Mini-Review

HNL Dimer- A Specific Biomarker of Neutrophil Activity - In Sepsis and Antibiotics Monitoring. A Mini-Review

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Abstract
Sepsis is a major burden to society and our health care systems. Early detection and treatment are keys to the improvement of the outcome of patients with sepsis with lower morbidity and mortality. We show in this review the potential usefulness, and superiority to current biomarkers, of plasma measurements of HNL Dimer in the monitoring of antibiotics in patients with sepsis. Future studies should be initiated to confirm this important aspect of sepsis management.

Keywords: Sepsis; Human Neutrophil Lipocalin (HNL)

Sepsis is a major burden to society and our health care systems. The global incidence is estimated by WHO to be about 50 million/year with 11 million deaths [1]. In the USA and most other developed countries sepsis is the costliest disease with estimated annual costs in the USA of about 24 billion $US [2] and in Great Britain of about 11 billion £ (WHITENET CHARITABLE TRUST The Cost of Sepsis Care in the UK Final Report, 2017). Early detection and treatment are keys to the improvement of the outcome of patients with sepsis with lower morbidity and mortality. Early detection relies on the demonstration of bacterial growth and partly on the detection of raised concentrations in blood of biomarkers such as CRP, Procalcitonin, Heparin-Binding Protein and others. However, none of these
biomarkers fulfil the expectations of the caring doctor since the discrimination between sepsis and other systemic inflammatory disease (SIRS) mostly is inadequate with great overlap in plasma concentrations and hence specificities and sensitivities in the range of 70-80% at the most [3-8]. Other important aspects of sepsis care are estimation of prognosis and the monitoring of adequate antibiotics treatment [9-13]. The latter is a great challenge since the septic patients very often are multi-sick having other complicating diseases that may affect biomarker levels. Also, the pathophysiology of sepsis is complex although the key player is the neutrophil granulocyte and its role in the eradication of the causing agent when the sepsis has a bacterial cause, which is the case in the vast majority of sepsis cases.

On activation neutrophils secrete a number of granule proteins, which can be measured in bodily fluids. One of these proteins is HNL (Human Neutrophil Lipocalin) also known as NGAL (Neutrophil Gelatinase Associated Lipocalin) and Lipocalin 2. HNL exists in several forms i.e. the monomeric form with a molecular weight of 22 kD, a dimeric form of 45 kD and as heteromeric forms at >90 kD of which one form is a complex between HNL and MMP9 (Gelatinase) [14, 15]. The origins of these forms are neutrophils and/or epithelial cells. The neutrophil produces and secretes all of these forms whereas epithelial cells only produce and secrete the monomeric form. Thus, the presence of the dimeric form in any human material, blood, urine, feces etc. indicates the presence and activation of neutrophils. We developed an HNL ELISA based on two of our monoclonal antibodies, a combination of antibodies that theoretically only recognizes epitopes in the dimeric form of HNL. This assumption was confirmed by the fact that the ELISA only detected the purified dimeric HNL and not at all purified monomeric HNL (Figure 1), hence the name HNL Dimer ELISA.

The HNL Dimer ELISA has been used in several studies to indicate the activation and involvement of neutrophils in the particular process. In a recent study on COVID-19 patients the assay was used to investigate the role of neutrophils in the development of AKI (Acute Kidney Injury) [16]. Indeed, the activities of neutrophils as measured by the HNL Dimer ELISA were closely related to the outcome of the COVID-19 patients in terms of severe AKI and survival. Thus, the relative risk of acquiring severe AKI was 8.99 (95% CI 2.91-31.81) times greater in those patients having plasma HNL Dimer above 13.2 μg/L at admission to the ICU as compared to patients with concentrations below the cut-off. The relative risk of 30-day mortality was 4.60 (95% CI 1.39–16.91) times higher in patients above that cut-off. A close relation of increasing plasma concentrations of HNL Dimer to increasing SOFA scores further emphasized the predictive potential of this biomarker in patients with SARS-CoV-2 infections.

In order to reflect the involvement of neutrophils in sepsis we measured plasma concentrations of HNL Dimer in the Bio-x [17, 18] and PEAK [19-21] studies. We show in figure 2 the distribution of HNL Dimer in plasma of Bio-x patients with sepsis at admission to the hospital but before antibiotics treatment. On average the concentrations were 5-10 times higher in patients with sepsis when compared to the concentrations found in healthy people. The distinction of plasma concentrations of HNL Dimer between bacterial sepsis and viral pneumonia/influenza is shown in the figure. The overlap in HNL Dimer concentrations was low, which is also reflected by the very high AUC ROC of 0.95 and the sensitivity and specificity given by the Youden index.
of 13.9 μg/L of 81% and 94%, respectively. The diagnostic performance, however, was very similar to plasma concentrations of PCT, but somewhat higher than those of CRP. The distinction between sepsis and other serious bacterial infections such as bacterial pneumonia is also shown in the figure. Here we see a somewhat greater overlap and translated into ROC analysis the AUCs were 0.81 and 0.80 for HNL Dimer and PCT, respectively. In this regard the performance of CRP was clearly inferior (0.65).

In the Bio-x study the results were from patients with microbiologically confirmed diagnosis only and samples were obtained before start of antibiotics treatment. These are important points in the evaluation of the diagnostic and predictive performances of the biomarkers, since the neutrophil biomarkers in particular, as will be shown below are very sensitive to antibiotics treatment and will normalize rapidly by adequate treatment. Therefore, the evaluation of the diagnostic or predictive performances days after start of antibiotics treatment will most likely underestimate the performances of such biomarkers. From the Bio-x study we conclude that the diagnostic performances of HNL-Dimer and PCT are quite similar when estimations are based on admission data and samples from antibiotics naïve patients. The PEAK study followed, twice daily during the length of their stay, a large number of patients (n=198) admitted to the ICU [19]. Of these patients, 144 were allocated to the infection-group of which 59% were diagnosed as having sepsis and the rest (41%) as having SIRS (Systemic Inflammatory Response Syndrome). The plasma concentrations of HNL Dimer were highly raised at admission day to the ICU in the whole cohort as compared to healthy people (p<0.0001), but with a significant difference (p=0.02) between patients with SIRS and sepsis. Similarly, the PCT concentrations were highly raised as compared to healthy people, but with a larger difference between SIRS and sepsis (p=0.0005). The SOFA score is a much-used score for the prediction of outcome of sepsis. As shown in figure 3 the plasma concentrations of dimeric HNL in sepsis patients with SOFA score of more than 7 at sepsis day 0 were significantly higher (p=0.03), whereas PCT and CRP concentrations were not statistically significantly related to the SOFA score. Thus, the association to outcome in sepsis of HNL Dimer seemed somewhat higher than for the other two biomarkers.

Thirty-one of the PEAK patients were monitored almost daily after adequate antibiotic treatment [19]. Figure 4 shows the rapid and significant reduction in plasma concentrations of HNL Dimer (p=0.0003) with near normal concentrations obtained already after 24 hours of adequate antibiotics treatment. For comparison, the changes in plasma concentrations of CRP, PCR, Calprotectin and HNL Total are also shown. Only PCT showed a significant reduction in concentrations (p=0.04), but not until after four days of treatment. The other biomarkers were unaltered or even increased after adequate antibiotics treatment. In figure 5 we show individual changes of HNL Dimer (n=21) and PCT (n=18) in the patients with paired samples before and day 2 after antibiotics treatment. Our results indicate that neutrophil activity is rapidly turned off by adequate antibiotics treatment and the eradication of the bacteria. Thus, the assay of plasma HNL Dimer concentrations may be a unique means of monitoring the success of antibiotics treatment in sepsis and probably also in other bacterial infectious diseases. The slower response of PCT to successful antibiotics treatment likely relates to the fact that the production of PCT is turned on in many different cells in the body of which some continue to produce PCT for some time after the inciting agent has
disappeared. This variability was indicated in Figure 5 in which PCT declined in almost half of the patients and increased dramatically in the other half as opposed to HNL Dimer that showed a more consistent decline after successful antibiotics treatment. We conclude from this minireview that plasma measurements of the dimeric form of HNL should be useful complements in the management of patients with sepsis and in particular in monitoring of successful and adequate antibiotics treatment of these patients. Thus, available data support further assessment of Dimeric HNL as a marker of adequate antibiotic initiation. The assay of the dimeric form of HNL also shows promise in the understanding of the role of neutrophils in serious infections such as SARS-CoV-2 and their role in organ damage of the kidney and the lung.

The specific detection of dimeric HNL by the HNL Dimer ELISA

![Graph showing the measurement of purified native HNL by the HNL Dimer ELISA. Only the dimeric form of HNL was detected.](image)

**Figure 1:** The measurement of purified native HNL by the HNL Dimer ELISA. Only the dimeric form of HNL was detected.
Figure 2: The left panel shows the plasma concentrations of dimeric HNL in sepsis as compared to the concentrations in plasma from patients with viral pneumonia/influenza, bacterial pneumonia and healthy non-infected controls. The top right panel shows ROC curves of sepsis vs viral pneumonia/influenza and the AUCs of the respective biomarker. The lower right panel shows the ROC curves of sepsis vs bacterial pneumonia and the AUCs of the respective biomarker. All samples were obtained before antibiotics treatment and the diagnosis was verified and confirmed by microbiological and clinical results.

Figure 3: The figure shows the relationship between HNL Dimer and PCT to SOFA score in patients with sepsis.
Plasma biomarkers in septic patients after adequate antibiotics treatment

**Figure 4:** The figure shows the kinetics of plasma biomarkers in septic patients treated with adequate antibiotics. The left panel shows the results of HNL Dimer. The upper middle panel shows the results of PCT, the upper right panel the results of HNL Total, the lower middle panel the results of CRP and the lower right panel the results of Calprotectin. Medians and 95% confidence intervals are shown.

**Figure 5:** The figure shows the change from before antibiotics to day 2 of adequate antibiotics treatment in HNL Dimer and PCT concentrations in plasma in patients with sepsis.
Conflict of Interest
Per Venge owns the worldwide patent of measuring HNL in infectious disease

References


