630431043
	0.64305394	0.84756223	0.596450189	1.863417854	1.283145592	IFIT1	1.011625524	1.604485968	1.367435517		1.941090719
· ^	0.736697087	0.682581224	0.31126961	1.665172295	1.096183673	IFIH1	0.51269938	0.677131803	0.614964862	1.089893378	1.129685142
A	0.692281103	0.79558963	0.385424280	1.8/312/003	1.186247482	IFI6	2.662452737	2.810609128	2.056178067	4.063842744	3.915749987
	0.591095576	0.0410/91/2	0.121590110	1.909370429	0.995265155	IF144	1.611197864	1.626112568	0.99882251	0.957829634	1.854207017
	0.003704033	0.700001750	0.307704030	1 2000000000	0.972962646	IFI35	0.85134324	0.832061752	1.008101155	1.493265385	1.379011343
	0.659221727	0.736402252	0.240490455	1,3000000007	1 196120001	HERC6	1.251223853	1.243896935	1.008493382	1.654659051	2.110535977
	0.030231737	0.730402332	0.427130492	1.712071407	1.100120991	HERC5	0.53783446	0.680100193	0.701848032	1.24698382	0.617116848
	0.846636567	0.00004577	0.540914105	1 882778237	1 30070175	HERC4	-0.841200614	-0.968971928	-0.650860266	-0.82631207	-0.744158812
	0.392090004	0.511404505	0.033043400	1 760147365	1.04455426	GBP5	1.032090025	0.697919258	0.687773578	1.953152755	1.675931943
	0.701113137	0.725097955	0.503321859	1.698767509	1 227847141	EPSTI1	1.020866224	1.373996773	1.261346301	1.795698135	1.302099731
-C18orf32	0.640708457	0.538833383	0.215656928	1.0507.07000	1 10988575	EIF4A1	0.293040687	0.729964388	0.659183743	1.214401196	0.904770093
OTOONOL	0.461682141	0.586630315	0.381422392	1 54011522	0.833372263	EGR1	1.760755155	-1.443962337	0.304496671	-4.962753888	3.197575567
4A	-1 762118555	-1 662711873	-2 220342188	-1 212212469	-1 836131361	DHX58	1.336670449	1.255929883	1.320958745	1.10218703	1.481351351
	-0.736550972	-0.789847617	-0.873204868	-1.5013996	-1.460412784	CXCL10	1.792154502	3.422134781	2.31101281	3.539210836	2.888757896
2	0.673680396	1.148968677	0.781523858	1.91730791	1.77574944	CD44	0.381670686	0.683512435	0.440690852	1.352904732	2.164353953
3	0.514993979	0.541902318	0.565066178	1.294655603	0.625377969	ATAD2B	-0.768274143	-1.011170596	-0.724555682	-2.526543754	-1.654225422
1	0.510614155	0.416092405	0.646466185	1.039847072	0.952878937	APAF1	-0.628435852	-0.7267511	-0.523891148	-3.304771213	-1.375874722
9	0.649206706	0.864351477	0.783614851	1.512848475	1.229128582	ADAR	0.504687791	0.43592216	0.561508709	-2.033047968	-0.277606608
3	0.468121416	0.7005429	0.472003684	0.923825114	0.863216931	ACTB	-0.409201064	0.233056082	0.163420159	0.533279959	0.402504285

(D)						
()	Genes	CD4+(Sheep)	CD8+(Sheep)	CD14+(Sheep)	CD21+(Sheep)	CD335+(Sheep)
	B2M	0.555014805	0.743559759	0.642736118	-3.166964292	0.176715854
	CLTC	-0.611987861	-0.247385808	-0.162945173	-3.778268513	-1.807783063
	DHX58	0.931844411	0.940998189	1.283793745	1.950240107	1.570444101
	EIF4B	0.201399468	0.326584257	0.285660754	-2.392207708	-0.350743731
	FAU	1.038610455	0.789887926	0.847207417	0.948749522	3.745174852
	IFI35	0.662547359	0.897738796	1.097915998	1.741171778	2.700369988
	IFI6	2.701745023	3.635058039	3.719327423	4.320482769	4.755508796
	IFIT3	0.951666611	1.648452647	2.122316917		
	ISG15	2.616943843	3.318903048	4.298601385	5.329151576	8.084652166
	MX1	1.070460774	0.953951731	2.117664923		
	MX2	1.263597474	0.954600131	1.503219659		
	RPL32	0.829700069	0.684574133	0.682085333	1.036138102	3.491284495
	RPS10	0.791819633	0.542729659	0.561397257	0.97346268	3.107929968
	RPS19	0.969410857	0.624354671	0.715343866	1.369822956	3.712340186
	RSAD2	1.000793076	1.63556357	3.011670856		
	SQSTM1	0.393547984	0.572013184	0.435207049	-3.779889764	-1.998615399
	STAT1	0.46995802	0.700672665	0.401877473	-4.255216254	-0.780305439
	SUN2	0.369178067	0.82537195	-0.326065822	-6.160606981	-2.114598903
	UBE2L6	1.343925734	1.17152532	0.823433926	2.322966095	2.144767883
	UBR5	-0.502610312	-0.536023915	-0.661688091	-2.962216415	-3.174126569
	XAF1	1.574601815	1.268233816	1.586611438	2.18494683	2.609327539
	ZNFX1	1.159804825	1.392658968	1.364107897	-1.80735843	1.043963877
	ISG20		2.792463239	3.201044467		3.123564522
	IFIT5		1.082622922	2.112629892		
	IFIT1		1.284923043	2.744670094		

	CD4		CD8		CD14		CE	021	CD335	
Gene	log₂FC from qRT-PCR	log₂FC from RNAseq								
DDX58	1.179333333	1.163823481	1.123333333	1.126889098	0.980666667	0.897711159			0.97	0.866777509
IFIT3	2.308666667	2.088394134	2.12	2.055511315	0.869802704	1.260277191	2.630251215	2.566300219	3.071333333	3.351404589
IRF7	2.005631332	1.975621957	1.945542755	2.041883882	1.051373011	1.39959564	1.936792165	2.008508899	2.627666667	2.423957195
ISG15	3.193989334	3.483488156	3.214456762	3.642171919	1.508623454	2.749599382	5.020430272	4.958535352	4.033333333	4.817152957
MX1	1.164756699	1.436277669	1.825847584	1.928714159	1.289272029	1.406726256	2.063278147	1.872160903	1.556666667	1.669321064

Table 1.A: log₂FC from RNAseq and qRT-PCR of PBMC subsets isolated from sungri/96 vaccinated Goat

Table 1.B: log₂FC from RNAseq and qRT-PCR of PBMC subsets isolated from sungri/96 vaccinated Sheep

	CD4		CD8		CD14		CD21		CD335	
Gene	log₂FC from qRT-PCR	log₂FC from RNAseq	log₂FC from qRT-PCR	log₂FC from RNAseq						
DDX58	1.311	1.4439822 94	2.00666666 7	1.88032193 2	1.80666666 7	1.63016686 2				
IFIT3	0.536779277	0.9516666 11	1.69666666 7	1.64845264 7	1.92333333 3	2.12231691 7				
IRF7	2.214341863	2.1746786 99	1.84333333 3	2.32051383 2	3.23333333 3	3.01743016 1			3.15333333 3	3.66900063 9
ISG15	2.781512248	2.6169438 43	2.79666666 7	3.31890304 8	3.12666666 7	4.29860138 5	4.14250017 8	5.32915157 6	5.64333333 3	8.08465216 6
MX1	1.34357254	1.0704607 74	1.01	0.95395173 1	2.18666666 7	2.11766492 3				