A brief, highly selective history of acute phase proteins as indicators of infection, inflammation and injury

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Abstract
There is an array of plasma protein alterations that occur in a wide variety of species, including humans in response to trauma, inflammation and infections, seemingly irrespective of etiologic agent. In numerous species, these plasma proteins are part of the innate immune response. In addition, it appears that a number of the plasma proteins in this array can be predictive of morbidity and/or mortality. We propose that based on historic use, selected acute phase proteins should be included in ongoing and future non-clinical and clinical studies to help us better understand disease progression in chronic, as well as acute diseases. In addition to assess if there is a relationship between vaccine-induced inflammation and degree of protection from live, attenuated or synthetic vaccines.

Keywords Acute phase proteins · Animal models of disease · Chronic disease · Covid-19 · C-reactive protein · Innate immunity · Morbidity and mortality · Trauma and sepsis · Vaccination

The present value of C-reactive protein as an indicator of the presence and severity of infection.

C-reactive protein (CRP) has long been a highly sensitive indicator of the presence and severity of infection, as well as efficacy of treatment in patients. A high-sensitivity C-reactive protein (hs-CRP) test, which is more sensitive than a standard test, also can be used to evaluate the risk of developing coronary artery disease.

In a recent study of 209 patients’ findings show that only C-reactive protein (CRP), in the multivariate analysis, was significantly associated with the progression to a severe case of COVID-19 (OR, 1.056; 95% CI, 1.025–1.089; p < 0.001). The data suggested that for every 1-unit increase in CRP level, the risk of developing severe events increased by about 5%. (Wang G, Wu C, Zhang Q, et al. (2020). This is consistent with the results of a review of 1896 survivors and 849 non-survivors cases of Covid-19 Sahu et al. (2020). In conjunction with eosinopenia, CRP can help triage patient patients presenting with fevers, sorting suspected COVID-19 patients from those with COVID-19-like symptoms. Lia et al. (2020).

The potential of proteomics in disease diagnosis and therapeutic monitoring

As the authors of a review on human plasma proteins state, “The human plasma proteome holds the promise of a revolution in disease diagnosis and therapeutic monitoring provided that major challenges in proteomics and related disciplines can be addressed.” (Anderson & Anderson, 2002). The authors list 289 proteins whose detection in plasma or serum is documented in the literature. These include enzymes released from tissue as a result of damage, enzyme inhibitors, a series of apolipoproteins, coagulation factors, complement components, immunoglobulins, cytokines and cytokine inhibitors, hormones, as well as albumin and a series of proteins that decrease or increase in response to inflammation, infection and injury. It is this last group that we will primarily focus on. The proteins in this latter group include albumin and transferrin which decrease during infection, inflammation and injury while alpha-1-acid glycoprotein, alpha-1-antitrypsin, alpha-2-macroglobulin, C-reactive protein, ceruloplasmin, haptoglobin, serum amyloid A and fibrinogen increase.
Acute phases proteins possible role in vaccination

A recent review describes a number of acute phase proteins and their importance in vaccination (Rafaat et al., 2020). These authors discuss in some detail the known and likely function of these proteins, a number of which are associated with response to vaccination. They argue that there is an association between the strength of the acute phase response and vaccination efficacy of key importance to human and veterinary medicine. They cite research in which the absence of interleukin-6, a potent inducer of acute phase proteins, increases the mortality of a live vaccine strain of F. tularensis in mice (Kurtz et al. (2013), unfortunately the plasma protein profile was not evaluated to see which proteins may have been altered.

As a separate part of the review, the authors discuss the acute phase protein response to adjuvants. The effects of adjuvants in rabbits indicated that CRP and fibrinogen expression levels were increased following the administration of adjuvant systems AS01 (monophosphoryl lipid A (MPL) and QS-21), AS03 (α-tocopherol and squalene in an oil-in-water (o/w) emulsion), AS15 (CpG 7909, monophosphoryl lipid, and QS-21) and DTPw (monophosphoryl lipid A) (Leroux-Roels, 2010). CRP levels increased 9–26-fold after injection of AS01, AS03, or AS15. Not all adjuvants cause an increase in inflammatory response (Destexhe et al. (2013) The use of other adjuvants such as aluminum phosphate and aluminum hydroxide did not affect CRP expression levels in the elderly or in the young following a diphtheria-tetanus-poliomyelitis-typhoid vaccine (Yousfi et al., 2005). Overall, these studies suggest the acute phase protein response in general, and that of CRP in particular, might be a marker for safety and perhaps efficacy in clinical and non-clinical vaccine trials. It would be interesting to test whether acute phase protein alterations after vaccination correlate with protection and/or antibody titer.

A non-clinical study of acute phase proteins in naïve and vaccinated rats

The notion of using acute phase proteins to monitor the response to vaccines, as well as the efficacy of vaccination is not new. Naïve rats inoculated ip with a live vaccine strain (LVS) of tularemia (Francisella tularensis) displayed decreases in serum zinc concentration and increases in acute phase proteins relative to increasing dose of microorganism. The vaccine strain, safe at a 4 log dose was lethal at 8 logs. The virulent strain (SCHU S4) was uniformly lethal at a 4 log dose in un-immunized rats. In contrast, rats immunized with 4 logs of the vaccine strain were protected when challenged with 4 or 8 logs of the virulent strain.

At the 4 log challenge dose, all vaccinated animals survived and there were no changes in serum zinc or acute phase concentrations. However, at the 8 log dose, though all animals survived, there were significant changes in serum zinc and acute phase proteins which persisted at least 4 days post challenge. Thus, by monitoring these biomarkers in naïve animals exposed to various dose of candidate vaccines and in immunized animals exposed to a range of doses of the etiologic agent, one may be able to select a dose that optimizes protection while minimizing the reaction to vaccination. (Powanda et al 1975)

It should also be noted that both LVS and SCHU S4 caused pyogranulomas in liver and spleen of nonimmune rats. Nonimmune rats given 10(4) SCHU S4 organisms did not survive beyond 72 h, but immune rats given challenge inoculum of 10(8) SCHU S4 organisms developed lesions and survived. Larger doses of LVS resulted in earlier onset of characteristic hepatitis and splenitis in nonimmune rats. Periportal lymphocytic infiltrates were present in the liver 48 h after SCHU S4 challenge inoculation of immune rats and 96 h after inoculation of LVS in nonimmune rats and were associated with intense macrophage aggregation. These changes indicate that the pathogenesis of tularemia is a result of the interdependency of the dose and virulence of the causative agent with the immune status of the host and that cellular immunity has a significant role in the response of the rat to tularemia. (Moe JB et al. (1975). The tissue damage associated with tularemia is related to it being able to infect the liver eliciting the innate immune response which includes changes in acute phase proteins. These data raise the question whether and to what degree is inflammation necessary for a live vaccine to effectively immunize? Is concomitant inflammation necessary for all vaccines, live, attenuated or artificial, such as the newly available mRNA vaccines for Covid-19, to effectively immunize? In some cases, adjuvants themselves can cause inflammation; again is inflammation required for maximum efficacy?

Use of acute phase proteins in testing anti-viral drugs

Acute phase proteins can also be used to test whether proposed therapies induce inflammation. Polyriboinosinic-polycytidylic acid complexes [poly(I)-poly(C)] are potent inducers of interferon in rodents and rabbits. In primates, however, these compounds are poor interferon inducers, apparently because the complex is rapidly degraded by plasma nucleases. A modified, nuclease resistant form of
poly(I)-poly(C) has therefore been developed for use in man. This modified poly(ICLC), induces interferon in chimpanzees and rhesus and cynomolgus monkeys. Poly(I)-poly(C) itself, in addition to inducing interferon, also causes fever and depressions in plasma zinc A study was conducted in rats to see whether there was a dose effect and whether significant changes in plasma proteins occurred. Repeated dose of 0.3 and 3.0 mg/kg poly(ICLC) were given over a 48 h period. Both 0.3 and 3.0 mg/kg poly(ICLC) caused increases in seromucoid (primarily consisting of alpha-1-acid glycoprotein) and haptoglobin up to five fold at 48 h. A 2 to three-fold increase in haptoglobin, and a dose dependent increase in alpha -2-macrofetoprotein. (Powanda et al., 1977).

**Some caveats about choice of models and relevance to humans**

While acute phase proteins may be used to assess the inflammation potential of a drug or vaccine, the animal model chosen will materially affect which plasma proteins are responders. (Watterson et al., 2009). For example, C-reactive protein and serum amyloid A are major reactants in humans, while in rats, it is alpha-2-macroglobulin. Both humans and rats have alpha-1-acid glycoprotein, haptoglobin and fibrinogen as moderate reactants, as does dog and mouse. (Watterson et al., 2009) This difference in major and moderate reactants has been observed in other species (Cray C, Zaias J, Altman et al. 1983). Moyer ED, Powanda MC (2004). Even in healthy chickens mRNA coding for alpha-1-acid glycoprotein (AGP), serum amyloid A (SAA), PIT54, C-Reactive protein (CRP) and Ovotransferrin (OVT) could be found in multiple extra hepatic tissues. The mRNA coding for CRP, OVT and SAA was detected in all 20 analyzed tissues with a higher expression in gastrointestinal tract, respiratory and lymphatic samples. ACP positive cells were present in the epithelium of the mucosal layer of gastrointestinal tract and kidney (Marques et al., 2017). These data suggest that with so many cell types containing mRNA for these acute phase proteins, the innate immune response to inflammation, infection and injury could be localized, as well as a systemic.

Though CRP in itself is a good marker for the presence and severity of disease, in a porcine model challenged with one of three bacterial agents, one parasitic and one viral agent, the use of a combination of acute phase proteins allowed the detection of disease with more sensitivity than any individual acute phase protein. (Heegard PMH et. al (2011). Use of multiple acute phase proteins in monitoring trauma-septic patients

In a study of trauma-septic patients indicated that select plasma proteins, α1-acid glycoprotein and ceruloplasmin, along with some amino acids and catabolites, allowed one to identify, up to 9 days before their demise, patients who would not survive with 99% certainty from a single plasma analysis profile (Moyer et al., 1981). There is a reprioritization of hepatic plasma protein release in trauma and sepsis wherein trauma patients who become septic have C-reactive protein, fibrinogen, ceruloplasmin, and α1-antitrypsin levels that continued to be elevated after the initial five posttrauma days, while transferrin, albumin, and α2-macroglobulin levels fell. This reprioritization response seems to be both a predictor of sepsis as well as a measure of the adequacy of the host response to trauma and sepsis (Sganga et al., 1985).

A small study of plasma protein changes in burned patients with and without the complication of infection indicated that the ratios of select plasma proteins, rather than the absolute values, provided a more effective discrimination among those patients who were burned (B) versus those with infection (BI) and versus those who were not only burned and infected, but also had complications requiring ventilatory support and/or some degree of renal impairment (BIC). Stepwise discriminant analysis indicated that the B, BI and BIC patients could be differentiated on the basis of α1-acid glycoprotein (AG) and haptoglobin (Hp) concentration in combination. When the data were ranked ordered and subjected to analysis of variance, the ratios of AG/Hp and AG/transferrin were found not only to discriminate B from BI patients, but also BIC from BI patients. (Powanda, MC, Moyer ED, Wilmore D et al. 1983; Moyer ED, Powanda MC 2004).

Many of these acute phase proteins appear to have significant roles in wound healing. Alpha-1-antitrypsin and alpha-2-macroglobulin accumulate at the site of injury presumably to prevent additional damage due to proteolytic enzymes released from already damaged tissue, as well as from phagocytic cells. Alpha-1-acid glycoprotein due to its high-carbohydrate content may act as an adhesive at the wound site. It also can inhibit phagocytosis and lymphocyte transformation and thus could protect the healing wound against the development of auto immunity. Ceruloplasmin transfers copper to cytochrome C oxidase vital to aerobic energy production, which along with glycolysis increases during wound healing. Ceruloplasmin and the copper it carries are also essential to collagen formation. Haptoglobin increases four to five times its normal plasma concentration following injury and is frequently found in inflammatory exudates. The principal biologic function of...
this protein appears to be the removal of free hemoglobin released by hemolysis. (Powanda and Moyer, 1981; Jan-
ciauskiene and T, Mahadeva R, 2011).

**Acute phase proteins as a monitor of chronic disease**

Might plasma proteins act as a biological dosimeter for cumulative stress, repeated infection, injury or inflammation and/or as an indicator of physiologic age? Data on the levels of acute phase proteins and plasma protein synthesis rates in adult vs. old rats (Papet et al., 2003) may prove instructive. Older rats (22 months) have increased levels of total proteins, even though albumin concentrations are lower, than adult (8 month) rats. There is also a doubling of fibrinogen, a 4.5-fold increase in α2-macroglobulin and a 35% increase in α1-acid glycoprotein concentration. There also is an increase in both the fractional and absolute synthesis rates for plasma proteins in the old rats, including for albumin, despite the decrease in its absolute concentration. Thymus weight decreased and spleen weight increased in old versus adult rats. These data are consistent with the “inflamm-aging” concept. (Franceschi et al., 2000).

Age and sex appear to profoundly affect plasma protein concentrations, at least in Gottigen minipigs. Blood samples were taken at 6, 16, 24 and 40–48 weeks. Significant changes by sex x age were seen in CRP, haptoglobin, pig major acute phase protein (PMAP) and albumin. Porcine α-1 acid glycoprotein (PAGP) displayed highly significant (p < 0.001) effects of age and gender. CRP, haptoglobin, pig major acute phase protein generally decreased rather than increased with age. Albumin increased with age. (Christofersen et al., 2015) Changing housing conditions (not specified) caused PAGP, CRP, PMAP and haptoglobin to increase (p < 0.05) and albumin to decrease (p < 0.05). Are the differences seen in this study versus that of (Papet et al., 2003) due to a difference in species, rat versus minipig? If so, which species better mimics the human response to age?

**Summary**

To sum up, despite their past use in non-clinical studies in a variety of species with various infectious and inflammatory agents, and in clinical studies, to predict morbidity and mortality, (Powanda, Beisel WR (2003) most acute phase proteins today are not extensively monitored. Considering the role of many of these plasma proteins in the innate immune response in numerous species, this may need reconsideration (Romo MR, Perez-Martinez D, Ferrer CC (2016). Perhaps, it is time to reassess whether it may be beneficial if widespread screening of many, if not all, of these proteins should be done in a variety of populations of different ages and defined health status. Were this done the following hypotheses could be tested.

If one, or more than one, plasma protein is used as a biomarker, will that provide a better understanding as to whether the existence and extent of inflammation following vaccination correlates with antibody titer, T cell mobilization and/or protection?

Can a constellation of plasma proteins either by absolute value, by ratios or by altered glycosylation be found that will act as a biological dosimeter with regard to cumulative stress, repeated infection, injury or inflammation? Would such biomarkers be of use in monitoring long-term Covid and help sort which patients might benefit from vaccination?

Can a constellation of plasma proteins either by absolute value, by ratios or by altered composition be found that will act as an indicator of physiologic age, as opposed to chronologic age? A comparison with changes in telomere length which alters with age (Shammas MA (2011) would be interesting.

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