Biocompatibility and Gene Expression Associated with Streptococcus Infection in Myocarditis

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Abstract Myocardial inflammation is a nonspecific response to various triggers such as viral and bacterial infection. This study was designed to examine the Streptococcus induction of myocarditis in rat and induction of CRP, SAP, LBP and cytokines on the transcriptional activity of the inflammation/related genes expression using a semi-quantitative RT-PCR in liver and heart tissues. Experimental myocarditis was established with intra-peritoneal injection of 10^8 CFU/ml of *Streptococcus faecalis* in three groups of rats and sacrificed after one, two and seven days post infection, respectively. Heart and liver functions were highly elevated while kidney functions were not affected. Histopathological examination for heart and liver showed damage effect for the bacterial infection. Expression pattern of CRP, SAP, LBP & IL6 genes were investigated, in liver these genes were highly expressed while some of them showed no expression pattern in heart tissues.

Keywords Myocarditis, Rat, *Streptococcus Faecalis*, C-Reactive Protein, Serum Amyloid P Component, Interleukin 6 & Lipopolysaccharide Binding Protein

1. Introduction

Myocarditis, a heterogeneous myocardial inflammatory disease, leads to both acute and chronic heart failure manifested by dilated cardiomyopathy. The etiology of myocarditis appears to be rather complicated involving idiopathic, infectious or autoimmune factors (Caforio and Iliceto, 2008). Bacteremia from any source may result in myocarditis, with the most common pathogens being meningococcus, streptococcus, and Listeria (Lori and Cooper, 2010). Myocarditis is an uncommon disease, usually difficult to diagnose because its clinical behavior may be entirely asymptomatic until sudden death. (Dec, 2002 & Jared et al., 2006) Myocarditis has been reported in up to 12% of young adults presenting with sudden death (Fabre & Sheppard, 2006) and is an important underlying etiology of other myocardial diseases (Puranik et al., 2005) such as dilated (Kawai, 1999) and arhythymogenic right ventricular cardiomyopathy (Calabrese et al., 2006). Research over the last 15 years has provided that there is unequivocal evidence for a local inflammation in the artery wall which is accompanied by a systemic low-grade inflammatory response. (Feinstein & Spagmulo, 1993 and Narula et al., 1999). Inflammatory response can be measured by a variety of inflammatory-biomarkers. The largest database so far has been accumulated for C-reactive protein (Kolz et al., 2008). C-reactive protein (CRP) is an acute phase reactant with roles in innate host defense, clearance of damaged cells, and regulation of the inflammatory response. Cytokines released from jeopardized tissues stimulate the liver to synthesize acute phase proteins, including C-reactive protein (CRP) and serum amyloid P component (SAP). IL-6 is the chief stimulator for the production of most acute phase proteins. (Kushner, 1982 and Gotschilich, 1989). Serum amyloid P component (SAP), the murine homologue to CRP, is a relatively low concentration constitutive serum protein in man. Paradoxically, in mouse SAP is the major acute phase reactant. (Jiang et al., 1995) Therefore, mRNA expression levels of CRP and SAP may provide sensitive markers of systemic acute phase response (Saber et al., 2009). Lipopolysaccharide binding protein (LBP), a key participant in the inflammatory response to infection, may be a useful marker for diagnosis and prognosis of patients with bacterial infections (Schumann et al., 1990). The present study sought to investigate the association and biocompatibility between C - reactive protein (CRP), Serum amyloid P component (SAP), Lipopolysaccharide binding protein (LBP) and cytokines exemplified by interleukin 6 (IL6) genes expression involved in the inflammatory pathway in a rat group experimentally infected with streptococcus bacteria.

2. Materials and Methods

2.1. Experimental animals
Ten weeks old male Swiss albino rats (n=40) weighing 150-200 g were maintained on a diet of *ad libitum* and tap water. These experiments have been approved by Ain Shams University committee on the use and care of Laboratory animals. After 2-weeks as a conditioning period, the animals were randomly distributed into 2 groups. One group (n=10) was treated as a control group and injected with saline. The other group (n=30) was treated as the experimental group, anesthetized with ketamine, and were intraperitoneal injected with a fresh culture of *Streptococcus faecalis*. Rats were sacrificed after 1, 2 and 7 days from infection with bacteria.

2.2. Bacterial Challenge

*Enterococcus faecalis* bacteria strain reference ATCC: 29212, supplied from the faculty of veterinary medicine, Cairo University, Giza, Egypt was used. The bacteria was grown for 18 hours at 37°C in glucose broth medium (Enriched media) and aliquots of the initial culture were inoculated into fresh media with 15 % glycerol for storing and sub cultured in fresh media for 18 hours prior to injection. The bacteria were cultured to approximately 10^8 CFU/ml. Estimations of cell density were determined by measuring the absorbance at 620 nm and verified by plate counts. Quantization of cultures was done by serial dilutions & plates were counted after overnight incubation with a bright light source. A fresh culture of bacteria was prepared (24 hours culture) and 1 ml of this culture was injected in the experimental rat groups.

2.3. Tissue Specific Expression of CRP, IL-6, LBP and SAP Genes by Semi-Quantitative RT-PCR Analysis

I) RNA extraction from tissue

RNA was extracted from rat livers and hearts according to the method of Nicholas, (2004) using BIOZOL BioFlux™ Reagent (ready to use) that was purchased from BioFlux (South San Francisco, U.S.A).

II) Reverse transcription

The extracted total RNA was then reverse transcribed with Oligo d(T) primer according to Revert Aid First Strand cDNA Synthesis Kit™ (purchased from Fermentas life science Co. (Invitorgen Corporation) (Van Allen Way, Carlsbad, Canada). Positive and negative control reactions were used to verify the results of the first strand cDNA synthesis steps, GAPDH gene RNA was used as positive control.

III) Polymerase chain reaction

The resulting cDNA was subjected to PCR for 35 cycles with respective primers designated from the sequence of the CRP, IL-6, LBP and SAP genes (table 1) using primer premier 5.0 software and were purchased from Invitrogen Corporation (Van Allen Way, Carlsbad, Canada). Dream Taq™ Green PCR Master Mix (Invitrogen Corporation) was used in the PCR. Products of PCR were then displayed on an appropriate agarose gel (2%) and examined for yield and specificity. Analysis of gel images was done using Gel analyzer Pro (version 3.1) software. PCR product for each gene was sequenced. The obtained nucleotide sequences were blasted on gene bank site (http://www.ncbi.nlm.nih.gov/) to ensure that the obtained PCR product is our target gene. The nucleotide sequence for each gene was analyzed using the web site for sequence manipulation suite (version 2): http://www.bioinformatics.org/sms2/ to characterize some target genes like the open reading frames, prediction and characterization for the protein sequences and the signal P 3.0 Sever using the expasy website (http://expasy.org/tools/) to predicts the presence and location of signal peptide cleavage sites in amino acid sequence. This method incorporates a prediction of cleavage sites and a signal peptide/non-signal peptide prediction based on a combination of several artificial neural networks and hidden Markov models.

<table>
<thead>
<tr>
<th>Gene &amp; its accession number</th>
<th>Primers(sense and antisense 5′→3′)</th>
<th>Annulling Temp.</th>
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<tbody>
<tr>
<td>CRP M83176</td>
<td>Sense: 5′-CGA AGG TTC AGC ATC TTC TC-3′</td>
<td>46.5°C</td>
</tr>
<tr>
<td></td>
<td>Antisense: 5′-CTG CAT TGA TCT GTT CTG GAG-3′</td>
<td></td>
</tr>
<tr>
<td>SAP</td>
<td>Sense: 5′-CTC AGA CAG ACC TCA ATC AG-3′</td>
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<tr>
<td></td>
<td>Antisense: 5′-TCA GCA ATA CCA GAG GAG GA-3′</td>
<td></td>
</tr>
<tr>
<td>IL-6</td>
<td>Sense: 5′-CCA GCC AGT TGC CTT CTT GGG A-3′</td>
<td>58°C</td>
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<tr>
<td></td>
<td>Antisense: 5′-GGC ATA GCA CAC TAG GTT TGC CGA-3′</td>
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<tr>
<td>LBP</td>
<td>Sense: 5′-GAG GCC GTC TCT CCA TCT-3′</td>
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<tr>
<td></td>
<td>Antisense: 5′-TCT GAG ATG GCA AAG TAG ACC-3′</td>
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</table>

IV) Histochemical diagnosis of myocardial infarction

Autopsy samples were taken from the liver and heart of rats in different groups and fixed in 10 % formal saline for 24 hours. Washing was done in tap water then serial dilutions of alcohol (methyl, ethyl and absolute ethyl) were used for dehydration. Specimens were cleared in X ylene and embedded in paraffin at 56°C in hot air oven for 24 hours. Paraffin bees wax tissue blocks were prepared for sectioning at 4 microns by slide microtome. The obtained tissue sections were collected on glass slides, deparaffinized and stained by hematoxylin and eosin stains (Banchroft et al; 1996) for histopathological examination through the light microscope.

3. Results

a) C-reactive protein gene expression

CRP gene was highly expressed by hepatocytes and heart after bacterial infection by 1 day, 2 days and 7 days (Fig. 1-A). In both tissues the bands molecular weight on agarose gel were 433 bp. While, there was no expression for CRP in the liver and heart of normal non-infected control group.
GADPH gene expression was used as +ve control in examining all the studied genes. Rat CRP gene/GADPH gene ratio was determined by densitometry (Fig.2), which was performed by measuring the photo stimulated luminescence values using gel analyzer pro version software.

**Figure 1.** A) The C-reactive protein (CRP) gene expression in accordance to GAPDH. (B) SAP gene expression in accordance to GAPDH (lane: 1) obtained by RT-PCR in liver (lane: 2, 3 & 4) & heart (6, 7 & 8) of infected groups with bacteria and control group (lanes 5 & 9) (M: DNA marker)

At seven days post-bacterial infection in liver, the rat CRP gene expression recorded the highest level compared to the control. However, after one and two days post infection, rat CRP gene expression levels was less than the level after seven days but still higher than that of control. In heart, rat CRP gene showed different expression pattern from that of liver post bacterial infection where at two days post infection in heart recorded the highest level of expression. However the rat CRP gene expression in heart was less than that of control in the three investigated intervals of time post bacterial infection. Moreover, rat CRP gene expression level in liver was higher than that of heart at all the stimulation periods. These different levels of expression were showed in fig. (3-A).

**Figure 2.** The relative density of expressed rat CRP gene bands in liver and heart tissues

b) Serum amyloid P component gene expression

SAP gene was expressed by hepatocytes in the three infected group of rats at different intervals of time of post infection. The bands molecular weight of SAP genes were 331bp. At two days post bacterial infection in liver, the rat SAP gene expression recorded its highest level compared to the control. However, after one and two days post infection, rat SAP gene expression levels was less than the level after two days but still higher than that of control. Heart tissue shows no expression for SAP gene as in normal control group. These different levels of expression were showed in fig. (3-B)).

c) Interleukin 6 gene expression

IL6 gene was expressed by hepatocytes in the three infected group of rats at different intervals of time of post infection Fig. (4-A). At one day post bacterial infection in liver, the rat IL6 gene expression recorded the highest level compared to the other intervals of time post infection but it still less than that of control. However, after two and seven days post infection, rat IL6 gene expression levels was less than the level after one day and less than that of control. In heart, rat IL6 gene showed different expression pattern from that of liver post bacterial infection where at the three intervals of time post infection in heart recorded the same level of expression. However the rat IL6 gene expression in heart was less than that of control in the three investigated intervals of time post bacterial infection. Moreover, rat IL6 gene expression level in liver was higher than that of heart at all the stimulation periods. These different levels of expression were showed in fig. (5-A).

**Figure 3.** (A) Effect of bacterial infection (1, 2 and 7 days) on expression of rat CRP / GAPDH genes in liver and heart. (B): Effect of bacterial infection (1, 2 and 7 days) on expression of rat SAP / GAPDH genes in liver and heart

**Figure 4.** (A) IL6 gene expression in accordance to GAPDH. (B) LBP gene expression in accordance to GAPDH (Lane: 1) obtained by RT-PCR in liver (2, 3 & 4) & heart (6, 7 & 8) of infected groups with bacteria and control group (lanes 5 & 9) (M: DNA marker)

**Figure 5.** (A) Effect of bacterial infection (1, 2 and 7 days) on expression of rat IL6 & GAPDH genes in liver and heart. (B) Effect of bacterial infection (1, 2 and 7 days) on expression of rat LBP & GAPDH genes in liver and heart
d) Lipopolysaccharide binding protein gene expression

LBP gene showed expression by hepatocytes after bacterial infection by 1 day, 2 days and 7 days fig. (4-B). LBP gene was also expressed in the heart tissue but only in two of the infected groups which are the 1 day and 2 days group and no expression in the 7 days group. The bands molecular weight of LBP genes was 540 bp according to the used marker. At one day post bacterial infection in liver, the rat LBP gene expression recorded the highest level compared to the control. However, after two and seven days post infection, rat LBP gene expression levels decreased and became less than that of control. In the heart, rat LBP gene showed different expression pattern from that of liver post bacterial infection where at two days post bacterial infection recorded the highest level of expression. However the rat LBP gene expression in heart was less than that of control at one and two days post bacterial infection. Moreover, rat LBP gene expression level in liver was higher than that of heart at all the stimulation periods. These different levels of expression were showed in fig. (5-B).

Analysis of Rat gene sequence

After detecting the PCR products for each target gene using the agarose gel electrophoresis, the PCR products were sequenced. Then the sequences were blasted on Gene bank using the BLAST tool to ensure that the visualized bands on gel were for our target gene. The analysis of rat target genes sequence was carried out using the web site for sequence manipulation suite (version 2) http://www.bioinformatics.org/sms2/, showed that CRP gene (Fig. (6) Consists of 531 bp with three open reading frames (ORF).

The ORF number 1 on the direct strand extends from base 79 to base 183, ORF number 2 on the direct strand extends from base 184 to base 288 and ORF number 3 on the direct strand extends from base 289 to base 381. Using the translation tool, nucleotide sequence of rat CRP gene was translated into 177 amino acids. The predicted protein molecular weight was 43.40 KDa and its isoelectric point was pH 5.07. Predicted signal p for the predicted protein sequence of CRP. The signalP-NN and SignalP-HMM results showed that the protein is non-secretory protein with signal peptide probability (0.001) and maximum cleavage site probability (0.00) between position 34 and 35.

Nucleotide sequence of rat SAP gene showed that SAP gene consists of 950 bp with three open reading frames (ORF). ORF number 1 on the direct strand extends from base 190 to base 291, ORF number 2 is on the direct strand extends from base 397 to base 696 where ORF number 3 is on the direct strand extends from base 772 to base 888.

Translation of SAP nucleotides into 316 amino acids predicted protein molecular weight as 77.07 KDa and its isoelectric point was pH 4.92. The predicted signalP-NN and SignalP-HMM for the predicted protein of SAP results showed that the protein is non-secretory protein with signal peptide probability (0.092) and maximum cleavage site probability (0.028) between position 57 and 58.

Analysis of rat IL6 gene sequence showed that IL6 gene consists of 1031 bp with three open reading frames (ORF). ORF number 1 is on the direct strand extends from base 7 to base 579, ORF number 2 is on the direct strand extends from base 658 to base 750 and finally the ORF number 3 is on the direct strand extends from base 775 to base 921. The nucleotide sequence of rat IL6 gene was translated into amino acid sequence using the translate tool. Using the translation tool, the predicted 343 amino acids of rat IL6 with its protein molecular weight was 85.18 KDa and its isoelectric point was pH 4.85. The signalP-NN result showed that a signal peptide was detected at the position from (1) to (13) with mean S value = 0.566 in which the most likely cleavage site is between position 13 and 14: (AAG-AC). The SignalP-HMM showed that the signal peptide probability = 0.947 and maximum cleavage site probability= 0.166 between position 20 and 21.

Analysis of rat LBP gene sequence showed that LBP gene consists of 632 bp with three open reading frames (ORF). ORF number 1 is on the direct strand extends from base 67 to base 171, ORF number 2 is on the direct strand extends from base 172 to base 264 and the ORF number 3 is on the direct strand extends from base 499 to base 630.

Translation of LBP revealed 110 amino acids predicted LBP protein molecular weight was 51.90 KDa and its isoelectric point was pH 4.95. The signalP-NN and SignalP-HMM results showed that the protein is non-secretory protein with signal peptide probability (0.076) and maximum cleavage site probability (0.009) between position 60 and 61.

Histopathological analyses

By examination of liver tissue from the infected rats and that of control group, it was observed that there was changes...
vary in severity and aggressiveness with the longer time after bacterial infection. In other words, after one day of infection, there was mild changes while after 2 days and 7 days, these changes become more aggressive and severe. (Fig.6)

Heart of infected rats showed focal hemorrhages in between the myocardial bundles after infection by one day while after bacterial infection by two days showed degeneration in the myocardial cells with focal hemorrhages in between. However, after seven days there was inflammatory cells infiltration in between myocardial cells (Fig.7).

4. Discussion

Myocardial inflammation is a nonspecific response to various triggers such as viral or bacterial infection, cardio toxic agents, catecholamine, infarction, or mechanical injury (Skouri et al., 2006). Myocarditis is a potentially life-threatening disease that primarily affects children and young adults with sometimes devastating consequences, including sudden death (Drory et al., 1991). Myocarditis is defined as: a process characterized by an inflammatory infiltrate of the myocardium, with necrosis or degeneration of adjacent myocytes not typical of the ischemic damage associated with coronary artery disease (Dominiczak et al., 2005). Bacteremia from any source may result in myocarditis, with the most common pathogens being meningococcus, streptococcus, and Listeria (Blauwet and Cooper, 2010).

More than a dozen different enterococcal species have been associated with human disease; the majority of human enterococcal infections are due to the species Enterococcus faecalis (Jett et al., 1994, Evans & Chinn (1947), Facklam & Collins (1989), Murray (1990) & Moellering (1992)).

The present study was concerned with bacterial myocarditis where it was carried out on four groups of rats, each group is composed of ten rats (n= 10). Three of the four groups were injected by Streptococcus faecalis intraperitoneally and the fourth group was treated as control group and was injected by saline. All rats in the experimental three groups developed myocarditis, this was proved, after sacrificing the animals on different intervals of time (after 1 day, 2 days and 7 days), through a series of biochemical, molecular and histological investigations. In this study we investigated numerous biochemical markers (Data not shown) for inflammation and gene expression pattern for group of inflammation related genes. These investigations were carried out on different intervals of time (one, two and seven days) after bacterial infection.

C-reactive protein is an acute-phase reactant protein synthesized primarily by liver cells in response to the proinflammatory cytokines including interleukin-6 and TNF-α (Castell et al., 1990). The major function of CRP by binding phosphocholine is to recognize some foreign pathogens and phospholipid constituents in damaged cells (Gotschlich and Edelman, 1967). It also activates the complement system and contributes to the induction of inflammatory cytokines and tissue factors in monocytes (Ballou and Kushner, 1992).

Inflammatory molecules as mildly raised C-reactive protein (CRP) are among the most extensively studied CVD biomarkers and is considered one underlying cause of CVD development and progression (Libby, 2000). Animal studies also support the proinflammatory and pro-atherogenic role of CRP, because administration of human CRP or over-expression of CRP in apolipoprotein E-deficient mice promotes the development of spontaneous atherosclerosis (Paul et al., 2004). CRP is mainly expressed by hepatocytes, but can be synthesized in other tissues (Black, Kushner & Samols, 2005 & Marnell, Mold & Du Clos, 2005), although the liver has been assumed as the only source of CRP synthesis and release, this protein has been demonstrated (Calabro, Willerson & Yeh, 2003) to be produced also in other districts, such as the vascular walls.

Mouse CRP (moCRP) is classified as a minor APR since its highest concentration was estimated at only~2µg/ml, as against levels of ≥300, µg/ml in humans that are undergoing an inflammatory response (Kushner, 1988 and Ballou & Kushner, 1992). HuCRP and moCRP are each encoded by a single-copy gene on chromosome 1 of both species as part of a syntenic group of genes (Yunis & Whitehead, 1990). Both genes have a very similar organization, with two exons and a single intron within the codon for the third residue of the mature protein (Lei et al., 1985 & Ohnishi et al 1988). The amino acid sequences from cDNA clones for human and mouse CRP show an identity of 71 % (Ohnishi et al., 1988 & Whitehead et al., 1990). The extent of the increase in moCRP seems likely to be a function of the gene structure, since the human CRP gene is expressed as a major APR in transgenic mice (Ciliberto et al., 1987).

In the current study, the injection of rats by Streptococcus faecalis induces a “priming” of the host immune response
characterized by increased expression of CRP genes in liver and heart tissues by 24 h after infection with a peak at 48 h when compared with the uninfected control rats that showed no expression for the gene. (Analysis of images to show the peak) Tissue-specific gene expression is regulated at the transcriptional initiation step through the recognition of promoter sites by transcription factors (Maniatis, Goodbourn, & Fisch, 1987). This general scheme for tissue-specific gene expression has been documented for many cell types, including pancreatic endocrine cells (Ohlssen & Edlund, 1986), hepatocytes (De Simone & Cortese, 1988), erythrocytes (Tsai et al., 1989) muscle cells (Lassar et al., 1989) and B cells (Muller et al., 1988 & Scheiderei et al., 1988). It seems likely therefore that the expression of the mouse acute-phase genes is regulated by liver-specific trans-acting factors and cis acting elements within the 5' flanking region of the genes (Namon & Richard, 1993). The pathogenesis and severity of both experimental and human myocarditis has been linked to the intramyocardial production of these cytokines, as well as that of interferon-! (IFN-y)(Lane et al., 1991, 1992, 1993).(70)

In rats infected with S. mansoni, a sudden increase in cytokine mRNA expression was observed only during the early stages of infection, during which the parasite passes through the lung. This result indicates that the immune response is associated to the parasitic infection pathway (Cetre et al., 1998).

IL-6, which is Th2 cytokine and the principal mediator of the acute phase response in rats, plays certain roles in immunity in vivo. However, the mechanism by which this cytokine operates has been reported to vary widely. The mechanisms inherent to responses to foreign antigens differ depending on the kinds of parasites, host and infection period. When we set a standard with rates of increase or decrease in cytokine expression in the spleen, it was determined to increase principally in the early stages of infection, and it was unbiased toward the Th1 or Th2 cytokine (KIM et al., 2007).

Interleukin-6 (IL-6), a proinflammatory cytokine, elicits pleiotropic effects on a variety of biological functions, including regulation of the immune response, hem-aptoiesis, inflammation and cellular differentiation (LE & Vlcek, 1989) & Hirano et al., 1990. However, IL-6 serves not only as an inducer of acute phase reactions but is also an important factor in the elicitation of cellular immune responses against infected cells and of mucosal humoral responses against re-infection (Romani et al., 1996). IL-6 is the chief stimulator for the production of most acute phase proteins (Gabay & Kushner, 1999). Recent studies have indicated the additional importance of cytokines in the pathogenesis of myocardial injury in viral myocarditis (Shioi et al. 1996). Evidence that IL-6 generates distinct signal only in the combination with IL-6 receptor in cardiac myocyte is based on the low expression level of IL-6 receptor (Saito et al. 1992; Yamazuchi-Takahara et al. 1995). Independently of high cholesterol or myocardial damage markers, IL-6 predicts future adverse cardiovascular events and reflects increased inflammatory activity in plaques and is therefore a strong marker of increased risk for mortality in coronary artery diseases (Lindmark et al., 2004 & Ridker et al., 2000). The cytokine IL-6 is synthesized, e.g., by mononuclear phagocytes, endothelial cells, and fibroblasts (Soderquist et al., 1995). IL-6 is known to be secreted from several cell types, including endothelial cells, macrophages, lymphocytes, and adipocytes, and exerts its biological actions through a complex yet well-defined fashion (Tsuchiyasu & Takashi, 2004). The most potent inducer of huCRP gene expression at the level of transcription is IL-6 (Ganapathi et al., 1988 & Ganter et al., 1989).

A recent study has shown increased myocardial IL-6 expression is associated with the progression of heart failure (Plenz, Song & Reichenberg et al., 1998). IL-6 mRNA is reportedly induced in the myocardium of a canine model of ischemia and reperfusion (Chandrasekar et al., 1999 & Kukiela et al., 1995).

In the present study, we investigate the expression pattern of IL-6 gene expression pattern in liver and heart tissues. We found that there was no abundance of IL-6 expression pattern in control non infected rat’s liver and cardiac tissue and that bacterial infection led to a marked increase in IL-6 gene expression in both liver and heat tissues detected after one, two and seven days post infection. Some investigators as Erodgan & Ongen, 1997, Mohler et al., 1997, Ridker et al., 2000 & Kanada et al., 2000 have suggested that IL6 enhances development of myocardial injury. Several mouse models have been used to analyze the direct relationship between IL-6 and infection. IL-6 deficient mice were reported to be susceptible to both microbial and viral infections (Lane et al., 1991). Elevated production of IL-6 has been reported in autoimmune and infectious diseases as well as traumatic injury.

Many other studies were in harmony with our results as the studies of (Lane et al., 1991, 1992, 1993) which support that the pathogenesis and severity of both experimental and human myocarditis has been lied to the intra-myocardial production of interleukin-1β (IL-1β) and interleukin-6 (IL 6), as well as that of interferon- IFN-y (Black et al., 2004) Also the studies of Hartmut et al., 1995 support our findings as they stated that cytokines may be released locally within the heart by vascular endothelial cells, fibroblasts, macrophages, and leukocytes in response to a variety of stimuli including ischemia-reperfusion, cardiopulmonary bypass, septic shock, heart transplant rejection, and myocarditis. The heart may also be exposed to cytokines released from other sites such as the liver or lungs in septic or inflammatory conditions (Gotschlich and Edelman, 1967).

Serum amyloid P component (SAP) is a member of pentaxins. Previous studies indicate that SAP exists in human atherosclerotic aortic intima and the plasma SAP levels are associated with cardiovascular disease. (Hirano et al., 1990) SAP is produced mainly in the liver and secreted into the blood circulation. The average plasma concentration of SAP is about 40 mg/l (160 nM) in human. (Puranik et al, 2005) In man and rabbit, one of the serum proteins that exhibit the highest proportional increase, from trace resting
levels to several hundred micrograms per ml, is CRP. SAP is a relatively low concentration constitutive serum protein in man. Paradoxically, in mouse it is SAP that is a 'major' acute phase reactant, increasing from resting levels of around 60, µg/ml or less to peak concentrations of several hundred micrograms/ml during acute inflammation [Gabay, 1999]. Although mouse CRP is poorly defined, it has been reported as a trace serum component [Shioi et al, 1996] that increases slightly in concentration during the acute-phase response [Saito et al, 1992]. These studies support our present study where by analysis of our PCR product images for both CRP and SAP genes expression in liver tissue, we found that the relative density for SAP genes bands was higher than that of CRP genes bands which gives a confirmed figure about that SAP represent the major acute phase protein in rat.

Numerous studies using primary mouse hepatocytes (Lin et al., 1990) have shown that the mouse SAP gene can be induced by the direct action of either IL-1 or IL-6. Thus, both CRP and SAP are usually considered to be class 1 acute-phase gene products.

Serum levels of SAP in mouse start to rise approximately 8 h after an acute-phase stimulus and peak at 24-48 h (Pepys, M.B., 1979) the serum half-life is 7-8 h and is independent of the SAP level (Baltz, Dyck & pepys, 1985). Increases are therefore proportional to the rate of synthesis and secretion of SAP rather than to changes in clearance. One of the most potent stimuli to SAP production is bacterial endotoxin. Recently it has been shown that this effect is mediated via certain cytokines produced by activated monocytes and macrophages (Le, Muller & Mortensen, 1982).

In this study, the data showed that there was no expression for SAP genes in normal uninfected rat’s liver while SAP expression markedly increase in liver tissue after the bacterial infection at one, two and seven days. Also, in the present study, no expressions for SAP genes were observed in heart tissue after bacterial infection when compared to the control. These findings are contradicted with those of Zhiqing et al, 2010 which indicates that SAP is produced by non-hepatic cells (Hirano et al, 1990).

In summary, our data demonstrate that in rats, during S. faecalis infection, the expression of SAP is rapidly and strongly in liver parenchyma but not in the heart during the bacterial induced acute phase response. In this regard, SAP might be referred to as an early intracellular acute phase protein. Taken together, these data point to an important role of SAP in cytoprotection of hepatocytes, which might become damaged during their intensive function under inflammatory conditions.

Indirect evidence for LBP participation in sepsis comes from clinical investigations showing that levels of LBP in several body fluids were increased significantly in patients with bacteremia, sepsis, major surgery and acute respiratory distress syndrome (ARDS), and were associated with clinical outcome. (Cermak et al., 2005). Lipopolysaccharide-binding protein (LBP) is a protein that is up-regulated and secreted in stress situations. Its exact role in modulating the cellular response to endotoxin remains unclear. As an acute phase response, protein LBP is especially interesting because of findings indicating a potential dual role in the regulation of LPS action on target cells: low baseline level concentrations of LBP have been shown to enhance LPS potency, whereas the high concentrations seen during the acute phase can decrease the suppressive effect of LPS by shutting it into the HDL clearance pathway. It has been demonstrated in a mouse model that the high serum concentrations of LBP are protective in the setting of sepsis and bacteremia [Yunis, and Whitehead, 1990]. LBP is present at low levels (ca. 100–200 ng/ml in the blood) under normal conditions whereas it is highly increased during infections (ca. 10–20 µg/ml) to neutralize LPS for the prevention of overwhelming inflammation (Zweigner et al., 2001).

Liver is considered to be the central organ regulating this acute phase response by releasing specific acute phase protein including LBP which has been defined as a class I acute phase protein. (Kushner, 1988). Previous data showed that LBP mRNA expression in liver and kidneys markedly increased after burns. (Marnell et al, 2005) LBP expression in several cell types was demonstrated in vitro to be dramatically up-regulated by exposure to pro-inflammatory cytokines including IL-1β, IL-6, and TNF-α. (Calabro et al, 2003)

Our data showed that there was no expression for LBP genes in normal uninfected rat’s liver while LBP expression markedly increases in liver tissue compared to GAPDH after the bacterial infection at one, two and seven days. These findings are in harmony with those of (Grace et al., 1994) and dog (Usui et al., 1998), which stated that LBP gene expression occur in tissues such as liver, kidney, spleen and lung was reported in rat. (Ciliberto et al, 1987) Also, this agree with the studies of Su et al., 1994 as they reported that rat LBP mRNA is expressed mainly in the liver and slightly in kidneys and lungs, and that its levels increase markedly in the liver, kidneys and lungs in bacterial infections (Maniatis et al., 1987). Rat and rabbit models as well as in-vivo experiments with primary human hepatocytes and human hepatoma cell lines demonstrated that LBP is a secretory class 1 acute-phase protein whose gene is transcriptionally activated by the acute-phase response element/signal transducer and activator of transcription 3 (APRF/STAT3) and other cytokine-inducible nuclear proteins [Ohnishi et al, 1988], where Our study indicated that the predicted signal p for the predicted protein sequence of rat LBP is a non-secretory protein.

There was no expression for LBP genes in normal uninfected rat’s heart in our study while we found that LBP genes expression pattern increased after 24 and 48 hours post infection and no expression for the gene was detected after seven days post infection. These findings can be supported by the findings of Dentener et al, 2000 which stated that the transcriptional regulation of the LBP gene is induced by interleukin (IL)-1 alone or synergistically by IL-1 and IL-6, leading to a maximal LBP concentration within 24-48 h after
stimulation, a response that can be strongly enhanced by tumor necrosis factor (TNF)-α and dexamethasone [Whitehead et al., 1990].

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