



THE UNIVERSITY *of* EDINBURGH

Edinburgh Research Explorer

Susceptibility to disease (tropical theileriosis) is associated with differential expression of host genes that possess motifs recognised by a pathogen DNA binding protein

Citation for published version:

Larcombe, SD, Capewell, P, Jensen, K, Weir, W, Kinnaid, J, Glass, EJ & Shiels, BR 2022, 'Susceptibility to disease (tropical theileriosis) is associated with differential expression of host genes that possess motifs recognised by a pathogen DNA binding protein', *PLoS ONE*, vol. 17, no. 1, e0262051.
<https://doi.org/10.1371/journal.pone.0262051>

Digital Object Identifier (DOI):

[10.1371/journal.pone.0262051](https://doi.org/10.1371/journal.pone.0262051)

Link:

[Link to publication record in Edinburgh Research Explorer](#)

Document Version:

Publisher's PDF, also known as Version of record

Published In:

PLoS ONE

General rights

Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy

The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.



RESEARCH ARTICLE

Susceptibility to disease (tropical theileriosis) is associated with differential expression of host genes that possess motifs recognised by a pathogen DNA binding protein

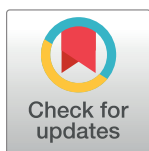
Stephen D. Larcombe^{1☯✉}, Paul Capewell^{1☯}, Kirsty Jensen², William Weir³, Jane Kinnaird¹, Elizabeth J. Glass², Brian R. Shiels^{1*}

1 Institute of Biodiversity, Animal Health, and Comparative Medicine, University of Glasgow, Glasgow, United Kingdom, **2** Division of Infection & Immunity, The Roslin Institute and R(D)SVS, University of Edinburgh, Edinburgh, United Kingdom, **3** School of Veterinary Medicine, University of Glasgow Veterinary School, University of Glasgow, Glasgow, United Kingdom

☯ These authors contributed equally to this work.

✉ Current address: Institute of Immunology and Infection, School of Biological Sciences, University of Edinburgh, Edinburgh, United Kingdom

* brian.shiels@glasgow.ac.uk



OPEN ACCESS

Citation: Larcombe SD, Capewell P, Jensen K, Weir W, Kinnaird J, Glass EJ, et al. (2022) Susceptibility to disease (tropical theileriosis) is associated with differential expression of host genes that possess motifs recognised by a pathogen DNA binding protein. PLoS ONE 17(1): e0262051. <https://doi.org/10.1371/journal.pone.0262051>

Editor: Gordon Langsley, Institut national de la santé et de la recherche médicale - Institut Cochin, FRANCE

Received: July 28, 2021

Accepted: December 15, 2021

Published: January 21, 2022

Peer Review History: PLOS recognizes the benefits of transparency in the peer review process; therefore, we enable the publication of all of the content of peer review and author responses alongside final, published articles. The editorial history of this article is available here: <https://doi.org/10.1371/journal.pone.0262051>

Copyright: © 2022 Larcombe et al. This is an open access article distributed under the terms of the [Creative Commons Attribution License](https://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Abstract

Background

Knowledge of factors that influence the outcome of infection are crucial for determining the risk of severe disease and requires the characterisation of pathogen-host interactions that have evolved to confer variable susceptibility to infection. Cattle infected by *Theileria annulata* show a wide range in disease severity. Native (*Bos indicus*) Sahiwal cattle are tolerant to infection, whereas exotic (*Bos taurus*) Holstein cattle are susceptible to acute disease.

Methodology/Principal findings

We used RNA-seq to assess whether *Theileria* infected cell lines from Sahiwal cattle display a different transcriptome profile compared to Holstein and screened for altered expression of parasite factors that could generate differences in host cell gene expression.

Significant differences (<0.1 FDR) in the expression level of a large number (2211) of bovine genes were identified, with enrichment of genes associated with Type I IFN, cholesterol biosynthesis, oncogenesis and parasite infection. A screen for parasite factors found limited evidence for differential expression. However, the number and location of DNA motifs bound by the TashAT2 factor (TA20095) were found to differ between the genomes of *B. indicus* vs. *B. taurus*, and divergent motif patterns were identified in infection-associated genes differentially expressed between Sahiwal and Holstein infected cells.

Data Availability Statement: RNA-seq data are publicly available at the NCBI GEO online repository under accession numbers GSM4824553–GSM4824563.

Funding: This work was funded by BBSRC: grant BB/L004739/1 and a BBSRC Strategic Programme grant (Control of Infectious Diseases: [BB/P013740/1]). The funder had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing interests: The authors have declared that no competing interests exist.

Conclusions/Significance

We conclude that divergent pathogen-host molecular interactions that influence chromatin architecture of the infected cell are a major determinant in the generation of gene expression differences linked to disease susceptibility.

Introduction

It is well established that the outcome of infection by a pathogen can vary between individuals from asymptomatic to fatal disease. How such variability is generated is of great importance, particularly for emerging viral disease such as COVID-19 and infectious disease of livestock. Adaptation of hosts that display no or mild clinical signs upon pathogen infection is thought to have arisen via competitive co-evolution [1,2]. In this process, mechanisms deployed by the pathogen to disrupt defence mechanisms are countered by the selection of genetic traits that attenuate this disruption. Over time this can lead to establishment of a general relationship where, upon infection, immunopathology is minimised to promote host survival and pathogen transmission. Hosts that have not co-evolved with a pathogen, however, are more susceptible to severe disease pathology when infected. Thus, host breeds that are generally tolerant of infection or susceptible to acute disease provide extremes of a susceptibility spectrum that can be exploited to investigate pathogen-host interactions that give rise to variable infection outcomes.

Tropical theileriosis, caused by the tick-borne apicomplexan parasite *Theileria annulata*, is a disease of great economic significance to livestock producers over large areas of the Old World [3,4]. It impacts both large and smallholding production systems with losses caused by mortality, reduced productivity and costs for treatment and control. *B. taurus* cattle show greater susceptibility to acute tropical theileriosis than *B. indicus* breeds native to regions endemic for *T. annulata* (reviewed in [5]). Susceptibility to the disease constrains the use of *B. taurus* in countries such as India, where a major crossbreeding programme aims to select for productive cattle tolerant to *T. annulata* infection [6]. While this has mitigated losses from acute disease, crossbred animals still show more pronounced signs of subclinical infection relative to native breeds [7,8]. Thus, strategies to improve disease tolerance of productive exotic breeds are needed but require greater understanding of how differential susceptibility to disease is conferred.

Previous work has investigated mechanisms of susceptibility to tropical theileriosis. Experimental challenge confirmed that infected Sahiwal (*B. indicus*) calves showed reduced measures of clinical pathology, relative to susceptible Holstein (*B. taurus*), including lymph node enlargement and fever [9]. A reduced pro-inflammatory response was observed for the Sahiwal calves, indicating that in a tolerant host immune pathology is limited, while in a susceptible one disease is caused by parasite-mediated dysregulation of the host immune response (reviewed in [2]).

During tropical theileriosis, sporozoites inoculated by a feeding tick rapidly invade leukocytes (of predominantly myeloid origin) and differentiate into the intracellular macroschizont stage [10]. Macroschizont formation is accompanied with transformation of the infected leukocyte into a cancer like cell. Infected cells proliferate in lymph nodes and metastasise throughout the lymphoid system to establish foci in the liver and lungs: death can occur via pneumonia like condition [11,12]. Based on the kinetics of disease it has been concluded that the major mechanism of tolerance is linked to the macroschizont infected leukocyte [9].

Moreover, five independent macroschizont infected cell lines derived from Sahiwal showed a reduced ability to transverse Matrigel compared to five Holstein lines, indicating that reduced invasiveness of the infected cell is linked to tolerance of infection [13].

Transformation of the bovine leukocyte by *Theileria* parasites occurs via constitutive activation of bovine transcription factors that act as inflammatory mediators and promote oncogenesis (for example, NF- κ B and AP1). Activation is accompanied with an extensive, irreversible reorganisation of the infected cell transcriptome [14] that alters the expression levels of many genes encoding proteins with the potential to promote oncogenesis or disrupt the immune response [15]. Thus, parasite-host interactions that activate and/or moderate inflammatory mediators in the transformed, infected cell are likely to be associated with disease susceptibility.

Only a handful of *Theileria*-encoded factors that translocate to the host cell compartment and modulate the infected leukocyte have been identified. TaPIN is a peptidyl-prolyl isomerase that was reported as secreted into the host cell by transforming *Theileria* species. It contributes to AP1 activation by destabilizing the FBW7 ubiquitin ligase that targets c-Jun, a key component of the AP1 heterodimer, for degradation [16]. Likewise, the Ta9 protein is located in the host cytoplasm and has recently been reported as an activator of AP1 [17]. Lastly members of the *TashAT* gene cluster are only found in transforming *Theileria* genomes, they are translocated to the host nucleus, possess AT hook domains homologous to those of mammalian HMGA proteins, bind DNA and modulate bovine gene expression [18–20]. It is unknown whether any of these candidate factors show altered expression in infected cells from tolerant breeds, but this could significantly influence the infected cell phenotype and potentially the severity of disease it generates.

To investigate whether differences in disease tolerance are mediated by pathogen-host interactions that modulate gene expression in the *Theileria*-transformed leukocyte, we have compared the transcriptome profile of the available set of *ex vivo* infected cell lines derived from Sahiwal vs. Holstein calves [13,21]. These low passage lines, although genetically diverse, offer the best infected cell lines available for comparative RNA-seq that relates to infection *in vivo*, since long term passage or cloning of infected cells results in attenuation of pathogenicity and a loss of parasite diversity [13,22]. The current work builds on an earlier microarray analysis, limited to 5000 immune related genes, that demonstrated differences in expression of genes encoding factors that regulate immune cell interaction [23]. The microarray study only covered the response to early infection events (first 72 h after invasion), and only a relatively small number of differentially expressed genes were identified (150 genes), reducing the power of pathway analysis. Moreover, a comparison of the host or parasite transcriptome of the Sahiwal and Holstein cell lines was not performed in the previous studies [13,21]. The results from our analysis show that infected cell lines from cattle breeds differentially susceptible to tropical theileriosis display significant differences in expression of bovine genes associated with parasite infection, innate immunity, cholesterol biosynthesis and oncogenesis. Furthermore, a screen for DNA motifs bound by the parasite encoded factor, TashAT2 identified differences in number and pattern between the genomes of *B. indicus* vs. *B. taurus*, which included infection-associated genes differentially expressed between Sahiwal and Holstein infected cell lines. These data provide a platform to characterise parasite-host interactions that have evolved to generate the wide range of disease outcomes that occur following infection.

Methods

Parasite infected cell lines and culture conditions

We used eleven infected cell lines, previously established *ex vivo* from the peripheral blood of cattle experimentally challenged with *T. annulata*. Six of these were derived from infected

Sahiwal calves (*B. indicus*) and five were derived from infected Holstein calves (*B. taurus*), as reported [13,21]. Both Sahiwal and Holstein animals were infected with sporozoites from the same parasite stock, *T. annulata* Hissar and cultured in RPMI-1640 medium (Sigma-Aldrich) supplemented with 10% FCS, 4 mM L-glutamine and 50 μ M β -mercaptoethanol at 37°C for a limited period (less than 10 passages). All cell lines were fully established based on the level of infection (>95% macroschizont infected cells). Clones were not generated in the original or current studies in order to retain the heterogeneity that naturally occurs upon infection *in vivo*, and because cloning and prolonged passage result in an attenuation of infected cell pathogenicity [13,22]. Thus, RNA was isolated from low passage cell lines, between passages 4–6, for Sahiwal and between passages 6–8, for Holstein. Cells were harvested by centrifugation (120 g for 10 min) after 48 h of culture. RNA was isolated using Trizol (Thermo Fisher Scientific) followed by purification using the RNeasy mini kit (Qiagen), including an on-column DNase digestion step, and using the suppliers' protocols. RNA quality and quantity were then assessed using a Nanodrop ND-1000 spectrophotometer (Thermo Fisher Scientific) and by gel electrophoresis.

RNA-seq

Between 160 and 260 μ g DNase-treated RNA was sequenced for each sample at the Centre for Genomic Research (CGR), Liverpool. Dual-indexed strand-specific RNA-seq libraries were made from total RNA, using NEBNext Poly(A) mRNA Isolation and Ultra Directional RNA Library Preparation kits (New England Biolabs). Sequencing was performed using one lane of Illumina HiSeq 4000 (Paired-end, 2 \times 150 bp sequencing, generating data from >280 M clusters per lane). Following sequencing, paired end reads were QCed and prepared for analysis at CGR: FASTQ files were trimmed for Illumina adapter sequences using Cutadapt version 1.2.1 [24] using option -O, so the 3' end of any reads which matched the adapter sequence for 3 bp or more were trimmed. The reads were further trimmed using Sickle version 1.200 [25], with a minimum window quality score of 20. Reads shorter than 20 bp after trimming were removed. Approximately 40 million paired end reads were generated from each sample.

Read mapping, differential expression analyses and pathway analyses

The trimmed FASTQ files for each sample were mapped to the *B. taurus* transcriptome: v UMD3.1 downloaded from Ensembl Biomart [26]. This strategy was used because of the greater fidelity of the *B. taurus* reference genome sequence relative to *B. indicus*, and on the premise that the two subspecies share a high level of sequence identity in mRNA coding regions of the genome. Bowtie2 [27] was used to map reads and generate SAM files, and the Python script "SAM2counts" [28] to generate reference sequence counts for each transcript. We collapsed reads mapping to multiple transcripts to the same Ensembl gene ID prior to analysis. Differential Expression analyses were performed on the count files using DESeq2 [29], to compare the five *B. taurus* samples to the six *B. indicus* samples. The *B. taurus* data values were used as the primary comparator, such that negative fold changes in gene expression are designated as lower expression in *B. taurus* vs. *B. indicus*, and positive fold changes are designated as higher in *B. taurus* vs. *B. indicus*. Genes with a false discovery rate (FDR) less than 0.1 (10%) were considered significantly differentially expressed between *B. indicus* relative to *B. taurus* infected cells lines. Following differential expression analyses, we used Ensembl Biomart to annotate the read files and Ingenuity Pathway Analysis (IPA) (Ingenuity Systems Inc., Redwood City, USA) to elucidate significantly modulated gene networks, disease pathways and potential effector molecules. We also mapped the RNA-seq reads to the *T. annulata*

genome [30], to assess whether parasite encoded genes are differentially expressed between the established macroschizont infected *B. taurus* vs. *B. indicus* cell lines.

Comparison with TBL20 infection associated dataset

To elucidate mechanisms that might underpin differences in breed susceptibility to *T. annulata* infection, we matched our data set of breed-associated differentially expressed genes with a data set of host cell genes whose differential expression was identified as associated with infection by *T. annulata*. The infection-associated data set was obtained by comparison of the transcriptome (obtained by microarray analysis) of uninfected (BL20) vs. macroschizont infected (TBL20) bovine lymphosarcoma cell lines, treated and not treated with the anti-*Theileria* drug buparvaquone. Full details can be found in [14,15]. As the two datasets used different systematic ID codes (Ensembl vs. Entrezgene), we first used Ensembl Biomart to filter the microarray BL20/TBL20 dataset by finding common Gene stable IDs. Since some genes were not found in this way we also used DAVID [31] and PANTHER [32] to generate alternative gene symbols to screen for overlap between gene lists independent of common gene IDs. To check for statistical significance, the representation factor was calculated as the number of genes in common \div expected number of genes; where the expected number of genes is given by: (number of genes in group 1 \times number of genes in group 2) \div number of genes in total. In our case we used 19,018 as the total number of genes as this is the number of unique identifiable bovine genes included on the microarray. The p value was then computed as a normal approximation of the exact hypergeometric probability [33].

Identification of infected host cell genes encoding predicted secreted or receptor proteins

Ensembl Biomart was used to generate the peptide sequences for all bovine genes classed as differentially expressed. A pipeline of publicly available resources was then used to perform *in silico* predictions. Signal P version 5.0 [34] generated a list of proteins with a predicted signal peptide (cutoff of >0.5 probability). Sequences for this new list were then screened for the presence of transmembrane domains or GPI anchors. This was performed using TMHMMserver v 2.0 [35] for transmembrane domains, and PredGPI [36] for GPI anchors. Any predicted proteins that scored positive for either motif was excluded from the list. The final list of proteins was obtained by screening against subcellular location prediction and gene ontology (GO) data available in the Genecards database version 4.14 [37]. Only proteins with a top confidence score as extracellular were retained in the final list. To generate a list of receptor proteins, we used gene ontology (GO) software to look for evidence of annotation as a receptor, or receptor-like domains in our data sets of differentially expressed genes. Thus, lists of annotations, protein family and classes were generated using Ensembl, DAVID, PANTHER and IPA. The data from each resource was collated, and we excluded any molecule that was not a confirmed or predicted receptor. Following receptor classification, the resulting gene list was curated by eye and nuclear receptors excluded, as we were primary interested in a cell surface location.

Validation of RNA-seq results by qRT-PCR

Validation of differential gene expression was carried out by qRT PCR, using RNA from the eleven *T. annulata* cell lines isolated on a different day than the samples used for RNA-seq. Primer pairs specific for 20 genes were designed (S1 Table) and qRT-PCR performed, as described previously [15]. Briefly, 500 ng of total RNA from each sample was used to synthesise cDNA, using the Affinity Script cDNA Synthesis Kit (Agilent Technologies) and Oligo-dT

as primer. 1 μ l cDNA for each sample was then used for qRT-PCR, using the Brilliant III Ultra-fast SYBR[®] Green qPCR Master mix (Agilent technologies) and the Stratagene Mx3005P system. Comparative quantitative analysis of gene expression across samples was performed using Stratagene MxPro Software. *REPS1* (RALBP1 associated Eps domain containing 1) was utilised as a house keeping gene, based on previous identification of constitutive expression in *Mycobacterium bovis* infected macrophage cells [38] and initial validation using RNA from the eleven *T. annulata* infected cell lines. The $\Delta\Delta$ CT method was then used to calculate fold-change (vs. the lowest expressed sample). The data were transformed on the \log_{10} scale to stabilize the variance and the mean group differences (Sahiwal vs. Holstein) were tested using Student's t tests.

Bioinformatic screen for DNA motifs bound by TashAT2

A consensus of AT rich motifs preferentially bound by the AT hook domain of recombinant TashAT2 was generated using previously published data [19]. This was used to generate a PROSITE pattern (TAAAT(1)N(4,6)T(1)A(3,4)T.) to search the *B. taurus* (UCD1.2) and *B. indicus* (Bos_indicus_1.0) genomes using FUZZNUC [39]. Bedtools [40] was used to find the intersections of identified motifs and introns and exons in the bovine genomes. Proximal upstream and downstream sequences in validated mRNA sequences not characterized as exons or introns were defined as untranslated regions (UTRs) and the intersection of motifs assessed. Sites without intersection were characterised as non-coding. Pathway enrichment of genes with TashAT2 binding motifs was assessed using PANTHER [32] and summary figures generated using CIRCOS [41].

Immunoblotting and immunofluorescence

Three of the independent Sahiwal infected cell lines (SA-C) and three Holstein lines (HA-C) were randomly selected, passaged 1X, cultured for 48 h and harvested, as described above. Cells were then washed 3X in phosphate buffered saline (PBS), resuspended in 20 μ L of PBS per 10^6 cells and an equal volume of 2X SDS-sample buffer added. Immunoblotting was performed using standard methodology, as described [42]. Antisera raised against the TA06460 fusion protein (the endoplasmic reticulum form of HSP90 (ER-HSP90)) was used at a dilution of 1:1500, as previously described [42]. For the Ta9 (TA15705) antigen, rat antisera were raised against a 35.7kDa fusion protein [43]. Antiserum production was carried out by the PTU/BS Unit at the Scottish National Blood Transfusion Service, Pentlands Science Park Edinburgh. The anti-Ta9 serum was used at a dilution of 1 in 1200.

Slide preparation, fixation and immunofluorescence were performed as described [15]. The cell lines used were the same as for immunoblotting and the EL24 antiserum specific for TashAT2 [44] was used at 1 in 250. Images were acquired with an Olympus BX60 microscope, SPOT camera and SPOTTM Advanced image software Version Mac: 4.6.1.26, using the matched exposure feature (Diagnostic Instruments).

Ethics statement

This study did not require ethical approval since it was conducted on *in vitro* cell lines established *ex vivo* from animals in a previous study where ethical approval was met (details available in [23]) and no further human or animal participation was needed.

Results

Early passage Holstein and Sahiwal infected cell lines display significant differences in their host transcriptome profile

Six Sahiwal and five Holstein lines infected with *T. annulata* (Hissar) were utilised for differential transcriptome analysis. As reported previously [9], under standard *in vitro* culture conditions, no significant difference in growth potential between the Sahiwal and Holstein lines was observed. RNA was isolated from each line and RNA-seq performed: the RNA-seq data are publicly available at the NCBI GEO online repository under accession numbers GSM4824553–GSM4824563. The results obtained indicate pronounced differences in host gene expression levels between *T. annulata* infected cells derived from Holstein or Sahiwal cattle. These differences were marked in terms of the large number of genes differentially expressed: 2211 genes (without duplicates) at $FDR < 0.1$ (adjusted), subsequently referred to as Holstein/Sahiwal differentially expressed (DE) dataset (H/S-DE). The full list of H/S-DE genes can be viewed in [S1 File](#). Of the 2211 DE genes there was an almost even split in genes displaying modulated expression associated with one breed relative to the other; thus, 1068 genes (48.3%) were expressed at lower levels in the Holstein infected lines compared to Sahiwal. Furthermore, analysis of the proportion of reads mapped to the *B. taurus* genome showed no statistical difference between the 6 Sahiwal relative to the 5 Holstein RNA-seq sample sets ([S2 Table](#)). These results support the premise that sequence identity between mRNA coding regions of the two genomes allow comparative analysis of gene expression using the *B. taurus* genome as the reference.

A demonstration that the H/S-DE data set represents differences in gene expression profiles between infected cells from Holstein vs. Sahiwal was provided by principal component analysis (PCA). Clustering of the samples with host type can clearly be seen in the plot shown in [Fig 1](#), where PC1 explained 35% of variation in the data, although a sizeable degree of variability

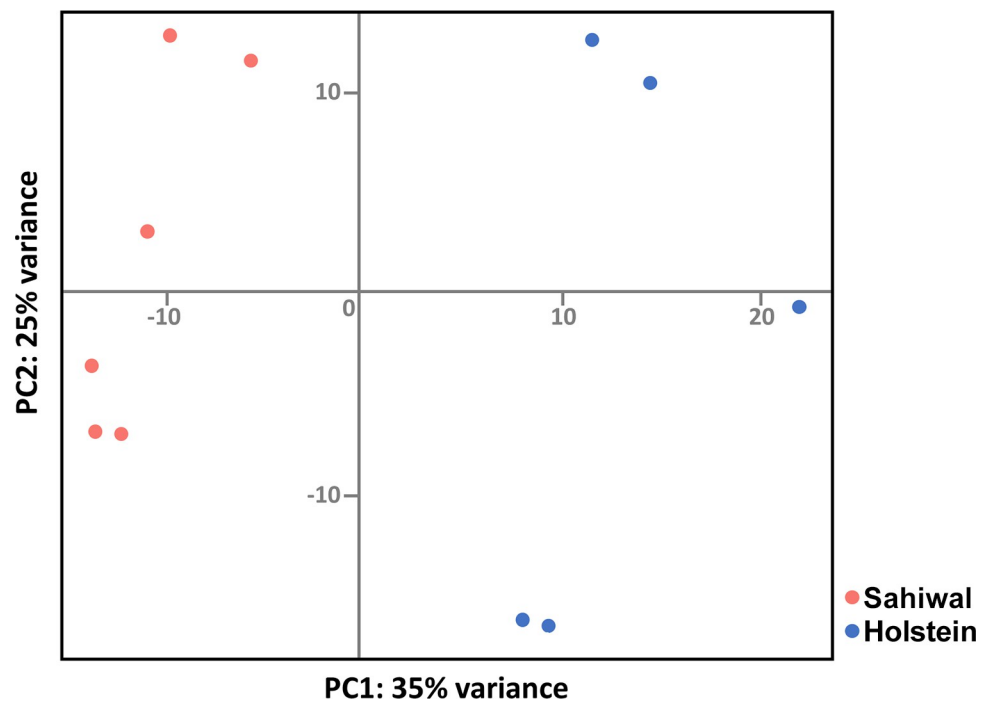


Fig 1. Clustering of samples by host breed. A PCA plot showing clustering based on the H/S-DE data set derived from five samples of Holstein (blue) or six samples of Sahiwal (red) cell lines infected with the *T. annulata* macroschizont.

<https://doi.org/10.1371/journal.pone.0262051.g001>

within each breed, that was more pronounced for the Holstein samples, was also detected (PC2 displaying 25% variance).

Indicative of the clear separation between the two breeds, a substantial proportion of genes in the data set showed a consistent difference between the breed types, with 439 of the 500 most statistically significant H/S-DE genes displaying differential expression across all Sahiwal vs. Holstein samples. Specific examples include the IFN stimulated gene (ISG) *MX1* expressed at lower levels in all Holstein samples relative to Sahiwal cells (-4.62 , \log_2); whereas the *FAT1* cadherin encoding gene, associated with TGF β signalling, was consistently expressed at a higher level for Holstein cell lines (7.49 , \log_2 ; [S1 File](#)).

Not unexpectedly, from the PCA, a sizeable number of genes indicated as differentially expressed between the breeds did not display a difference that was consistent across all six Sahiwal vs. five Holstein samples. A good example being the gene encoding pro-inflammatory cytokine IL-6, designated as significantly higher in Holstein (1.34 \log_2), where two Holstein samples showed lower expression values than the two highest values recorded for the Sahiwal cell lines ([S1 File](#)). Thus, genes of this type are indicated as generally differentially expressed between the two breeds but show a more substantial level of variable expression within a breed, and as reported previously [21] this appeared to be more marked for the Holstein sample set.

[Table 1](#) shows the top 19 genes displaying greatest higher or lower expression (fold change \log_2) in Holstein (*B. taurus*) relative to Sahiwal (*B. indicus*) infected leukocytes. A number of these genes possess gene ontology (GO) annotation indicating they could play a role in determining phenotypic differences between Holstein and Sahiwal macroschizont infected cells. Thus, for genes with higher relative levels of expression in Holstein infected cells: Slit Guidance Ligand 2 (*SLIT2*), functions in cancer and leukocyte chemotaxis/infiltration [45,46]; Ubiquitin D (*UBD/FAT10*), operates in protein modification/degradation and activation of pro-inflammatory mediators (reviewed in [47]); *GULP1*, is involved in phagocytosis [48] and associated with emphysema [49], a clinical symptom of acute theileriosis [50]. For the genes designated as expressed lower in Holstein infected cells relative to Sahiwal, those encoding proteins associated with a tumour suppressor/inhibitor of metastasis function (e.g. *ANXA8*; *PAX6*; *COL4A1* and *ADAMTS18*) or associated with the innate immune/ISG response were of particular interest. Of the 68 IFN type I response genes [51] identified in the H/S-DE data set, 56 displayed lower expression in Holstein derived infected cells relative to Sahiwal ([S1 File](#)).

IPA reveals breed-associated modulation of gene expression linked to innate immunity, cholesterol biosynthesis and oncogenesis

Differential susceptibility to theileriosis could arise via pre-existing breed associated differences in gene expression between infected (and uninfected) cells or only become manifest after infection of the bovine leukocyte. Therefore, we applied Ingenuity Pathway Analysis (IPA) on both the full data set of 2211 H/S-DE genes and a more limited data set of Infection Associated (IA) genes. To identify breed associated differences in gene expression that are induced by infection, we overlapped the H/S-DE data set with a set of bovine genes demonstrated to display altered expression in *T. annulata*, macroschizont infected *B. taurus* derived lymphosarcoma (TBL20) cells relative to uninfected (BL20) cells [14]. Of the 2211 H/S-DE genes, 517 overlapped and were indicated as the infection associated Holstein/Sahiwal data set (IA-H/S); see [S2 File](#) for the full data set. The representation factor for the 517 genes was 1.4 and the p value was $< 4.2 \times 10^{-15}$; indicating statistical significance for the obtained overlap ([Fig 2](#)). Strikingly the most significant canonical pathways identified by IPA for both the full ([S3 File](#)) and infection associated data set ([S4 File](#)) were virtually identical, with involvement in innate immunity, cholesterol biosynthesis and oncogenesis highlighted.

Table 1. Top 19 (by Log₂ fold change) down- and up-regulated genes in *B. taurus* vs. *B. indicus* *T. annulata* infected cell lines.

Gene	log ₂ FC	lfcSE	Padj	Gene symbol	Gene description
ENSBTAG00000021565	-26.131	3.038	<0.001	PRSS2	serine protease 2
ENSBTAG00000002335	-11.108	1.021	<0.001	HAND1	heart and neural crest derivatives expressed 1
ENSBTAG00000012849	-9.151	1.622	<0.001	COL4A1	collagen type IV alpha 1 chain
ENSBTAG00000009842	-8.637	1.417	<0.001	CRYM	crystallin mu
ENSBTAG00000031355	-8.014	0.780	<0.001	LOC529196	C-C chemokine receptor type 1-like
ENSBTAG00000046377	-7.793	0.992	<0.001	LOC100336807	
ENSBTAG0000001010	-7.450	1.462	<0.001	ADAMTS18	ADAM metalloproteinase with thrombospondin type 1 motif 18
ENSBTAG00000046412	-7.448	1.056	<0.001	LOC512617	acyl-CoA dehydrogenase family member 10
ENSBTAG00000004561	-6.888	1.114	<0.001	PAX6	paired box 6
ENSBTAG00000038064	-6.865	0.457	<0.001	LOC614531	ras-related GTP-binding protein A
ENSBTAG00000005857	-6.751	1.312	<0.001	SLC6A1	solute carrier family 6 member 1
ENSBTAG00000027782	-6.529	0.998	<0.001	OR5K1	olfactory receptor, family 5, subfamily K, member 1
ENSBTAG00000008129	-6.455	1.668	0.002	CLSTN3	calsyntenin 3
ENSBTAG00000047616	-6.311	1.719	0.005	ZNF114	zinc finger protein 114
ENSBTAG00000018499	-5.923	1.758	0.011	ANXA8L1	annexin A8-like 1
ENSBTAG00000035110	-5.918	2.049	0.038	DEUP1	deuterosome assembly protein 1
ENSBTAG00000019428	-5.765	1.473	0.002	CCR1	chemokine (C-C motif) receptor 1
ENSBTAG00000018367	-5.733	1.347	<0.001	CD6	CD6 molecule
ENSBTAG00000014628	-5.624	0.852	<0.001	OAS2	2'-5'-oligoadenylate synthetase 2
ENSBTAG00000018694	8.087	1.629	<0.001	ACSS3	acyl-CoA synthetase short chain family member 3
ENSBTAG00000044185	7.729	3.016	0.075	SOX6	SRY-box 6
ENSBTAG00000005108	7.513	1.576	<0.001	SLIT2	slit guidance ligand 2
ENSBTAG00000025803	7.506	1.809	<0.001	INSYN1	inhibitory synaptic factor 1
ENSBTAG00000020657	7.490	0.865	<0.001	FAT1	FAT atypical cadherin 1
ENSBTAG00000003345	7.149	1.769	0.001	FAT4	FAT atypical cadherin 4
ENSBTAG00000037649	7.111	1.212	<0.001	VIPR2	vasoactive intestinal peptide receptor 2
ENSBTAG00000015905	7.013	1.136	<0.001	ARHGAP32	Rho GTPase activating protein 32
ENSBTAG00000025398	6.927	1.707	0.001	LOC504548	ubiquitin D
ENSBTAG00000004510	6.777	1.310	<0.001	SARDH	sarcosine dehydrogenase
ENSBTAG00000047336	6.726	2.294	0.034	LOC100847115	thyrotropin-releasing hormone receptor-like
ENSBTAG00000031802	6.597	1.335	<0.001	SPATA16	spermatogenesis associated 16
ENSBTAG00000014354	6.519	1.246	<0.001	FXYD6	FXYD domain containing ion transport regulator 6
ENSBTAG00000019340	6.436	1.438	<0.001	PCDH9	protocadherin 9
ENSBTAG00000021526	6.296	1.994	0.020	RPRM	reprimin, TP53 dependent G2 arrest mediator homolog
ENSBTAG00000027246	6.247	1.388	<0.001	UBD	ubiquitin D
ENSBTAG00000020385	6.216	1.408	<0.001	ALDH1B1	aldehyde dehydrogenase 1 family member B1
ENSBTAG00000016525	6.179	1.898	0.015	ITGA1	integrin subunit alpha 1
ENSBTAG00000007141	6.159	1.438	<0.001	GULP1	GULP PTB domain containing engulfment adaptor 1

<https://doi.org/10.1371/journal.pone.0262051.t001>

Innate immunity pathways

Enriched pathways linked to innate immunity in the infection-associated data set were “Activation of Interferon by Cytosolic Pattern Recognition Receptors” and “Interferon signalling” and these pathways were also highly enriched by the analysis of the full H/S-DE data set. Other related significantly enriched pathways included “Role of RIG1 Like Receptors in Antiviral Innate Immunity” and, “NF-κB Activation by Viruses” (full data set only).

IPA indicates the activation state of canonical pathways via the calculated Z-score, a negative Z score predicting lower activation potential, while a positive score indicates higher

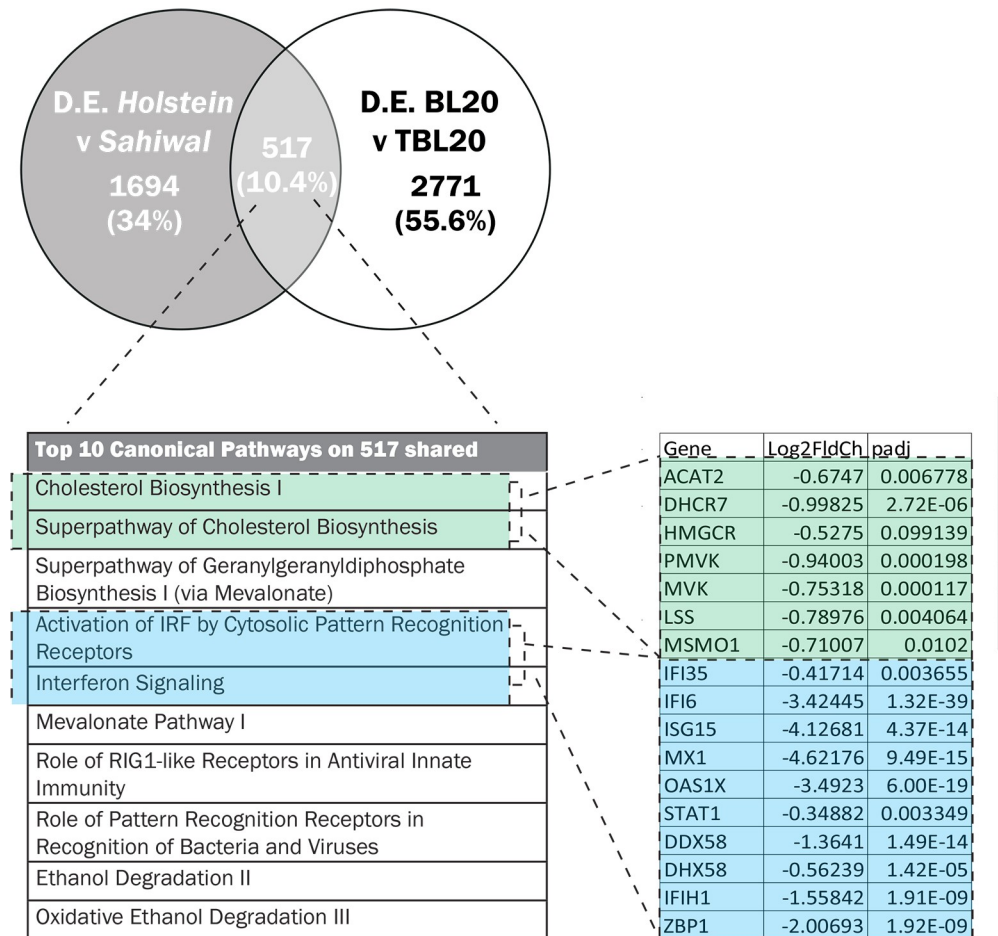


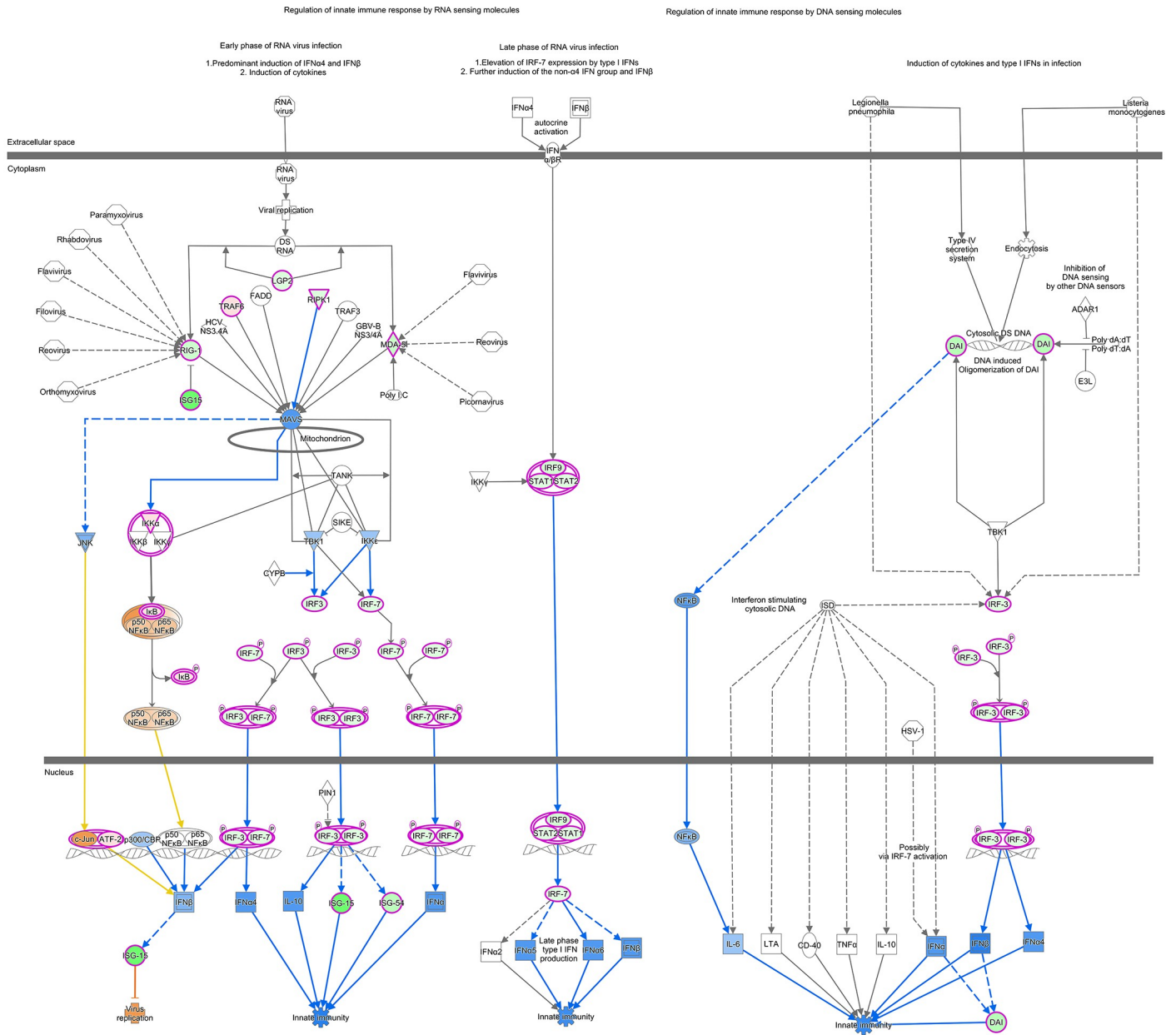
Fig 2. Overlap of genes modulated between Holstein and Sahiwal breeds of infected cell and *Theileria* infection of BL20 cells. (A) Venn diagram of overlapping genes, (B) the top IPA canonical pathways and (C) the modulation of infected-associated genes found in enriched pathways linked to cholesterol biosynthesis or the IFN response.

<https://doi.org/10.1371/journal.pone.0262051.g002>

potential. For pathways linked to innate immunity highlighted above, all except NF- κ B showed lower activation status in Holstein relative to Sahiwal infected cells (Z score range > -1.3 to -2.8). This finding is illustrated for “Activation of Interferon by Cytosolic Pattern Recognition Receptors” (Fig 3) and “Interferon Signalling Pathways” (Fig 4) generated by IPA for the H/S-DE data set. As would be expected, these two pathways overlap, with lower transcript levels in Holstein infected cells for STAT1, STAT2, IRF9 and the ISG targets of their transcription factor complex. Furthermore, all identified infection-associated ISG genes (*MX1*, *MX2*, *OAS1Y/Z*, *OAS2*, *ISG15*, *IFI6*, *IFI44* and *RSAD2*) showed lower expression values in Holstein infected cells relative to Sahiwal (Fig 2; S2 File).

Nodes, colours, shapes and lines in pathway are as described in legend to Fig 3.

IPA for disease or molecular function also predicted lower innate immune function for Holstein infected cells. As shown in Table 2, the three most significant disease or functions identified for the full H/S-DE data set that are predicted to be lower in Holstein compared to Sahiwal infected cells were “Antiviral response”; “Immune response of cells” and “Antimicrobial response”. Furthermore, the IPA function that predicts activation or inhibition of upstream regulators identified inhibition of cytokines, transcription factors and compounds

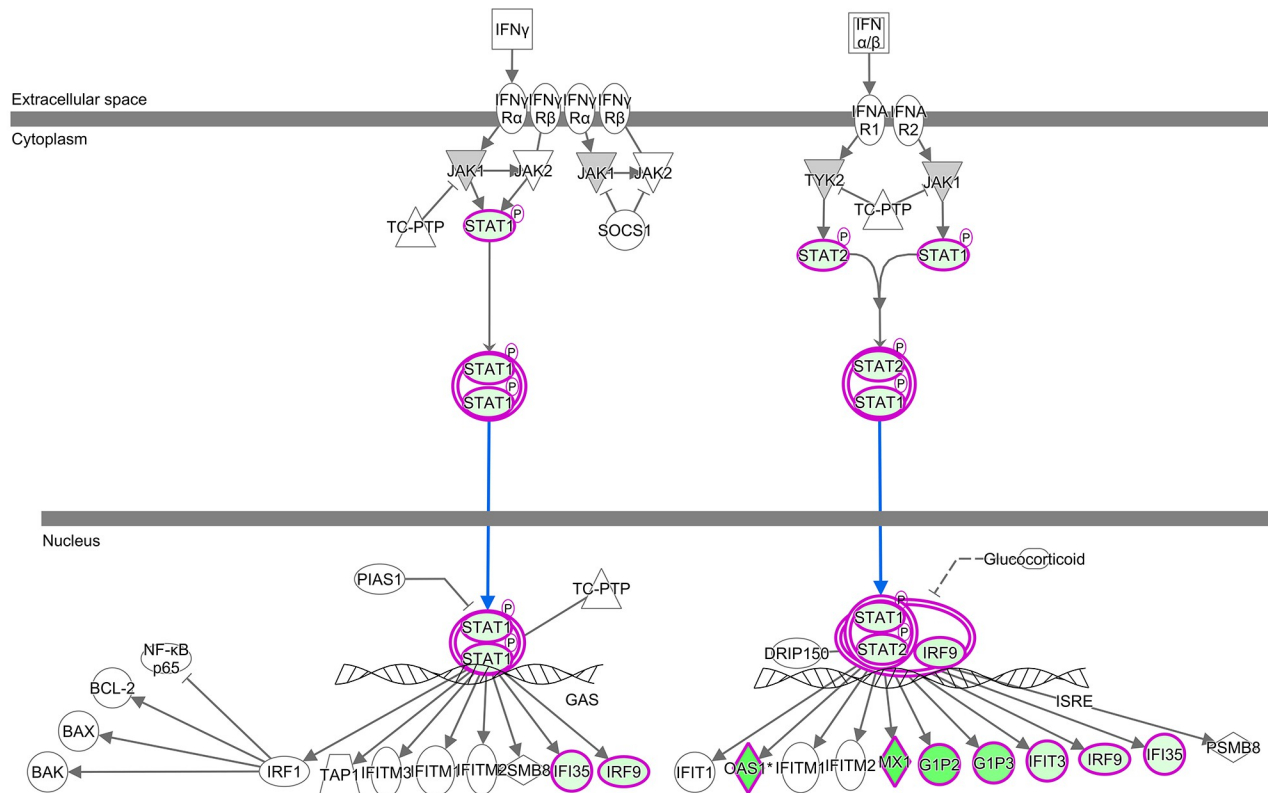


© 2000-2020 OMAGEN. All rights reserved.

Fig 3. IPA canonical pathway, “Activation of IRF by Cytosolic Pattern Recognition Receptors”. The nodes represent genes/molecules/complexes in a pathway, and the lines and arrows between nodes indicate known relationships from the Ingenuity Knowledge Base. Nodes with purple outline indicate molecules that were measured as differentially expressed in our dataset, with the intensity of coloured infill indicating the level of up (red) or down (green) regulation of Holstein relative to Sahiwal. The blue- and orange-coloured molecules and lines are predicted activation states generated by the Molecular Activity Predictor function in Ingenuity Pathway Analysis (IPA). Blue colour indicates a predicted inhibition, and orange a predicted activation state in Holstein relative to Sahiwal. Yellow lines indicate relationships where our findings are inconsistent with the state of the downstream molecule. Broad lines with explanatory text beside the pathway indicate the cellular location of molecules in the pathway. The molecules in the pathway are given shapes that indicate their functional class (Nested Circle/Square = Group/Complex, Horizontal ellipse = Transcriptional Regulator, Vertical Ellipse = transmembrane receptor, Vertical Rhombus = enzyme, Square = Cytokine/Growth Factor, Triangle = Kinase, Vertical Ellipse = Transmembrane Receptor, Circle = other). The edges between molecules are also differentiated to indicate the type of relationship between them. Solid lines are direct relationships and dashed lines are indirect.

<https://doi.org/10.1371/journal.pone.0262051.g003>

associated with the interferon response in Holstein relative to Sahiwal infected cells (negative Z scores >2). Very similar results were obtained for both the full and IA data sets (Table 3 & S3 Table).



© 2000-2020 QIAGEN. All rights reserved.

Fig 4. IPA canonical pathway, “Interferon signalling”.

<https://doi.org/10.1371/journal.pone.0262051.g004>

Cholesterol biosynthesis

The top three scoring IPA canonical pathways for the infection-associated data set of 517 genes (Fig 2, S4 File) were all linked to cholesterol biosynthesis, with negative Z scores or reduced relative expression values for pathway genes in Holstein infected cells.

Similar results were obtained using the full H/S-DE data set with “Superpathway of Cholesterol Biosynthesis” ranked as the second top canonical pathway and the highest negative Z score recorded (S3 File). Inhibition of the SREBF2 transcription factor, an important positive regulator of genes involved in cholesterol biosynthesis [52], was also predicted (Z score -3.48, P value 6.66 E-06), and relative *SREBF2* expression levels were lower for Holstein (*B. taurus*) infected cells (S1 File). Elevated expression of genes involved in cholesterol biosynthesis is associated with oncogenesis/metastasis via the PI3K/AKT pathway [53,54], which is activated in *Theileria* infected cells [55] and enriched by IPA in both the full H/S-DE and IA data sets (S3 and S4 Files). Therefore, prediction of elevated PI3K/AKT signalling together with lower expression of cholesterol biosynthesis pathway genes in Holstein infected cells was unexpected. However, in other studies decreased expression of cholesterol biosynthesis genes has been associated with activation of Ras/Erk signalling and a more malignant cancer cell phenotype [56].

Oncogenesis

IPA showed a clear association of the full Holstein vs. Sahiwal data set of DE genes with oncogenesis/neoplasia. Thus, the canonical pathways “Retinoic Acid Mediated Apoptosis

Table 2. IPA for disease pathways and functions enriched in data set of differentially expressed genes between Holstein and Sahiwal *T. annulata* infected cell lines.

Category	Diseases or Functions Annotation	p-value	Predicted Activation State	Activation z-score	# Molecules
Antimicrobial Response, Inflammatory Response	Antiviral response	<0.001	Decreased	-3.341	48
Inflammatory Response	Immune response of cells	<0.001	Decreased	-2.725	84
Antimicrobial Response, Inflammatory Response	Antimicrobial response	<0.001	Decreased	-3.107	56
Neurological Disease	Progressive neurological disorder	<0.001	Decreased	-2.157	133
Cancer, Organismal Injury and Abnormalities, Renal and Urological Disease	Urinary tract cancer	<0.001	Decreased	-2.236	240
Cancer, Organismal Injury and Abnormalities, Renal and Urological Disease	Renal cancer	<0.001	Decreased	-2.236	150
Cell Death and Survival	Cell death of lymphoma cell lines	<0.001	Decreased	-3.104	36
Lipid Metabolism, Small Molecule Biochemistry, Vitamin and Mineral Metabolism	Synthesis of cholesterol	<0.001	Decreased	-2	18
Cell Death and Survival	Apoptosis of lymphoma cell lines	<0.001	Decreased	-2.81	29
Cell Death and Survival	Cell death of macrophage cancer cell lines	<0.001	Decreased	-2.735	11
Cell Death and Survival	Cell death of connective tissue cells	<0.001	Decreased	-2.034	75
Cell-To-Cell Signalling and Interaction, Inflammatory Response	Response of phagocytes	0.002	Decreased	-2.498	32
Cell-To-Cell Signalling and Interaction, Embryonic Development	Response of embryonic cell lines	0.002	Decreased	-3.111	12
Cell-To-Cell Signalling and Interaction	Response of myeloid cells	0.003	Decreased	-2.466	31
Cell-To-Cell Signalling and Interaction, Inflammatory Response	Immune response of phagocytes	0.003	Decreased	-2.787	29
Embryonic Development, Organismal Development	Development of body trunk	<0.001	Increased	3.647	153
Infectious Diseases	Viral Infection	<0.001	Increased	3.15	220
Infectious Diseases	Infection by RNA virus	<0.001	Increased	3.075	114
Cancer, Organismal Injury and Abnormalities	Development of malignant tumour	<0.001	Increased	2.984	786
Infectious Diseases	Infection of Mammalia	0.001	Increased	2.978	42
Cellular Movement	Cell movement of melanoma cell lines	0.002	Increased	2.715	21
Infectious Diseases	Replication of Herpesviridae	<0.001	Increased	2.449	15
Cancer, Organismal Injury and Abnormalities	Incidence of tumour	<0.001	Increased	2.428	815
Infectious Diseases	Infection of cells	<0.001	Increased	2.331	97
Cancer, Organismal Injury and Abnormalities	Malignant solid tumour	<0.001	Increased	2.277	1292
Infectious Diseases	Replication of Murine herpesvirus 4	0.003	Increased	2.236	5
Cardiovascular System Development and Function, Embryonic Development, Organ Development, Organismal Development, Tissue Development	Cardiogenesis	<0.001	Increased	2.198	79
Cancer, Organismal Injury and Abnormalities, Reproductive System Disease	Tumorigenesis of reproductive tract	<0.001	Increased	2.19	459
Cancer, Organismal Injury and Abnormalities, Reproductive System Disease	Female genital neoplasm	<0.001	Increased	2.19	459
Cancer, Endocrine System Disorders, Organismal Injury and Abnormalities, Reproductive System Disease	Ovarian tumour	<0.001	Increased	2.19	205
Cancer, Endocrine System Disorders, Organismal Injury and Abnormalities	Endocrine gland tumour	<0.001	Increased	2.183	1024
Cancer, Gastrointestinal Disease, Organismal Injury and Abnormalities	Digestive organ tumour	<0.001	Increased	2.161	1130
Cancer, Organismal Injury and Abnormalities	Frequency of tumour	<0.001	Increased	2.152	801
Cancer, Haematological Disease, Organismal Injury and Abnormalities	Hematologic cancer	0.002	Increased	2.088	329
Cancer, Gastrointestinal Disease, Hepatic System Disease, Organismal Injury and Abnormalities	Liver tumour	<0.001	Increased	2.017	556

<https://doi.org/10.1371/journal.pone.0262051.t002>

Table 3. Top activated or repressed upstream regulators predicted by IPA from gene targets within the H/S-DE data set.

Upstream Regulator	Molecule Type	Predicted Activation State	Activation z-score	p-value of overlap	Genes in dataset (Number of regulators from data in network)
IFNA2	cytokine	Inhibited	-5.284	<0.001	108 (11)
IRF7	transcription regulator	Inhibited	-5.147	<0.001	137 (13)
Interferon alpha	group	Inhibited	-5.024	<0.001	180 (13)
IRF3	transcription regulator	Inhibited	-4.863	<0.001	120 (12)
PRL	cytokine	Inhibited	-4.788	<0.001	241 (10)
poly rI:rC-RNA	biologic drug	Inhibited	-4.586	<0.001	292 (13)
IFNL1	cytokine	Inhibited	-4.577	<0.001	374 (15)
IFN Beta	group	Inhibited	-4.506	<0.001	233 (11)
STAT1	transcription regulator	Inhibited	-4.412	<0.001	318 (12)
Ifnar	group	Inhibited	-4.245	<0.001	181 (14)
IFNG	cytokine	Inhibited	-4.078	<0.001	367 (14)
IRF1	transcription regulator	Inhibited	-3.994	<0.001	317 (13)
CpG ODN 2006	chemical reagent	Inhibited	-3.846	<0.001	244 (11)
IFNB1	cytokine	Inhibited	-3.806	<0.001	286 (13)
IRF5	transcription regulator	Inhibited	-3.763	<0.001	357 (17)
IFNA1/IFNA13	cytokine	Inhibited	-3.538	<0.001	103 (10)
TGM2	enzyme	Inhibited	-3.516	<0.001	
SREBF2	transcription regulator	Inhibited	-3.482	<0.001	250 (5)
TRIM24	transcription regulator	Activated	4.884	<0.001	68 (6)
MAPK1	kinase	Activated	4.26	<0.001	106 (8)
PNPT1	enzyme	Activated	4.123	<0.001	260 (7)
NKX2-3	transcription regulator	Activated	3.788	<0.001	
INSIG1	other	Activated	3.628	<0.001	190 (5)
ACKR2	G-protein coupled receptor	Activated	3.606	<0.001	233 (5)
PTGER4	G-protein coupled receptor	Activated	3.286	<0.001	363 (10)
MYC	transcription regulator	Activated	3.202	<0.001	231 (2)
SIRT1	transcription regulator	Activated	3.184	0.0582	
SOCS1	other	Activated	3.18	0.015	
MFSD2A	transporter	Activated	3.148	<0.001	
IL1RN	cytokine	Activated	3.048	<0.001	362 (11)
LEPR	transmembrane receptor	Activated	3	0.219	
Mek	group	Activated	2.992	0.475	
MMP3	peptidase	Activated	2.828	0.351	
IRF8	transcription regulator	Activated	2.819	0.016	
JQ1	chemical reagent	Activated	2.779	0.055	
USP18	peptidase	Activated	2.772	<0.001	72 (9)
POR	enzyme	Activated	2.729	<0.001	
IKZF3	transcription regulator	Activated	2.72	0.005	
HMGA1	transcription regulator	Activated	2.718	0.337	

<https://doi.org/10.1371/journal.pone.0262051.t003>

Signalling”, “Death Receptor Signalling” and “Induction of Apoptosis by HIV1” showed negative Z scores of -2.8, -2.3 and -1, respectively (S3 File): predicting lower capacity for apoptosis/cell death in Holstein infected cells. In contrast, “FAT10 (UBD) Cancer Signalling Pathway” (Z score +2.6), “Small Cell Lung Carcinoma” (Z score +1.89), “Non-Small Cell Lung Cancer Signaling” (Z score +2.36), “PI3K Signalling in B lymphocytes” (Z score + 0.8) and “Wnt/Ca

+ Pathway” (Z score +0.7) predicted elevated pathway activation for Holstein infected cells. “FAT10 cancer signalling” with the higher expression of *UBD (FAT10)* predicted to promote carcinogenesis via NF- κ B and TGFB is illustrated (S1 Fig). In addition, four of the top fifteen entries with a negative Z score in the disease processes or molecular function analysis (Table 2) were placed in “Cell Death and Survival”, while the majority (twelve) of the top twenty activated (positive Z score) disease processes/functions were in “Cancer” or “Cellular movement”.

Canonical pathways associated with oncogenesis displaying a positive Z score were not identified in analysis of the infection associated-H/S data set. However, infection associated genes known to function in oncogenesis were identified as displaying higher expression in Holstein infected cells compared to Sahiwal. These genes include *UBD (FAT10)* and *SLIT2* (see Table 1 and S1 Fig). *PRUNE2* (B-cell CLL/lymphoma 2 and adenovirus E1B 19 kDa interacting family) and *CLU*, encoding Clusterin a secreted chaperone, were also identified and have been shown to operate in cell death, cell transformation, tumour progression and neurodegenerative disorders [57–59].

Taken together the results of the IPA analysis indicate for Holstein infected cells relative to Sahiwal a higher oncogenic potential, but lower innate immunity and cholesterol biosynthesis potential. And for all three of these processes, a clear link to genes whose expression is altered by parasite infection of leukocytes was established

Differential expression of genes encoding predicted secreted/receptor proteins

In our dataset of H/S-DE genes, we found 83 predicted to encode secreted/extracellular proteins, with 28 of these identified as infection associated (S5 File). The most significantly elevated of these genes was *SLIT2*. Genes encoding *CLU*, *SPARC* and *PLAU*, were also higher more than 1fold, \log_2 in Holstein cells and modulated by parasite infection. *SPARC* (cysteine rich acidic matrix associated protein) promotes both metastasis [60] and degradation of p53 [61], which also occur in *Theileria* infected leukocytes [13,62]. *PLAU* encodes a secreted urokinase associated with cancer progression/metastasis via upregulation by the AT hook factor, *HMGA1* [63]. Infection associated genes encoding secreted proteins that showed lower relative expression > 1 fold, \log_2 in Holstein cells were *ADMATS18*, *C1QTNF6* (complement C1q tumour necrosis related protein: anti-inflammatory), *SPOCK2* (ERK signalling pathway, pro-cancer), *TNFSF10* (cytokine that induces apoptosis in transformed cells) and *CCL5* (chemokine involved in inflammation and immuno-regulation; chemo-attractant for blood monocytes and memory T helper cells). A number of non-infection associated genes that encode secreted proteins with potential to influence the phenotype of the infected cell and/or cells of the host immune system were also identified (e.g. *IL9*, *CXCL2*, *CLEC3B* and *ADAMTS19*). Therefore, differential stimulation of and interaction with cells of the haematopoietic system by factors secreted by Holstein relative to Sahiwal infected cells is likely.

Numerous genes encoding cellular receptors were also present in the H/S DE data set (see S6 File). Infection associated receptor encoding genes include *CCR1* (encoding a receptor for *CCL5*), *CRLF2* (cytokine receptor like factor 2), *IL10RA*, *TLR2*, *CD2*, *CD48* and *IL7R*. Non-infection associated genes encoding receptors included *ROBO2*; encoding the receptor for the secreted *SLIT2* factor (both receptor and factor expressed higher in infected Holstein cells), *TLR7*, *CD4*, *TRAF1* (TNF receptor associated factor 1), *IL27RA*, *IL17RD* and *IRAK1* (Interleukin-1 receptor-associated kinase 1). Therefore, Holstein infected cells are predicted to possess the potential to react differently than Sahiwal, in response to a large array of receptor ligands, including chemokines, cytokines and growth factors.

qRT-PCR validation of differential expression of candidate genes

To confirm the validity of the RNA-seq data, qRT-PCR analysis was performed. Seventeen genes were selected based on their representative RNA-seq profiles across the H/S-DE data set. The results are summarized in Table 4 and shown for a subset of the genes in Fig 5. *SLIT2* and *PRUNE2* represent infection-associated genes where expression was assessed as higher by RNA-seq in Holstein relative to Sahiwal infected cells. qRT-PCR confirmed this profile (Table 4), with statistically significant higher mRNA levels for both genes, with on average 62.1- and 171.6-fold higher values for *SLIT2* and *PRUNE2* from Holstein infected cell lines, respectively. *IFI44*, *ISG15*, *MX1*, *RSAD2*, *SAMD9* and *TNFSF10* were chosen as they are infection-associated genes that show lower expression levels in Holstein infected cells compared to Sahiwal cells. In all six cases the qRT-PCR supported this profile with a statistically significant lower mean expression level in Holstein cells of 8.9-fold (*IFI44*), 22.2-fold (*ISG15*), 17.5-fold (*MX1*), 6.1-fold (*RSAD2*), 5.9-fold (*SAMD9*) and 3.4-fold (*TNFSF10*), respectively (Table 4, Fig 5A & 5B). *ADAMTS20*, *NECTIN3* and *WLS* were selected to represent non-infection associated genes with higher expression in Holstein cells. The qRT-PCR analysis confirmed this profile with statistically significant higher relative expression of 2.0 fold (*ADAMTS20*), 9.0-fold (*NECTIN3*) and 5.4-fold (*WLS*) (Table 4, Fig 5C & 5D) in Holstein cells compared to Sahiwal cells. The reciprocal profile was also validated with *HERC5*, *IFI6*, *CGAS* (*MB21D1*) and *OAS1Y*. Statistically significant lower expression of 3.3 -fold (*HERC5*), 6.0-fold (*IFI6*), 2.2-fold (*CGAS*) and 10.5-fold (*OAS1Y*) was found in Holstein infected cells compared to Sahiwal cells (Table 4). Thus, for the majority of genes in the RNA-seq data set, the trend of expression predicted is likely to be accurate. Two further genes encoding the pro-inflammatory cytokines, IL6 and IL23A, designated as significantly higher in the H/S DE data set but which showed non-consistent differential expression across the eleven samples, were tested. The results indicated that although the mean expression values were 1.6-fold (*IL6*) and 3.6-fold (*IL23A*) higher

Table 4. qRT-PCR validation of differential expression of a subset of H/S-DE genes.

Gene	Gene Symbol	Sahiwal	Holstein	P value
ADAM Metallopeptidase With Thrombospondin Type 1 Motif 20	ADAMTS20	1.3 ± 0.1	2.6 ± 0.4	0.011
Cyclic GMP-AMP Synthase	CGAS	2.8 ± 0.2	1.2 ± 0.1	<0.001
HECT And RLD Domain Containing E3 Ubiquitin Protein Ligase 5	HERC5	3.7 ± 0.6	1.1 ± 0.1	<0.001
Interferon Alpha	IFNA	1.7 ± 0.3	2.1 ± 0.4	n.s.
Interferon Alpha Inducible Protein 6	IFI6	11.8 ± 1.1	2.0 ± 0.4	0.001
Interferon Beta 1	IFNB1	2.3 ± 0.4	5.7 ± 1.1	0.006
Interferon Beta 3	IFNB3	5.1 ± 1.4	2.9 ± 0.7	n.s.
Interferon Induced Protein 44	IFI44	50.0 ± 8.4	5.6 ± 1.4	0.002
Interleukin 6	IL6	2.8 ± 0.6	4.5 ± 1.8	n.s.
Interleukin 23 Subunit Alpha	IL23A	1.6 ± 0.2	5.8 ± 3.8	n.s.
ISG15 Ubiquitin Like Modifier	ISG15	110.8 ± 26.3	5.0 ± 1.8	0.001
MX Dynamin Like GTPase 1	MX1	191.2 ± 14.4	10.9 ± 8.1	0.005
Nectin Cell Adhesion Molecule 3	NECTIN3	5.4 ± 2.0	48.3 ± 13.8	0.018
2'-5'-Oligoadenylate Synthetase 1	OAS1	231.0 ± 23.5	22.0 ± 17.9	0.001
Prune Homolog 2 With BCH Domain	PRUNE2	2.0 ± 0.6	335.3 ± 172.9	0.018
Radical S-Adenosyl Methionine Domain Containing 2	RSAD2	153.4 ± 35.1	25.1 ± 19.2	0.013
Sterile Alpha Motif Domain Containing 9	SAMD9	8.5 ± 1.5	1.4 ± 0.1	0.001
Slit Guidance Ligand 2	SLIT2	2.1 ± 0.5	127.4 ± 65.0	0.007
TNF Superfamily Member 10	TNFSF10	12.5 ± 1.6	3.7 ± 0.9	0.009
Wnt Ligand Secretion Mediator	WLS	1.2 ± 0.1	6.7 ± 1.7	0.008

<https://doi.org/10.1371/journal.pone.0262051.t004>

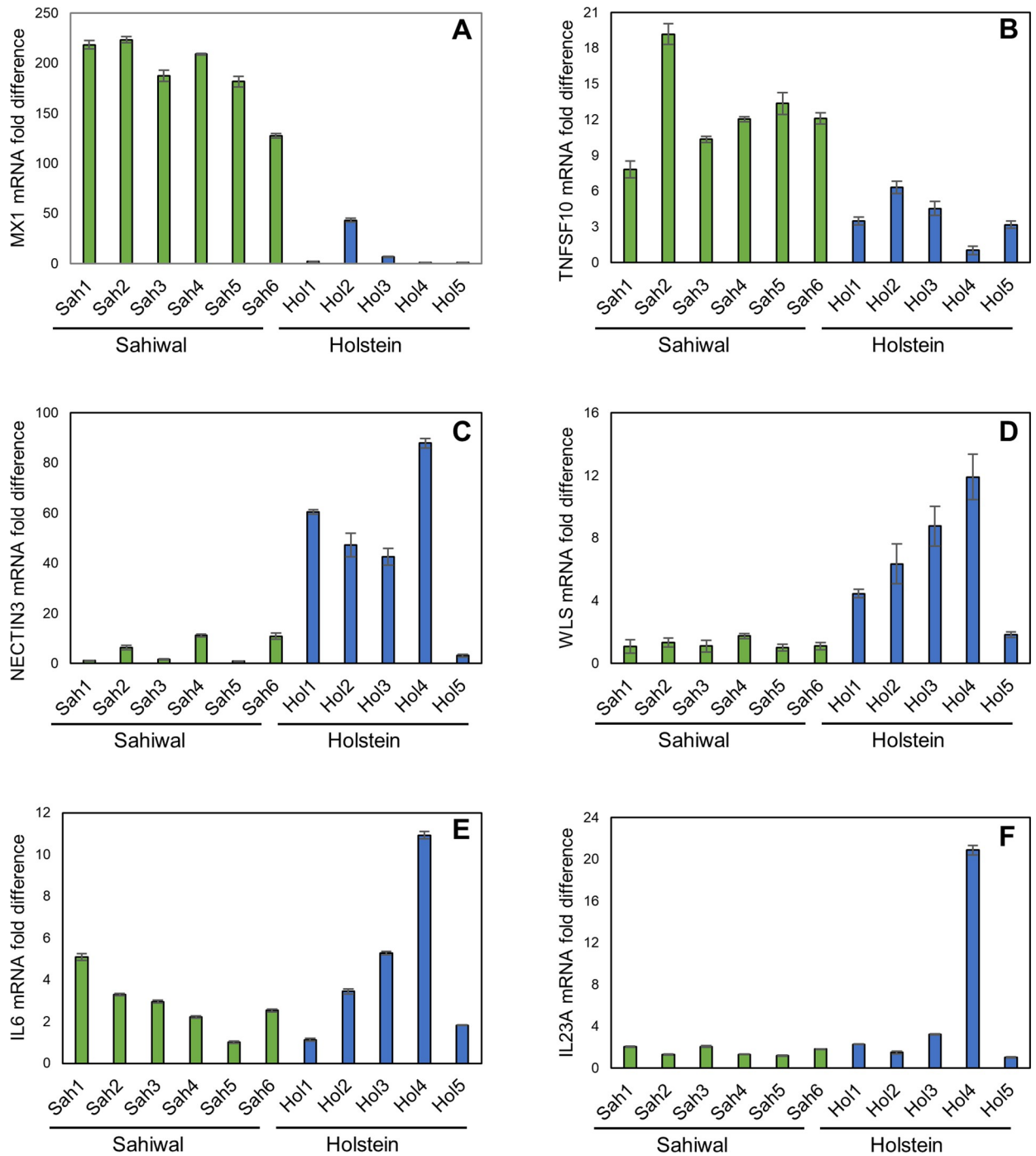


Fig 5. qRT-PCR validation of candidate H/S-DE genes. Panels show results for representative genes. (A) *MX1*; (B) *TNFSF10*; (C) *NECTIN3*; (D) *WLS*; (E) *IL6*; (F) *IL23A*. Within each panel Y axis shows mRNA fold difference (calculated vs. the lowest expressed sample); X axis designates sample number and breed; Hol1-5, Holstein; Sah1-6, Sahiwal.

<https://doi.org/10.1371/journal.pone.0262051.g005>

for Holstein infected cells relative to Sahiwal, variability within the data precluded validation of a significant difference (Table 4, Fig 5E & 5F).

IPA analysis of our RNA-seq datasets predicted marked differences between Holstein and Sahiwal infected cells in pathways linked to activation of and response to type I IFN. However,

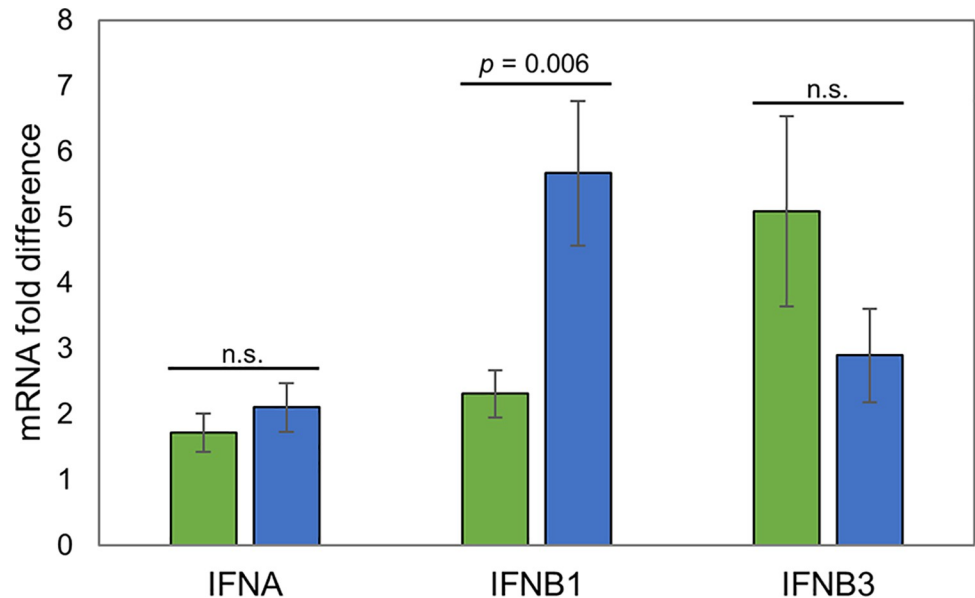


Fig 6. qRT-PCR analysis of INF genes: *IFNA*, *IFNB1* and *IFNB3*. The Y-axis shows mRNA fold difference (calculated vs. the lowest expressed sample): Green infill designates, Sahiwal; blue infill, Holstein. Degree of significance denoted above bar.

<https://doi.org/10.1371/journal.pone.0262051.g006>

no type I IFN genes were identified as differentially expressed in the H/S-DE dataset. In an attempt to elucidate whether a type I IFN could be associated with differential expression of ISGs, the expression of *IFNB1*, *IFNB3* and *IFNA* was investigated by qRT-PCR. No significant difference in *IFNA* and *IFNB3* mRNA levels were detected between Holstein and Sahiwal derived infected cells. However, there was a significant difference in *IFNB1* expression. Unexpectedly, on average 2.5-fold higher expression of *IFNB1* was detected in Holstein infected cells compared to Sahiwal cells (Table 4, Fig 6).

Screen for differential expression of parasite transformation candidates and motifs bound by TashAT2 in *B. indicus* vs. *B. taurus* genomes

In total, 109 *T. annulata* encoded genes were identified as significantly differentially expressed between Holstein and Sahiwal infected cell lines (S7 File). Compared to host genes, however, the parasite genes showed markedly smaller magnitudes of change in expression level; a maximum \log_2 fold change of 2.6 compared to -26.1 for the bovine gene set. A large number of the parasite genes were annotated as hypothetical or encode proteins with a predicted function/location that cannot be linked to transformation of the infected cell or host immune response. Moreover, there was limited evidence for altered expression of genes encoding candidate modulators of host cell phenotype (TaPIN, Ta9 and TashATs). Thus, gene *TA15705*, encoding the Ta9 immuno-dominant antigen [17], was the only one identified as differentially expressed (0.84 \log_2 fold higher in Holstein, third most significant). Further analysis of Ta9 expression was performed by immunoblot. The result failed to indicate significant elevation in Ta9 protein levels across three of the Holstein lines relative to three Sahiwal (S2 Fig). A similar result was obtained for TashAT2 translocated to the host nucleus of the infected cell [44]. Immunofluorescence failed to detect a consistent difference in reactivity against the infected cell nucleus between all three Holstein lines relative to Sahiwal (S3 Fig).

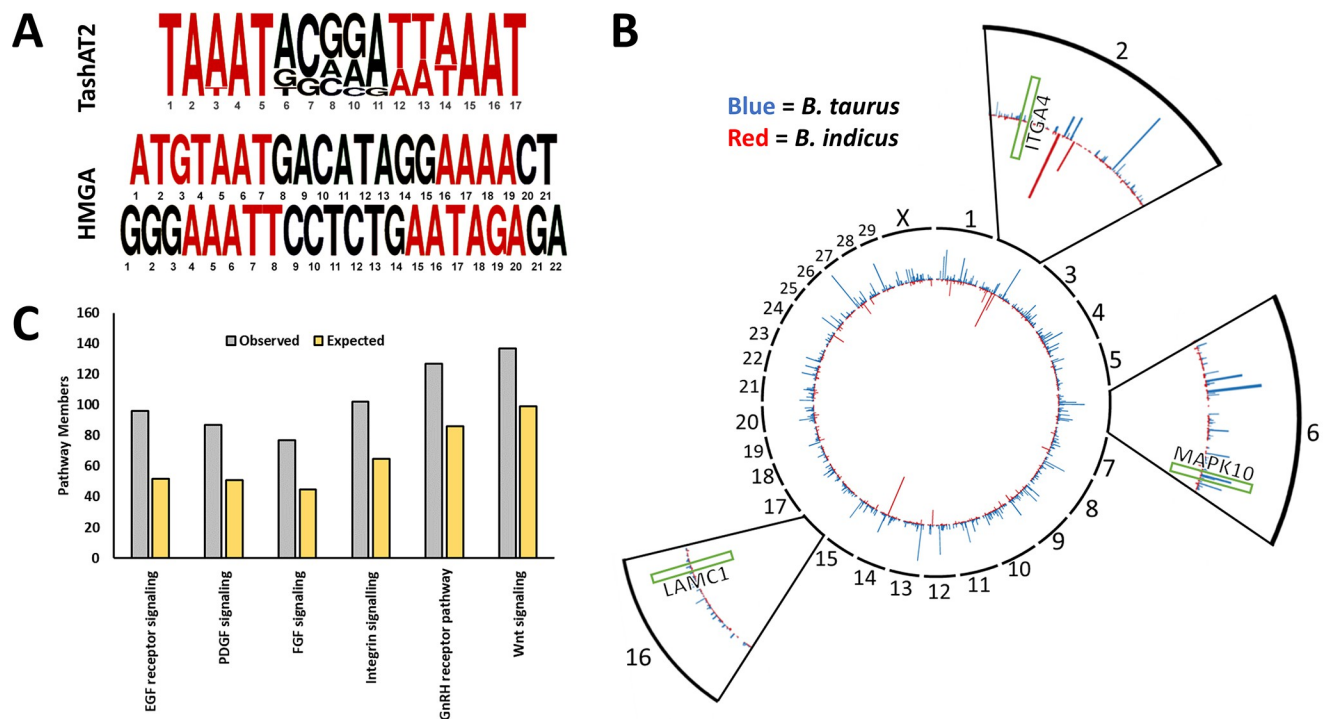


Fig 7. Analysis of DNA motifs bound by the parasite encoded TashAT2 factor in the genomes of *B. indicus* vs. *B. taurus*. (A) Consensus motif of the DNA region bound by parasite TashAT2 in the genome of *B. indicus* vs. *B. taurus* compared to the binding motifs of mammalian HMGA [19]. AT rich regions bound by AT hooks found in both TashAT2 and HMGA are shown in red. (B) A CIRCOS summary plot with automatic binning showing the number of TashAT2 binding motifs found in genes across the *B. indicus* (red) and *B. taurus* (blue) genomes. Genes with the same number of motifs are excluded to demonstrate the pattern of differences between each breed. Zoomed views are shown for chromosomes 2, 8 and 16 without automatic binning. The approximate regions of genes in the “EGF” and “Integrin signalling” pathways that possess different numbers of TashAT2 binding motifs and differ in expression between infected Holstein and Sahiwal breeds are indicated in green (*ITGA4*, *MAPK10* and *LAMC1*). (C) Overrepresented PANTHER pathways in the cohort of bovine genes with FDR adjusted p value < 0.05 after a Fisher’s Exact test. Blue is the observed members of a pathway in the dataset, whereas red indicates the expected number if the distribution was random.

<https://doi.org/10.1371/journal.pone.0262051.g007>

Polymorphism of a secreted parasite factor, or diversity of the host target it interacts with, could differentially modulate host cell gene expression in the absence of differential expression, and this model has been predicted for TashAT proteins. Thus, for TashAT2/3, polymorphism has been identified in the number and pattern of their AT hook DNA binding domains [64]. The AT hook domain of TashAT2/3 shows similarity to that of mammalian HMGA proteins, suggesting potential overlap in target DNA motifs, and it is known that the TashAT2 domain recognises motifs rich in AT (Fig 7A) [19]. Our IPA upstream analysis predicted activation of HMGA1 (Table 3), and HMGA1 has been shown to lower expression of cholesterol biosynthesis genes [55], one of the most enriched pathways in our data sets. Therefore, we assessed whether the pattern of motifs bound by TashAT2 show diversity in the genomes of *B. indicus* relative to *B. taurus* and whether they might be enriched in our IA-H/S data set. Our initial screen identified 167,151 incidences of the target motifs in the *B. taurus* and 149,107 in the *B. indicus* genome (S8 File). The majority of the motif patterns in both genomes were located outside of gene coding regions ($\approx 56\%$ and $\approx 65\%$ in *B. taurus* and *B. indicus* respectively), while $\approx 42\%$ were within introns in *B. taurus* and $\approx 35\%$ in *B. indicus*. Approximately 2% of patterns were in upstream or downstream UTRs and less than 1% were within exons in each breed (S8 File). As motif sites located within gene coding regions are more likely to be cis-acting elements and their associated targets more easily inferred, we used the cohort of motifs located in

UTRs, introns and exons for a more detailed comparison (S9 File). In *B. taurus* 11,983 genes featured at least one TashAT2 binding motif compared to 9,516 in *B. indicus* and there were substantial differences in the pattern and number of TashAT2 binding motifs between breeds (Fig 7B; S9 File). There were 7,575 shared genes that possessed at least one TashAT2 binding motif and it was evident that for certain genes (including *SLIT2*, *PRUNE2*, *LITAF* and *NECTIN3*) the number of motifs differed between *B. indicus* and *B. taurus* genomes. Subsequent PANTHER analysis of the shared genes data set showed an enrichment in genes involved in “EGF (PI3K/AKT, ERK) receptor signalling”, “PGDF signalling”, “Wnt signalling pathway”, “FGF signalling”, “Gonadotropin-releasing hormone (GnRH)” and “Integrin signalling” (FDR adjusted $p < 0.05$; Fig 7C). Comparison with IPA on the H/S-DE gene set showed overlap with “ERK/MAPK signalling”, “PI3K/AKT signalling”, “Wnt/Ca⁺ pathway and “Integrin signalling” (S3 File). Additionally, of the 517 transcripts previously noted to be altered by parasite infection of BL20 cells and differ in expression between infected Holstein and Sahiwal cells, 232 (45%) feature TashAT2 binding motifs. This is significantly more than would be expected by chance under a binomial distribution in which $\approx 26\%$ of annotated features in the bovine genome possess a binding motif ($p < 0.001$). *SLIT2*, *LITAF*, *PRUNE2* and genes involved in “Integrin signalling” showed altered numbers of TashAT2 motifs (S10 File) and, together with “PI3K/AKT signalling”, integrin signalling genes were significantly enriched by IPA in the Infection Associated-H/S data set (S4 File).

Discussion

Understanding how differential susceptibility to infectious disease is conferred is of great importance. Such understanding is needed to highlight at risk groups, inform therapeutic regimes and develop strategies for breeding productive but infection tolerant livestock. To gain insight on how variable infection responses are generated, we have utilised two cattle breeds representing the ends of a susceptibility spectrum to tropical theileriosis, caused by *Theileria annulata*. The Sahiwal breed (*B. indicus*) has evolved in tandem with the pathogen and in general, displays mild symptoms of disease when infected, whereas Holstein (*B. taurus*) are generally susceptible to acute disease. Based on the premise that tolerance has most likely evolved via competitive molecular interactions between pathogen and host that lead to pathology, we focused on identifying gene expression differences between low passage *T. annulata* macroschizont infected cell lines from tolerant Sahiwals and susceptible Holsteins, and screened for parasite factors that could generate these differences.

RNA-seq analysis identified a large number of genes displaying a significant difference in expression level between Holstein and Sahiwal infected cells (the H/S-DE data set). In order to replicate, as closely as possible, the situation *in vivo* and to prevent loss of virulence through cloning, this data set was generated from uncloned, low passage infected cell lines derived from 11 different animals. PCA analysis of the data set showed that although the two breed types clearly separated there was an inherent level of variability for samples within a breed. The result was not unexpected. It has been shown that gene expression levels, including cytokine encoding genes, vary within breed for both Holstein and Sahiwal infected cell lines [13,21,42], and gene expression markers associated with virulence show appreciable variability [65]. Such within breed variability could be generated by: a) genetic differences between individual cattle, with variance likely to be higher for the Holstein (genetically unrelated) relative to the Sahiwal cattle (full and 1/2 siblings) [9]; b) genetic differences in parasite genotype represented by infected cell lines, because although all cell lines are infected by Hissar, this strain contains multiple parasite genotypes [65]; c) epigenetic differences generated during establishment of cell lines *in vivo* or *in vitro* and d) variable host cell type composition between infected

cell lines. We propose that the gene expression profile of infected cells may be unique for each individual animal and that this influences the wide variability in infection outcome between and within “tolerant” and “susceptible” breeds [9,66]. To validate this premise, analysis of multiple pairs of infected and uninfected cells from a range of “tolerant” and “susceptible” breeds is required.

Despite the expectation of variable expression profiles, a large number of genes in our H/S-DE data set displayed expression differences that were consistent across all Sahiwal vs. Holstein samples (see denoted genes, [S2 File](#)); while the remainder, as indicated above, showed a higher level of variability between individuals across the breeds (for example see *IL-6*). The results support the conclusion that there is a major influence of breed type on the phenotype of the macroschizont-infected leukocyte linked to disease susceptibility [2,5,13].

IPA was performed on both the full H/S-DE data set and the more limited set (IA-H/S) of genes whose expression is induced by parasite infection of the host leukocyte. Strikingly the most significantly enriched pathways for both data sets were the same. However, previous studies indicate that a number of infection-associated genes differentially expressed between breeds have been missed by our analysis, and there are at least several candidates. Thus, the *TGFB2* gene, that is induced by infection and expressed at a higher level in Holstein infected cells [13], was present in our H/S-DE data set (elevated 3.5 log₂ fold) but was not identified as infection associated. *ICAM1* was reported as differentially expressed between Sahiwal and Holstein infected cells [23] but was not highlighted as significantly different in our study. The reasons for these false negatives are not conclusively known, but most likely relate to differences in bovine cells used to establish changes linked to infection and methodology for identification of differential gene expression.

The pathway showing the most consistent and significant modulation across our data sets was type 1 IFN. Modulation was most evident as lower relative expression of ISG in Holstein infected cells (82% of identified ISGs) but was also recorded for genes encoding pattern recognition receptors (DDX58 [RIG-1], ZBP1 [DAI], IFIH1 [MDA-5] and CGAS) and transcription factors (STAT1, STAT2, IRF3, IRF7 and IRF9). Notably, for many ISG genes differential expression was consistent across all Holstein vs. Sahiwal samples. Validation of gene expression differences was obtained by qRT-PCR for several ISG genes (*MX1*, *OAS1Y*, *ISG15*, *RSAD2* [Viperin], *IFI6*, *IFI44* and *HERC5*), and a number of ISG genes (*HERC6*, *ISG15*, *IFI6*, *IFI44*, *IFIH1*, *IFI35*, *MX1*, *MX2*, *OAS2* and *ZBP1*) were identified as modulated by parasite infection, supporting the previous report on *ISG15* [20]. The function of ISG genes in innate immunity has been studied predominantly using viral systems (reviewed in [67,68]). Limited information regarding the function of ISGs against intracellular protozoa is available. Nevertheless, it is reasonable to propose that, by acting as tumour suppressors or regulators of immune effector cells, ISGs could modulate the phenotype of the *Theileria*-infected leukocyte and influence infection outcome [69–74]. Modulation of ISG expression between cattle breeds could also be of relevance to other pathogens. An ISG response is generated in infected cells against Foot and Mouth Disease Virus [75] and *B. indicus* cattle are known to be less susceptible to FMDV infection than *B. taurus* [76].

While IPA predicted modulation of type I IFN production associated with tolerance, there was no direct evidence for this in the RNA-seq data set. This may have been due to difficulty in mapping RNA-seq reads to specific IFN genes within the large type I IFN family [77]. However, contrary to the IPA network prediction, qRT-PCR indicated significantly higher levels of *IFNB1* expression in the Holstein cell lines, supporting previous data demonstrating expression of *IFNB* in infected leukocytes [78]. How differential expression of *IFNB1* is generated between the infected cell lines is currently unclear but could occur via lower expression of genes involved in cholesterol biosynthesis recorded for Holstein infected cells (see [Fig 5](#)). A

reduction in cholesterol biosynthesis has been linked to induction of *IFNB* expression and inflammatory disease [52,79] and is implicated in the induction of an inflammatory cytokine storm in COVID-19 [80]. It appears, therefore, that tolerance to tropical theileriosis is associated with a higher relative ISG expression in Sahiwal infected cells; while disease susceptibility of Holstein may be linked to elevated levels of *IFNB1* cytokine production and a stronger propensity to stimulate an inflammatory response. Whether these events are associated with the massive pulmonary oedema following metastasis of infected cells to the lungs [12] requires investigation. Parallels with mechanisms conferring susceptibility to other infectious diseases are of interest.

IPA also identified several processes linked to oncogenesis. This included a prediction of greater metastasis potential for Holstein infected cells (via, for example elevated *FAT10* [*UBD*], *TGFB2*, *ITGA1*, *PLAU* and *SPARC* gene expression), supporting and validating the previous work demonstrating a role for *TGFB2* in stimulating elevated metastasis of Holstein relative to Sahiwal infected cell lines [13]. Previous KEGG pathway analysis of microarray data led to postulation that disease susceptibility is linked to an aberrant interaction with cells of the immune response that promotes immune dysfunction [23]. Major differences in expression of genes associated with immune cell interaction (such as receptors, chemokines and cytokines), highlighted in our current analysis, supports this premise. Thus, reduced oncogenic potential and lower aberrant immune cell activation could account for the less pronounced lymph node enlargement observed in infected Sahiwal relative to Holstein cattle [9].

A major aim of this study was to investigate potential pathogen-host interactions that generate gene expression differences between infected cells from tolerant vs. susceptible breeds. One potential interaction previously considered is that the type of cell preferentially infected could differ between *B. indicus* and *B. taurus* breeds. Analysis with antibodies against cell type markers indicated that myeloid cells were the predominant infected cell type infected for both breeds [81], although evidence was presented for some macroschizont infected NK cells in Sahiwal lines. Our RNA-seq data support these findings. For genes encoding the myeloid lineage markers, *CCR2*, *CCR3*, *CD13*, *CD11*, *CD14*, *CD34*, *CD36*, *CD9* and *CD38*, none were identified as differentially expressed. For genes encoding the *CD8* and *CD2* markers previously identified on some Sahiwal infected cells [81], *CD8* was not identified as significant in our data set. *CD2* showed higher expression levels associated with Sahiwal line, although in three Holstein lines the level of expression was close to that of one of the Sahiwal lines. In contrast, the gene encoding the *CD4* marker found on T cells and myeloid cells showed significantly lower expression values in all Holstein cell lines ($-4.6 \log_2$). The results imply that either the profile of cell types infected can vary to a degree between *B. indicus* and *B. taurus*, or following transformation of predominantly myeloid cells, the expression of cell surface molecules is differentially modulated. Previous analysis of infected, purified myeloid cells [23] also highlighted lower expression of *CD4* in Holstein relative to Sahiwal cells, and together with alteration of surface marker expression following infection of the BL20 line [14,15], indicates the latter possibility as most likely.

Interactions that modulate gene expression between Sahiwal and Holstein infected cells are likely to involve pathogen modulators of host cell gene expression. No conclusive evidence of this for known parasite candidates was detected. While the gene encoding *Ta9* reported to activate *AP1* [17] was scored as significantly expressed at a higher level ($0.84 \log_2$ fold) in Holstein infected leukocytes, this was not validated at the protein level by immunoblot. IPA, however, did identify *HMG1A* as a potential regulator of differential gene expression between the two breeds. Mammalian *HMGAs* act as architectural transcription factors and bind patterns of AT rich motifs to alter chromatin structure (both locally and globally) [82]. *HMG1A* factors regulate gene expression primarily during development [83], but also in neoplasia [84,85] and

genes targeted by NF- κ B [86]. Given that TashAT2 is known to encode a DNA binding domain with homology to HMGAs, binds a related AT rich motif pattern (Fig 7A) and was detected in the host nucleus of both Sahiwal and Holstein infected cells (S3 Fig), we screened the available *B. indicus* and *B. taurus* genomes for the presence of motifs bound by TashAT2. The results clearly demonstrated a difference in the predicted pattern of these motifs (Fig 7B). Thus, genes in both genomes uniquely display a predicted TashAT2 bound motif, while for genes that shared possession of motifs, differences in motif number were frequently detected. Within this cohort of genes there are many that could influence the infected cell phenotype. For example, there is enrichment for genes that function in “Epidermal Growth Factor signalling” and overlap with pathways highlighted by IPA of the H/S-DE data set (“ERK”, “PI3K/AKT”, “Integrin signalling”). PI3K/AKT signalling has previously been linked to mechanisms involved in transformation of the *Theileria*-infected leukocyte [55,87] and is modulated by mammalian HMGA2 proteins [84]. Integrin signalling is well established in oncogenesis and cellular interaction [88,89], and integrin genes are targets for modulation by HMGA [85]. It was also demonstrated that 45% of infection-associated genes differentially expressed between Sahiwal and Holstein infected cell lines feature at least one putative TashAT2 binding motif, which is more than expected by chance ($p < 0.001$). “PI3K/AKT” and “Integrin signalling” were significant in IPA of the IA-H/S data set (S4 File) and several infection-associated genes involved in integrin signalling, possess differences in the number of TashAT2 binding motifs between the two genomes (LAMC1, ACTA2, ITGA4 and ITGB5). The majority of integrin genes are expressed at a lower level in Sahiwal infected cells relative to Holstein. Reduced expression of ITGA4 via PI3K signalling has been linked previously to a loss of infected cell virulence [87].

Based on the above results, and on the premise that AT hook DNA binding proteins act as base composition readers to mould the structure of the epigenome [90], we propose the following model. Differences in the pattern of AT rich DNA motifs between the genomes of tolerant vs. susceptible cattle breeds (and individuals), when bound by pathogen TashAT factors located in the host cell nucleus, give rise to alternative chromatin architectures. These architectures then allow variable accessibility of host transcription factors activated by infection, resulting in differential modulation of gene expression in the macroschizont infected cell. Given the known association between the infected cell and pathology, and the polymorphism identified in the TashAT2/3 DNA binding domain [64], this model could explain the considerable variability in disease severity between and within breeds that occur upon infection [9,66]. Validation of differential TashAT2 factor binding to sites in host DNA by chromatin immunoprecipitation, and further investigation of divergent pathogen-host interactions that have evolved to modulate host cell gene expression will provide understanding of how differential susceptibility to disease has arisen, and inform strategies aiming to breed productive livestock tolerant to infection.

Supporting information

S1 Fig. IPA canonical pathway, “FAT10 cancer signalling”. The nodes represent genes/molecules/complexes in a pathway, and the lines and arrows between nodes indicate known relationships from the Ingenuity Knowledge Base. Nodes with purple outline indicate molecules that were measured as differentially expressed in our dataset, with the intensity of coloured infill indicating the level of up- (red) or down- (green) regulation of Holstein relative to Sahiwal. The blue and orange coloured molecules and lines are predicted activation states generated by the Molecular Activity Predictor function in Ingenuity Pathway Analysis. Blue colour indicates a predicted inhibition, and orange a predicted activation state in Holstein relative to

Sahiwal. Broad lines with explanatory text beside the pathway indicate the cellular location of molecules in the pathway. The molecules in the pathway are given shapes that indicate their functional class (Nested Circle/Square = Group/Complex, Horizontal ellipse = Transcriptional Regulator, Vertical Ellipse = transmembrane receptor, Vertical Rhombus = enzyme, Square = Cytokine/Growth Factor, Triangle = Kinase, Vertical Ellipse = Transmembrane Receptor, Circle = other). The edges between molecules are also differentiated to indicate the type of relationship between them. Solid lines are direct relationships and dashed lines are indirect.

(TIF)

S2 Fig. Immunoblot carried out with protein extracts prepared from 3 independent Sahiwal lines compared to 3 Holstein lines. Sahiwal samples are denoted SA, SB, SC; Holstein denoted HA, HB, HC. Protein size markers are indicated on the right (kDa). **A.** Extracts probed with Rat anti-Ta9 (*TA15705*) at 1/1200 dilution. Ta9 was detected at a variable size of 43-46kDa, which is similar to that described previously for the polymorphic Ta9 antigen [17,43]. **B.** As a control, the same blot was reprobbed with Rabbit antiserum raised against constitutively expressed ER HSP90 (*TA06470*) at 1/1500 dilution. The Ta9 and HSP90 reactive proteins are denoted by arrow.

(TIF)

S3 Fig. Immunofluorescence assay carried out on cells prepared from 3 independent Sahiwal lines compared to 3 Holstein lines. Sahiwal samples are denoted SA, SB, SC; Holstein denoted HA, HB, HC. Cells were reacted with antiserum specific for TashAT2 (EL24) and images obtained using matched exposures. Bar = 7 μ m.

(TIFF)

S1 Table. Details of oligonucleotide primers used in qRT-PCR. F and R denote forward and reverse primers respectively. *Primers designed based on Ensembl transcript.

(DOCX)

S2 Table. Number of RNA-seq reads mapped on to *B. taurus* genome are not statistically different. Rows denote RNA-seq sample set derived from 6 Sahiwal (S1-6) or 5 Holstein (H1-5) infected cell lines. Columns denote the % of reads mapped to the *B. taurus* genome. Summary denotes mean % of reads mapped, standard deviation and no significant difference between means.

(DOCX)

S3 Table. Top upstream regulators with predicted activated or inhibited activation states in infection-associated H/S-DE genes. In total 655 upstream regulators were found with a p-value of < 0.05; including 123 with a significantly predicted activation state (activated or inhibited, based on a z-score >2, or <-2). Only the top 20 activated and top 20 inhibited regulators based on z-score are presented in the table.

(DOCX)

S1 File. Excel spread sheet of 2211 H/S-DE genes (<0.1 FDR). Columns designate: Ensemble Gene ID; base mean of RNA-seq counts across all samples; \log_2 fold change between mean of Holstein vs. Sahiwal sample counts; lfcSE (Standard Error of the log fold change); stat (Wald statistic); p value (unadjusted p value for the Wald test); padj (Benjamini-Hochberg adjusted p-value for significance of the Wald test); DESeq2 normalized RNA-seq counts for each of the six Sahiwal samples (Sah 1–6) and five Holstein (Hol 1–6); the gene name; an alternative gene name, when applicable; whether RNA-seq counts were consistent across all Sahiwal samples vs. Holstein; Interferon associated genes, as identified in the study of Liu et al. [50] or by GO

[37]; description of protein/factor encoded by the gene. Yellow highlight designate genes also modulated in *Theileria* infected (TBL20) cells.

(XLSX)

S2 File. Excel spread sheet of the 517 genes in H/S-DE that overlap with genes designated as infection associated (modulated in TBL20 vs. BL20). Columns designate: Gene id; DESeq normalized base mean of counts in A (Sahiwal) samples; base mean counts in B (Holstein) samples; \log_2 fold change between mean of Holstein vs. Sahiwal sample counts; lfcSE (Standard Error of the log fold change); stat (Wald statistic); pvalue (unadjusted p value for the Wald test); padj (Benjamini-Hochberg adjusted p-value for significance of the Wald test; the gene name; description of protein/factor encoded by the gene.

(XLSX)

S3 File. Excel spread sheet of the 100 most significant canonical pathways designated as enriched by IPA for H/S-DE data set. Columns designate: The enriched canonical pathway; the $-\log$ p value; ratio of the number of genes from the list that maps to the pathway divided by the total number of genes that map to pathway; the (activation) Z-score; the gene symbol for molecules in H/S-DE data set present in the pathway. Blue highlight denotes pathways with a Z score >2 , indicating repression in Holstein vs. Sahiwal infected cells; red highlight indicates activation (Z-score >1.5).

(XLSX)

S4 File. Excel spread sheet of the 100 most significant canonical pathways enriched by IPA from the data set of 517 overlapped infection-associated H/S-DE genes. Columns designate: The enriched canonical pathway; the $-\log$ p value; the ratio of the number of genes from the list that maps to the pathway divided by the total number of genes that map to pathway; the (activation) Z-score; the gene symbol for molecules in H/S DE data set present in the pathway. Blue highlight denotes pathways with an activation score (Z score >2) indicating repression in Holstein vs. Sahiwal infected cells.

(XLSX)

S5 File. Excel spread sheet of genes in H/S-DE data set predicted to encode secreted proteins. Columns designate: Ensemble Gene ID; DESeq normalized base mean of counts in all samples; \log_2 fold change between mean of Holstein vs. Sahiwal sample counts; lfcSE (Standard Error of the log fold change); stat (Wald statistic); pvalue (unadjusted p value for the Wald test); padj (Benjamini-Hochberg adjusted p-value for significance of the Wald test; the gene name; description of protein/factor encoded by the gene; prediction of location based on gene ontology data in Genecards [35]; Yellow highlight designate genes also modulated in *Theileria* infected (TBL20) cells.

(XLSX)

S6 File. Excel spread sheet of genes in H/S-DE data set predicted to encode receptors. Columns designate: Ensemble Gene ID; Deseq2 normalized base mean for RNA-seq counts for all Sahiwal and Holstein samples; the \log_2 fold change between Holstein and Sahiwal samples for that gene; lfcSE (Standard Error of the log fold change); stat (Wald statistic); pvalue (unadjusted p value for the Wald test); padj (Benjamini-Hochberg adjusted p-value for significance of the Wald test; the gene name; description of protein/factor encoded by the gene; Yellow highlight designate genes also modulated in *Theileria* infected (TBL20) cells.

(XLSX)

S7 File. Excel spread sheet of 109 parasite (*T. annulata*) genes (<0.1 FDR). Columns designate: TA number (T. annulata gene ID); DESeq normalized base mean of RNA-seq counts

across all samples; designation of whether expression is higher (up) or lower (down) in Holstein (*B. taurus*) samples relative to Sahiwal (*B. indicus*); DESeq normalized base mean of counts in A (Sahiwal) samples; base mean counts in B (Holstein) samples; \log_2 fold change between mean of Holstein vs. Sahiwal sample counts; lfcSE (Standard Error of the log fold change); stat (Wald statistic); pvalue (unadjusted p value for the Wald test); padj (Benjamini-Hochberg adjusted p-value for significance of the Wald test; description of protein/factor predicted to be encoded by the gene; location of gene on *T. annulata* genome; genomic sequence ID.

(XLS)

S8 File. Excel spread sheet of nucleotide motifs bound by TashAT2 in genome of *B. taurus* and *B. indicus*. Columns designate chromosome; motif start position; motif end position; genomic feature linked to motif position (non-coding, intron, exon, untranslated region (UTR)); gene product associated with motif and genomic feature, if applicable.

(XLSX)

S9 File. Excel spread sheet of nucleotide motifs bound by TashAT2 in gene coding (RNA) regions of *B. taurus* and *B. indicus* genomes. The data sheets list motifs identified in *B. taurus* genes, motifs identified in *B. indicus* genes and motifs in genes shared by *B. taurus* and *B. indicus*. Columns indicate: Gene identifier, number of motifs identified in that gene region; and for shared genes: B. number of motifs identified in *B. taurus* gene; C. number of motifs identified in *B. indicus* gene.

(XLSX)

S10 File. List of genes in integrin signalling pathway that bear different numbers of nucleotide motif bound by TashAT2 in *B. taurus* and *B. indicus* genomes. Green highlight designates genes that display differential expression between Sahiwal and Holstein infected cells.

(DOCX)

Acknowledgments

We would like to thank the Centre for Virus Research, University of Glasgow for providing access to the IPA software.

Author Contributions

Conceptualization: Elizabeth J. Glass, Brian R. Shiels.

Data curation: Stephen D. Larcombe, Paul Capewell, William Weir.

Formal analysis: Paul Capewell, Kirsty Jensen, Jane Kinnaird, Brian R. Shiels.

Funding acquisition: William Weir, Elizabeth J. Glass, Brian R. Shiels.

Investigation: Stephen D. Larcombe, Kirsty Jensen, Brian R. Shiels.

Methodology: Stephen D. Larcombe, Paul Capewell, Kirsty Jensen, William Weir.

Project administration: Brian R. Shiels.

Supervision: Brian R. Shiels.

Validation: Stephen D. Larcombe, Kirsty Jensen, Jane Kinnaird.

Writing – original draft: Stephen D. Larcombe, Paul Capewell, Brian R. Shiels.

Writing – review & editing: Kirsty Jensen, William Weir, Jane Kinnaird, Elizabeth J. Glass, Brian R. Shiels.

References

1. Betts A, Gray C, Zelek M, MacLean RC, King KC. High parasite diversity accelerates host adaptation and diversification. *Science*. 2018; 360:907–911. <https://doi.org/10.1126/science.aam9974> PMID: 29798882
2. Glass EJ, Crutchley S, Jensen K. Living with the enemy or uninvited guests: functional genomics approaches to investigating host resistance or tolerance traits to a protozoan parasite, *Theileria annulata*, in cattle. *Vet Immunol Immunopathol*. 2012; 148:178–89. <https://doi.org/10.1016/j.vetimm.2012.03.006> PMID: 22482839
3. Brown C, Hunter A, Luckins A. Diseases caused by protozoa. *Handbook of Animal Diseases in the Tropics*. 4th Ed. Cambridge UK: Cambridge University Press; 1990.
4. Minjauw B, McLeod A. Tick-borne Diseases and Poverty. The Impact of Ticks and Tick-borne Diseases on the Livelihoods of Small Scale and Marginal Livestock Owners in India and Eastern and Southern Africa. Research Report, DFID Animal Health Programme. Centre for Tropical Veterinary Medicine, University of Edinburgh. 2003; 116.
5. Glass EJ, Jensen K. Resistance and susceptibility to a protozoan parasite of cattle—gene expression differences in macrophages from different breeds of cattle. *Vet Immunol Immunopathol*. 2007; 120(1–2):20–30. <https://doi.org/10.1016/j.vetimm.2007.07.013> PMID: 17727964
6. Singh CV. Cross-breeding in Cattle for Milk Production: Achievements, Challenges and Opportunities in India—A Review. *Adv Dairy Res*. 2016; 4:158.
7. Kolte SW, Larcombe SD, Jadhao SG, Magar SP, Warthi G, Kurkure NV, et al. PCR Diagnosis of Tick-Borne Pathogens in Maharashtra State, India Indicates Fitness Cost Associated with Carrier Infections Is Greater for Crossbreed Than Native Cattle Breeds. *PLoS One*. 201; 12(3):e0174595. <https://doi.org/10.1371/journal.pone.0174595> PMID: 28358861
8. Larcombe SD, Kolte SW, Ponnudurai G, Kurkure N, Magar S, Velusamy R, et al. The impact of tick-borne pathogen infection in Indian bovines is determined by host type but not the genotype of *Theileria annulata*. *Infect Genet Evol*. 2019; 75:103972. <https://doi.org/10.1016/j.meegid.2019.103972> PMID: 31344487
9. Glass EJ, Preston PM, Springbett A, Craigmile S, Kirvar E, Wilkie G, et al. *Bos taurus* and *Bos indicus* (Sahiwal) calves respond differently to infection with *Theileria annulata* and produce markedly different levels of acute phase proteins. *Int J Parasitol*. 2005; 35(3):337–47. <https://doi.org/10.1016/j.ijpara.2004.12.006> PMID: 15722085
10. Glass EJ, Innes EA, Spooner RL, Brown CG. Infection of bovine monocyte/macrophage populations with *Theileria annulata* and *Theileria parva*. *Vet Immunol Immunopathol*. 1989; 22(4):355–68. [https://doi.org/10.1016/0165-2427\(89\)90171-2](https://doi.org/10.1016/0165-2427(89)90171-2) PMID: 2516673
11. Hulliger L, Wilde KH, Brown CG, Turner L. Mode of multiplication of *Theileria* in cultures of bovine lymphocytic cells. *Nature*. 1964; 203:728–30. <https://doi.org/10.1038/203728a0> PMID: 14207267
12. Irvin AD, Morrison WI. Immunopathology, immunology and immuno- prophylaxis of Theileria infections. In: Soulsby EJJ, editor. *Immune Responses in Parasitic Infections: Immunology, Immunopathology and Immunoprophylaxis vol III Protozoa* Baton Rouge, Florida: CRC Press Inc; 1987. 223–274.
13. Chaussepied M, Janski N, Baumgartner M, Lizundia R, Jensen K, Weir W, et al. TGF- β 2 Induction Regulates Invasiveness of *Theileria*-Transformed Leukocytes and Disease Susceptibility. *PLoS Pathog*. 2010; 6(11):e1001197. <https://doi.org/10.1371/journal.ppat.1001197> PMID: 21124992
14. Kinnaird JH, Weir W, Durrani Z, Pillai SS, Baird M, Shiels BR. A Bovine Lymphosarcoma Cell Line Infected with *Theileria annulata* Exhibits an Irreversible Reconfiguration of Host Cell Gene Expression. *PLoS One*. 2013; 8(6):e66833. <https://doi.org/10.1371/journal.pone.0066833> PMID: 23840536
15. Durrani Z, Weir W, Pillai S, Kinnaird J, Shiels B. Modulation of activation-associated host cell gene expression by the apicomplexan parasite *Theileria annulata*. *Cell Microbiol*. 2012; 14(9):1434–54. <https://doi.org/10.1111/j.1462-5822.2012.01809.x> PMID: 22533473
16. Marsolier J, Perichon M, DeBarry JD, Villoutreix BO, Chluba J, Lopez T, et al. *Theileria* parasites secrete a prolyl isomerase to maintain host leukocyte transformation. *Nature*. 2015; 520(7547):378–82. <https://doi.org/10.1038/nature14044> PMID: 25624101
17. Unlu AH, Tajeri S, Bilgic HB, Eren H, Karagenc T, Langsley G. The secreted *Theileria annulata* Ta9 protein contributes to activation of the AP-1 transcription factor. *PLoS One*. 2018; 13(5):e0196875. <https://doi.org/10.1371/journal.pone.0196875> PMID: 29738531

18. Hayashida K. et al. Comparative genome analysis of three eukaryotic parasites with differing abilities to transform leukocytes reveals key mediators of *Theileria*-induced leukocyte transformation. *mBIO* 2012; 3(5),e00204–12. <https://doi.org/10.1128/mBio.00204-12> PMID: 22951932
19. Swan DG, Stern R, McKellar S, Phillips K, Oura CA, Karagenc TI, et al. Characterisation of a cluster of genes encoding *Theileria annulata* AT hook DNA-binding proteins and evidence for localisation to the host cell nucleus. *J Cell Sci.* 2001; 114:2747–2754. PMID: 11683409
20. Oura CA, McKellar S, Swan DG, Okan E, Shiels BR. Infection of bovine cells by the protozoan parasite *Theileria annulata* modulates expression of the ISGylation system. *Cell Microbiol.* 2006; 8(2):276–88. <https://doi.org/10.1111/j.1462-5822.2005.00620.x> PMID: 16441438
21. McGuire K, Manuja A, Russell GC, Springbett A, Craigmile SC, Nichani AK, et al. Quantitative analysis of pro-inflammatory cytokine mRNA expression in *Theileria annulata*-infected cell lines derived from resistant and susceptible cattle. *Vet Immunol Immunopathol.* 2004; 99:87–98. <https://doi.org/10.1016/j.vetimm.2004.01.003> PMID: 15113657
22. Graham SP, Brown DJ, Vatanserver Z, Waddington D, Taylor LH, Nichani AK, et al. Proinflammatory cytokine expression by *Theileria annulata* infected cell lines correlates with the pathology they cause in vivo. *Vaccine.* 2001; 19(20–22):2932–44. [https://doi.org/10.1016/s0264-410x\(00\)00529-6](https://doi.org/10.1016/s0264-410x(00)00529-6) PMID: 11282205
23. Jensen K, Paxton E, Waddington D, Talbot R, Darghouth MA, Glass EJ. Differences in the transcriptional responses induced by *Theileria annulata* infection in bovine monocytes derived from resistant and susceptible cattle breeds. *Int J Parasitol.* 2008; 38(3–4):313–25. <https://doi.org/10.1016/j.ijpara.2007.08.007> PMID: 17949724
24. Martin M. Cutadapt removes adapter sequences from high-throughput sequencing reads. *Embnet. J.* 2011; 17,10–12.
25. Joshi NA, Fass JN. (2011). Sickel: A sliding-window, adaptive, quality-based trimming tool for FastQ files (Version 1.33). Available from: <https://github.com/najoshi/sickle>.
26. Zerbino DR, Achuthan P, Akanni W, Amode MR, Barrell D, Bhari J, et al. Ensembl 2018 *Nucleic Acids Res.* 2018; 46(D1):D754–D761. <https://doi.org/10.1093/nar/gkx1098> PMID: 29155950
27. Langmead B, Salzberg S. Fast gapped-read alignment with Bowtie 2. *Nature Methods.* 2012; 9:357–359. <https://doi.org/10.1038/nmeth.1923> PMID: 22388286
28. Buffalo V. sam2counts.py—convert SAM mapping results to reference sequence counts. Available from: <https://github.com/vsbuffalo/sam2counts>.
29. Love MI, Huber W, Anders S. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. *Genome Biology.* 2014; 15:550. <https://doi.org/10.1186/s13059-014-0550-8> PMID: 25516281
30. Pain A, Renaud H, Berriman M, Murphy L, Yeats CA, Weir W, et al: Genome of the host-cell transforming parasite *Theileria annulata* compared with *T. parva*. *Science* 2005; 309:131–133. <https://doi.org/10.1126/science.1110418> PMID: 15994557
31. Huang DW, Sherman BT, Lempicki RA. Systematic and integrative analysis of large gene lists using DAVID Bioinformatics Resources. *Nature Protoc.* 2009; 4(1):44–57. <https://doi.org/10.1038/nprot.2008.211> PMID: 19131956
32. Mi H, Huang X, Muruganujan A, Tang H, Mills C, Kang D, et al. PANTHER version 11: expanded annotation data from Gene Ontology and Reactome pathways, and data analysis tool enhancements. *Nucleic Acids Res.* 2017; 45(D1):D183–D189. <https://doi.org/10.1093/nar/gkw1138> PMID: 27899595
33. Land J. Statistical significance of the overlap between two groups of genes. Available from: http://nemates.org/MA/progs/overlap_stats.html.
34. Almagro Armenteros JJ, Tsirigos KD, Sønderby CK, Petersen TN, Winther O, Brunak S, et al. SignalP 5.0 Improves Signal Peptide Predictions Using Deep Neural Networks. *Nat Biotechnol.* 2019; 37(4):420–423. <https://doi.org/10.1038/s41587-019-0036-z> PMID: 30778233
35. Krogh A, Larsson B, von Heijne G, Sonnhammer EL. Predicting transmembrane protein topology with a hidden Markov model: application to complete genomes. *J Mol Biol.* 2001; 305(3):567–580. <https://doi.org/10.1006/jmbi.2000.4315> PMID: 11152613
36. Pierleoni A, Martelli PL, Casadio R. PredGPI: A GPI-anchor Predictor. *BMC Bioinformatics.* 2008; 9:392. <https://doi.org/10.1186/1471-2105-9-392> PMID: 18811934
37. Weizmann Institute of Science. GeneCards. Available from: <https://www.genecards.org>.
38. Jensen K, Gallagher IJ, Johnston N, Welsh M, Skuce R, Williams JL, et al. Variation in the Early Host-Pathogen Interaction of Bovine Macrophages with Divergent *Mycobacterium bovis* Strains in the United Kingdom. *Infect Immun.* 2018; 86(3):e00385–17. <https://doi.org/10.1128/IAI.00385-17> PMID: 29263113

39. Rice P., Longden I., Bleasby A. EMBOSS: the European Molecular Biology Open Software Suite Trends Genet. 2000; 16:276–277. [https://doi.org/10.1016/s0168-9525\(00\)02024-2](https://doi.org/10.1016/s0168-9525(00)02024-2) PMID: 10827456
40. Quinlan Aaron R., and Hall Ira M. "BEDTools: a flexible suite of utilities for comparing genomic features." Bioinformatics 2010; 26.6: 841–842. <https://doi.org/10.1093/bioinformatics/btq033> PMID: 20110278
41. Krzywinski Martin, et al. "Circos: an information aesthetic for comparative genomics." *Genome research* 2009; 19.9:1639–1645. <https://doi.org/10.1101/gr.092759.109> PMID: 19541911
42. Kinnaird JH, Singh M, Gillan V, Weir W, Calder ED, Hostettler I, et al. Characterization of HSP90 isoforms in transformed bovine leukocytes infected with *Theileria annulata*. *Cell Microbiol.* 2017; 19(3): e12669. <https://doi.org/10.1111/cmi.12669> PMID: 27649068
43. Bilgic HB, Karagenc T, Bakirci S, Shiels B, Tait A, Kinnaird J, et al. Identification and Analysis of Immunodominant Antigens for ELISA-Based Detection of *Theileria annulata*. *PLoS One.* 2016; 11(6): e0156645. <https://doi.org/10.1371/journal.pone.0156645> PMID: 27270235
44. Swan DG, Phillips K, Tait A, Shiels BR. Evidence for localisation of a *Theileria* parasite AT hook DNA-binding protein to the nucleus of immortalised bovine host cells. *Mol Biochem Parasitol.* 1999; 101(1–2):117–29. [https://doi.org/10.1016/s0166-6851\(99\)00064-x](https://doi.org/10.1016/s0166-6851(99)00064-x) PMID: 10413048
45. Chaturvedi S, Yuen DA, Bajwa A, Huang YW, Sokollik C, Huang L, et al. Slit2 prevents neutrophil recruitment and renal ischemia-reperfusion injury. *J Am Soc Nephrol.* 2013; 24(8):1274–87. <https://doi.org/10.1681/ASN.2012090890> PMID: 23766538
46. Zhang QQ, Zhou DL, Lei Y, Zheng L, Chen SX, Gou HJ, et al. Slit2/Robo1 signalling promotes intestinal tumorigenesis through Src-mediated activation of the Wnt/ β -catenin pathway. *Oncotarget.* 2015; 6(5):3123–35. <https://doi.org/10.18632/oncotarget.3060> PMID: 25605242
47. Aichem A, Groettrup M. The ubiquitin-like modifier FAT10 in cancer development. *Int J Biochem Cell Biol.* 2016; 79:451–461. <https://doi.org/10.1016/j.biocel.2016.07.001> PMID: 27393295
48. Faralli JA, Desikan H, Peotter J, Kanneganti N, Weinhaus B, Filla MS, et al. Genomic/proteomic Analyses of Dexamethasone-Treated Human Trabecular Meshwork Cells Reveal a Role for GULP1 and ABR in Phagocytosis. *Mol Vis.* 2019; 25:237–254. PMID: 31516309
49. Datta S, Nam HS, Hayashi M, Maldonado L, Goldberg R, Brait M, et al. Expression of GULP1 in Bronchial Epithelium Is Associated with the Progression of Emphysema in Chronic Obstructive Pulmonary Disease. *Respir Med.* 2017; 124:72–78. <https://doi.org/10.1016/j.rmed.2017.02.011> PMID: 28284325
50. Oryan A, Namazi F, Sharifiyazdi H, Razavi M, Shahriari R. Clinicopathological Findings of a Natural Outbreak of *Theileria annulata* in Cattle: An Emerging Disease in Southern Iran. *Parasitol Res.* 2013; 112(1):123–7. <https://doi.org/10.1007/s00436-012-3114-4> PMID: 22968949
51. Liu SY, Sanchez DJ, Aliyari R, Lu S, Cheng G. Systematic identification of type I and type II interferon-induced antiviral factors. *Proc Natl Acad Sci U S A.* 2012; 109(11):4239–44. <https://doi.org/10.1073/pnas.1114981109> PMID: 22371602
52. Pombo JP, Sanyal S. Perturbation of Intracellular Cholesterol and Fatty Acid Homeostasis During Flavivirus Infections. *Front Immunol.* 2018; 9:1276. <https://doi.org/10.3389/fimmu.2018.01276> PMID: 29915602
53. Baenke F, Peck B, Miess H, Schulze A. Hooked on fat: the role of lipid synthesis in cancer metabolism and tumour development. *Dis Model Mech.* 2013; 6(6):1353–63. <https://doi.org/10.1242/dmm.011338> PMID: 24203995
54. Ehmsen S, Pedersen MH, Wang G, Terp MG, Arslanagic A, Hood BL, et al. Increased Cholesterol Biosynthesis Is a Key Characteristic of Breast Cancer Stem Cells Influencing Patient Outcome. *Cell Rep.* 2019; 27(13):3927–3938.e6. <https://doi.org/10.1016/j.celrep.2019.05.104> PMID: 31242424
55. Heussler VT, Küenzi P, Fraga F, Schwab RA, Hemmings BA, Dobbelaere DA. The Akt/PKB Pathway Is Constitutively Activated in *Theileria*-transformed Leucocytes, but Does Not Directly Control Constitutive NF-kappaB Activation. *Cell Microbiol.* 2001; 3(8):537–50. <https://doi.org/10.1046/j.1462-5822.2001.00134.x> PMID: 11488815
56. Treff NR, Derek Pouchnik D, Dement GA, Britt RL and Reeves R. High-mobility group A1a protein regulates Ras/ERK signalling in MCF-7 human breast cancer cells. *Oncogene* 2004; 23:777–785. <https://doi.org/10.1038/sj.onc.1207167> PMID: 14737112
57. Salameh A, Lee AK, Cardó-Vila M, Nunes DN, Efsthathiou E, Staquicini FI, et al. PRUNE2 is a human prostate cancer suppressor regulated by the intronic long noncoding RNA PCA3. *Proc Natl Acad Sci U S A.* 2015; 112(27):8403–8. <https://doi.org/10.1073/pnas.1507882112> PMID: 26080435
58. Koltai T. Clusterin: a key player in cancer chemoresistance and its inhibition. *Onco Targets Ther.* 2014; 7:447–56. <https://doi.org/10.2147/OTT.S58622> PMID: 24672247

59. Sansanwal P, Li L, Sarwal MM. Inhibition of intracellular clusterin attenuates cell death in nephropathic cystinosis. *J Am Soc Nephrol*. 2015; 26(3):612–25. <https://doi.org/10.1681/ASN.2013060577> PMID: 25071085
60. Chen J, Wang M, Xi B, Xue J, He D, Zhang J, et al. SPARC is a key regulator of proliferation, apoptosis and invasion in human ovarian cancer. *PLoS One*. 2012; 7(8):e42413. <https://doi.org/10.1371/journal.pone.0042413> PMID: 22879971
61. Fenouille N, Puissant A, Tichet M, Zimniak G, Abbe P, Mallavialle A, et al. SPARC functions as an anti-stress factor by inactivating p53 through Akt-mediated MDM2 phosphorylation to promote melanoma cell survival. *Oncogene*. 2011; 30(49):4887–900. <https://doi.org/10.1038/onc.2011.198> PMID: 21685937
62. Hayashida K, Hattori M, Nakao R, Tanaka Y, Kim JY, Inoue N, et al. MDM2 regulates a novel form of incomplete neoplastic transformation of *Theileria parva* infected lymphocytes. *Mol Biochem Parasitol*. 2010; 174(1):8–17. <https://doi.org/10.1016/j.molbiopara.2010.06.005> PMID: 20540970
63. Resmini G, Rizzo S, Franchin C, Zanin R, Penzo C, Pegoraro S, et al. HMGA1 regulates the Plasminogen activation system in the secretome of breast cancer cells *Sci Rep*. 2017; 7(1):11768. <https://doi.org/10.1038/s41598-017-11409-4> PMID: 28924209
64. Weir W., Karagenc T, Baird M., Tait A. and Shiels B.R. Evolution and diversity of secretome genes in the apicomplexan parasite *Theileria annulata*. *BMC Genomics*. 2010; 11:42. <https://doi.org/10.1186/1471-2164-11-42> PMID: 20082698
65. Hall R, Ilhan T, Kirvar E, Wilkie G, Preston PM, Darghouth M, et al. Mechanism(s) of attenuation of *Theileria annulata* vaccine cell lines. *Trop Med Int Health*. 1999; 4: A78–84. <https://doi.org/10.1046/j.1365-3156.1999.00454.x> PMID: 10540315
66. Preston PM, Brown CG, Bell-Sakyl L, Richardson W, Sanderson A. Tropical theileriosis in *Bos taurus* and *Bos taurus* cross *Bos indicus* calves: response to infection with graded doses of sporozoites of *Theileria annulata*. *Res Vet Sci*. 1992; 53(2):230–43. [https://doi.org/10.1016/0034-5288\(92\)90115-i](https://doi.org/10.1016/0034-5288(92)90115-i) PMID: 1439213
67. Schneider WM, Chevillotte MD, Rice CM. Interferon-stimulated genes: a complex web of host defenses. *Annu Rev Immunol*. 2014; 32:513–45. <https://doi.org/10.1146/annurev-immunol-032713-120231> PMID: 24555472
68. Wong MT, Chen SS. Emerging roles of interferon-stimulated genes in the innate immune response to hepatitis C virus infection. *Cell Mol Immunol*. 2016; 13(1):11–35. <https://doi.org/10.1038/cmi.2014.127> PMID: 25544499
69. Bogunovic D, Boisson-Dupuis S, Casanova JL. ISG15: leading a double life as a secreted molecule. *Exp Mol Med*. 2013; 45:e18. <https://doi.org/10.1038/emm.2013.36> PMID: 23579383
70. Helbig KJ, Beard MR. The role of viperin in the innate antiviral response. *J Mol Biol*. 2014; 426(6):1210–9. <https://doi.org/10.1016/j.jmb.2013.10.019> PMID: 24157441
71. Kim JC, Ha YJ, Tak KH, Roh SA, Kwon YH, Kim CW, et al. Opposite functions of GSN and OAS2 on colorectal cancer metastasis, mediating perineural and lymphovascular invasion, respectively. *PLoS One*. 2018; 13(8):e0202856. <https://doi.org/10.1371/journal.pone.0202856> PMID: 30148861
72. Kuriakose T, Kanneganti TD. ZBP1: Innate Sensor Regulating Cell Death and Inflammation. *Trends Immunol*. 2018; 39(2):123–134. <https://doi.org/10.1016/j.it.2017.11.002> PMID: 29236673
73. Numajiri Haruki A, Naito T, Nishie T, Saito S, Nagata K. Interferon-inducible antiviral protein MxA enhances cell death triggered by endoplasmic reticulum stress. *J Interferon Cytokine Res*. 2011; 31(11):847–56. <https://doi.org/10.1089/jir.2010.0132> PMID: 21992152
74. Xu D, Zhang T, Xiao J, Zhu K, Wei R, Wu Z, et al. Modification of BECN1 by ISG15 plays a crucial role in autophagy regulation by type I IFN/interferon. *Autophagy*. 2015; 11(4):617–28. <https://doi.org/10.1080/15548627.2015.1023982> PMID: 25906440
75. Medina GN, Segundo FD, Stenfeldt C, Arzt J, de Los Santos T. The Different Tactics of Foot-and-Mouth Disease Virus to Evade Innate Immunity. *Front Microbiol*. 2018; 9:2644. <https://doi.org/10.3389/fmicb.2018.02644> PMID: 30483224
76. Singh R, Deb R, Singh U, Raja TV, Alex R, Kumar S, et al. Heterozygosity of the SNP (rs136500299) of ITGB6 receptor gene possibly influences susceptibility among crossbred bull to foot and mouth disease infection. *Virus disease*. 2015; 26:48–54. <https://doi.org/10.1007/s13337-015-0249-9> PMID: 26436121
77. Walker AM, Roberts RM. Characterization of the bovine type I IFN locus: rearrangements, expansions, and novel subfamilies. *BMC Genomics*. 2009; 10:187. <https://doi.org/10.1186/1471-2164-10-187> PMID: 19393062
78. Sager H, Brunschweiler C, Jungi TW. Interferon production by *Theileria annulata*-transformed cell lines is restricted to the beta family. *Parasite Immunol*. 1998; 20(4):175–82. <https://doi.org/10.1046/j.1365-3024.1998.00141.x> PMID: 9618728

79. Perucha E, Melchiotti R, Bibby JA, Wu W, Frederiksen KS, Roberts CA, et al. The cholesterol biosynthesis pathway regulates IL-10 expression in human Th1 cells. *Nat Commun.* 2019; 10(1):498. <https://doi.org/10.1038/s41467-019-08332-9> PMID: 30700717
80. Lee W, Ahn JH, Park HH, Kim HN, Kim H, Yoo Y, et al. COVID-19-activated SREBP2 disturbs cholesterol biosynthesis and leads to cytokine storm. *Signal Transduct Target Ther.* 2020; 5(1):186. <https://doi.org/10.1038/s41392-020-00292-7> PMID: 32883951
81. Preston PM, Dargouth M, Boulter NR, Hall FR, Tall R, Kirvar E, et al. A dual role for immunosuppressor mechanisms in infection with *Theileria annulata*: well-regulated suppressor macrophages help in recovery from infection; profound immunosuppression promotes non-healing disease. *Parasitol Res.* 2002; 88:522–534. <https://doi.org/10.1007/s00436-002-0613-8> PMID: 12107474
82. Ozturk N, Singh I, Mehta A, Braun T, Barreto G. HMGA proteins as modulators of chromatin structure during transcriptional activation. *Front Cell Dev Biol.* 2014; 2:5. <https://doi.org/10.3389/fcell.2014.00005> PMID: 25364713
83. Vignali R, Marracci S. HMGA Genes and Proteins in Development and Evolution. *Int. J. Mol. Sci.* 2020; 21:654. <https://doi.org/10.3390/ijms21020654> PMID: 31963852
84. Wang X, Wang J Wu. Emerging roles for HMGA2 in colorectal cancer. *Translational Oncology* 2021; 14:100894 <https://doi.org/10.1016/j.tranon.2020.100894> PMID: 33069103
85. Reeves R, Edberg DD, Li Y. Architectural transcription factor HMGI(Y) promotes tumour progression and mesenchymal transition of human epithelial cells. *Mol Cell Biol.* 2001; 21(2):575–94. <https://doi.org/10.1128/MCB.21.2.575-594.2001> PMID: 11134344
86. Jørn Henriksen^{1,2}, Marianne Stabell^{1,2}, Meza-Zepeda Leonardo A^{1,2}, Lauvrak Silje AU¹, Kassem Moustapha³ and Myklebost Ola. Identification of target genes for wild type and truncated HMGA2 in mesenchymal stem-like cells. *BMC Cancer* 2010; 10:329. <https://doi.org/10.1186/1471-2407-10-329> PMID: 20576167
87. Haidar M, Whitworth J, Noé G, Liu WQ, Vidal M, Langsley G. TGF- β 2 induces Grb2 to recruit PI3-K to TGF-RII that activates JNK/AP-1-signaling and augments invasiveness of *Theileria*-transformed macrophages. *Sci Rep.* 2015; 5:15688. <https://doi.org/10.1038/srep15688> PMID: 26511382
88. Neri P, Ren L, Azab AK, Brentnall M, Gratton K, Klimowicz AC, et al. Integrin β 7-mediated regulation of multiple myeloma cell adhesion, migration, and invasion. *Blood.* 2011; 117(23):6202–13. <https://doi.org/10.1182/blood-2010-06-292243> PMID: 21474670
89. Lamb CA, O'Byrne S, Keir ME, Butcher EC. Gut-Selective Integrin-Targeted Therapies for Inflammatory Bowel Disease. *J Crohns Colitis.* 2018; 12(2):S653–S668. <https://doi.org/10.1093/ecco-jcc/jjy060> PMID: 29767705
90. Quante T, Bird A. Do short, frequent DNA sequence motifs mould the epigenome? *Nat. Rev. Mol. Cell Biol.* 2016; 17:257–262. <https://doi.org/10.1038/nrm.2015.31> PMID: 26837845