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Journal Title: Cytokine
Volume: Volume 60, Number 1
Publisher: Elsevier: 12 months | 2012-10-01, Pages 249-256
Type of Work: Article | Post-print: After Peer Review
Publisher DOI: 10.1016/j.cyto.2012.06.016
Permanent URL: https://pid.emory.edu/ark:/25593/s9f0r

Final published version: http://dx.doi.org/10.1016/j.cyto.2012.06.016

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Accessed June 14, 2020 5:49 PM EDT
Tumor necrosis factor –308 polymorphism (rs1800629) is associated with mortality and ventilator duration in 1057 Caucasian patients

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Abstract

Purpose—Management of sepsis in critically ill patients remains difficult and requires prolonged intensive care. Genetic testing has been proposed as a strategy to identify patients at risk for adverse outcome of critical illnesses. Therefore, we wished to determine the influence of heredity on predisposition to poor outcome and on duration of ventilator support of intensive care unit (ICU) patients.

Methods—A study was conducted from July 2001 to December 2005 in heterogeneous population of patients from 12 US ICUs represented by the Genetic Predisposition to Severe Sepsis (GenPSS) archive. In 1057 Caucasian critically ill patients with SAPS II probability of survival of >0.2 in the US, six functional single nucleotide polymorphisms in relation to inflammatory cytokines and innate immunity (rs1800629, rs16944, rs1800795, rs1800871, rs2569190, and rs909253) were evaluated in terms of mortality and ventilator free days.

Results—The AA homozygote of TNF(–308) (rs1800629) was most over-represented in the deceased patient group (P = 0.015 with recessive model). The carriage of the TNF(–308)* AA genotype showed significantly higher odds ratio of 2.67(1.29–5.55) (P = 0.008) after adjustment with the covariates. However, the presence of 1, 2, or 3 acute organ dysfunctions was larger prognostic factors for the adverse outcome (OR(95%CI) = 2.98(2.00–4.45), 4.01(2.07–7.77), or 19.95(4.99–79.72), P < 0.001 for all). Kaplan–Mayer plot on ventilator duration of TNF(–308)* AA patient significantly diverged from that of TNF(–308) (GG + GA) ((AA v GG + GA), Adjusted HR(95%CI) = 2.53(1.11–5.79) with Cox regression, P = 0.028).
Conclusions—TNF(−308)* AA is significantly associated with susceptibility to adverse outcome and to longer ventilator duration. Therefore, heredity likely affects both predisposition to ICU prognosis as well as the resource utilization.

Keywords
Genetic Predisposition to Disease; Genetic testing; Sepsis; Cytokines; Ventilators; mechanical

1. Introduction

Sepsis is life-threatening and the leading cause of multiple organ failure [1], and overcoming this formidable enemy is challenging even with the present critical care, which has been highly advanced. Sepsis is also a costly public health problem [2], therefore early prediction, detection and treatments are considered to be very important to decrease the expense of medical resources as well as the mortality of septic patients. Needless to say, various factors, including age, gender, chronic health, physiologic derangement, treatments, diagnostic strategies, and complications affect the outcome of severe sepsis. However, few studies have ever formally evaluated strengths of these determinants to predict susceptibility to and also the outcome of severe sepsis and the association among these co-factors.

On the other hand, evidence of genetic influences with respect to predisposition to various critical illnesses has started to garner attention, and susceptibility to severe sepsis has been considered to be attributed to both heritable and comorbid factors [3–5]. A number of studies have been conducted to evaluate associations between genetic variations and susceptibility to sepsis or outcome of septic patients under a similar concept to that study [3–6]. However, most of those genetic researches have been statistically underpowered, and control patients have been inappropriately selected for the purpose of determination of the genetic influence *bona fide* on susceptibility to diseases and on the patients’ outcome [7–9].

In an attempt to solve the problems of these population association studies, we have already performed a collaborative case-control study with respect to the susceptibility to severe sepsis in a larger, heterogeneous population of patients from 12 USA ICUs represented by the Genetic Predisposition to Severe Sepsis (GenPSS) archive [10]. In that project, we used the Project IMPACT™ (PI) critical care clinical database information management system developed by the Society of Critical Care Medicine (SCCM) [11]. We have previously reported the methodology in detail, and a single nucleotide polymorphism (SNP) in the first intron of LTA at +252(LTA(+252)) was found to influence predisposition to severe sepsis, a predisposition that is modulated by gender and age [10]. However, no clinical outcome in relation to genetic polymorphisms has been shown in the article. The present work represents a continuation of this population association study to address the outcome using the six candidate SNPs which have been widely assumed to be associated with systemic inflammation [10].

Here, we performed logistic regression analyses for ranking by regression of age, chronic health, physiologic derangement, treatments, complications and the genotypes all on the outcomes of severely ill patients with the same PI database, which was utilized in our previous study [10]. Because critical illnesses often cause acute respiratory failure which requires mechanical ventilation, genetic conditions were additionally assessed with respect to the ventilator duration, i.e. the ICU resource utilization.
2. Materials and methods

2.1. Design and data sources

We conducted a retrospective analysis of outcome and ventilator duration in patients with severe sepsis in a heterogeneous, large population in the US. In the present study, we examined whether genetic conditions affect the outcome of critical illnesses and how large the effect is relative to acquired conditions. All participating clinical centers utilized the PI database information management system developed by SCCM [11] and managed by Tri-Analytics, Inc. (http://www.trianalytics.com/, accessed 11 November 2011) and Cerner Corporation (Kansas City, MO). PI is an administrative database designed for critical care units across all disciplines [11].

2.2. Patient selection and definitions of severe sepsis

The diagnosis of severe sepsis was made when a patient met the criteria proposed by the American College of Chest Physicians/SCCM Consensus Conference [10,12,13]. A previously validated approach was applied to the identification of severe sepsis involving the co-occurrence of International Classification of Diseases, Ninth Revision (ICD-9) codes for a bacterial or fungal infectious process and acute organ dysfunction [10].

During the 4 years since Jul 2001, there were more than 200,000 patients among 120 centers in the PI database. The GenPSS archive includes critically ill patients from 12 intensive care units across the US, and the archive contains a total of 854 severely septic patients with genetic material. The patients with SAPS (simplified acute physiology score) II probability of survival [14] of <0.20, i.e. poor chance of survival, are thought to be too severe to evaluate their genetic effect on their clinical outcome; therefore we excluded the severest patient population from the 1708 patients (Fig. 1). Also, the distribution of the polymorphisms assessed in this study (vide infra) was different among African Americans and Caucasians, and these polymorphisms identify different haplotypes in different ethnic groups (data not shown). Therefore, analyses were stratified on presented only for Caucasians. The study population of the critically ill patients is shown in Table 1. Even with the non-random elimination of the severe patient population, SAPS II probability of survival [14] was similar as the actual survival rate (Table 1). Selected treatments for critical illnesses were also evaluated with respect to outcome of the ICU patients in GenPSS archive. Because the patient characteristics are largely diverse, the purpose of the treatments even with the same drug, e.g. steroids, might vary in the critical care settings. And several treatments and procedures have been proven not to be associated with the clinical outcome with our preliminary analyses (data not shown). Therefore, the number of treatments as confounders was strictly limited.

2.3. Definitions of ventilator free days

Ventilator free days (VFDs) were defined as the number of days between successful weaning from mechanical ventilation and 28 day after study enrollment by a formula by Schoenfeld and Bernard [15]. The formulae are as follows:

- \( VFDs = 0 \): If the patient dies before 28 days.
- \( VFDs = (28 - x) \): If the patient is successfully weaned from mechanical ventilation within 28 days, where \( x \) is the number of days spent receiving mechanical ventilation.
- \( VFDs = 0 \): If the patient requires mechanical ventilation for 28 days or more.
2.4. ICD-9. Codes used to identify pre-existing conditions, acute organ dysfunction, and sequelae

Preexisting conditions (cardiovascular diseases, diabetes mellitus, chronic obstructive pulmonary diseases (COPD), malignant neoplasm, trauma, and obesity) and comorbidities during ICU stay (disseminated intravascular coagulation (DIC), deep venous thrombosis (DVT)) were defined in accordance with the ICD-9 codes as follows. 393–429, Pre-existing cardiovascular diseases; 250, Diabetes mellitus; 490–495, COPD; 140–159, 161–165, 170–208, 230–234, Malignant neoplasm; 800–848, 850–854, 860–879, 881–887, 890–897, 900–904, 925, 928, 929, 950–959, Trauma; 278.00, 278.01, Obesity; 286.6, 286.7, 286.9, DIC; 451–454, DVT. We adapted ICD-9 based classification of acute organ dysfunction from a stuff by Weycker et al. [16].

2.5. Genotyping

Detailed information about sample collection, patient confidentiality, and genotyping methods has been described previously [10]. Genomic DNA for genotyping was isolated from these blood stains. First, we amplified the target region of DNA by polymerase chain reaction (PCR) with primers specific for the sequences of each of our six selected markers. The selected markers include SNPs present before the first codon of TNF at −308 (rs1800629), IL1B at −511 (rs16944), IL6 at −174 (rs1800795), IL10 at −819 (rs1800871), and CD14 at −159 (rs2569190) (TNF−308), IL1B−511, IL6−174, IL10−819, and CD14−159), and in the first intron of lymphotoxin-α (formerly known as TNF-β) at +252 (rs909253) (LTA+252)). These six polymorphisms were selected from 22 candidate markers related to systemic inflammation to infection, i.e. sepsis, based on our preliminary analysis of a separate pilot patient cohort [10]. We focused on genes whose products had a known functional role in the inflammatory response and selected the same six candidate SNPs which were evaluated at the former GenPSS study [10].

We introduced a single nucleotide extension strategy for genotyping using dideoxy terminator nucleotides, or, the template-directed, dye-terminator incorporation assay with fluorescence polarization detection (TDI-FP) methods [10]. The PCR primers and the conditions for the six SNPs are shown in Appendix 1. Also, the TDI-FP oligonucleotide probes, dye-terminator combinations, and reaction conditions are shown in Appendix 2.

2.6. Statistical analysis

We performed a multivariable logistic regression analysis applying a forced entry method with SPSS software (SPSS Inc, Chicago, IL) in order to assess risk factors, including heritable and acquired conditions, which are considered to be associated with outcome of critical illnesses. We additionally evaluated treatments’ effects against several critical illnesses by the logistic regression. Usages of mechanical ventilation, acute renal replacement therapy, vasopressin, blood products administration, and enteral feeding for the patients were queried by Microsoft Access database. Ventilator durations in the ICU patients with regard to the common allele presence of a selected SNP were also assessed by Cox regression analysis using the statistical software program SPSS (SPSS Inc., Chicago, IL).

The Hardy–Weinberg equilibrium for the population distribution of the variant alleles was determined according to the approach described by Guo and Thompson [17]. Allelic chi-squares were examined for each SNP. We considered the differences significant with respect to mortality and VFD at a full scan permutation of the correlation/trend test $P$-value of $p \leq 0.05$. The multiple logistic regression model was established to evaluate the independent relationship between a SNP variant and the presence of the potential confounders in relation to the outcome of critical illnesses. Odds ratios (ORs) were calculated in 95% confidence interval (CI) and all P values were two-tailed, with statistical significance defined by $P \leq 0.05$.

Cytokine. Author manuscript; available in PMC 2013 October 01.
Hazard ratios (HRs) for Cox regression analysis of ventilator durations were also calculated and all P values of these multivariable regression analyses were with statistical significance defined by P ≤ 0.05. Adjustment for multiple testing was not performed due to the exploratory hypothesis generating nature of the study. All of both heritable and acquired covariates included in the models were indicated in Table 1.

The protocol, which includes waiver of informed consent, was approved by the Human Studies Committee at the Washington University School of Medicine. The GenPSS project design meets the confidentiality and specimen access controls required of a typical NIGMS Human Genetic Cell Repository (http://ccr.coriell.org/Sections/Collections/NIGMS/?SsId=8, accessed 11 November 2011).

3. Results

3.1. Population characteristics

To investigate the association between heritable and acquired factors of the critically ill in the heterogeneous US population, we analyzed a total of 1057 Caucasian ICU patients from the GenPSS archive and genotyped for the six SNPs. The baseline characteristics of the genotyped patients are presented in Table 1. Patients with severe sepsis were 55.3% of total population and male gender was 52.6%. Median age (range) was 66.0 (18–90) years old and their ventilator free days was 17.3 (0.4) (median, SE)). SAPS II probability of survival (mean (SD)) [14] was 70.4 (29.6) and the actual in-hospital survival rate was also 70.4%. Cardiovascular disease was the most prominent pre-existing conditions appearing in the patient cohort according to the ICD-9 entry at admission. Enteral feeding was performed for 37.1% of the patients. Mechanical ventilation was introduced for 49.5% of the patients and the number of acute organ dysfunction (mean (SD)) was 0.81 (0.76).

3.2. Genotypic distributions of the six inflammation gene polymorphisms and the outcome of the critically ill in the genotype categories

Genotype call rate of the six SNPs was 91.5–99.0%, although the genotypic distribution in LTA(+252) diverged from Hardy-Weinberg equilibrium (HWE) in the studied subjects (P < 0.001; Table 2). The TNF(−308)* AA homozygote showed the highest in-hospital mortality (48.7%) among the 18 genotypes evaluated and recessive model of TNF(−308) and IL10(−819) showed statistically significant association with outcome of the critically ill (P = 0.015 and P = 0.040, respectively with correlation/trend test). As expected, given their collocation on chromosome six, TNF(−308) and LTA(+252) were in linkage disequilibrium (LD correlation R = 0.502, D = 0.808, P < 0.001).

3.3. Ranking by the regression of the prognostic factors on critical illnesses

We performed the ranking of heritable and acquired risk factors for severe sepsis (Fig. 2). The more organ dysfunctions the septic patients obtained, the higher mortality they had, hence presence of three organ dysfunctions was the highest prognostic factor for the adverse outcome (OR(95%CI) = 19.96(4.99–79.72), P < 0.001). Gastrointestinal bleeding was also the significant risk factors for deterioration (OR(95% CI) = 7.64(1.76–33.12), P = 0.007).

As we expected, TNF(−308)* AA genotype carriers resulted in having worse outcome than the other genotype with recessive model after normalization by the selected heritable and acquired factors (OR(95% CI) = 2.67(1.29–5.55), P = 0.008). Hence, there is evidence that TNF(−308) is significantly associated with adverse outcome both on its own and also when adjusted for potential confounders including the number of organ failures, gastrointestinal bleeding as well as severe sepsis occurrence. IL10(−819)* TT genotype carriers were also
susceptible to poor outcome than the other \textit{IL10} genotype group, although the genetic influence was weaker (OR(95% CI) = 1.89(1.00–3.57), \( P = 0.050 \), Fig. 2).

3.4. Genotypic distributions of the six inflammation gene polymorphisms and the ventilator free days of the critically ill in the genotype categories

Six hundred and twenty-five out of the 1057 Caucasian patients received ventilatory support (Appendix 3). The \( TNF(-308)* \) AA homozygote again showed the shortest ventilator free days of 5.56 (1.95) (mean(SE)) among the 18 genotypes evaluated (Fig. 3) and recessive model of \( TNF(-308) \) showed marginally significant association with the ventilator free days of the critically ill \( (P = 0.050 \) with the correlation/trend test, Fig. 3, Appendix 3). However, this results might be affected by the worse outcome of \( TNF(-308)* \) A presences of the critically ill patient population (Table 2). Therefore, we performed a further evaluation on their ventilator duration by Cox regression analysis because correction for potential confounding factors due to baseline imbalance of covariates indicated in Table 1 was thought to be important.

3.5. \( TNF(-308) \) SNP in relation to ventilator duration of the critically ill

Fig. 4 shows the cumulative frequency curve on ventilator duration of the ICU patients by \( TNF(-308)* \) genotype (AA v non-AA). The \( TNF(-308)* \) AA genotype significantly diverged from the other genotypes (Adjusted HR(95%CI) = 2.53(1.11–5.79) with Cox regression, \( P = 0.028 \)). This deviation meant that \( TNF(-308)* \) AA genotypes needed longer mechanical ventilation among this acute phase, in the 28 days after the start of mechanical ventilation, of their ICU stay. Therefore, genetic testing for this \( TNF(-308) \) SNP is thought to be useful for predicting the predisposition to longer ventilation, i.e. longer occupation of ICU resources.

4. Discussion

Genomic approaches to study injury and critical illness still have high hurdles, unique to these methods, regarding resources, experimental design, and data analysis. Hence, it is difficult to determine which factors largely affect the outcome of the multifactorial disorders in clinical settings. Over the period of history, \( TNF(-308) \) has held center stage of critical care genomics since McGuire et al. reported the association between the SNP and susceptibility to cerebral malaria \cite{18}. In the present study, we derived associations between \( TNF(-308) \) or \( IL10(-819) \) and inhospital mortality with univariate genetic association analyses (Table 2). Thereafter, only the influence of \( TNF(-308) \) on both clinical outcome (Fig. 2) and ventilator duration (Fig. 4) was validated with multivariate analysis.

Recently meta-analysis on the \( TNF(-308) \) was performed and an association between development of sepsis and \( TNF(-308)* \) A presence was recognized, although \( TNF(-308)* \) A was not associated with sepsis mortality except for Asian population \cite{3}. In this regard, immunosuppression due to severe sepsis \cite{19} is considered to be predisposed to cytomegalovirus (CMV) reactivation \cite{20}. Patients with severe sepsis who developed active CMV infection were found to have significantly prolonged mechanical ventilation \cite{20}, therefore the longer ventilator duration of \( TNF(-308)* \) AA carriers in the present study might have been caused partially by the CMV reactivation. On the other hand, a recent article evaluated with an Asian population insisted that APACHE II score combined with \( TNF(-308) \) genotyping was more accurate than APACHE II score alone in predisposing to clinical outcomes \cite{21}. It is widely recognized that there is a substantial genotypic difference in \( TNF(-308) \) among various racial populations \cite{22}. As part of this trend, the present study is one of the largest population genetic studies in relation to \( TNF(-308) \) with
Caucasian patient population, and as we expected, the association between TNF(−308) and outcome of the critically ill in this large and heterogeneous population was also recognized. Our previous report did not reveal any association between TNF(−308) and susceptibility to severe sepsis in the US patients of the GenPSS archive while LTA(+252) influenced predisposition to the disease [10]. Because the two markers on the same chromosome TNF(−308) and LTA(+252) were in a linkage disequilibrium, diplotype of these two SNPs may have a stronger association with clinical outcome of critically ill patients. Actually, a forward stepwise logistic haplotype trend regression test [23] for occurrence of severe sepsis on the two marker haplotype composed of TNF(−308) and LTA(+252) markers determined that the best predictor for severe sepsis was the (G, G) haplotype with a P-value of 6.47e-06 [10], even though the sepsis risk allele for the TNF(−308) marker is generally the minor A allele [3,24,25]. TNF(−308)* A shows high TNF producer phenotype [26], and an important role of TNF in controlling M.tuberculosis and many other intracellular infections in humans has been recognized [27]. Therefore, high TNF production does not always lead to vulnerability to sepsis but potentially deterioration of the inflammatory disorder due to extreme hypercytokinemia [28]. On the contrary, Foster et al. demonstrated that histone methylation inhibits TNF gene transcription [29], hence the epigenetic modification of chromatin and DNA is thought to be an important level of gene regulation in SIRS and sepsis pathophysiology. As the results, inflammatory gene silencing, i.e. endotoxin tolerance in sepsis could lead to deleterious effects on ICU patients [30]. In this regard, the rarity of the mutant allele homozygotes of TNF(−308) in both the US populations is noteworthy. Moreover, there was not even a single TNF(−308)* AA individual in the Japanese ICU patients [21,24,31]. The genetic effect on outcome of critical illnesses is thought to be substantial among most racial groups, and this may strongly select against mutant homozygote individuals and act to reduce the TNF(−308)* A allele frequency.

The present report begs the question of whether it is useful to introduce genetic testing in an acute phase of critical care illnesses. The answer might be ‘yes’, but only in selected and limited situations. Genetic testing has least utility in the severest patients who typically die soon after admission. Their care would not likely be affected by their genotypes. Hence, genetic testing for the patients with SAPS II PS of <0.2 in ICU settings, whom we eliminated in the present study, would incur expense with little promise of benefit. On the other hand, the genetic information may be useful in ‘milder’ critically ill patients to more rapidly institute appropriate treatments. TNF(−308)* AA homozygotes needed longer duration of mechanical ventilation for the other genotypes, TNF(−308)* (GG + GA), in their 28 ICU days (Fig. 4). This information might serve as a useful reference suggesting earlier tracheostomy and planning for long-term ventilator support for TNF(−308)* AA homozygotes.

The present study has several important limitations. First, 22 genetic markers of more than millions of genetic polymorphisms were evaluated preliminarily and only six were further analyzed in detail. Needless to say, sepsis pathophysiology is complex web of interactions and the disease advances through many physiologic pathway, i.e. inflammation, innate immunity, adoptive immunity, coagulation–fibrinolysis [25,32], which are closely involved in various types of cell death of vital organs [33]. However, even among these limited SNPs, we found statistically significant association between the genetic factors and mortality as well as ventilator duration. Hence, biological pathways related to TNF(−308) are suggested to affect both outcome of critical illness and use of ICU resources. Secondly, the LTA(+252) was deviated from HWE in the study population. There was evidence to suggest that the association between the outcome and these markers was not driven by departures from HWE. The TNF(−308) and LTA(+252) were shown in previous studies to be associated with both sepsis and respiratory failure [10,34]. The samples were not randomly drawn from...
a population but instead were taken from patients admitted to the ICU (Fig. 1, Table 1). The
likelihood of cases and controls suffering from either sepsis, respiratory failure or both is
greater than would normally be found in the general “healthy” population, plausibly
accounting for the departures from HWE as discussed at our previous GenPSS study [10].
Longevity of ventilator duration is thought to be affected by susceptibility to severe sepsis
[35]. Therefore, it is also striking that 7NKR(−308) was associated with longer mechanical
ventilation even though this SNP did not affect sepsis occurrence [10]. A linkage
disequilibrium between 7NKR(−308) and LTA(+252) might have caused the association
between 7NKR(−308) and longer mechanical ventilation as well as susceptibility to poor
outcome. Lastly, the assignment of race is known to be error-prone. We used the
assignments recorded in the PI database. These self-assignments were made by patients and
families at the time of admission.

The GenPSS project launched in 2001, and we selected more than 20 functional and
candidate polymorphisms related to systemic inflammation [10] and the genes whose
products had a known functional role in the inflammatory response had been focused. These
SNPs were analyzed with TDI-FP, a strategy for high-throughput, cost-effective genotyping
of samples [10]. With current technologies, information on up to more than a million unique
sequence variants can be provided on a single chip with lower cost. Actually,
oligonucleotide arrays can be used to identify presence of specific alleles in individuals, and
a CYP chip and the microarray instrumentation system have been commercially available in
the pharmacogenomics field (FDA K042259; http://www.fda.gov/, accessed 16 April 2012).
This sort of DNA chip-based genotyping technique is more suitable for profiling of the
genetic background of a particular individual than the present genotyping assay, i.e. TDI-FP,
in the critical care field [36].

Genetic testings, to ultimately investigate the inherent order of biology, is thought to
compete with the usefulness of biomarkers in prediction and prevention. One advantage of
genetic testings is that their information remains unchanged in any clinical conditions in
contrast to a variety of biomarkers from biological samples. Therefore, physicians will be
able to perform prepared response to each biological reaction to stress on the basis of
genomic markers in the critical care settings. Furthermore, the classic central dogma of
molecular biology states that “DNA makes RNA makes protein”, and functional genomics
explores to determine the significance of the observed changes in gene expression [37]. In
terms of this concept, multiple-gene biomarkers, e.g. cDNA microarrays, are likely to be
more sensitive and discriminating than single genes. Reaping the maximum fruit from these
technologies, Ashley et al. lately provided an approach to comprehensive analysis of a
human genome in a defined clinical context [38]. They assessed whole-genome genetic risk,
focusing on variants in genes that are associated with Mendelian disease, novel and rare
variants across the genome, and variants of pharmacogenomic importance [38]. This kind of
whole-genome assessment for sepsis and the critical illnesses may come true in near future.

5. Conclusions

Heredity likely affects both predisposition to severe inflammation and to poor outcome of
the disease. The specific effects remain opaque. Further analyses of haplotypes in candidate
regions, e.g. MHC region including 7NF and LTA in sepsis, and genome-wide analysis for
the critical illnesses may prove to be a more productive approach. Although the genetic
influences are likely overwhelmed by comorbid factors and acute illness in individual cases,
population studies continue to suggest biological pathways that affect both critical illness
and use of ICU resources.
Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

This research was supported by NIH/GM (062809) and the Pfizer Travel Fellowship 2006 by Japanese Association for Acute Medicine. The funding sources had no role in study design, collection, analysis, and interpretation of data.

Abbreviations

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<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>ICU</td>
<td>intensive care unit</td>
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<tr>
<td>SNP</td>
<td>single nucleotide polymorphism</td>
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<td>GenPSS</td>
<td>Genetic Predisposition to Severe Sepsis</td>
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<td>SAPS</td>
<td>simplified acute physiology score</td>
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<td>TNF</td>
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<td>template-directed, dye-terminator incorporation assay with fluorescence polarization detection</td>
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<td>HWE</td>
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<td>LD</td>
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<td>SIRS</td>
<td>systemic inflammatory response syndrome</td>
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References


**Appendix A. Supplementary material**

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.cyto.2012.06.016.
Fig. 1.
Flow chart of patients’ selection in the GenPSS project (GenPSS, Genetic Predisposition to Severe Sepsis).
Fig. 2.
Multivariable logistic regression model for detecting the incident effect of the TNF(−308) and IL10(−819) SNPs on the outcome of critical illnesses. COPD, chronic obstructive pulmonary diseases; VAP, ventilator associated pneumonia; RRT, renal replacement therapy; DIC, disseminated intravascular coagulation; DVT, deep venous thrombosis. GI; gastrointestinal, TNF(−308), SNP at position −308 nucleotides 5′ of the first exon of tumor necrosis factor; IL10(−819), SNP at position −819 nucleotides before the first exon of IL-10.
Diamons to the right of the vertical line indicate an increased risk of death from critical illnesses in patients carrying each covariate. The diamonds represent the odds ratios from the logistic regression analyses. Horizontal lines through the diamonds represent 95% confidence intervals (CIs).
Ventilator free days of the critically ill in genotype categories of the six single nucleotide polymorphisms VFD, ventilator free day; SNP, single nucleotide polymorphism; TNF(−308), SNP at position −308 nucleotides 5′ of the first exon of tumor necrosis factor; LTA(+252), SNP at position 252 site of lymphotoxin-α; IL1B(−511), SNP at position −511 nucleotides 5′ of the first exon of interleukin-1β; IL6(−174), SNP at position −174 nucleotides before the first exon of IL-6; IL10(−819), SNP at position −819 nucleotides before the first exon of IL-10, CD14(−159), SNP at position −159 nucleotides before the first exon of CD14. Y-axes for all graphs show the ventilator free days of the critically ill in each genotype category. The TNF(−308)* AA homozygote showed the shortest ventilator free days of 5.6 (2.0) (mean (SE)) among the 18 genotypes evaluated (P = 0.050 with recessive model of the correlation/trend test; AA v (GG + GA), see Appendix 3).

Fig. 3.
Fig. 4.
Kaplan–Meier analysis of 28-day ventilator duration of the critically ill by \( TNF(-308) \) genotype (AA v non-AA). \( TNF(-308) \), SNP at position \(-308\) nucleotides 5′ of the first exon of tumor necrosis factor. Statistical significance was assessed using the Cox regression analysis, censored cases; ventilator removal because of decease.
Table 1
Baseline characteristics of the study population.

<table>
<thead>
<tr>
<th></th>
<th>Genotyped critically ill patients (n = 1057)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (median, range)</td>
<td>66, 18–90</td>
</tr>
<tr>
<td>Male sex; n (%)</td>
<td>556 (52.6)</td>
</tr>
<tr>
<td>Survivors; n (%)</td>
<td>744 (70.4)</td>
</tr>
<tr>
<td>Severe sepsis; n (%)</td>
<td>585 (55.3)</td>
</tr>
<tr>
<td>SAPS II prob. of survival (mean ± SD, range)</td>
<td>70.4 ± 29.6, 20.0–100</td>
</tr>
<tr>
<td>Ventilator free days (mean ± SE, range)</td>
<td>17.3 ± 0.4, 0–28</td>
</tr>
<tr>
<td><strong>Pre-existing conditions; n (%)</strong></td>
<td></td>
</tr>
<tr>
<td>Cardiovascular diseases</td>
<td>960 (90.8)</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>751 (71.1)</td>
</tr>
<tr>
<td>COPD</td>
<td>702 (66.4)</td>
</tr>
<tr>
<td>Malignant neoplasm</td>
<td>449 (42.5)</td>
</tr>
<tr>
<td>Trauma</td>
<td>71 (6.7)</td>
</tr>
<tr>
<td>Obesity</td>
<td>402 (38.0)</td>
</tr>
<tr>
<td><strong>Treatments/procedures; n (%)</strong></td>
<td></td>
</tr>
<tr>
<td>Renal replacement therapy</td>
<td>86 (8.1)</td>
</tr>
<tr>
<td>Vasopressin</td>
<td>36 (3.4)</td>
</tr>
<tr>
<td>Blood product administrations</td>
<td>151 (14.3)</td>
</tr>
<tr>
<td>Enteral feeding</td>
<td>392 (37.1)</td>
</tr>
<tr>
<td><strong>Acute organ dysfunctions and sequelae; n (%)</strong></td>
<td></td>
</tr>
<tr>
<td>Respiratory</td>
<td>523 (49.5)</td>
</tr>
<tr>
<td>Cardiovascular</td>
<td>93 (8.8)</td>
</tr>
<tr>
<td>Renal</td>
<td>154 (14.6)</td>
</tr>
<tr>
<td>Hematologic</td>
<td>40 (3.8)</td>
</tr>
<tr>
<td>Neurologic</td>
<td>36 (3.4)</td>
</tr>
<tr>
<td>Hepatic</td>
<td>14 (1.3)</td>
</tr>
<tr>
<td>DIC</td>
<td>32 (3.0)</td>
</tr>
<tr>
<td>DVT</td>
<td>68 (6.4)</td>
</tr>
<tr>
<td>VAP</td>
<td>55 (5.2)</td>
</tr>
<tr>
<td>GI bleeding</td>
<td>11 (1.0)</td>
</tr>
<tr>
<td><strong>Number of acute organ dysfunctions (mean ± SD)</strong></td>
<td>0.81 ± 0.76</td>
</tr>
</tbody>
</table>

SAPS, simplified acute physiology score; COPD, chronic obstructive pulmonary diseases; DIC, disseminated intravascular coagulation; DVT, deep venous thrombosis; VAP, ventilator associated pneumonia; GI, gastrointestinal, SE, standard error; SD, standard deviation.
Table 2

Genotypic distributions of the six SNPs and mortality for the group of the critically ill patients in each genotype category.

<table>
<thead>
<tr>
<th>SNP locus</th>
<th>Genotype: n (genotypic distribution (%))</th>
<th>$P$ values of additive model on mortality</th>
<th>$P$ values of recessive model on mortality</th>
<th>$P$ values of dominant model on mortality</th>
<th>Genotype call rate</th>
<th>Hardy–Weinberg equilibrium test $P$ value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$TNF$ (−308)</td>
<td>GG: 661 (68.4%) GA: 269 (27.8%) AA: 37 (3.83%)</td>
<td>0.091</td>
<td>0.015&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.318</td>
<td>0.915</td>
<td>0.147</td>
</tr>
<tr>
<td>$LTA$ (+252)</td>
<td>AA: 451 (43.1%) AG: 433 (41.4%) GG: 162 (15.5%)</td>
<td>0.824</td>
<td>0.573</td>
<td>0.928</td>
<td>0.990</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>$IL1B$ (−511)</td>
<td>CC: 464 (47.2%) CT: 408 (41.5%) TT: 111 (11.3%)</td>
<td>0.655</td>
<td>0.748</td>
<td>0.419</td>
<td>0.930</td>
<td>0.141</td>
</tr>
<tr>
<td>$IL6$ (−174)</td>
<td>GG: 358 (34.4%) GC: 512 (49.2%) CT: 171 (16.4%)</td>
<td>0.251</td>
<td>0.400</td>
<td>0.312</td>
<td>0.985</td>
<td>0.595</td>
</tr>
<tr>
<td>$IL10$ (−819)</td>
<td>CC: 587 (59.2%) CT: 355 (35.8%) TT: 49 (4.94%)</td>
<td>0.231</td>
<td>0.040&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.597</td>
<td>0.938</td>
<td>0.618</td>
</tr>
<tr>
<td>$CD14$ (−159)</td>
<td>CC: 269 (27.5%) CT: 485 (49.6%) TT: 224 (22.9%)</td>
<td>0.729</td>
<td>0.882</td>
<td>0.491</td>
<td>0.925</td>
<td>0.849</td>
</tr>
</tbody>
</table>

SNP, single nucleotide polymorphism; $TNF$ (−308), SNP at position −308 nucleotides 5' of the first exon of tumor necrosis factor-α; $LTA$ (+252), SNP at position 252 site of lymphotoxin-α; $IL1B$ (−511), SNP at position −511 nucleotides 5' of the first exon of interleukin-1β; $IL6$ (−174), SNP at position −174 nucleotides before the first exon of IL-6; $IL10$ (−819), SNP at position −819 nucleotides before the first exon of IL-10; $CD14$ (−159), SNP at position −159 nucleotides before the first exon of CD14.

<sup>a</sup>$TNF$ (−308)*AA and $IL10$ (−819)*TT genotypes were observed at a frequency higher in non-survivors than in survivors ($P=0.015$ and 0.040, respectively with correlation/trend test).