Lipopolysaccharide-induced acute phase response: New views on trace metal plasticity in hepatic inflammation

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The term "acute phase" was originally introduced by Avery and colleagues to refer to serum obtained from patients who were acutely ill with infectious disease and which contained C-reactive protein (CRP) [1,2]. The acute phase reaction (APR) is a systemic reaction of the organism to local or systemic disturbances in its homeostasis caused by infection, tissue insult, trauma, or immunological disorders initiated and regulated by a large subset of cytokines. Most strikingly, the pattern of hepatic protein synthesis is drastically altered resulting in an increase of some plasma proteins. Generally, proteins whose concentration may increase by as much as 25% during the first week of injury or infection are classified as positive APR reactants [3]. At the same time, the production of several other proteins, i.e. negative APRs, is compensatory reduced. The measurement of positive and negative APR reactants as disease sensor is clinically useful because they are sensor of inflammatory disease [4,5]. The application of bacterial lipopolysaccharides (LPS), also known as endotoxin or lipoglycan, is a well-established experimental model for inducing APR in rodents [6]. Although the sensitivity toward LPS differs by several orders of magnitude between humans and mice, the quantitative and qualitative pattern of most APR reactants and associated clinical symptoms are similar [7,8].

Previously it was demonstrated that zinc deficiency enhances the APR, inflammatory response, and expression of APR reactants in vitro and in vivo through up-regulation of the JAK-STAT3 pathway [9]. Similarly, hypophosphatohemia clinically provoked stronger APR in critically ill patients by favouring the development of complications by impairing high-energy substrate availability for host defense and other cell functions [10]. Contrarily, severe iron deficiency blunted the response to LPS by blunting the activation of the NF-κB pathway, induction of NF-xB responsive genes in macrophages and affecting hepcidin expression in hepatocytes [11]. In addition, modulator effects on ARP in cultured hepatocytes were described for selenium [12,13]. All these studies suggest that the status of diverse metals or metalloids impact the APR amplitude. However, a detailed analysis of trace metal kinetics after application of LPS was not reported yet.

In previous studies, my group optimized laser ablation inductively coupled mass spectrometry (LA-ICP-MS)-based protocols for trace metal imaging in liver sections from experimental Wilson's disease models and clinical samples of patients suffering from this inherited disorder [14-16]. These protocols have multi-element capability and allow the quantification of a large variety of metals and metalloids within the tissue with high spatial resolution. In brief, LA-ICP-MS is based on the ablation of tissue material by use of a finely focused laser beam. Subsequently, the ablated material is transported into the inductively coupled plasma source of a mass spectrometer using an inert carrier gas stream. There it is vaporized, atomized and ionized and the different elements are separated according to their mass/charge ratio [17]. By use of this methodology, we showed that copper age-dependently accumulated in livers and brains of Atp7b deficient mice and simultaneously induced imbalances in hepatic iron and zinc content suggesting that the LA-ICP-MS technology is powerful and innovative in displaying complex changes in metal composition within a tissue [16].

The LA-ICP-MS technology was used here to analyse hepatic alterations in trace metals after LPS challenge. The endotoxin was injected into C57BL/6 wild type mice (n=3 per group) via the intraperitoneal route at 1 mg/kg body weight using standard operating procedures [6]. Control animals received an identical volume of vehicle (i.e. 0.9% NaCl). After one and six hours, the animals were sacrificed and the livers removed and snap frozen. 30 µm thick tissue cryosections for LA-ICP-MS measurements of tissue were prepared and analyzed essentially under conditions reported before [14,15]. Images of element distributions were generated in Microsoft Excel and visualized with the help of the Excel Laser Ablation Imaging (ELAI) visualization tool as described before [18]. Standards were prepared from homogenized liver tissue that was spiked with varying concentrations of standard salt solutions.

While in this analysis, the hepatic concentration of manganese (Mn), magnesium (Mg), iron (Fe), copper (Cu), and zinc as images by LA-ICP-MS showed no gross alterations after LPS challenge, the content of phosphorus (P) showed a significant increase already 2 hours after LPS injection that sustained for over 6 hours (Figure 1). The increase in P corroborated with increased hepatic potassium (K) and a subsequent decrease in hepatic sodium (Na).

In my view this preliminary pilot study demonstrates that LPS challenge in mice leads to an orchestrated program affecting hepatic concentrations of P, K and Na. Possibly, priming with LPS first provokes a redistribution of systemic P into the liver followed by a subsequent relocalisation of K acting as a positive counter-ion to restore the overall electric charge. Excess influx of K might be compensated in a second step by an efflux of Na.

It is tempting to speculate that the redistribution of P from the periphery into the liver might be a necessity to initiate the different APR signalling cascades majorly activated by phosphorylation of...
Figure 1. Element measurement by LA-ICP-MS in normal and inflamed mouse liver. Murine liver specimens were isolated two or six hours after LPS injection. Animals that received an equal volume of 0.9% NaCl served as controls. The concentration and distribution of Na, P, K, Mn, Mg, Fe, Cu, Zn, and C was imaged by LA-ICP-MS. Light microscopic (LM) photographs of the respective liver specimens at a magnification of 4x are depicted for orientation. The determined element concentrations are visualized in a colour scale and are given in µg/g liver tissue. C used for normalization is given in %. More information about sample preparation, measurement, and data analysis are given elsewhere [16].
transcriptions factors rapidly resulting in production of acute phase proteins [6,19]. Alternatively, the observed pattern of element influx and efflux is secondarily provoked by diverse subsets of inflammatory cells infiltrating the inflamed tissue after endotoxemia [20].

To sum up, this study has shown that LPS injection in mice provokes significant element distributions within the liver. This set of experiments is far beyond to clarify the significance and the exact biological mechanisms leading to respective redistributions. However, the measurements depicted in Figure 1 indicate that the LA-ICP-MS technology allows documenting and quantifying respective changes.

In future studies, I and my research team will in more detail address the precise kinetics and biological consequences of metal alterations within hepatic tissue after challenge with diverse inflammatory triggers. Possibly such "new views on trace metal plasticity in hepatic inflammation" investigations will help to develop new therapeutic avenues for treatment of overshooting APRs.

**Comment on animal manipulation**

All mice used in this study received humane care and were housed according to the guidelines of Institutional Animal Care and Use Committees and in accordance with governmental requirements.

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