

# *Brucella melitensis* and *Mycobacterium tuberculosis* depict overlapping gene expression patterns induced in infected THP-1 macrophages

Masoudian, M.<sup>1</sup>; Derakhshandeh, A.<sup>2</sup> and Ghahramani Seno, M. M.<sup>3\*</sup>

<sup>1</sup>Ph.D. Student in Biotechnology, Department of Pathobiology, School of Veterinary Medicine, Shiraz University, Shiraz, Iran; <sup>2</sup>Department of Pathobiology, School of Veterinary Medicine, Shiraz University, Shiraz, Iran; <sup>3</sup>Department of Basic Sciences, Faculty of Veterinary Medicine, Ferdowsi University of Mashhad, Mashhad, Iran and Institute of Biotechnology, Ferdowsi University of Mashhad, Mashhad, Iran

\*Correspondence: M. M. Ghahramani Seno, Department of Basic Sciences, Faculty of Veterinary Medicine, Ferdowsi University of Mashhad, Mashhad, Iran and Institute of Biotechnology, Ferdowsi University of Mashhad, Mashhad, Iran. E-mail: mgseno@um.ac.ir

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## Summary

Pathogens infecting mammalian cells have developed various strategies to suppress and evade their hosts' defensive mechanisms. In this line, the intracellular bacteria that are able to survive and propagate within their host cells must have developed strategies to avert their host's killing attitude. Studying the interface of host-pathogen confrontation can provide valuable information for defining therapeutic approaches. Brucellosis, caused by the *Brucella* strains, is a zoonotic bacterial disease that affects thousands of humans and animals around the world inflicting discomfort and huge economic losses. Similar to many other intracellular dwelling bacteria, infections caused by *Brucella* are difficult to treat, and hence any attempt at identifying new and common therapeutic targets would prove beneficial for the purpose of curing infections caused by the intracellular bacteria. In THP-1 macrophage infected with *Brucella melitensis* we studied the expression levels of four host's genes, i.e. *EMP2*, *ST8SIA4*, *HCP5* and *FRMD5* known to be involved in pathogenesis of *Mycobacterium tuberculosis*. Our data showed that at this molecular level, except for *FRMD5* that was downregulated, the other three genes were upregulated by *B. melitensis*. *Brucella melitensis* and *M. tuberculosis* go through similar intracellular processes and interestingly two of the investigated genes, i.e. *EMP2* and *ST4SIA8* were upregulated in THP-1 cell infected with *B. melitensis* similar to that reported for THP-1 cells infected with *M. tuberculosis*. At the host-pathogen interaction interface, this study depicts overlapping changes for different bacteria with common survival strategies; a fact that implies designing therapeutic approaches based on common targets may be possible.

**Key words:** *Brucella melitensis*, *EMP2*, *FRMD5*, *HCP5*, *ST8SIA4*

## Introduction

Multicellular organisms such as mammals have evolved sophisticated and complex mechanisms to make themselves immune to invasion by a huge number of opportunists they encounter on a daily basis. On the other hand, for survival, many microbes are either completely dependent on other living organisms or at least part of their life cycle must involve taking advantage of the resources available from the other organisms. Therefore, to survive, these microorganisms have had to develop strategies to at least suppress their prospective host's defensive mechanisms. In this context, the intracellular pathogenic bacteria not only effectively evade the host's various components of immune system, but have evolved mechanisms to use the cells meant to kill them as hiding niches or as dwellings (Imbuluzqueta *et al.*, 2010). Moreover, since many antibiotics either do not enter the cells or their concentrations inside the cells do not reach a therapeutic level, infections caused by the intracellular dwelling bacteria are either very difficult to treat or are refractory to many treatment procedures (Carryn *et al.*, 2002).

*Brucella* genus includes ten gram-negative species of which *B. abortus*, *B. melitensis*, *B. suis*, and *B. canis* are known to affect humans (Grilló *et al.*, 2012; He, 2012). Human infections with *B. abortus*, *B. melitensis* and *B. suis*, which are considered as category B priority pathogens by the US Center for Disease Control (CDC), are more commonly reported, affecting 500,000 people around the world per year (He, 2012). Moreover, brucellosis is a cause of huge economic losses worldwide; for example, it is estimated to lay a burden of 600 million dollars on Latin America's economy (Barbier *et al.*, 2011).

*Brucella* as an intracellular pathogen has developed sophisticated mechanisms to subvert its host defensive strategies rendering them ineffective. Inside the macrophage cells, *Brucella* is protected and trafficked within a membrane-bound compartment called *Brucella*-containing vacuole (BCV) (Roop II *et al.*, 2009). It is known that factors from the bacteria either directly, by changing the BCV properties, or indirectly, by controlling and changing the expression of various host factors, help the intracellular pathogen survive within the host cells (Roop II *et al.*, 2009). In this line, it has been

shown that the transcriptome of macrophages infected with a pathogenic strain of *B. melitensis* is significantly different from that of the cells infected with the attenuated strain which is cleared more easily by the macrophages (Wang *et al.*, 2011). It is expected that this difference should maintain the factors that help the pathogenic strain survive within the macrophage cells.

In a recent interesting work, using macrophages differentiated from THP-1 cells. Kumar *et al.* (2010) used RNAi to target all host genes, one by one, and studied the consequences with regard to the changes in replication capacity of the intracellular pathogen *M. tuberculosis*. These investigators identified 74 host genes whose downregulation resulted in reduced pathogen propagation within the cells, irrespective of the *Mycobacterium* strain they tested. Further, these investigators demonstrated that upon infection with *M. tuberculosis* the expression of certain genes, whose earlier downregulation was shown to result in lower *M. tuberculosis* replication capacity are indeed induced; a fact that may indicate this intracellular pathogen directs creation of a tolerating or permissive milieu within the host cell. In order to verify whether similar changes happen when the cells encounter different intracellular pathogens, a subset of the aforementioned genes were selected and their expression behaviors were studied using qPCR in differentiated THP-1 cells infected with *B. melitensis*. Previous research works show that *Brucella* and *Mycobacterium* survive within macrophages by limiting the fusion of lysosomes with the phagosomes harboring the bacteria (Yuk *et al.*, 2012; Mostowy, 2013; Huang and Brummell, 2014). Therefore, for therapeutic purposes it would be interesting to verify whether, irrespective of pathogen type, common pathways may be triggered by these intracellular pathogens to be used as common drug target(s).

## Materials and Methods

### Candidate gene selection

Using RNAi to knockdown genes in differentiated THP-1 cells infected with *M. tuberculosis*, Kumar *et al.* (2010) identified a list of 74 genes whose downregulation restricted *M. tuberculosis* propagation irrespective of the infecting bacterial strain. These investigators further showed that a subset of these genes are upregulated upon THP-1 infection with *M. tuberculosis*, suggesting a mechanism by which *M. tuberculosis* induces expression of host factors that favor its survival and propagation within the host cells. Having in mind the goal of pinpointing possible therapeutic targets, from the list of 74 genes identified by Kumar *et al.* (2010) we chose four genes, i.e. *EMP2* (epithelial membrane protein 2) and *ST8SIA4* (2-8 sialyl transferase), *HCP5* (HLA complex P5), *FRMD5* (FERM domain 5) that, based on available knowledge did not seem essential for macrophage function and their downregulation had resulted in over 50% reduction in *M. tuberculosis* propagation within THP-1 cells.

### Internal control gene selection for qPCR

Based on the data from Mae *et al.* (2010) that indicated *ACTB* and *RPL37A* as the most stable genes upon THP-1 differentiation among 21 other potential internal control genes and the data from Cao *et al.* (2012) that showed *PPIB* and *PGK1* as the most stable genes during THP-1 stimulation with LPS, we chose *RPL37A* and *PPIB* as potential normalizers for this work. Subsequently, based on the expression values for these two genes in our THP-1 cells, *PPIB* was selected to normalize the qPCR data.

### Cell culture, macrophage differentiation and infection

The THP-1 human monocytic cell line was obtained from the Persian Type Culture Collection (Pasteur Inst., Iran). *Brucella melitensis* was obtained from the Microbiology Lab at the School of Veterinary Medicine, Shiraz University, Shiraz, Iran. This bacterium had been isolated from an aborted sheep fetus and biochemically characterized in the aforementioned lab.

The THP-1 cells were maintained and cultured in RPMI 1640 (GIBCO) supplemented with 10% fetal bovine serum (FBS), 50  $\mu$ M 2-mercaptoethanol, 5  $\mu$ M L-glutamine and 100 IU/100  $\mu$ g/ml penicillin/streptomycin at 37°C in a tissue culture incubator. The media were refreshed every 2-3 days and the cells were split before the cell number reached  $1 \times 10^6$ /ml. When the passage number reached 10, a new frozen vial was used to start a fresh culture. On each round of experiment, THP-1 cells were counted in the presence of trypan blue to evaluate cell viability and six tissue culture dishes (35 mm) were seeded with  $5 \times 10^5$  cells in 2 ml complete RPMI containing 5 ng/ml phorbol 12-myristate 13-acetate (PMA) (Cat # P 1585, SIGMA). The following day, when the differentiated cells had become adherent as confirmed by light microscopy, the cells were washed thrice gently with warm RPMI without antibiotic. The bacteria (*B. melitensis*) were grown to late logarithmic phase ( $OD_{600} = 1$ ) in Tryptic Soy Broth (Merck, Germany), centrifuged at 12000 RPM for 10 min and resuspended in warm RPMI 1640 without antibiotic. For the test group, the media containing the bacteria were added to three cultures at the multiplicities of infection (MOI) of 10. The infected plates were centrifuged at 500 g for 10 min and were subsequently transferred to a tissue culture incubator. After 4 h of incubation at 37°C, the cells were gently washed thrice with complete RPMI containing 100  $\mu$ g/ml gentamicin followed by 2 h of incubation in the same medium to kill extracellular bacteria. The cells were subsequently washed with complete RPMI containing 10  $\mu$ g/ml gentamicin and were incubated in the same medium until used to extract RNA at ~20 h post infection. The remaining cultures were used as controls and hence were treated exactly as the infected cultures were, but without being exposed to the bacteria. Each round of experiment contained three replicates for each condition (i.e. infected and uninfected), and the whole experiment was also independently repeated two more times.

**Table 1:** Specifications of the primers used in this work

Gene	Primers sequences	Tm (°C)	Product size (bps)
<i>EMP2</i>	F: CTGTGTGTCATGATTGCGGC	60.18	158
	R: ATCATGCCGCTGATGAAGGT	59.82	
<i>ST8SIA4</i>	F: CTGGGGCAACCAGGACTTTC	60.90	143
	R: ATGAGTTGCGTCTCCTGGTG	60.04	
<i>HCP5</i>	F: AAGGAGAGTTGATCAAGGCCG	60.07	145
	R: AGGCCCTACTTCTCTCAGGC	60.69	
<i>FRMD5</i>	F: CAATGAGCGAGTAGCTGTG	59.35	142
	R: TCCTCCTCAATGCTGCAGGT	61.20	
<i>PPIB</i>	F: AGATGTAGGCCGGGTGATCT	60.11	152
	R: CTCCGCCCTGGATCATGAAG	60.25	

### RNA isolation and cDNA synthesis

Cells were used to extract total RNA using Qiagen RNeasy plus Mini kit (Cat # 74134 Qiagen, Germany) that included a built-in on-column DNase treatment step. The extracted RNAs were qualitatively and quantitatively assessed by, respectively, running on agarose gel and spectrophotometry. The extracted RNAs were stored at -80°C until used.

For cDNA synthesis PrimeScript kit (perfect real time) (Cat # RR037A Takara, Japan) and the provided instruction was used. For each sample, 300 ng RNA with 25 picomole oligodT and 50 picomole random hexamer in 10 µL final volume was reverse transcribed. The cDNA product was diluted one in half and transferred to -20°C in aliquots until used.

### Primer design and qPCR experiment

The primers were designed using Primer3 software (Untergasser *et al.*, 2012) (Table 1). All primer pairs were designed to have similar melting temperatures and were checked for not having common SNPs using UCSC genome browser (Karolchik *et al.*, 2003). The primers were synthesized by Bioneers Ltd. (South Korea). The primers were tested by running a melt curve analysis and also running the related PCR products on agarose gel to confirm the products size. In order to determine the best matching primer set concentrations, a primer optimization procedure was conducted for each set of the primer sets (Table 2) and the efficiencies of amplification were also determined by standard curve method.

**Table 2:** Primer optimization data. The best performing concentrations for each primer sets as determined in our work are presented

Primer set	Optimum concentration (nM)
<i>EMP2</i>	F:400 / R:600
<i>FRMD5</i>	F:400 / R:600
<i>HCP5</i>	F:400 / R:600
<i>PPIB</i>	F:600 / R:300
<i>ST8SIA4</i>	F:600 / R:400

Quantitative PCR was conducted on a MiniOpticon instrument (BioRad, USA) using the SYBR Premix Ex *Taq II* (Cat # RR820Q, Takara, Japan) according to the

instructions provided by the manufacturer. qPCR cycling conditions were as follows: 30 s pre-incubation at 95°C, 40 cycles of 95°C/5 s and 58°C/30 s. After completion of the amplification step a melting curve analysis was also conducted.

### Statistical analysis

The differences in expression values between test and control conditions for each gene were statistically analyzed using REST software (<http://rest.gene-quantification.info>) (Pfaffl *et al.*, 2002). A cut-off of P<0.05 was considered as significant.

### Results

#### Reference gene selection

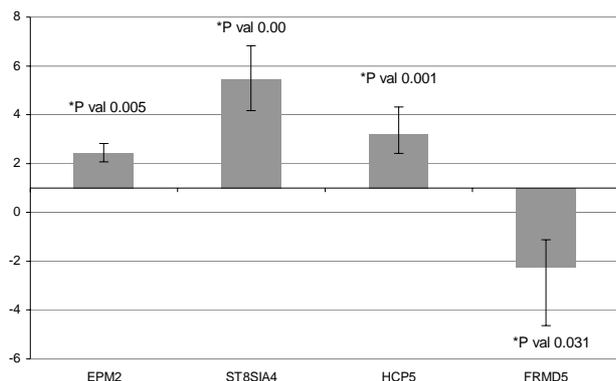
As explained in the materials and methods section, based on published data (Maeß *et al.*, 2010; Cao *et al.*, 2012) *PPIB* and *RPL37A* genes were initially selected as potential reference genes for the current work. A qPCR experiment was conducted using the primer sets amplifying *PPIB* or *RPL37A* and after analyzing the expression values for these two genes, *PPIB* was chosen over *RPL37A* for this work. This is because *RPL37A* was highly expressed in THP-1 cells and *PPIB* expression level was within the same range as the other four experimental genes.

#### Differential regulation of *EMP2*, *ST8SIA4*, *HCP5* and *FRMD5* in THP-1 macrophages infected with *Brucella melitensis*

qPCR data for *EMP2*, *ST8SIA4*, *HCP5* and *FRMD5* showed that these genes are significantly dysregulated upon THP-1 infection with *B. melitensis* (Fig. 1). *EMP2* (Fold change (FC) = 2.402, P=0.005), *ST8SIA4* (FC = 5.421, P=0.00) and *HCP5* (FC = 3.18, P=0.001) are upregulated while *FRMD5* (FC = -2.222, P=0.031) is downregulated when the THP-1 macrophages are infected with *B. melitensis*.

### Discussion

The characteristics of eukaryotic cells and their intracellular conditions are governed by their functional



**Fig. 1:** qPCR data on genes changed by THP-1 macrophages infected with *B. melitensis*. The qPCR analysis of four genes in THP-1 macrophages infected with *B. melitensis* showed that compared to the uninfected cells, three of them were significantly upregulated while one gene was significantly downregulated. Bars show standard error of mean

genomics. Therefore, it is rational to expect that intracellular dwelling parasites must have developed strategies to change their host properties to their advantage by inducing changes in the functional genomic aspects of their host. In this line, Kumar *et al.* (2010) took an interesting approach to interrogate the aspects of the transcriptome of THP-1 macrophages that influence the persistence of the infection caused by *M. tuberculosis*. These investigators showed that many hundreds of the host's genes were involved in the process of development and persistence of the infection, and therefore manipulating these factors may be helpful in maintaining the infection under control. Interestingly, they verified that several of the genes that had earlier shown their downregulation by RNAi suppressed the progress of the infection were indeed upregulated in the infected cells, implying *M. tuberculosis* induces changes in the host's transcriptome in favour of its survival and replication. Here, we chose *B. melitensis* with similar life cycle to that of *Mycobacterium* regarding its persistence in phagosomes and changes to the levels of expression of four genes, i.e. *EMP2*, *ST8SIA4*, *HCP5* and *FRMD5* taken from Kumar's study were determined in the infected THP-1 macrophages. The goal here was to verify whether similar alterations in gene transcription may be induced by these two different intracellular pathogens. Our data confirmed a similar pattern in expression changes in two of the selected genes, i.e. *EMP2* and *ST8SIA4* between THP-1 cells infected with *B. melitensis* and those infected with *M. tuberculosis* as reported by Kumar *et al.* (2010). Based on the previous studies, *B. melitensis* and *M. tuberculosis* use similar strategies to survive within the host by limiting the fusion of phagosomes harbouring them with lysosomes (Arenas *et al.*, 2000; Knodler *et al.*, 2001; Yuk *et al.*, 2012). Therefore, the genes commonly changed in THP-1 cells infected with either *B. melitensis* or *M. tuberculosis* may be involved in some stages of normal phagosome maturation such as fusion with lysosomes.

Kumar *et al.* (2010) showed that *EMP2* is not only

upregulated when THP-1 cells are infected with *M. tuberculosis*, but its downregulation by RNAi restricts *M. tuberculosis* replication within these cells. Our data also showed that this gene is upregulated in THP-1 cells infected with *B. melitensis* (Fig. 1). *EMP2* is a transmembrane protein and is a member of the GAS3/PMP22 subfamily that is involved in the delivery of certain proteins such as integrin isoform to the cell surface (Wadehra *et al.*, 2005; Shimazaki *et al.*, 2009). *EMP2* is reported to be highly expressed at membrane sites of Chlamydia infection and blockage of this protein using anti-*EMP2* antibody reduces the infectability of the genital tract of mouse model by Chlamydia (Shimazaki *et al.*, 2009). Similarly, it has been shown that the HEC-1A cell line with reduced expression of *EMP2* is refractory to infection by Chlamydia, but overexpression of this protein in this cell line makes it amenable to infection (Shimazaki *et al.*, 2007). In the RNAi experiment conducted by Kumar *et al.* (2010), it was demonstrated that *EMP2* downregulation limits the propagation capacity of *M. tuberculosis* within THP-1 cells. These investigators designed their experiment so that they targeted the gene after *M. tuberculosis* had entered the cells. Therefore, this design ruled out any possible influence from reduced THP-1 cells infectability by lower surface expression of *EMP2* and suggested that this gene may be active somewhere in post-infection stages of *Mycobacterium* life cycle. Moreover, they showed that the expression level of *EMP2* gene was, indeed, upregulated after THP-1 cells had been infected with *Mycobacterium*, indicating positive influence on *Mycobacterium* survival and propagation from *EMP2* after infection. Interestingly, it has been shown that macrophages infected with *Mycobacterium* and activated with  $\text{INF}\gamma$  show enhanced killing activity (Gutierrez, 2004; Herbst, 2011) while  $\text{INF}\gamma$  stimulation of cells results in downregulation of *EMP2* expression (Kumar *et al.*, 2010). Therefore, it seems reasonable to assume that  $\text{INF}\gamma$  accomplishes its antibacterial effect at least partly through modulating the expression of a variety of genes, such as *EMP2*, to overtake the suppression state already induced by some bacteria.

*ST8SIA4* encodes for a protein that is involved in the synthesis of polysialic acid (Nakayama *et al.*, 1995). This gene that is overexpressed in whole blood compared to its average expression in all tissues shows substantial expression in immune related cells such as monocytes, neutrophils and B and T cells (Safran *et al.*, 2010; Montague *et al.*, 2014; Uhlén *et al.*, 2015). In our experiment *ST8SIA4* is upregulated in THP-1 macrophages infected with *B. melitensis* as it was also reported to be overexpressed in THP-1 macrophages infected with *M. tuberculosis* (Kumar *et al.*, 2010). It has also been shown that upon infection with some viruses such as the SARS-associated coronavirus, the infected macrophages endure overexpression of *ST8SIA4* (Peiris and Cheung, 2009). Downregulation of *ST8SIA4* by RNAi lowers the capacity of THP-1 macrophages in supporting propagation of intracellular *M. tuberculosis* (Kumar *et al.*, 2010). As mentioned earlier  $\text{INF}\gamma$

stimulation of the macrophages infected with *M. tuberculosis* increases the capacity of the infected cells in overtaking the suppression state induced by the bacteria (Gutierrez *et al.*, 2004; Kumar *et al.*, 2010; Herbst *et al.*, 2011). Similar to the case for *EMP2*, the cells stimulated with INF $\gamma$  undergo a downregulation in *ST8SIA4* expression levels as well (Kumar *et al.*, 2010). Moreover, the notions that for survival within macrophages, *Brucella* intercepts membrane trafficking from endoplasmic reticulum to Golgi apparatus (Atluri *et al.*, 2011), and that *ST8SIA4* is localized to Golgi apparatus (Angata and Fukuda, 2014) may suggest some other molecular mechanisms for the involvement of this gene in pathogenesis of *Brucella*.

Our data show that *HCP5* gene is upregulated in THP-1 cells infected with *B. melitensis*. *HCP5* codes for a non-coding RNA that is structurally related to human endogenous retroviruses (HERVs) group of transposable elements (Kulski and Dawkins, 1999). *HCP5* locus is positioned within the MHC class I region and the sequence shows some homology to retroviral *pol* gene. Some genomic variations in this locus have been associated with viral load in HIV infection, and hence with delay in development of AIDS (Fellay *et al.*, 2007), as well as with susceptibility to development of herpes zoster (Crosslin *et al.*, 2015).

Based on our data, in THP-1 macrophages infected with *B. melitensis* the expression level of *FRMD5* gene is downregulated by over two-fold (Fig. 1). Except for some data on the association of *FRMD5* and tumor progression (Wang *et al.*, 2012) little information is available on the possible function of *FRMD5* in development and progression of infection. It has been shown that *FRMD5* shares 51% identity with another family member, i.e. *FRMD3* (Susana *et al.*, 2013) and that *FRMD3* overexpression induces apoptosis (Haase *et al.*, 2007). Therefore, if a similar pro-apoptotic function could be attributed to *FRMD5*, the notion that infection of human monocytes with *B. suis* inhibits apoptosis (Gross *et al.*, 2000) can best be used to envision the interrelationship between bacterial infection, *FRMD5* downregulation and apoptosis. Otherwise, in infected cells apoptosis would limit the bacteria propagation and persistence within the host. On the other hand, Kumar *et al.* (2010) demonstrate positive influence from *FRMD5* downregulation by RNAi on bacterial replication within THP-1 macrophages. Though it may be argued that the level of suppression may have an influence, the scientific reason for this apparent contradiction remains to be explored.

Taken together, based on the previous studies and the data from this work it is obvious that intracellular pathogenic bacteria influence their host's transcriptome for their own benefit. Therefore, taking approaches to revert the changes induced by these pathogens can prove beneficial for prophylactic and/or therapeutic purposes. Moreover, though the changes induced by the bacteria can be type specific, our data shows that the bacteria with similar life style such as *Mycobacterium* spp. and *Brucella* spp. direct overlapping molecular events that

can be considered as common therapeutic targets.

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## Conflict of interest

The authors declare no conflict of interest.

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