Serum Markers Associated with Severity and Outcome of Hantavirus Pulmonary Syndrome

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**Background.** Hantavirus pulmonary syndrome (HPS) is caused by Andes virus (ANDV) and related hantaviruses in the Americas. Despite a fatality rate of 40%, the pathogenesis of HPS is poorly understood and factors associated with severity, fatality, and survival remain elusive.

**Methods.** Ninety-three ANDV-infected HPS patients, of whom 34 had a fatal outcome, were retrospectively studied. Serum levels of cytokines and other inflammation-associated markers were analyzed using multiplex immunoassay and enzyme-linked immunosorbent assay. Associations with disease severity, fatal outcome, and survival were identified using logistic regression.

**Results.** HPS patients exhibited increased serum levels of markers associated with inflammation, intestinal damage, and microbial translocation compared to controls. Patients with fatal outcome displayed higher levels of interleukin (IL) 6, IL-10, interferon-γ, soluble tumor necrosis factor-related apoptosis-inducing ligand, and intestinal fatty acid–binding protein (I-FABP) than survivors. Levels of complement factor 5/5a were higher in survivors compared with fatal cases. IL-6 and I-FABP, the latter a marker for intestinal damage, were by multivariate analyses identified as independent markers associated with disease severity (odds ratio [OR], 2.25; 95% confidence interval [CI], 1.01–5.01) and fatal outcome (OR, 1.64; 95% CI, 1.01–2.64), respectively.

**Conclusions.** HPS patients displayed a multifaceted, systemic inflammatory response, with IL-6 and I-FABP as independent markers of disease severity and fatality, respectively.

**Keywords.** hantavirus pulmonary syndrome; cytokines; microbial translocation; intestinal fatty acid–binding protein; IL-6.

Hantaviruses comprise a family of negative-sense, single-stranded RNA viruses belonging to the Bunyaviridae order. Certain members of the Hantaviridae family cause severe disease in humans: hantavirus pulmonary syndrome (HPS) in the Americas, and hemorrhagic fever with renal syndrome (HFRS) in Europe and Asia [1]. HPS is primarily caused by Andes virus (ANDV) in Argentina and Chile, and Sin Nombre virus in North America [1]. The first recognized HPS outbreak occurred in the Four Corners region of the United States in 1993 [2], and soon after, HPS was discovered also in South America [3]. Hantaviruses are continuing to cause sporadic outbreaks throughout the Americas. For example, the 2012 outbreak in Yosemite National Park, California, received worldwide attention [4]. With a fatality rate of 35%–40%, HPS represents one of the deadliest infectious human diseases [1, 5]. Yet, no US Food and Drug Administration (FDA)–approved vaccine or specific treatment exists.

Hantaviruses are normally transmitted to humans through inhalation of virus-contaminated rodent excreta [1], making HPS and HFRS zoonotic infectious diseases. While hantaviruses are in general not transmitted between humans, several cases of interhuman ANDV transmission have been reported [1, 6]. HPS patients initially present with flulike symptoms, such as fever and headache, as well as gastrointestinal symptoms including abdominal pain and diarrhea [7–9]. These symptoms are soon followed by a rapidly progressing pulmonary edema, often putting patients in life-threatening respiratory failure, and in the most severe cases cardiogenic shock leading to death [1, 8].

While the mechanisms behind HPS pathogenesis are poorly understood, hantavirus-induced immune responses have been suggested to drive immunopathology [1]. Immune responses upon hantavirus infection include strong inflammatory responses...
responses displayed by increased levels of proinflammatory cytokines and vigorous natural killer (NK) cell, CD8 T-cell, and B-cell responses [10–15]. However, studies providing more in-depth knowledge on how immune responses during hantavirus infection might affect the disease outcome are lacking, hampering the development of HPS interventions and treatments.

Here, we retrospectively characterized the systemic inflammatory response in the largest cohort of HPS patients analyzed to date, including 93 HPS patients, among whom 34 had a fatal outcome. With the aim of identifying correlates of fatal outcome and survival, we further identified serum markers associated with the respective outcomes, using logistic regression. Our findings strengthen previous observations [9, 14], showing that HPS patients display high levels of proinflammatory cytokines in general. We here present findings that delineate inflammatory responses in HPS patients with fatal vs nonfatal outcome. Most importantly, we show that markers of intestinal damage and microbial translocation are increased during HPS, and that multiple serum markers, including interleukin (IL) 6 and intestinal fatty acid–binding protein (I-FABP), are associated with severity and fatal outcome.

MATERIALS AND METHODS

Study Design and Patients

Serum samples from 93 HPS patients with confirmed ANDV infection were included in the study. Samples were collected for diagnostic purposes at the Hantavirus National Reference Laboratory in Buenos Aires, Argentina, during 2010–2016, and stored at –80°C until analysis. Ethical approval was obtained from the institutional ethical committee at Administración Nacional de Laboratorios e Institutos de Salud “Dr Carlos G. Malbrán” in Buenos Aires, Argentina. HPS was confirmed by enzyme-linked immunosorbent assay (ELISA) detecting ANDV-specific immunoglobulin M and immunoglobulin G antibodies, as previously described [16], or by quantitative reverse-transcription polymerase chain reaction detecting ANDV RNA. In addition, serum from 33 community-matched controls was collected under informed consent.

Multiplex Immunoassay

A custom 15-plex Magnetic Luminex Screening assay (R&D Systems) was used for measurement of IL-1β, IL-2, IL-6, IL-10, IL-12 (p70), IL-15, IL-18, tumor necrosis factor (TNF), interferon gamma (IFN-γ), B-cell activating factor (BAFF), complement factor 5 (C5)/complement factor 5a (C5a), vascular endothelial growth factor (VEGF), soluble TNF-related apoptosis-inducing ligand (sTRAIL), granzyme A, and granzyme B according to the manufacturer’s instructions. Serum was diluted 1:2 prior to analysis.

Enzyme-Linked Immunosorbent Assay

Ferritin levels were measured in serum diluted 1:1000, using a commercially available human ferritin ELISA kit according to the manufacturer’s instructions (Abcam). Serum levels of lipopolysaccharide-binding protein (LBP), soluble CD14 (sCD14), I-FABP, and soluble CD25 (sCD25) were measured using DuoSet ELISA development kits (R&D Systems) according to the manufacturer’s guidelines. Serum was diluted 1:4000 for LBP, 1:2000 for sCD14, 1:5 for I-FABP, and 1:10 for sCD25 in ready-to-use ELISA diluent (Mabtech).

Statistical Analyses

Statistical analyses were performed using Graph Pad Prism version 7 and Stata version 13 software. Mann–Whitney U test was used for comparisons between 2 groups and Kruskal–Wallis test was used for comparisons of >2 groups. Spearman rank correlation coefficient was used for examining associations between serum markers. Principal component analysis (PCA) was performed for visualization of possible clusters, using the R package “scatterplot3d.” Serum marker concentrations used in the heatmap were standardized and clustered within each group.

Logistic regression analyses, adjusted for sex, age, and day of sampling, were performed using disease severity or fatal outcome as outcome and serum markers as predictors. The univariate analyses were carried out using standard logistic regression. The multivariate analyses were performed using penalized logistic regression (“ridge”). Ordinal logistic regression was used both for univariate and multivariate associations. Concentrations of IL-1β, IL-2, IL-6, IL-12, TNF, IFN-γ, VEGF, granzyme A, granzyme B, and I-FABP were log-transformed before analysis with PCA and logistic regression. Samples with concentrations below the detection limit were assigned a positive value smaller than the lowest detectable limit for that serum marker.

RESULTS

Study Cohort

Diagnostic serum samples from 93 HPS patients were collected at median of 6 days after reported symptom debut (range, 1–23 days). Two of the patients were diagnosed upon discharge and were thus excluded from logistic regression analyses. The HPS patients consisted of 27 females and 66 males at a mean age of 36 years (range, 7–85 years). Of the 93 patients, 34 had a fatal outcome. Patient characteristics are summarized in Table 1. Clinical data including white blood cell (WBC) count, platelet count, and hematocrit were available for a subset of the cohort (Table 1). In general, the HPS patients had increased WBC count and decreased platelet count, while the mean hematocrit was within normal range (Table 1). Patients were divided into 4 different severity groups, as previously described [13]: I, prodromal symptoms without respiratory involvement (n = 3); II, mild to moderate respiratory compromise without
hemodynamic compromise (n = 37); III, severe respiratory insufficiency with hemodynamic compromise (n = 19); IV, severe respiratory insufficiency with refractory-to-treatment hemodynamic compromise and a final fatal outcome (n = 34). The control group consisted of 21 women and 12 men, with a mean age of 37 years (range, 24–69).

**HPS Is Characterized by a Multifaceted Inflammatory Response**

To characterize the systemic inflammatory response during HPS, levels of inflammation-associated serum markers were measured in HPS patients and controls, using multiplex immunoassay and ELISA. For initial statistical analyses, HPS patients were divided into 3 groups, depending on the day post symptom debut at which they had been sampled (days 1–4, n = 38; days 5–10, n = 38; days 11–23, n = 17). Serum levels of IL-1β, IL-6, IL-10, IL-18, TNF, IFN-γ, BAFF, C5/C5a, sCD25, granzyme A, and granzyme B were all significantly increased in all HPS groups, compared to controls (Figure 1; Supplementary Table 1). Levels of IL-12, IL-15, and ferritin were significantly higher in HPS patients sampled on days 1–4 and days 5–10, compared with controls, whereas IL-2 levels were significantly elevated only in HPS patients sampled at days 1–4 post symptom debut (Figure 1; Supplementary Table 1). Levels of sTRAIL and VEGF did not differ between HPS patients and controls (Figure 1; Supplementary Table 1). Together, these findings confirm previous studies showing that HPS patients display high levels of proinflammatory cytokines and IL-10 [9, 14]. In addition, these data show that HPS patients have increased systemic levels of BAFF, C5/C5a, granzyme A, granzyme B, and ferritin.

**Markers Associated With Intestinal Damage and Microbial Translocation Are Increased During HPS**

A potential driver of inflammation is microbial translocation (MT), referring to the leakage of bacteria or bacterial products from the gastrointestinal tract into the systemic circulation [17]. Given the systemic inflammation and gastrointestinal symptoms of HPS patients [7, 8], we were interested to investigate if markers of MT would be increased during HPS. LBP and sCD14 have been widely used as surrogate markers of MT [17]. Moreover, I-FABP, often measured in the context of MT, is a systemic marker for intestinal epithelial cell injury [18]. Serum levels of sCD14, LBP, and I-FABP were measured in HPS patients and controls. Levels of sCD14 and LBP were elevated in all groups of HPS patients compared to controls (Figure 2A and 2B; Supplementary Table 1). I-FABP levels were increased in HPS patients sampled at days 1–4 and days 5–10 post symptom debut (Figure 2C; Supplementary Table 1). Levels of sCD14 and LBP in HPS patients were positively correlated (ρ = .54, P < .0001) (Supplementary Figure 1). Furthermore, levels of sCD14 and I-FABP were positively correlated to levels of the inflammation-associated cytokines IL-6, IL-10, IL-12, IL-15, TNF, and IFN-γ (Supplementary Figure 1). Pairwise correlations between all serum markers are summarized in Supplementary Figure 1. Taken together, these data indicate that intestinal damage and MT may occur during HPS.

**Levels of Multiple Serum Markers Differ Between Fatal and Nonfatal HPS**

To get an overview of the differences in responses between fatal and nonfatal HPS cases and controls, data were summarized using PCA and heatmap analysis (Figure 3A and 3B). PCA...
separated fatal HPS cases from nonfatal cases, albeit not as marked as total HPS cases from controls (Figure 3A).

Next, we sought to identify the specific serum markers discriminating patients with fatal vs nonfatal outcome. As only 1 of the patients who died was sampled later than day 10 (sampled at day 13) post symptom debut, groupwise comparisons were performed on patients sampled at days 1–4 (fatal, n = 19; nonfatal, n = 19) and days 5–10 (fatal, n = 14; nonfatal, n = 23) post symptom debut. Serum levels of IL-6 and I-FABP were significantly higher in patients with fatal compared to nonfatal outcome, both in patients sampled on days 1–4 and those sampled on days 5–10 post symptom debut (Figure 3C; Supplementary Table 2). In patients sampled on days 1–4, levels of C5/C5a were lower in patients with fatal outcome (Figure 3C; Supplementary Table 2). In patients sampled on days 5–10, higher levels of IL-10, IFN-γ, and sTRAIL were observed in fatal compared

Figure 1. Cytokines and inflammatory markers are increased in serum of patients with hantavirus pulmonary syndrome (HPS). Levels of serum markers in HPS patients, sampled either at days 1–4 (n = 38), days 5–10 (n = 38), or days 11–23 (n = 17) post symptom debut, and controls (n = 29–33). Kruskal–Wallis test; median values. *P < .05; **P < .01; ***P < .001; ****P < .0001. Abbreviations: BAFF, B-cell activating factor; C5, complement factor 5; C5a, complement factor 5a; IFN-γ, interferon gamma; IL, interleukin; sCD25, soluble CD25; sTRAIL, soluble TNF-related apoptosis-inducing ligand; VEGF, vascular endothelial growth factor.
with nonfatal cases (Figure 3C; Supplementary Table 2). No significant differences were found for IL-1β, IL-2, IL-15, IL-18, TNF, BAFF, VEGF, sCD25, ferritin, granzyme A, granzyme B, sCD14, or LBP (Figure 3C; Supplementary Table 2). Together, these findings show that levels of several serum markers differ between HPS patients with fatal compared with nonfatal outcome, suggesting that certain responses may be associated with HPS severity and fatality.

IL-6 and I-FABP Are Associated With Severe Disease and Fatal Outcome

The observed differences in serum marker levels between fatal and nonfatal HPS prompted us to further investigate for possible associations with disease severity and outcome. Univariate and multivariate regression analyses adjusting for sex, age, and day of sampling were performed, using severity and fatal outcome as dependent variables.

We first explored associations to the disease severity. Univariate analyses showed that increased levels of IL-1β, IL-6, IL-15, IL-18, ferritin, granzyme B, and I-FABP were associated with increased odds of severe disease, whereas C5/C5a was associated with decreased odds of severe disease (Figure 4A; Supplementary Table 3). Next, to identify serum markers that were independently associated with severe disease, multivariate analyses were performed. Interestingly, these analyses showed that IL-6 was the only marker independently associated with increased odds of severe disease (Figure 4B; Supplementary Table 3). The multivariate analyses further showed that increased levels of C5/C5a and BAFF were significantly associated with decreased odds of more severe disease (Figure 4B; Supplementary Table 3).

The patient outcome data allowed for additional logistic regression analyses, using fatal outcome as dependent variable. In this univariate regression model, IL-6, IL-15, IFN-γ, ferritin, granzyme B, and I-FABP were identified as markers associated with increased odds of fatal outcome (Figure 4C; Supplementary Table 3). In contrast, C5/C5a was associated with decreased odds of fatal outcome (Figure 4C; Supplementary Table 3). Interestingly, multivariate analyses identified I-FABP as the only marker that was independently associated with increased odds of fatal outcome (Figure 4D; Supplementary Table 3). In conclusion, univariate analyses revealed associations between multiple serum markers and HPS severity and fatality, while the multivariate analyses highlighted IL-6 and I-FABP as independent factors associated with increased odds of severe disease and fatal outcome, respectively.

DISCUSSION

While an adequate immune response is essential for proper clearance of pathogens, a deregulated, exaggerated immune response can give rise to immunopathology. HPS is an acute, severe disease with a high fatality rate associated with an excessive immune response [1]. Still, correlates of fatality and survival are poorly known, impeding the development of treatments. Better understanding of pathological immune responses could aid in the development of treatments for hantavirus infection and other viral hemorrhagic fevers. Here, we characterized the inflammatory response in a unique cohort of HPS patients and identified correlates of fatality and survival.

Our findings, demonstrating a broad and strong inflammatory response in HPS patients (Figure 1), extend previous observations [9, 14]. Moreover, we demonstrate increased levels of BAFF, granzyme A, granzyme B, C5/C5a, and ferritin during HPS. We also observed increased levels of IL-1β, which is in contrast to 2 previous studies, in which no differences were found compared to healthy controls [9, 14]. While we did not observe elevated levels of VEGF and sTRAIL, increased levels of these markers have been shown in previous studies of HPS patients [14, 19].

The higher levels of C5/C5a observed in patients with nonfatal outcome (Figure 3), together with the associations found...
Figure 3. Multiple serum markers distinguish fatal from nonfatal hantavirus pulmonary syndrome. A, Principal component (PC) analysis of patient groups and controls plotted against PC1–PC3. B, Heatmap displaying the standardized serum concentration of each marker in each patient. Colors depict high (red) or low (black) concentration. C, Levels of serum markers in patients with fatal and nonfatal HPS sampled on days 1–4 (fatal, n = 19; nonfatal, n = 19) and days 5–10 (fatal, n = 14; nonfatal, n = 23) post symptom debut. Mann–Whitney U test; median values. *P < .05; **P < .01; ***P < .001. Abbreviations: BAFF, B-cell activating factor; C5, complement factor 5; C5a, complement factor 5a; I-FABP, intestinal fatty acid–binding protein; IFN-γ, interferon gamma; IL, interleukin; LBP, lipopolysaccharide-binding protein; ns, not significant; PC, principal component; sCD14, soluble CD14; sCD25, soluble CD25; sTRAIL, soluble TNF-related apoptosis-inducing ligand; TNF, tumor necrosis factor; VEGF, vascular endothelial growth factor.
between increased C5/C5a levels and decreased odds of severe disease and fatal outcome (Figure 4B and 4C), suggest a possible role for the complement system in HPS pathogenesis. Likewise, our data showing an association between the B-cell–stimulating cytokine BAFF and decreased odds of severe HPS (Figure 4B) indicate a potential protective effect of BAFF. Further characterization of serum markers associated with survival during HPS could provide important leads on HPS pathogenesis and be useful in the development of therapies.

Strong cytotoxic cell responses have been repeatedly reported during human hantavirus infection [10–12] and in vitro, hantavirus-infected endothelial cells have been shown to activate NK cells via IL-15 [20]. The observed increases in serum levels of IL-2, IL-12, IL-15, and IL-18 in HPS patients (Figure 1) [14] suggest that hantavirus infection triggers secretion of a broad repertoire of cytokines that may activate cytotoxic lymphocytes. Moreover, the finding of increased systemic granzyme A and granzyme B levels (Figure 1) likely reflects high cytotoxic activity of these cells. The increased odds of severe disease and fatal outcome associated with increased levels of IL-15 and granzyme B (Figure 4A and 4C) may suggest that severely ill HPS patients are predisposed to mount stronger cytotoxic cell responses. Taken together, our data strengthen the view of HPS as a disease characterized by highly activated immune responses and inflammation [1].

Microbial translocation has been suggested as a driver of immune activation and inflammation during human immunodeficiency virus, hepatitis B virus, hepatitis C virus, and dengue virus infection [21–23]. Here, we demonstrate increased serum levels of the MT surrogate markers sCD14 and LBP during HPS (Figure 2A and 2B). These data indicate that MT may occur during HPS. We also observed elevated levels of I-FABP (Figure 2C), suggesting injury of the intestinal epithelial cells of HPS patients. This raises the question as to whether intestinal damage could be associated with the gastrointestinal symptoms frequently observed in HPS patients [7, 8]. During Ebola virus disease, gastrointestinal symptoms are also common and have been associated with a fatal outcome [24]. Interestingly, I-FABP levels were higher in HPS patients with fatal compared with nonfatal outcome (Figure 3C) and in multivariate analysis, I-FABP was the only marker (out of 20) that was independently associated with increased odds of fatal outcome (Figure 4D). This suggests that intestinal damage marks a life-threatening stage of HPS. Possible causes of intestinal cell injury during HPS, causing subsequent leakage of I-FABP to the circulation, remain elusive. Several lines of evidence indicate that cells of the gastrointestinal tract could be infected during hantavirus infection. For instance, viral antigen can be detected in the small intestine and feces of bank voles, the natural host of HFRS-causing Puumala virus (PUUV) [25]. Furthermore, PUUV antigen has been detected in stomach and colon of HFRS patients [26]. As hantaviruses inhibit apoptosis [27], it is unlikely that systemic I-FABP levels would arise as a consequence of virus-induced cell death of infected intestinal epithelial cells. In support of this view, it was recently shown that human intestinal epithelial cells are susceptible to infection with PUUV in vitro and
that infection does not cause cell death [28]. Possible causes of
intestinal damage during HPS could be intestinal ischemia or
bystander killing by cytotoxic cells or via soluble factors such
as TNF [29, 30]. Unraveling the factors responsible for causing
intestinal injury during HPS has the potential to reveal import-
ant and novel aspects of hantavirus-induced pathology.

The proinflammatory cytokines IL-1β, IL-6, and IL-15 as well
as ferritin and granzyme B were all associated with a more severe
disease in univariate analyses (Figure 4A). This suggests that
certain inflammatory pathways might be more active in patients
with severe HPS. In multivariate analyses, controlling for inter-
marker relationships, IL-6 was identified as the only marker
that was independently associated with the disease severity of
HPS (Figure 4B). These findings extend previous studies, in
which increased IL-6 levels were suggested to be associated
with the severity of HPS and HFRS [9, 31]. In addition, during
Ebola virus disease, dengue virus infection, and Crimean–
Congo hemorrhagic fever, elevated levels of IL-6 have been
linked to disease severity [32–34], suggesting that this is a com-
mon characteristic of viral hemorrhagic fevers. Contemplating
these findings, an interesting parallel can be drawn to cytokine
release syndrome (CRS), a toxic immune reaction upon can-
cer treatment with chimeric antigen receptor–modified T cells,
during which IL-6, IL-10, and IFN-γ are the main cytokines
during which IL-6 levels were suggested to be associated
with severe pulmonary syndrome in Argentina. Virology 1996;
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In conclusion, this study provides novel insights into the
inflammatory response during fatal and nonfatal HPS. Our
results identify distinct inflammation-associated serum mark-
ers associated with severity and fatality in HPS. Importantly,
multivariate analyses identified IL-6 as a marker associated with
disease severity whereas I-FABP, a marker of intestinal damage,
was shown to be associated with fatality. It is tempting to spec-
ulate that these markers may contribute toward prediction of
disease severity in HPS.

Supplementary Data
Supplementary materials are available at The Journal of Infectious
Diseases online. Consisting of data provided by the authors to
benefit the reader, the posted materials are not copyedited and
are the sole responsibility of the authors, so questions or com-
ments should be addressed to the corresponding author.

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