Responses of Bovine Innate Immunity to *Mycobacterium avium* subsp. *paratuberculosis* Infection Revealed by Changes in Gene Expression and Levels of MicroRNA

Michela Malvisi¹²*, Fiorentina Palazzo³, Nicola Morandi¹, Barbara Lazzari¹⁴, John L. Williams¹⁵, Giulio Pagnacco², Giulietta Minozzi²

¹ Parco Tecnologico Padano, Lodi, Italy, ² Department of Veterinary Medicine, University of Milan, Milan, Italy, ³ Faculty of Bioscience and Technology for Food, Agriculture and Environment, University of Teramo, Teramo, Italy, ⁴ Institute of Agricultural Biology and Biotechnology, National Research Council, Lodi, Italy, ⁵ School of Animal and Veterinary Sciences, University of Adelaide, Roseworthy, Australia

* michela.malvisi@yahoo.com

Abstract

Paratuberculosis in cattle is a chronic granulomatous gastroenteritis caused by *Mycobacterium avium* subsp. *paratuberculosis* (MAP) which is endemic worldwide. In dairy herds, it is responsible for huge economic losses. However, current diagnostic methods do not detect subclinical infection making control of the disease difficult. The identification of MAP infected animals during the sub-clinical phase of infection would play a key role in preventing the dissemination of the pathogen and in reducing transmission. Gene expression and circulating microRNA (miRNA) signatures have been proposed as biomarkers of disease both in the human and veterinary medicine. In this paper, gene expression and related miRNA levels were investigated in cows positive for MAP, by ELISA and culture, in order to identify potential biomarkers to improve diagnosis of MAP infection. Three groups, each of 5 animals, were used to compare the results of gene expression from positive, exposed and negative cows. Overall 258 differentially expressed genes were identified between unexposed, exposed, but ELISA negative and positive groups which were involved in biological functions related to inflammatory response, lipid metabolism and small molecule biochemistry. Differentially expressed miRNA was also found among the three groups: 7 miRNAs were at a lower level and 2 at a higher level in positive animals vs unexposed animals, while 5 and 3 miRNAs were respectively reduced and increased in the exposed group compared to the unexposed group. Among the differentially expressed miRNAs 6 have been previously described as immune-response related and two were novel miRNAs. Analysis of the miRNA levels showed correlation with expression of their target genes, known to be involved in the immune process. This study suggests that miRNA expression is affected by MAP infection and play a key role in tuning the host response to infection.
The miRNA and gene expression profiles may be biomarkers of infection and potential diagnostic of MAP infection earlier than the current ELISA based diagnostic tests.

Background

*Mycobacterium avium* subspecies *paratuberculosis* (MAP) is the causal agent of paratuberculosis (paraTB) or Johne’s disease in cattle, a chronic granulomatous gastroenteritis [1, 2]. ParaTB is endemic worldwide and occurs primarily in ruminants, including cattle, sheep, goats, and farmed deer. However, the disease has been reported in non-ruminants, such as wild rabbits [3], foxes and stoats [4] and in primates such as mandrills and macaques [5, 6].

Johne’s disease causes substantial economic losses in dairy herds through lost productivity [7]. A link between MAP and Crohn’s disease in humans has been suggested [8, 9], although the causal role of MAP has not yet been proven [10, 11] and the association remains controversial [11]. However, this possible risk to human health has increased interest in the disease and has made the need to improved diagnosis more pressing.

In cattle, the disease starts with the slow development of intestinal lesions in infected animals, a proportion of which become clinically ill two to six years after infection [12]. The disease progresses in four stages, which start with the silent phase, followed by subclinical, clinical and advanced phases [13]. Cows in the subclinical stages can be classified as low, moderate and high shedders, on the basis of the number bacteria that can be detected by fecal culture [14]. The subclinical stage is immunologically characterized by a protective Th1 immune response, and an elevated level of IFN-γ [15]. The progression of infection and appearance of clinical disease is associated with the shift from a Th1 to a non-protective Th2-mediated humoral response in the late subclinical phase [15].

The identification of infected animals at an early subclinical stage is critical to avoid transmission via the oral-fecal route and the dissemination of the pathogen. Infection mainly occurs in young calves, which are most susceptible, while adult cows are more resistant to infection [16]. The clinical phase is characterized by untreatable diarrhea, progressive weight loss, decreased milk production and ultimately death [17]. Currently infection is detected by an ELISA test to detect serum antibodies against MAP, or PCR of feces to detect the presence of the bacterium. However, antibodies are only present late in infection and detection of the bacterium by PCR has only moderate sensitivity when low shedders are tested [18, 19]. The fecal bacterial culture [20], does not identify subclinical cases [13, 21] and can give false positives when environmental MAP passes through uninfected animals or false negatives due to intermittent shedding.

There are no effective treatments for Johne’s disease, and vaccine efficacy as protective tool in paratuberculosis prevention is still debatable [22]. Therefore, the detection and isolation of animals in the early stages of infection can play a key role in Johne’s disease eradication. It is therefore important to develop a diagnostic test for animals during the early stages of infection, before they start shedding and spreading the disease.

Understanding host-pathogen interaction and disease responses has improved with the availability of high throughput—omics technologies. Studies have progressed from the analysis of candidate genes or loci associated with MAP susceptibility [23–25] to the investigation of the whole transcriptome. Expression microarrays have been used to analyze gene expression related to MAP infection in Holstein-Friesian orally-inoculated calves [26, 27]. Degradation of miRNA-targeted mRNA is now a well-known mechanism of post-transcriptional regulation of
gene expression in plants and animals [28], although the levels of miRNAs and transcript abundance are poorly correlated [29]. Nevertheless, circulating miRNAs, contained in extracellular compartments such as plasma [30] or exosomes [31], have been identified as markers for human diseases, and are used in the diagnosis of some cancers [32], cardiovascular diseases [33], disorders of the immune system [34], neurodegenerative disease [35] and diabetes [36]. Significantly, miRNAs have been suggested as diagnostic markers of human tuberculosis infections [37]. In the veterinary field, specific miRNA patterns have been associated with viral diseases [38, 39] and gram-positive bacterial infections [40]. Although the mechanism leading the bacteria-induced miRNA expression changes has yet to be fully understood, both pathogenic and commensal bacteria have been shown to affect miRNA expression in the host [41]. Differentially expressed miRNAs have been suggested as diagnostic markers of MAP infection, but experimentally infected Holstein-Friesian calves did not show specific miRNA signatures, at least during the early latent period of MAP infection when diagnosis would be most useful [42]. In the present study, the mRNA and circulating miRNA expression were investigated in Holstein cattle positive and negative for MAP by ELISA to gain insights into gene regulatory networks related to Johne's disease in clinical and preclinical phases of the disease.

Results
Differentially expressed genes and quantitative RT-PCR validation of the sequencing data

RNAseq data from whole blood of 5 infected (PP), 5 exposed (NP) and 5 negative (NN) animals identified 12,366 genes, 258 of which were differentially expressed in the three comparisons: 162 genes were differentially expressed (DE) comparing PP vs NN, 94 genes for NP vs NN and 2 genes for PP vs NP. In this study Log$_2$ fold change above 1 or below -1 and 0.05 as false discovery rate (FDR) threshold were chosen to select DE genes. The complete lists of DE genes for the three comparisons are provided in supporting information files (S1, S2 and S3 Tables). Fifty-four DE genes were shared by PP vs NN and NP vs NN with the same trend of expression (Fig 1). All of animals were from the same breed and they have been carefully selected to be uniform for age, physiologic and health status but they differed for the ELISA test result. Thus the similarities found in the gene expression when the groups were compared with the control suggest that exposed animals may have been incubating disease in the preclinical stage. In addition, some animals of the NP group showed expression values similar to the PP group. A similar number of DE genes had increased vs decreased levels of expression in each comparison. The Log$_2$ fold expression changes ranged from -5.0 to 3.6 for PP vs NN, -2.4 to 2.9 for NP vs NN and -2.7 to 1.7 in the PP vs NP comparisons. The Pearson correlation coefficient for RNA-Seq vs RT-qPCR data for TTYH3, LOC617313, ZNF467 and IDO1 which were DE in NP and PP group vs NN was 0.945 (p<0.01). (Fig 2).

Differential expression of miRNAs

The miRNA-Seq data was used to identify known, and to predict novel miRNAs. Overall, 645 novel miRNAs were identified by the Mirdeep2 software [43].

Two samples (one PP and one NP) were excluded due to low read numbers and analysis for differential expression of miRNAs was conducted on the remaining 20 samples.

miRNAs with an FDR ≤0.05 were considered as differentially expressed. The comparison PP vs NN identified DE of 8 known and 1 novel miRNA. Among these, 7 had lower expression in the PP group compared with the NN group, with the novel miRNA 14_7917 having the
greatest decrease (-6.54 Log$_2$ FC), and 2 known miRNAs had higher expression in the PP group compared with the NN group (Table 1).

Four miRNAs, bta-mir-19b, bta-mir-19b2, bta-mir-1271 and novel miRNA 14_7917, which showed a differential expression between PP and NN groups were also differentially expressed between NP vs NN groups with the same trend of expression. Other differentially expressed miRNAs between the NP and NN groups, which were not DE between PP and NN, were bta-mir-24-1, bta-mir-24-2, bta-mir-378c and Novel:27_25982 (Table 2).

Two miRNAs, bta-miR-19b and bta-miR-19b2, had lower levels of expression in the PP group vs the NP group with -0.9 Log$_2$ Fold Change (Log$_2$ FC) (Table 3), while having reduced expression in PP vs NN and also showing reduced expression in the NP vs NN group, although to a lesser extent.

Most of the DE miRNAs have previously described in other species as having a role in immunity and inflammation. miRNAs involved in inflammation and immunological pathways and their target genes are summarized in Table 4.

Target prediction and validation of regulatory miRNAs

The correlation of expression between predicted targets and miRNA expression levels revealed that expression levels of 6 miRNAs, bta-mir-19b, bta-mir-19b-2, bta-mir-1271, bta-mir-32,
bta-mir-7857 and the Novel:14_7917, were inversely correlated with the level of mRNA expression observed in the PP and NP groups compared to the NN group (Table 5).

Bta-mir-19b and bta-mir-19b-2 share the same targets and therefore overall 12 genes were identified which showed expression consistent with being targets of miRNAs with differential level of expression, 10 of which were in the PP group and 2 in NP group compared with the NN group. Bta-mir-19b and bta-mir-19b2 were differentially expressed in both positive and exposed groups, although apparently affecting the expression of different target genes in the two groups. HIC1, TBC1D8 and IMPDH1 had the same sequence motif and were targeted by both bta-mir-19b and bta-mir-19b-2 in the PP group, while ZBTB4 was targeted by bta-mir-19b and bta-mir-19b-2 in the NP group. None of the other differentially expressed genes were

![Fig 2. Validation of RNA-Seq data with qRT-PCR assays.](image)

**Fig 2. Validation of RNA-Seq data with qRT-PCR assays.** Relative quantification was calculated with respect to the control group, after normalization with ACTB and GAPDH expression levels.

doi:10.1371/journal.pone.0164461.g002

| Comparison | miRNA name  | Chromosome | Log₂FC* | PValue  | FDR    | Mature sequence       |
|------------|-------------|------------|---------|---------|--------|-----------------------|
| PP vs NN   | bta-mir-19b | 12         | -1.38   | <0.001  | <0.001 | UGUGCAAAUCCAGCAAAACUGA |
| PP vs NN   | bta-mir-19b-2 | X | -1.40   | <0.001  | <0.001 | UGUGCAAAUCCAGCAAAACUGA |
| PP vs NN   | bta-mir-1271 | 7          | -0.88   | <0.001  | <0.001 | CUUGGCAACCUGUAUGACUCUCA |
| PP vs NN   | bta-mir-100 | 15         | -6.28   | <0.001  | 0.009  | AACCGGUAGAUGCAACUUGUG |
| PP vs NN   | bta-mir-301a | 19        | -2.38   | <0.001  | 0.003  | CAGUGCAAIAGUUAUGUCAAAGCAU |
| PP vs NN   | bta-mir-32  | 8          | -2.73   | <0.001  | 0.008  | UAUUGGCAAGACUAGUGUGCAU |
| PP vs NN   | bta-mir-6517 | 10        | 1.25    | <0.001  | 0.001  | UCAGGGCGUGACACUACCACG |
| PP vs NN   | bta-mir-7857 | 23       | 1.88    | <0.001  | <0.001 | AUUGCCAGUGUGGCGAAGAUGC |
| PP vs NN   | Novel:14_7917 | 14       | -6.54   | 0.001   | 0.041  | AGCAGCAUGUGAAGGGCUCCUGA |

* Log₂ Fold Change in the positive group (PP) when compared with the unexposed group (NN).

doi:10.1371/journal.pone.0164461.t001
Table 2. Differentially expressed miRNAs in NP vs NN comparison.

| Comparison  | miRNA name   | Chromosome | Log2FC* | PValue  | FDR     | Mature sequence     |
|-------------|--------------|------------|---------|---------|---------|---------------------|
| NP vs NN    | bta-mir-19b  | 12         | -0.48   | <0.001  | 0.001   | UGGCUAAUCAUGCAAAACUGA |
| NP vs NN    | bta-mir-19b-2| X          | -0.51   | <0.001  | 0.003   | UGGCUAAUCAUGCAAAACUGA |
| NP vs NN    | bta-mir-1271 | 7          | -0.72   | <0.001  | <0.001  | CUUGGCCCUAUUGCAUGCAAG |
| NP vs NN    | bta-mir-24-1 | 8          | 0.94    | <0.001  | <0.001  | UGGCCUACUGACGCUAGAAG |
| NP vs NN    | bta-mir-24-2 | 7          | 0.89    | <0.001  | <0.001  | UGGCUAGACGCUAGCAAG |
| NP vs NN    | bta-mir-378c | 24         | 1.44    | <0.001  | 0.017   | ACUGGAGCUAGAGCAAGUA |
| NP vs NN    | Novel:14_7917| 14         | -5.54   | <0.001  | 0.009   | ACGAGCAGGUGACGCGGUCUGA |
| NP vs NN    | Novel:27_25982| 27        | -5.26   | 0.001   | 0.041   | CCGGUACUGACGCGACCCGA |

* Log2 Fold Change in the exposed group (NP) when compared with the unexposed group (NN)

Table 3. miRNAs expression in PP compared with NP animals.

| Comparison  | miRNA name   | Chromosome | Log2FC* | PValue  | FDR     | Mature sequence     |
|-------------|--------------|------------|---------|---------|---------|---------------------|
| PP vs NP    | bta-mir-19b  | 12         | -0.91   | <0.001  | 0.002   | UGGCUAAUCAUGCAAAACUGA |
| PP vs NP    | bta-mir-19b-2| X          | -0.89   | <0.001  | 0.006   | UGGCUAAUCAUGCAAAACUGA |

* Log2 Fold Change in the positive group (PP) when compared with the exposed group (NP)

Table 4. Differentially expressed miRNAs with inflammatory and immunological functions in other species.

| DE miRNA | Species | Tissue                        | Disease                  | Expression | Target         | Target function          | Reference |
|----------|---------|-------------------------------|--------------------------|------------|----------------|--------------------------|----------|
| mir-19b  | Mouse   | mir-17-92 deficient CD4+ T cells | Decreased                | PTEN       | Inhibition of PI3K-AKT-mTOR | [103]    |
| mir-19b-2| Human   | Mucosal tissue                | Crohn's disease          | Decreased  | n.d.           | n.d.                     | [104]    |
| mir-100  | Mouse   | Lungs                         | Viral infection          | Increased  | n.d.           | n.d.                     | [38]     |
| mir-301a | Mouse   | Lung                          | Viral infection          | Decreased  | n.d.           | n.d.                     | [38]     |
| mir-32   | Cow     | Mammary epithelial cells      | S. uberis infection      | Decreased  | n.d.           | n.d.                     | [40]     |
| mir-24-2 | Human   | PBMC                          | Tuberculosis             | Increased  | n.d.           | n.d.                     | [61]     |
| mir-378  | Human   | Serum                         | Tuberculosis             | Increased  | MAPA1          | n.d.                     | [67]     |
| mir-1271 | Human   | Platelets                     | Ulcerative colitis       | Increased  | n.d.           | n.d.                     | [68]     |
targets of more than one miRNA. bta-mir-1271 had significantly lower levels in both PP and NP groups compared to the NN group. TTYH3, which is a target for mir-1271, had a higher level of expression in PP and NP vs NN.

A novel miRNA, Novel:14_7917, showed a lower expression in PP vs NN animals. Putative targets of this miRNA (FAM78A, ARL2 and AP2A1) showed higher levels of expression in the PP vs NN animals.

**Ingenuity Pathway Analysis**

From the 258 DE genes among the three comparisons 174 could be mapped in the Ingenuity Knowledge Base database [44], 119 in PP vs NN animals and 55 in the NP vs NN animals.

The Ingenuity pathway analysis (IPA) of DE genes from the PP vs NN comparison identified biological functions related to inflammatory disease (27 molecules), inflammatory response (36 molecules), cellular movement (37 molecules), lipid metabolism (17 molecules), small molecules biochemistry (29 molecules), cell-to-cell signaling and interaction (31 molecules), and immune cell trafficking (27 molecules). In particular, 16 DE genes were found to be involved in inflammatory response, 7 in lipid metabolism and 12 in small molecule biochemistry associated functions.

IPA identified "Eicosanoid Signaling" and "Leukotriene Biosynthesis" as the top canonical pathways in both PP vs NN and NP vs NN comparisons (Table 6). In addition, the analysis of DE genes between PP and NN groups were associated with "G-Protein Coupled Receptor Signaling" and "Granulocyte Adhesion and Diapedesis" canonical pathways, while DE genes from the NP vs NN comparison were enriched "EIF2 Signaling" and "Tryptophan Degradation to 2-amino-3-carboxymuconate Semialdehyde".
Analysis of the IPA knowledge database for upstream regulators of the DE genes identified 3 significant predicted regulators that differed between PP and NN animals, namely LEPR (z-score = 2, activated), IL-12 complex (z-score = -2.410, inhibited) and IL-2 (z-score = -2.950, inhibited). In the NP vs NN comparison one upstream regulator gene, MYCN, was predicted to be activated (z-score 2.219).

Functions and networks associated with DE genes in MAP infected or exposed dairy cows

The DE genes observed in PP or NP cows versus NN, unexposed controls, were investigated for involvement in immunity-related functions. IPA analysis of gene expression profiles revealed networks associated with immune response and lipid metabolism. Gene networks defined for DE genes in the PP vs NN comparison were: "Lipid Metabolism, Small Molecule Biochemistry, Inflammatory Response" (score 45, focus molecules 22), "Inflammatory Disease, Respiratory Disease, Dermatological Diseases and conditions" (score 45, focus molecules 22), "Cell-To-Cell Signaling and Interaction, Cellular Growth and Proliferation, Hematological System Development and Function" (score 26, focus molecules 15), and "Cell Cycle, Digestive System Development and function, Cancer" (score 23, focus molecules 13). DE genes detected in the NP vs NN comparison were associated with several networks having a role in: "Inflammatory Response, Cellular Movement, Respiratory Disease" (score 46, focus molecules 20), "Cell Cycle, Nervous System Development and Function, Lipid Metabolism" (score 24, focus molecules 12), "Cell Signaling, Molecular Transport, Vitamin and Mineral Metabolism" (score 22, focus molecules 11) and "Cancer, Organismal Injury and Abnormalities, Molecular Transport" (score 21, focus molecules 11). The relationships, direct and indirect among the DE genes of PP vs NN and NP vs NN comparisons, are shown in Figs 3 and 4 respectively.

The representative network for the DE genes in the PP vs NN groups showed the relationships among the DE genes, as well as the predicted upstream and downstream effects of activation or inhibition molecules. The network includes ALOX5 (arachidonate 5-lipoxygenase), ALOX15 (arachidonate 15-lipoxygenase), IDO1 (indoleamine 2,3-dioxygenase 1), IL1RL1 (IL-1 receptor-like 1), IL5RA (IL-5 receptor, alpha), SOCS2 (suppressor of cytokine signaling 2) among the down-regulated genes, while IL23R (IL-23 receptor) is one of the up-regulated genes in the network. From the IPA pathway analysis of genes associated with inflammatory response, genes with lower expression in the NP vs NN group, included BHLHE40, CAMK2N1, CFB, Table 6. Top canonical pathways associated with DE genes in PP vs NN and NP vs NN comparisons.

| Ingenuity Canonical Pathways | Molecules | DE gene set |
|-----------------------------|-----------|-------------|
| Eicosanoid Signaling         | ALOX15, LTC4S, ALOX5, CYSLTR2 | PP vs NN |
| Leukotriene Biosynthesis     | LTC4S, ALOX5 | PP vs NN |
| G-Protein Coupled Receptor Signaling | P2RY14, ADORA3, CHRM3, ADRB3, PRKCG | PP vs NN |
| Granulocyte Adhesion and Diapedesis | VCAM1, IL1RL1, MMP14, HRH4 | PP vs NN |
| Leukotriene Biosynthesis     | LTA4H, LTC4S | NP vs NN |
| EIF2 Signaling               | RPS15, RPS23, RPL26, RPL37 | NP vs NN |
| Eicosanoid Signaling         | LTA4H, LTC4S | NP vs NN |
| Tryptophan Degradation to 2-amino-3-carboxymuconate Semialdehyde | IDO1 | NP vs NN |

doi:10.1371/journal.pone.0164461.t006
Fig 3. The most relevant network associated with DE genes of the PP vs NN comparison. Up-regulated (pink) and down-regulated (green) genes involved in lipid metabolism, small molecules biochemistry and Inflammatory response functions. Different shapes indicate different function of gene products: squares for cytokines, horizontal rectangles for ligand-dependent nuclear receptor, vertical rectangles for G-protein coupled receptors, vertical rhombus for enzymes, horizontal rhombus for peptidase, triangles for phosphatase, inverted triangle for kinase, vertical ovals for transmembrane receptors, horizontal ovals for transcription regulators, trapezius for transporters, double circles for complexes and circles for others.

doi:10.1371/journal.pone.0164461.g003

CLNK, ESM1, FUCA1, GZMB, HRH4, IDO1, IGFBP7, IL1RL1, RTF and SULF2, while ACE, CSPG4, DCHS1, LTC4S, NOTCH3, TPT1 showed increased expression.

Discussion

Early diagnosis of MAP is a big challenge for the control and the eradication of MAP from infected herds. The current study integrated the analysis of miRNA and mRNA expression data in order to relate gene expression with disease progression in cattle. Naturally infected adult cows, healthy cows from an infected herd and unexposed cows were compared to assess changes in the transcriptome associated with MAP infection, which may be useful for early diagnosis.
The analysis showed that expression of key inflammation and immune-related genes ALOX5 and ALOX15 was reduced in the PP group when compared with the NN group (S1 Table). The reduced expression of these genes has previously been described in calves 6 months after experimental inoculation with MAP [27]. The production of the enzyme 15-lipoxygenase, encoded by the ALOX15 gene, is induced by IL-4 in monocytes and macrophages during Th2 type response [45]. Mice deficient in 12/15-LOX have increased susceptibility to chronic toxoplasmosis which is associated with reduced production of IL-12 and gamma interferon (INF-γ).
Thus, reduced expression of this gene is associated with a modified immune inflammatory response and may be related to the long term persistence of MAP in infected animals. IPA includes ALOX5 and ALOX15 within a gene network enriched for molecules associated with inflammatory response. This network encompasses several genes found to have lower expression in the PP vs NN group, including IL1RL1, IDO1, IL5RA, SOCS2, while IL23R showed increased expression (Fig 3).

IL23R and SOCS2 are involved in cell-mediated immune response. SOCS2 is a regulator of Th2 response, and homozygous knockout mice have a higher Th2 immune response after helminth antigen stimulation compared with wild-type mice [47]. IL-23 is produced by macrophages and dendritic cells and is a regulator of inflammation and of Th1 response. IL-23 is over expressed in the ileal mucosa of sheep with paratuberculosis [48]. These findings suggest that the enhanced expression of IL-23R in infected animals, stimulates the Th1 immune response. However, the reduced SOCS2 expression, may inhibit responses that are unfavourable to the intracellular pathogens survival, and promote a Th2 response.

IDO1 expression was lower in the PP than NN as well as in the NP vs NN group (S2 Table) and has been found to have increased expression in the ileum, jejunum and ileum draining lymph nodes of MAP infected sheep [49]. IDO expression is known to be induced by IFN-γ [49, 50] and lower expression may be associated with the shift towards a Th2 response, typical of the JD progression. The shift from Th1 to Th2 response would also be stimulated by the decrease of SOCS2 expression in the PP group. IPA includes IDO1 within a gene network enriched for genes associated with inflammatory response in both PP and NP group. This network included additional genes found to have lower expression in the NP vs NN group, such as IL1RL1 (Fig 4).

The reduced expression level of IL1RL1 in the PP vs NN group was accompanied by an increase in IL23R. However, in NP vs NN the expression of IL1RL1 was reduced although the expression of IL23R was unchanged. Increased IL1RL1 expression is associated with the activation of Th2 response [51], while reduced IL1RL1 may maintain the Th1 response. A Th1 response is required for pathogen elimination and IL23R up-regulation occurred only in the positive group indicating a further effort for the pathogen clearance.

Predicted upstream regulators of DE genes in MAP infected or exposed dairy cows compared to control group

The analysis of the regulators of DE genes from the PP vs NN comparison revealed the LEPR trans-membrane receptor was activated in the PP group, while the IL-12 complex and IL-2 regulators were predicted to be inhibited. GZMB, IDO1, KLRC1, KLRD1 and SOCS2 are known to be regulated by the IL-12 complex and had reduced expression in MAP infected animals. In contrast, the MIF gene, which codes for the multifunctional cytokine 'macrophage migration inhibitory factor', and which is known to be down-regulated by the IL-12 complex [52], showed higher expression in the PP vs NN group. IL-12 cytokine has been shown to induce INF-γ production in primary CD4+T cells [53, 54], which is essential for the activation of macrophages and pathogen clearance. Furthermore, it is known that there is a synergic effect of IL-12 and IL-18 in the enhancement of the NK cells cytotoxicity and Th1 cell differentiation [55]. CD4+ Th1 cells are thought to play a key role in the protection against mycobacteria by production of IFN-γ, which activates infected macrophages to kill the intracellular bacteria [56]. The benefit of IFN-γ as anti-mycobacterial has been demonstrated in patients with IL-12 deficiency [57]. The reduction of IL-12 expression observed in the PP group may have led to a reduction of immune response against MAP. These observations support the hypothesis that the regulation of the host IL-12 response may be a way for MAP to prolong its survival in
monocyte cells. Similarly, inhibition of IL-2 regulators was observed in the PP vs NN group. The decline in T-cell function in advanced mycobacterial infections in mice has been associated with impaired production of IL-2 leading to a decline in proliferation of activated T-cells and the loss of IFN-γ [58]. Taken together, these observations suggest the suppression of macrophages immune response and the suppression of T-cells recognition and effector functions, may contribute to the prolonged survival of MAP in macrophages. Indeed, the association of mycobacterial infections with the decline in the cellular immune function of the host is well known [59].

Expression levels of RPS23, RPS15, RPL37, RPL26 and IGFBP7 in NP animals predicted the activation of transcriptional regulator MYCN. MYC prevents the activation of Nramp1 in mouse macrophages through the competition with IRF-8, leading to an impaired killing of intra-phagosomal pathogens [60]. Activation of MYCN in the NP group, which were clinically healthy and in which MAP was not detectable but which had been exposed to pathogen and may have bene incubating disease, indicates that MAP is able to prevent the activation of macrophages in order to ensure its intra-cellular survival.

**miRNAs as potential biomarkers of MAP infection**

Some of the miRNAs found to be DE between the NN and PP groups have been described as having a role in immune function and inflammation following bacterial or viral infection. Expression of mir-100 was at lower levels in the PP vs the NN group in the present study (Table 1), although increased levels of mir-100 have been described in mouse lung after viral infection [38] and even in hemocytes from gram-negative infected shrimps [61]. However, reduced mir-100 expression has been observed in a bovine mammary epithelial cell line after the infection with streptococcus uberis [40], indicating that the miRNA response is associated with gram-positive bacterial infection. Mir-301a is an activator of NF-kB [62] and was under-expressed in PP vs NN groups. A decreased expression of mir-301a has been associated with both viral [38] and mycobacterial infection [63]. Levels of mir-32 were lower in PP vs NN groups in the present study while previous studies on bovine and human pulmonary tuberculosis reported the overexpression of mir-32 in infected alveolar macrophages [64] and CD4+ T cells [65]. Inconsistencies in miRNA expression between the mixed cell populations found in PBMC versus single cell types, specifically CD4+ T cells, has been observed [65], as well as in miRNA signatures from different biological fluids [63]. These differences may reflect different cell functions, or changes in expression levels only in cells that are actively infected with MAP.

Among differentially expressed miRNAs between the PP and NN groups bta-mir-19b, bta-mir-19b-2, bta-mir-1271 and Novel:14_7917 have previously been found to be involved in human tuberculosis and in inflammatory bowel disease. Interestingly, reduced expression of mir-19b and mir-19b-2 has been observed respectively in serum [66] and sputum [63] of tuberculosis affected human vs healthy patients. Furthermore, mir-19b was reported to have low levels of expression in mucosal tissue of Crohn’s affected patients when compared to healthy controls [67]. In the present study exposed animals (NP group) had an increased level of expression of bta-mir-24-1, bta-mir-24-2, bta-mir-378c compared with the NN group (Table 2) and higher variability in the NP group compared to the other two groups. Increased expression of mir-24-2 [68] and mir-378 [69], has been reported in serum of human patients with pulmonary tuberculosis and mir-378 increased expression was also reported in platelets of ulcerative colitis human patients [70] when compared to the healthy controls. In the present study, while mir-24-1, mir-24-2 and mir-378c levels were the same in NN and PP groups of cows, increased levels were observed in the exposed (NP) group. Exposed animals were from a herd with high MAP infection rate and most likely were in contact with the pathogen.
However, the infection status of each NP individual is not known and it is possible that animals were infected at different stages of disease progression. This would explain the high variability of the expression levels observed in the exposed group compared with the other two groups for bta-mir-19b, bta-mir-19b2, bta-mir-301a and bta-mir-32 that were all differentially expressed between the PP vs NN groups. It should also be noted that not all the animals infected with MAP progress to clinical disease, and it is unclear whether some animals are able to develop an immune response that controls or eliminates the pathogen [71]. The differentially expressed miRNAs observed here may be involved in regulating early immune response and may play a role in controlling disease progression.

Target genes and their role in immune function

The expression levels of genes potentially targeted by the differentially expressed miRNAs related to immune function was investigated and the effect of such regulation is summarized in S4 Table. HIC1, TBC1D8 and IMPDH1 were over expressed in the PP group in comparison with the NN control groups and had the same target sequence motif for both bta-mir-19b and bta-mir-19b-2, which had lower expression in the PP group. None of the other differentially expressed genes was targeted by more than one miRNA (Table 5).

Interestingly, ZBTB44 (under expressed in the PP group) and ZBTB4 (over expressed in the NP group) belonged to the same zinc finger and BTB domain-containing family of the BTB-ZF and POK (POZ and Krüppel) families [72]. miRNAs belonging to the human miR-17-92 cluster, including mir-19b, which had lower expression in NP vs NN, have been described as targeting ZBTB4 [73] which indeed had higher expression levels in the NP group. Experimentally infected human macrophages showed lower levels of ZBTB4 in phagosomes infected with an attenuated Mycobacterium avium subsp. hominissuis strain, compared with macrophage infected with the more virulent wild-type mycobacteria [74]. The attenuated strain, unlike the wild-type strain, was not able to replicate in macrophages or prevent phagosome-lysosome fusion. In the present study, the high expression of ZBTB4 in the NP cows may suggest a response of macrophages to the pathogen invasion. ZBTB44, which had lower expression in PP vs NN groups, carries the target motif for bta-mir-7857, which had higher levels in PP vs NN.

These observations indicate that levels of key response genes in the macrophage are being regulated by RNA interference mechanisms and not changes in transcription per se.

TTYH3, which is targeted by bta-mir-1271, had higher expression in both PP and NP groups compared to the unexposed NN group. TTYH3 codes for a member of the tweety family proteins, which are calcium-dependent chloride channels that allow the passage of the chloride anion through the cell membrane for the stabilization of the membrane potential, transport, and cell volume regulation. Anionic channels play a key role in the maintenance of peripheral blood mononuclear cell functions, in particular those connected with innate immune response: NADPH oxidase activity, phagocytosis, and trans-endothelial migration [75]. Increased expression of TTYH2, a parologue of TTYH3, has been observed in the whole blood of Holstein cows infected with Mycobacterium avium subsp. paratuberculosis [76]. The results presented here confirm the increased expression levels of this gene family in PP animals compared with the NN group, suggesting the involvement of TTYH3 in the response to MAP challenge.

The serologically positive (PP) animals were from the same herd as the exposed serologically negative (NP) animals of the same age and sex. Patterns of miRNA and mRNA expression observed between the two groups suggests that the status was different, although expression of bta-mir-19b and bta-mir-19b2 differed in both NP and PP groups in comparison with the NN control group (Table 3), suggesting that the exposed group was responding, possibly to latent MAP infection. Interestingly different target genes of the miRNAs were differentially expressed.
in the PP vs NN or in the NP vs NN groups. One of the genes targeted by both bta-mir-19b and bta-mir-19b2 is HIC1 (Hypermethylated In Cancer 1), which showed an increased expression only in the PP group which is consistent with the decrease in the miRNAs levels. HIC1 controls the expression of toll-like receptor 2, which enhances the NF-kB related axis [77] and pro-inflammatory cytokine secretion. NF-kB activation, and an enhanced production of inflammatory mediators, has been observed in macrophages and other cells types after Mycobacterium tuberculosis infection, as well as in intestinal epithelial cells after mycobacteria challenge [78]. This suggests that the host immune response to the infection is being carefully regulated in order to avoid the uncontrolled release of inflammatory mediators.

Mir-19a, has been shown to target IMPDH1, decreasing the related protein expression level in breast cancer cells [79]. The target prediction analysis carried out here also suggests that bta-mir-19b and bta-mir-19b2, which had lower levels in PP vs NN animals, target IMPDH1, which is consistent with the increased IMPDH1 level in PP vs NN groups. Interestingly, although the level of mir-19 was also lower in the NP vs NN animals, the expression of IMPDH1 was not appreciably higher in NP vs NN groups. IMPDH1 codes for an inosine 5’-monophosphate dehydrogenase enzyme, which is essential for purine synthesis in lymphocytes, catalyzing the conversion of IMP to GMP [80]. IMPDH inhibition has anti-proliferative effects in human monocytes in vitro [81], and is associated with reduced pro-inflammatory cytokines, nitric oxide and lactate dehydrogenase production in a murine macrophage cell line [82]. It has also been shown that IMPDH inhibitors can reduce IFN-γ mRNA levels after allogeneic stimulation in mice [83]. Thus IMPDH activity seems to be essential to the normal proliferation, pro-inflammatory functions and activation of macrophages. The higher expression levels detected in PP animals vs NN controls, suggest the activation of cellular immune response upon MAP infection.

The present study identified a novel miRNA, Novel:14_7917, the expression of which was lower in PP vs NN animals. Predicted targets for this miRNA included FAM78A, ARL2 and AP2A1, which is consistent with the higher expression seen in PP vs NN. AP2A1 and ARL2 are involved in the movement of vesicles through the cellular membrane. AP2A1 is involved in endocytosis [84–86], while ARL2 plays a role in both the regulation of tubulin folding and microtubule remodeling [87], which is essential for phagosome maturation [88]. Thus, the enhanced expression of these genes observed in the PP group may be associated with increased intracellular traffic due to phagocytosis related to the uptake of MAP.

The transcriptome and miRNA analysis indicated that mechanisms of innate and acquired immunity were affected, which may influence the status of the infecting mycobacterium. These data suggest a balance between increased and suppressed immune mechanisms, reflecting opposing action of the host response to clear infection and the strategy used by the pathogen to ensure survival within the host cells. The miRNA profiles and levels of predicted gene targets were consistent, that is increased miRNA was associated with a decrease in target genes and vice versa, suggesting that expression of these genes was regulated at the level of RNA interference not at the level of transcription. The status of animals in the NP group was not known however the analysis suggested that animals in this group were responding to the pathogen and may have been at an earlier stage of infection. In particular, the NP group showed reduced levels of bta-mir-19b, bta-mir-19b-2 and bta-mir-1271 and an increased expression of target genes which was observed in the PP vs NN groups. This pattern could indicate that signs of infection can be detected prior to sero-conversion and hence prior to the current ELISA diagnostic test. However, more work will be necessary to confirm this and test the specificity of the response. In conclusion, examining the transcriptome and miRNA profiles can focus analyses on specific pathways, in this case identifying host pathogen interactions. This facilitated the
identification of miRNA and gene candidates for diagnostic tools to detect animals infected with MAP at an early stage of disease than the current ELISA test.

**Methods**

**Animal resource**

Holstein dairy cows were selected from two herds, one MAP free (verified by ELISA and absence of clinical cases) and the other positive for MAP (verified by ELISA, fecal culture and presence of clinical cases). From the positive herd, 5 positive (PP) cows based on ELISA test results and 5 ELISA negative, potentially exposed (NP) animals were chosen. All animals were 4 to 5 years old, had a body condition score (BCS) of 3 and were at 170 to 190 days in milk (DIM) when sampled. Positive subjects were confirmed by fecal culture. Five negative non-exposed control animals (NN) were selected from the negative herd, matched for age, lactation, BCS and DIM with the positive and exposed animals. For the miRNA-Seq experiment, in addition to these animals, 7 negative cows from the same negative herd were added to enlarge the control group.

**Sample preparation, RNA extraction and quality control**

Samples used were taken during obligatory routine animal sanitary controls by an authorized veterinarian. No ethical approval was required, in compliance with the European Directive 2010/63/UE and the Italian regulation D. Lgs n. 26/2014. Whole blood was collected into PAXgene® Blood RNA Tubes (PreAnalytiX GmbH) and kept at room temperature for at least two hours before freezing at -20°C for 24 hours before being transferred at -80°C until processing, as recommended by the manufacturer. Total RNA was extracted using the PAXgene® Blood miRNA Kit (PreAnalytiX GmbH) according to the manufacturer's protocol. Total RNA was eluted in a final volume of 80 μL. RNA concentration was measured by NanoDrop™ 1000 spectrophotometer (Thermo Scientific) and RNA integrity was assessed with an Agilent 2200 TapeStation system (Agilent Technologies).

**RNA-Seq library preparation and sequencing**

Libraries were prepared with the Illumina Truseq RNA sample prep kit (Illumina Inc., USA) following manufacturer's protocol and the size and yield evaluated using an Agilent TapeStation 2200. Libraries were then quantified with an ABI9700 qPCR instrument using the KAPA Library Quantification Kit in triplicate, according to the manufacturer's protocol (Kapa Biosystems, Woburn, MA, USA) and then normalized to 10 nM as recommended by Illumina for cluster generation on the Hiseq2000. Fifteen libraries were prepared and equimolar amounts of 5 samples were mixed before NaOH denaturation. Each of the pools was run in a lane of a Hiseq Flowcell. The Illumina Truseq PE cluster kit v3 was used to generate clusters on the grafted Illumina Flowcell and the hybridized libraries were sequenced on a Hiseq2000 with a 100 cycles of paired-end sequencing module using the Truseq SBS kit v3 (Illumina Inc., USA).

**miRNA-Seq library preparation and sequencing**

For each sample, 5 μl of RNA were used to prepare a library with the TruSeq SmallRNA kit (Illumina Inc., USA) following the manufacturer's instructions. In order to minimize primer dimers formation, half of the TruSeq Small RNA sample reagents were used, followed by 11 PCR cycles of PCR to amplify the library. 10 μl of unique indexed libraries were pooled and DNA fragments from 140 to 160 bp (the length of miRNA inserts plus the 3’ and 5’ adaptors)
were selected from a Pippin 3% Agarose Dye free Gel cassette (BluPippin, Sage Science, MA, USA), which were then recovered in 40 μL of Pippin elution buffer. Fragments were purified by Qiagen MinElute PCR Purification kit (Qiagen, CA, USA). The indexed libraries were quantified as described above. Two pools of 11 sample libraries were prepared and 10 μL of each pool at a final concentration of 2 nM were used in a lane for 50bp Single-Read sequencing for a total of 2 lanes of an Illumina HiSeq2000.

RNA-Seq data analysis
Preliminary quality control of raw reads was carried out with FastQC software v0.11.2 [89]. Illumina raw sequences were trimmed using Trimmomatic [90] and PCR primers and Illumina adapter sequences were removed. Minimum base quality 15 over a 4 base sliding window was required, then only sequences longer than 36 nucleotides were included in the downstream analysis. Reads which successfully passed trimming were mapped against the Bos taurus UMD3.1.68 reference genome sequence, using STAR [91] aligner to obtain BAM alignment files. The BAM files were sorted and indexed using Samtools [92]. In order to quantify counts for each sample a list of genes and relative abundance of mapping reads were extracted using htsq-count [93]. These count files were used for downstream statistical analysis.

miRNAseq data analysis
Trimming and quality control were performed as described for RNA-Seq analysis, with the difference that a 15 nucleotide minimum sequence length was required. Reads which passed the quality control were used for novel small-RNA discovery using Mirdeep2 [43]. Both “cow” and “human” known small-RNAs (mature and precursors) were downloaded from MirBase [94] and used as support datasets to help the discovery process. Absolute positions of novel miRNAs on the bovine reference genome were retrieved by BLAST [95]. Counts of all the known and the novel miRNAs were used to quantify expression levels for each sample using Mirdeep2. This pipeline produced a list of small-RNA IDs and the relative abundance of mapping reads (counts) for each sample which was used in the downstream statistical analysis.

Expression Analysis of mRNAs and miRNAs
Statistical analyses to compare mRNA and miRNA expression profiles were performed using the "R" statistical environment edgeR [96], vegan [97] and gplots [98] packages. Genes that had at least one count per million in at least 3 samples were included in the gene expression analysis. A general linear model was used in the edge R Package to generate lists of mRNAs and miRNAs with statistically significant different expression among the three comparisons: PP vs NN, NP vs NN and PP vs NP. Differentially expressed mRNAs and miRNAs were defined as having a False Discovery Rate (FDR) below 0.05.

Real-time RT-qPCR for RNA-Seq validation
To validate the RNA-Seq gene expression data RT-qPCR was performed on 4 genes selected from those which were differentially expressed in both PP vs NN and NP vs NN comparisons and had a FDR<0.01. Three hundred ng of total RNA were reverse transcribed into cDNA using the QuantiTect Reverse Transcription Kit (Qiagen), RT-qPCR was performed using 5 ng cDNA, and 250 nM each primer in GoTaq qPCR Master Mix (Promega). The reaction mixtures were incubated in 384-wells plates at 95°C for 10 min, followed by 45 cycles of 95°C for 15 seconds and 62°C for 1 min using a CFX 384 Real-Time PCR Detection System (Bio-rad...
Laboratories, USA). All reactions were performed in triplicate and “no-template” controls were included. Following amplification, a melting curve analysis was performed to verify the specificity of the reactions. GAPDH and ACTB were chosen as the reference genes, as they were stably expressed in the RNAseq data, and assayed in the same samples to normalize the data. In order to determine the efficiency and the dynamic range of the reaction, for each primer pair a standard curve was constructed from triplicate assays for 4 dilutions from 30 ng to 0.03 ng of pooled cDNA samples. Primers for Real-time RT-qPCR are listed in Table 7 and were designed using Primer-BLAST (NCBI) or deduced by the literature. The relative expression level of each selected gene was calculated according to the Pfaffl method [99] and was reported as normalized fold expression relative to the control (NN cows). Log$_2$ fold-change data of RT-qPCR and RNA-Seq analyses were compared to validate results. The Pearson correlation coefficient between the two analyses was calculated using IBM Statistical Package for Social Sciences software (SPSS, ver. 21; IMB SPSS Inc., Chicago, IL, USA) and differences were considered as statistically significant if the p-Value was <0.05.

**Functional Analysis of gene expression data**

The RNA-Seq differentially expressed (DE) gene lists obtained for each comparison (PP vs NN, NP vs NN, PP vs NP) were submitted to the Qiagen Ingenuity Pathway Analysis (IPA; Ingenuity Systems Inc., USA). The DE genes were filtered for an FDR ≤0.05 and a fold change ≥1 or ≤-1 and were used as input in three separate data sets. Each gene was mapped to its corresponding gene object in the Ingenuity Knowledge Base. In the Core Analysis, IPA assigns functional information and biological relevance by analyzing RNA expression data in the context of known biological response and regulatory networks. In addition, canonical pathways, biological functions, and networks were investigated to identify major biological pathways associated with MAP infection. The upstream regulators were investigated based on an activation z-score, which was considered as significant when below -2 (inhibited) or above 2 (activated).

**Prediction of miRNA target genes and correlation with mRNA data**

Due to the uncertainty associated with the prediction of miRNA target genes [40], two approaches were used. For known miRNAs only the gene targets identified by both methods were considered in the subsequent analysis. miRDB software [101] was used with default
threshold settings for the initial prediction, based on miRNA-target matching the miRNA seed region and 3’ UTR mRNAs. The method was applied for both known and novel differentially expressed miRNAs and a threshold of 90 was set to select a panel of potential target genes. Predicted targets of known miRNAs were then analyzed with the TargetScan software v7.0 [102]. RNA-Seq expression levels were investigated for the genes that were shared between the two lists. Predicted gene targets of the novel miRNAs were obtained only from the miRDB prediction software, because target identification was done directly from the novel miRNA sequences. Targets which in RNA-Seq results had FDR < 0.05 and opposite pattern of expression with respect to related miRNAs were considered as putatively affected by miRNA expression.

Supporting Information

S1 Table. Differentially expressed genes in the PP subjects when compared with the NN group. Genes with an FDR < 0.05 and a Log₂ fold change (Log₂FC) < -1 or > 1 were considered as differentially expressed.

(SDOCX)

S2 Table. Differentially expressed genes in the NP subjects when compared with the NN group. Genes with an FDR < 0.05 and a Log₂ fold change (Log₂FC) < -1 or > 1 were considered as differentially expressed.

(SDOCX)

S3 Table. Differentially expressed genes in the PP subjects when compared with the NP group. Genes with an FDR < 0.05 and a Log₂ fold change (Log₂FC) < -1 or > 1 were considered as differentially expressed.

(SDOCX)

S4 Table. Effects of gene regulation by miRNAs on the immune functions.

(SDOCX)

Author Contributions

Conceptualization: JLW GM.
Formal analysis: MM BL.
Funding acquisition: JLW.
Investigation: MM FP.
Methodology: JLW GM.
Project administration: GM.
Resources: FP NM.
Supervision: JLW GM.
Validation: MM.
Visualization: MM GM.
Writing – original draft: MM GP GM.
Writing – review & editing: MM JLW GP GM.
References

1. Olsen I, Sigurjónsdóttir G, Djenne B. Paratuberculosis with special reference to cattle. Vet Q. 2002; 24: 12–28. doi: 10.1080/01652176.2002.9695120 PMID: 11918176

2. Whittington RJ, Sergeant ESG. Progress towards understanding the spread, detection and control of Mycobacterium avium subsp. paratuberculosis in animal populations. Aust Vet J. 2001; 79: 267–278. PMID: 1149414

3. Greig A, Stevenson K, Henderson D, Perez V, Hughes V, Pavlik I, et al. Epidemiological study of paratuberculosis in wild rabbits in Scotland. J Clin Microbiol. 1999; 37: 1746–1751. PMID: 10325318

4. Beard PM, Daniels MJ, Henderson D, Pirie A, Rudge K, Buxton D, et al. Paratuberculosis infection of nonruminant wildlife in Scotland. J Clin Microbiol. 2001; 39: 1517–1521. doi: 10.1128/JCM.39.4.1517-1521.2001 PMID: 11283080

5. McClure HM, Chiodini RJ, Anderson DC, Swenson RB, Thayer WR, Coutu JA. Mycobacterium paratuberculosis infection in a colony of stump-tailed macaques (Macaca arctoides). J Infect Dis. 1987; 155: 1011–1019. PMID: 3559275

6. Zwick LS, Walsh TF, Barbiers R, Collins MT, Kinsella MJ. Paratuberculosis in a mandrill (Papio sphinx). J Vet Diagn Invest. 2002; 14: 326–328.

7. Garcia AB, Shalloo L. Invited review: The economic impact and control of paratuberculosis in cattle. J Dairy Sci. 2015; 98: 5019–5039. doi: 10.3168/jds.2014-9241 PMID: 26074241

8. Bull TJ, McMinn EJ, Sidi-Boumedine K, Skull A, Durkin D, Neil P, et al. Detection and verification of Mycobacterium avium subsp. paratuberculosis in fresh ileocolonic mucosal biopsy specimens from individuals with and without Crohn’s disease. J Clin Microbiol. 2003; 41: 2915–2923. doi: 10.1128/JCM.41.7.2915-2923.2003 PMID: 12843021

9. Ghadiali AH, Strother M, Naser SA, Manning EJ, Sreevatsan S. Mycobacterium avium subsp. paratuberculosis strains isolated from Crohn’s disease patients and animal species exhibit similar polymorphic locus patterns. J Clin Microbiol. 2004; 42: 5345–5348. doi: 10.1128/JCM.42.11.5345-5348.2004 PMID: 15528739

10. Greenstein RJ. Is Crohn’s disease caused by a Mycobacterium? Comparisons with leprosy, tuberculosis, and Johne’s disease. Lancet Infect Dis. 2003; 3: 507–514. PMID: 12901893

11. Mendoza JL, Lana R, Díaz-Rubio M. Mycobacterium avium subspecies paratuberculosis and its relationship with Crohn’s disease. World J Gastroenterol. 2009; 15: 417–422. PMID: 19152445

12. Khare S, Nunes JS, Figueiredo JF, Lawhorn SD, Rossetti CA, Gull T, et al. Early phase morphological lesions and transcriptional responses of bovine ileum infected with Mycobacterium avium subsp. paratuberculosis. Vet Pathol. 2009; 46: 717–728. doi: 10.1354/vp.08-VP-0187-G-FL PMID: 19276052

13. Gilardoni LR, Paolicchi FA, Mundo SL. Bovine paratuberculosis: a review of the advantages and disadvantages of different diagnostic tests. Rev Argent Microbiol. 2012; 44: 201–215. PMID: 23102470

14. Crossley BM, Zagmutt-Vergara FJ, Frycock TL, Whitlock RH, Gardner IA. Fecal shedding of Mycobacterium avium subsp. paratuberculosis by dairy cows. Vet Microbiol. 2005; 107: 257–263. doi: 10.1016/j.vetmic.2005.01.017 PMID: 15863285

15. Roussey JA, Steibl J, Coussemens PM. Regulatory T cell activity and signs of T cell unresponsiveness in bovine paratuberculosis. Front Vet Sci. 2014; 1: 20. doi: 10.3389/fvets.2014.00020 PMID: 26664919

16. Mitchell RM, Medley GF, Collins MT, Schukken YH. A meta-analysis of the effect of dose and age at exposure on shedding of Mycobacterium avium subspecies paratuberculosis (MAP) in experimentally infected calves and cows. Epidemiol Infect. 2012; 140: 231–246. doi: 10.1017/S0950268811000689 PMID: 21524342

17. Whittington RH, Buergelt C. Preclinical and clinical manifestations of paratuberculosis (including pathology). Vet Clin North Am Food Anim Pract. 1996; 12: 345–356. PMID: 8828109

18. Wells SJ, Collins MT, Faaberg KS, Wees C, Tavornpanich S, Petruini KR, et al. Evaluation of a rapid fecal PCR test for detection of Mycobacterium avium subsp. paratuberculosis in dairy cattle. Clin Vaccine Immunol. 2006; 13: 1125–1130.

19. Clark DL Jr, Koziczkowski JJ, Radcliff RP, Carlson RA, Ellingon J. Detection of Mycobacterium avium subspecies paratuberculosis: comparing fecal culture versus serum enzyme-linked immunosorbent assay and direct fecal polymerase chain reaction. J Dairy Sci. 2008; 91: 2620–2627. doi: 10.3168/jds.2007-0902 PMID: 18565921

20. Collins MT. Diagnosis of paratuberculosis. Vet Clin North Am Food Anim Pract. 1996; 12: 357–371. PMID: 8828110
21. Sockett DC, Thomas CB. Evaluation of four serological tests for bovine paratuberculosis. J. Clin. Microbiol. 1992; 30: 1134–1139. PMID: 1583110
22. Hines ME, Turmquist SE, Ilha MR, Rajeef S, Jones AL, Whittington L, et al. Evaluation of novel oral vaccine candidates and validation of a caprine model of Johnes’s disease. Front Cell Infect Microbiol. 2014; 4: 26. doi: 10.3389/fcimb.2014.00026 PMID: 24624365
23. Pinedo PJ, Buergelt CD, Donovan GA, Melendez P, Morel L, Wu R, et al. Candidate gene polymorphisms (BoIFNG, TLR4, SLC11A1) as risk factors for paratuberculosis infection in cattle. Prev Vet Med. 2009; 91: 189–196. doi: 10.1016/j.prevetmed.2009.05.020 PMID: 1952022
24. Minozzi G, Buggiotti L, Stella A, Strozzi F, Luini M, Williams JL. Genetic loci involved in antibody response to Mycobacterium avium ssp. paratuberculosis in cattle. PLoS One. 2010; 5: e11117. doi: 10.1371/journal.pone.0011117 PMID: 20559561
25. Settles M, Zanella R, McKay SD, Schnabel RD, Taylor JF, Whitlock R, et al. A whole genome association analysis identifies loci associated with Mycobacterium avium subsp. paratuberculosis infection status in US holstein cattle. Anim Genet. 2009; 40: 655–662. doi: 10.1111/j.1365-2052.2009.01896.x PMID: 19422364
26. David J, Barkema HW, Mortier R, Ghosh S, le Guan L, De Buck J. Gene expression profiling and putative biomarkers of calves 3 months after infection with Mycobacterium avium subspecies paratuberculosis. Vet Immunol Immunopathol. 2014; 160: 107–117. doi: 10.1016/j.vetimm.2014.04.006 PMID: 24841487
27. David J, Barkema HW, le Guan L, De Buck J. Gene expression profiling of calves 6 and 9 months after inoculation with Mycobacterium avium subspecies paratuberculosis. Vet Res. 2014; 45: 96. doi: 10.11186/s13567-014-0096-5 PMID: 25294045
28. Huntzinger E, Izaurralde E. Gene silencing by microRNAs: contributions of translational repression and mRNA decay. Nat Rev Genet. 2011; 12: 99–110. doi: 10.1038/nrg2936 PMID: 21245828
29. Bartel DP. MicroRNAs: target recognition and regulatory functions. Cell. 2009; 136: 215–233. doi: 10.1016/j.cell.2009.01.002 PMID: 19167326
30. Turchinovich A, Weiz L, Langheinz A, Burwinkel B. Characterization of extracellular circulating microRNA. Nucleic Acids Res. 2011; 39: 7223–7233. doi: 10.1093/nar/gkr254 PMID: 21609964
31. Konishi H, Ichikawa D, Arita T, Otsuji E. Microarray Technology and Its Applications for Detecting Plasma microRNA Biomarkers in Digestive Tract Cancers. Methods Mol Biol. 2016; 1368: 99–109. doi: 10.1007/978-1-4939-3136-1_8 PMID: 26614071
32. De Rosa S, Indolfi C. Circulating microRNAs as Biomarkers in Cardiovascular Diseases. EXS 2015; 106: 139–149. doi: 10.1007/978-3-0348-0955-9_6 PMID: 26608202
33. Heegaard NH, Carlsen AL, Skovgaard K, Heegaard PM. Circulating Extracellular microRNA in Systemic Autoimmunity. EXS 2015; 106: 171–195. doi: 10.1007/978-3-0348-0955-9_8 PMID: 26608204
34. Grasso M, Piscopo P, Crestini A, Confalonieri A, Denti MA. Circulating microRNAs in Neurodegenerative Diseases. EXS. 2015; 106: 151–169. doi: 10.1007/978-3-0348-0955-9_7 PMID: 26608203
35. Farr RJ, Joglekar MV, Hardikar AA. Circulating microRNAs in Diabetes Progression: Discovery, Validation, and Research Translation. EXS. 2015; 106: 215–244. doi: 10.1007/978-3-0348-0955-9_10 PMID: 26608206
36. Ueberbürger B, Kohns M, Mayatepek E, Jacobse M. Are microRNAs suitable biomarkers of immunity to tuberculosis? Mol Cell Pediatr. 2014; 1: 8. doi: 10.1186/s40348-014-0008-9 PMID: 26567102
37. Choi EJ, Kim HB, Baek YH, Kim EH, Pascua PN, Park SJ, et al. Differential microRNAs expression following infection with a mouse-adapted, highly virulent avian H5N2 virus. BMC Microbiol. 2014; 14: 252. doi: 10.1186/s12866-014-0252-0 PMID: 25266911
38. Li X, Zhu L, Liu X, Sun X, Zhou Y, Lang Q, et al. Differential expression of microRNAs in porcine parvovirus infected porcine cell line. Virol J. 2015; 12: 128. doi: 10.1186/s12985-015-0339-4 PMID: 26290078
39. Lawless N, Foroushani AB, McCabe MS, O’Farrelly C, Lynn DJ. Next generation sequencing reveals the expression of a unique miRNA profile in response to a gram-positive bacterial infection. PLoS One 2013; 8: e57543. doi: 10.1371/journal.pone.0057543 PMID: 23472090
40. Staedel C, Darfeuille F. MicroRNAs and bacterial infection. Cell Microbiol. 2013; 15: 1496–1507. doi: 10.1111/cmi.12159 PMID: 23795564
41. Farrell D, Shaughnessy RG, Britton L, MacHugh DE, Markey B, Gordon SV. The Identification of Circulating miRNA in Bovine Serum and Their Potential as Novel Biomarkers of Early Mycobacterium
avium subsp. paratuberculosis Infection. PLoS One 2015; 10: e0134310. doi: 10.1371/journal.pone.0134310 PMID: 26218736

43. Friedländer MR, Mackowiak SD, Li N, Chen W, Rajewsky N. miRDeep2 accurately identifies known and hundreds of novel microRNA genes in seven animal clades. Nucleic Acids Res. 2012; 40: 37–52 doi: 10.1093/nar/gkr688 PMID: 21911355

44. Ingenuity Knowledge Base—Qiagen. Available: http://www.ingenuity.com/science/knowledge-base

45. Conrad DJ, Kuhn H, Mulkins M, Highland E, Sigal E. Specific inflammatory cytokines regulate the expression of human monocye 15-lipoxygenase. Proc Natl Acad Sci U S A. 1992; 89: 217–221. PMID: 1729692

46. Middleton MK, Zukas AM, Rubinstein T, Kinder M, Wilson EH, Zhu P, et al. 12/15-lipoxygenase-dependent myeloid production of interleukin-12 is essential for resistance to chronic toxoplasmosis. Infect Immun. 2009; 77: 5690–5700. doi: 10.1128/IAI.00560-09 PMID: 19822654

47. Knosp CA, Carroll HP, Elliott J, Saunders SP, Nel HJ, Amu S, et al. SOCS2 regulates T helper type 2 differentiation and the generation of type 2 allergic responses. J Exp Med. 2011; 208: 1523–1531. doi: 10.1084/jem.20101167 PMID: 21646394

48. Gossner AG, Venturina VM, Peers A, Watkins CA, Hopkins J. Expression of sheep interleukin 23 (IL23A, alpha subunit p19) in two distinct gastrointestinal diseases. Vet Immunol Immunopathol. 2012; 150: 118–122.

49. Plain KM, de Silva K, Earl J, Begg DJ, Purdie AC, Whittington RJ. Indoleamine 2,3-dioxygenase, tryptophan catabolism, and Mycobacterium avium subsp. paratuberculosis: a model for chronic mycobacterial infections. Infect Immun. 2011; 79: 3821–3832. doi: 10.1128/IAI.05204-11 PMID: 21730087

50. Taylor MW, Feng GS. Relationship between interferon-gamma, indoleamine 2,3-dioxygenase, and tryptophan catabolism. FASEB J. 1991; 5: 2516–2522. PMID: 1907934

51. Joshi AD, Oak SR, Hartigan AJ, Finn WG, Kunkel SL, Duffy KE, et al. Interleukin-33 contributes to both M1 and M2 chemokine marker expression in human macrophages. BMC Immunol. 2010; 11: 52. doi: 10.1186/1471-2172-11-52 PMID: 20958987

52. Chaiyaroj SC, Rutta AS, Muenthaisong K, Watkins P, Na Ubol M, Looaarueswan S. Reduced levels of transforming growth factor-beta1, interleukin-12 and increased migration inhibitory factor are associated with severe malaria. Acta Trop. 2004; 89: 319–327. PMID: 14745458

53. Barbulescu K, Becker C, Schlaak JF, Schmitt E, Meyer zum Büschenfelde KH, Neurath MF. IL-12 and IL-18 differentially regulate the transcriptional activity of the human IFN-gamma promoter in primary CD4+ T lymphocytes. J Immunol. 1998; 160: 3642–3647. PMID: 9558063

54. Okamura H, Tsutsi H, Komatsu T, Yutsudo M, Hakura A, Tanimoto T, et al. Cloning of a new cytokine that induces IFN-gamma production by T cells. Nature. 1995; 378: 88–91. doi: 10.1038/378088a0 PMID: 7477296

55. Flynn JL, Chan J, Triebold KJ, Dalton DK, Stewart TA, Bloom BR. An essential role for interferon gamma in resistance to Mycobacterium tuberculosis infection. J Exp Med. 1993; 178: 2249–2254. PMID: 7504064

56. Alten-Koltunoff M, Goren S, Nousbeck J, Fong CG, Sher A, Ozato K, et al. Innate immunity to intraphagosomal pathogens is mediated by interferon regulatory factor 8 (IRF-8) that stimulates the expression of macrophage-specific Nramp1 through antagonizing repression by c-Myc. J Biol Chem. 2008; 283: 2724–2733. doi: 10.1074/jbc.M707704200 PMID: 18045875

57. Zhu F, Wang Z, Sun B-Z. Differential expression of microRNAs in shrimp Marsupenaeus japonicus in response to Vibrio alginolyticus infection. Dev Comp Immunol. 2015; 55: 76–79. doi: 10.1016/j.dci.2015.10.012 PMID: 26483347

58. Ma X, Becker Buscaglia LE, Barker JR, Li Y. MicroRNAs in NF-κB signaling. J Mol Cell Biol. 2011; 3: 159–166. doi: 10.1093/jmcb/mjr007 PMID: 21502305

59. Yi Z, Fu Y, Ji R, Li R, Guan Z. Altered microRNA signatures in sputum of patients with active pulmonary tuberculosis. PLoS One 2012; 7: e43184. doi: 10.1371/journal.pone.0043184 PMID: 22900099

Transcriptomic Profiling in Bovine Paratuberculosis
64. Vegh P, Magee DA, Naipas NC, Bryan K, McCabe MS, Browne JA, et al. MicroRNA profiling of the bovine alveolar macrophage response to Mycobacterium bovis infection suggests pathogen survival is enhanced by microRNA regulation of endocytosis and lysosome trafficking. Tuberculosis. 2015; 95: 60–67. PMID: 25692199

65. Fu Y, Yi Z, Li J, Li R. Deregulated microRNAs in CD4+ T cells from individuals with latent tuberculosis versus active tuberculosis. J Cell Mol Med. 2014; 18: 503–513. doi: 10.1111/jcmm.12205 PMID: 24373112

66. Xu Z, Zhou A, Ni J, Zhang Q, Wang Y, Lu J, et al. Differential expression of miRNAs and their relation to active Tuberculosis Tuberculosis 2015; 95: 395–403. doi: 10.1016/j.tube.2015.02.043 PMID: 25936536

67. Wu F, Zhang S, Dassopoulos T, Harris ML, Bayless TM, Meltzer SJ, et al. Identification of microRNAs associated with ileal and colonic Crohn’s disease. Inflamm Bowel Dis. 2010; 16: 1729–1738. doi: 10.1002/ibd.21267 PMID: 20848482

68. Liu Y, Jiang J, Cao Z, Yang B, Cheng X. Modulation of T cell cytokine production by miR-144* with elevated expression in patients with pulmonary tuberculosis. Mol Immunol. 2011; 48: 1084–1090. doi: 10.1016/j.molimm.2011.02.001 PMID: 21367459

69. Zhang X, Guo J, Fan S, Li Y, Wei L, Yang X, et al. Screening and Identification of Six Serum micro-RNAs as Novel Potential Combination Biomarkers for Pulmonary Tuberculosis Diagnosis. PLoS ONE. 2013; 8: e61076. doi: 10.1371/journal.pone.0081076 PMID: 24349033

70. Duttagupta R, DiRenzo S, Jiang R, Bowers J, Gollub J, Kao J, et al. Genome-wide maps of circulating microRNA biomarkers for ulcerative colitis. PLoS One. 2012; 7: e31241. doi: 10.1371/journal.pone.0031241 PMID: 22359580

71. Rideout BA, Brown ST, Davis WC, Gay JM, Giannella RA, Hines ME, et al. Diagnosis and Control of Johne’s Disease. Committee on Diagnosis and Control of Johne’s Disease. Washington: National Academy Press; 2003.

72. Costoya J. Functional analysis of the role of POK transcriptional repressors. Brief Funct Genomic Proteomic. 2007; 6: 8–18. doi: 10.1093/bfgp/elm002 PMID: 17384421

73. Kim K, Chadalapaka G, Pathi SS, Jin UH, Lee JS, Park YY, et al. Induction of the Transcriptional Repressor ZBTB4 in Prostate Cancer Cells by Drug-induced Targeting of microRNA-17-92/106b-25 Clusters. Mol Cancer Ther. 2012; 11: 1852–1862. doi: 10.1158/1535-7163.MCT-12-0181 PMID: 22752225

74. Jha SS, Danelishvili L, Maser J, Li YJ, Moric I, Vogt S, et al. Virulence-related Mycobacterium avium subsp hominisuis MAV_2928 gene is associated with vacuole remodeling in macrophages. BMC Microbiol. 2010; 1: 100.

75. Moreland JG, Davis AP, Bailey G, Nauseef WM, Lamb FS. Anion channels, including ClC-3, are required for normal neutrophil oxidative function, phagocytosis, and transendothelial migration. J Biol Chem. 2006; 281: 12277–12288. doi: 10.1074/jbc.M511030200 PMID: 16522634

76. Shin M, Park H, Shin SW, Jung M, Im YB, Park H, et al. Whole-Blood Gene-Expression Profiles of Cows Infected with Mycobacterium avium subsp. paratuberculosis Reveal Changes in Immune Response and Lipid Metabolism. J. Microbiol. Biotechnol. 2015; 25: 255–267. PMID: 25248984

77. Janeckova L, Pospichalova V, Fafilek B, Vojtechova M, Tureckova J, Dobes J, et al. HIC1 Tumor Suppressor Loss Potentiates TLR2/NF-κB Signaling and Promotes Tissue Damage-Associated Tumorigenesis. Mol Cancer Res. 2015; 13: 1139–1148. doi: 10.1158/1535-7186.MCR-15-0033 PMID: 25934696

78. Tripathi P, Aggarwal A. NF-κB transcription factor: a key player in the generation of immune response. Curr Sci. 2006; 90: 519–531.

79. Ouchida M, Kanzaki H, Ito S, Hanafusa H, Jitsumori Y, Tamaru S, et al. Novel Direct Targets of miR-19a Identified in Breast Cancer Cells by a Quantitative Proteomic Approach. PLoS One 2012; 7: e44095. doi: 10.1371/journal.pone.0044095 PMID: 22952885

80. Gu JJ, Tolin AK, Jain J, Huang H, Santiago L, Mitchell BS. Targeted Disruption of the Inosine 5′-Monophosphate Dehydrogenase Type I Gene in Mice. Mol Cell Biol. 2003; 23: 6702–6712. doi: 10.1128/MCB.23.18.6702-6712.2003 PMID: 12944494

81. Waters RV, Webster D, Allison A. Mycophenolic acid and some antioxidants induce differentiation of monocytic lineage cells and augment production of the IL-1 receptor antagonist, Ann N.Y. Acad Sci. 1993; 696: 185–196. PMID: 8109827

82. Jonsson CA, Carlsten H. Mycophenolic acid inhibits inosine 5-monophosphate dehydrogenase and suppresses production of pro-inflammatory cytokines, nitric oxide, and LDH in macrophages. Cellular Immunol 2002; 216: 93–101.
83. Lui SL, Ramassar V, Urmson J, Halloran PF. Mycophenolate mofetil reduces production of interferon-dependent major histocompatibility complex induction during allograft rejection, probably by limiting clonal expansion. Transplant Immunol. 1998; 6: 23–32.

84. Hussain KM, Leong KLJ, Ng MM-L, Chu JJH. The Essential Role of Clathrin-mediated Endocytosis in the Infectious Entry of Human Enterovirus 71. J Biol Chem. 2011; 286: 309–321. doi: 10.1074/jbc.M110.168468 PMID: 20956521

85. Traub LM. Common principles in clathrin-mediated sorting at the Golgi and the plasma membrane. Biochim Biophys Acta. 2005; 1744: 415–437. doi: 10.1016/j.bbamcr.2005.04.005 PMID: 15922462

86. Mousavi SA, Malerød L, Berg T, Kjeken R. Clathrin-dependent endocytosis. Biochem J. 2004; 377: 1–16. doi: 10.1042/BJ20031000 PMID: 14505490

87. Tian G, Thomas S, Cowan NJ. Effect of TBCD and its regulatory interacto r Arl2 on tubulin and microtubule integrity. Cytoskeleton (Hoboken). 2010; 67: 706–714.

88. Harrison RE, Grinstein S. Phagocytosis and the microtubule cytoskeleton. Biochem Cell Biol. 2002; 80: 509–515. PMID: 12440692

89. FastQC software. Babraham Institute—Babraham Bioinformatics. Available: http://www.bioinformatics.babraham.ac.uk/projects/fastqc/.

90. Bolger AM, Lohse M, Usadel B. Trimmomatic: A flexible trimmer for Illumina Sequence Data. Bioinformatics. 2014; 30: 2114–2120. doi: 10.1093/bioinformatics/btu170 PMID: 24695404

91. Dobin A, Davis CA, Schlesinger F, Drenkow J, Zaleski C, Jha S, et al. STAR: ultrafast universal RNA-seq aligner. Bioinformatics. 2013: 29: 15–21. doi: 10.1093/bioinformatics/bts635 PMID: 23104886

92. Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, et al. The Sequence alignment/map (SAM) format and SAMtools. Bioinformatics. 2009; 25: 2078–2079. doi: 10.1093/bioinformatics/btp352 PMID: 19505949

93. Anders S, Pyl PT, Huber W. HTSeq—a Python framework to work with high-throughput sequencing data. Bioinformatics. 2015; 31: 166–169. doi: 10.1093/bioinformatics/btp616 PMID: 25260700

94. Kozomara A, Griffiths-Jones S. miRBase: integrating microRNA annotation and deep-sequencing data. Nucleic Acids Res. 2011; 39: 152–157.

95. Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. Basic local alignment search tool. J Mol Biol. 1990; 215: 403–410. doi: 10.1016/S0022-2836(05)80360-2 PMID: 22317112

96. Robinson MD, McCarthy DJ, Smyth GK. edgeR: a Bioconductor package for differential expression analysis of digital gene expression data. Bioinformatics. 2010; 26: 139–140. doi: 10.1093/bioinformatics/btp152 PMID: 19910308

97. vegan—R package. Available: http://vegan.r-forge.r-project.org

98. gplots—R package. Available: http://www.rdocumentation.org/packages/gplots versions/3.0.1

99. Pfaffl MW. A new mathematical model for relative quantification in real-time RT-PCR. Nucleic Acids Res. 2001; 29: e45. PMID: 11328886

100. Robinson TL, Sutherland IA, Sutherland J. Validation of candidate bovine reference genes for use with real-time PCR. Vet Immunol Immunopath. 2007; 115: 160–165.

101. Wong N, Wang X. miRDB: an online resource for microRNA target prediction and functional annotations. Nucleic Acids Res. 2015; 43: 146–152.

102. TargetScan Human—Search for predicted microRNA targets in mammals. Available: http://www.targetscan.org/

103. Liu SQ, Jiang S, Li C, Zhang B, Li QJ. miR-17-92 cluster targets phosphatase and tensin homology and Ikaros Family Zinc Finger 4 to promote TH17-mediated inflammation. J Biol Chem. 2014; 289: 12446–12456. doi: 10.1074/jbc.M114.550723 PMID: 24644282

104. Jiang S, Li C, Olive V, Lykken E, Feng F, Sevilla J, et al. Molecular dissection of the miR-17-92 cluster’s critical dual roles in promoting Th1 responses and preventing inducible Treg differentiation. Blood. 2015; 116: 5487–5487.

105. Li X, Zhu L, Liu X, Sun X, Zhou Y, Lang Q, et al. Differential expression of micromas in porcine parvovirus infected porcine cell line. Virol J. 2015; 12: 128. doi: 10.1186/s12985-015-0359-4 PMID: 26290078

106. Huang L, Liu Y, Wang L, Chen R, Ge W, Lin Z, et al. Down-regulation of miR-301a suppresses pro-inflammatory cytokines in Toll-like receptor-triggered macrophages. Immunology. 2013; 140: 314–322. doi: 10.1111/imm.12139 PMID: 23808420