Gender-Specific Effects on Immune Response and Cardiac Function after Trauma Hemorrhage and Sepsis

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Summary
Background: Studies in human as well as animal models indicate a gender-specific responsiveness of the immune and organ systems with regard to shock, trauma, and sepsis. Methods: A literature review was performed. Results: Cell-mediated immune responses and cardiovascular functions are suppressed in males following trauma hemorrhage, whereas they are maintained or even enhanced in females in the proestrus state of the estrus cycle. Experimental studies have demonstrated that divergent immune responses in males and females following adverse circulatory conditions are mediated by the gender-specific hormones testosterone and estrogen. Several clinical trials, however, failed to demonstrate a significant association of gender and inflammatory response. This may be explained by the heterogeneity of the population in terms of their hormonal status at the time of injury. Conclusions: With regard to the underlying mechanisms, receptors for sex hormones have been identified on various immune cells, suggesting direct effects of these hormones on immune function. Alternatively, indirect effects of sex steroids such as changes in cardiovascular responses or androgen- and estrogen-synthesizing enzymes might contribute to gender-specific immune responses. Clinical studies suggest that sex hormones, such as dehydroepiandrosterone, modulate the function of peripheral blood mononuclear cells also following abdominal surgery. Thus, sex hormones, receptor antagonists, and sex steroid-synthesizing enzymes might be useful in the future for modulating the complex immune responses after trauma hemorrhage and sepsis.

The first two authors contributed equally to this work.

Schlüsselwörter
Geschlecht · Geschlechtshormone · Hämorrhagischer Schock · Immunsuppression · Immunmodulation

Zusammenfassung

The first two authors contributed equally to this work.
Introduction

Differences between women and men can be identified at many levels of the immune response, and may affect its outcome. Clinical and experimental studies have demonstrated gender-specific humoral and cell-mediated immune responses (fig. 1). For example, the incidence of autoimmune diseases, such as systemic lupus erythematosus (SLE) (female to male ratio of 9:1), Hashimoto’s thyroiditis, rheumatoid arthritis, and primary biliary cirrhosis, differs significantly between the sexes [1–4]. Moreover, higher levels of circulating plasma antibodies have been found in females with autoimmune disease compared to male patients [5]. While it is plausible that some sex-linked genes may contribute to the genetic predisposition for autoimmune disease processes, other likely culprits for this gender bias are male and female sex hormones [2, 6]. Administration of testosterone in female F1 NZB/NZW mice, for example, prevented the development of lupus erythematosus [7]. Further support for the notion that male and female sex steroids differently affect autoimmune disease processes comes from studies showing lower androgen levels and increased levels of active estrogen metabolites in women with SLE compared to age-matched healthy female controls [8, 9]. In addition, not only humoral but also cell-mediated immune responses appear to exhibit sexual dimorphism. As early as 1898, Calcolzari [10] reported a connection between reproduction and immunology when he found the thymus of rabbits castrated before sexual maturity to be larger than that of male controls. Moreover, a shorter skin allograft rejection time has been reported in females compared to males [11]. The aforementioned studies demonstrate that male and female sex steroids modulate humoral and cell-mediated immune functions and thereby affect several disease processes. In this regard, gender has also been identified as an important factor in the human body’s response to injury and sepsis: in a huge series of 681,000 patients utilizing the US National Trauma Data Bank, Haider et al. [12] demonstrated a significantly lower mortality as well as a decreased complication rate in female compared to male trauma patients. Male trauma patients also have a higher risk for the development of sepsis and multiple organ failure [13–16] and a significantly lower survival rate (31 vs. 74%) in sepsis [17].

This article will review the epidemiological and clinical data published on this subject, and discuss why some clinical studies failed to demonstrate a difference between the two sexes in the response to trauma hemorrhage and sepsis. Pathophysiological findings that may explain sexual dimorphism are described, and based on these findings novel therapeutic approaches targeted at the influence of sex hormones on the immune response to injury and sepsis are presented. To this end, a MEDLINE search was performed using the terms (sex hormones OR gender) AND (shock OR trauma OR hemorrhage OR sepsis). Experimental studies in animals as well as clinical trials and retrospective analyses were included if they examined the influence of either gender or sex hormones on survival or other applicable outcome parameters in one or more of the following clinical situations or experimental models thereof: cardiocirculatory shock, severe trauma, hemorrhage, or sepsis. Redundant studies were excluded, and additional explanatory evidence was included as deemed necessary by the authors.

Clinical Evidence for Gender Differences in Trauma, Shock, and Sepsis

An analysis of the National Trauma Data Bank, comprising more than 150,000 patients involved in blunt and penetrating trauma, observed an association between gender and mortality among blunt trauma patients, especially those aged ≥50 years [18, 19], while other studies concluded that female gender was not associated with decreased mortality if patients were adequately stratified in a multivariate analysis considering independent risk factors, including the injury severity score (ISS) or patient age [20] (table 1).

Offner et al. [21] studied 545 trauma patients older than 15 years with an ISS higher than 15 and prospectively detected those with survival of more than 48 h. They revealed that male gender was associated with significantly increased risk of severe infectious complications after trauma, and this correlation was most significant following injuries of moderate severity. Gender differences in B and T lymphocyte function, T helper cell counts, and natural killer cell counts have been hypothesized as mechanisms on a cellular and molecular level [22]. Furthermore, increased serum interleukin(IL)-6 and procalcitonin levels in male compared to female patients have been suggested as humoral mechanisms [23].

Sepsis syndrome is a manifestation of an acute infection and is characterized by a generalized inflammatory response [13]. In an experimental ex vivo setting of septic shock, when peripheral blood mononuclear cells (PBMC) were incubated with lipopolysaccharide (LPS), the release of tumor necrosis factor(TNF)-α from these cells after 6 h was significantly higher in samples of healthy males compared to that from female volunteers [24]. This increased cytokine release capacity in male patients was also observed after blunt injuries with an ISS ≥16, and was additionally associated with an increased susceptibility to sepsis compared to female trauma victims [23]. Coyle et al. [25] prospectively evaluated the effects of intravenous application of LPS in 72 healthy volunteers (48 male, 24 female), and found increased core temperature and decreased mean arterial blood pressure in men compared to woman. However, no differences were observed regarding host immune responses to LPS (white blood cell count, cortisol or cytokine level) between males and females. Although several clinical trials indicated gender-specific differences in clinical outcome after trauma, animal studies, in which the experimental conditions are generally better con-
Gender-Specific Immune Response

...of the population studied in relation to their hormonal state [18].

**Gender-Specific Effects on Immune Function after Experimental Trauma Hemorrhage and Sepsis**

Several experimental studies have shown that immune functions are depressed in males as well as ovariectomized and aged females after trauma hemorrhage and sepsis, and remain depressed despite fluid resuscitation [14, 26]. In this respect, female rats showed a higher resistance towards disturbances of the microcirculation following trauma or intestinal ischemia [27]. Moreover, proestrus female mice exhibit increased levels of proinflammatory Th1 lymphokines, such as IL-2 or interferon(IFN)-γ, compared to male mice following trauma hemorrhage [28]. On the other hand, anti-inflammatory Th2 lymphokines, such as IL-10, are decreased in females compared to males after trauma hemorrhage [29]. A gender-dependent immune response has also been described in experimental models of sepsis with survival rates of females significantly higher than those of males (60 vs. 25%) [30]. One explanation for the observation that females are more resistant towards septic challenges may be an increased early expression of inflammatory cytokines such as IL-1 [31]. Interestingly, the immune response to trauma and hemorrhage in females depends on the hormonal status at the time of injury: a more pronounced immune reaction is found during proestrus as compared to diestrus, potentially corresponding with enhanced levels of estrogen and pregnenolone [26, 32, 33]. In clinical studies, the maximum expression of proinflammatory cytokines has been observed in the follicular phase of the menstrual cycle [34–36]. Sex-specific immune reactions have also been shown in the thymus, at the location of T cell lymphopoiesis [37, 38]. In female mice, thymocytes and lymphocytes show an enhanced reaction to antigens compared to thymocytes and lymphocytes from male mice [38], and a reduced apoptosis rate in models of trauma and hemorrhage, respectively [37]. An increased apoptotic rate in males may represent an attempt to decrease autoreactive T cells after trauma and hemorrhage [39]. With respect to sex hormones, studies in castrated mice that develop SLE [40] showed that the application of androgens delayed the onset of SLE. Interestingly, thymectomy abolished the protective effects of administered androgens on the induction of SLE [40]. Nevertheless, the relevance of some of these studies is narrowed by limitations of their study design; in older trials, mainly young male animals were utilized. More recent studies on shock and trauma also took into account the effects of age, gender, and sex hormones, showing advantages for young female animals. Further studies are required to clarify whether these advantages in morbidity and mortality might be neutralized in older female animals to the level of aged males.

**Gender-Specific Effects on Cardiovascular Function under Stressful Conditions**

After trauma hemorrhage, an impaired cardiac performance has been observed in male animals, leading to a generalized organ hypoperfusion [41, 42]. In this respect, cardiovascular and hepatic function was reduced in males and estrus females, but not in proestrus females, following trauma and

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**Table 1. Multi-institutional studies on the outcome of trauma patients depending on gender**

<table>
<thead>
<tr>
<th>Study, year [ref.]</th>
<th>Age group, years</th>
<th>ISS</th>
<th>n</th>
<th>Mortality rate male vs. female</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Morris et. al., 1990 [78]</td>
<td>all ≤25</td>
<td>199,737</td>
<td>–</td>
<td>RR*: 0.7–2.0</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>Wohltmann et al., 2001 [79]</td>
<td>all ≤25</td>
<td>20,261</td>
<td>8 vs. 7%</td>
<td>n.s.</td>
<td></td>
</tr>
<tr>
<td>Gannon et al., 2002 [20]</td>
<td>18–65 ≤25</td>
<td>22,332</td>
<td>7 vs. 5%</td>
<td>&lt;0.002</td>
<td></td>
</tr>
<tr>
<td>George et al., 2003 [18]</td>
<td>≥20</td>
<td>155,691</td>
<td>6.3 vs. 4.5%</td>
<td>&lt;0.001</td>
<td></td>
</tr>
</tbody>
</table>

*Relative risk of mortality for males. n.s. = Not significant.
resuscitation [43]. Interestingly, studies showed that castration of male rats 2 weeks before the induction of trauma hemorrhage led to maintenance of myocardial function [44]. Estradiol, on the other hand, has been found to improve cardiovascular and hepatic function following cardiocirculatory shock [28, 45]. As a potential mechanism, an increased blood volume in females following trauma hemorrhage could explain the observed gender specificities in immune and organ functions [46]. A study analyzing regional myocardial contractile function in a magnetic resonance imaging-based multiparametric strain analysis indicated that there is a significantly larger circumferential and longitudinal left ventricular strain in normal female volunteers, indicating a better functional cardiac reserve in women [47]. Similarly, Sarikouch et al. [48] observed increased end-diastolic and end-systolic left ventricular volumes in female patients and decreased biventricular ejection fractions in males late after repair of tetralogy of Fallot. The exact underlying mechanism remains unclear, nonetheless, and should be further investigated.

The Role of Gender-Specific Hormones in Trauma Hemorrhage and Sepsis

Gender-specific immune reactions are regulated on a hormonal level. The responsible hormones are mainly released from the gonads, and secondarily from the thymus and the hypothalamus and the pituitary glands (table 2) [49–55]. In experimental trauma hemorrhage, prior castration and depletion of male sex hormones increased the release of splenic and peritoneal macrophage cytokines [51, 53–56]. Interestingly, preconditioning of female mice with 5-dihydrotestosterone (DHT) over 2 weeks before trauma hemorrhage caused a reduction in splenic and peritoneal macrophage function similar to males [50, 51]. In contrast, female sex steroids exhibit an increase in cell-mediated immune reactions [57–59]. Elevated systemic levels in proestrus females play a pivotal role in the immunocompetence following trauma hemorrhage [58] maintained by splenocytes and splenic macrophages [60, 61]. Moreover, application of 17β-estradiol was associated with a significantly increased survival rate in sepsis [61]. Although Sperry et al. [62] failed to show differences between pre- and postmenopausal human females after injury and hemorrhagic shock, the protective role of female sex steroids is further underlined by experimental data indicating an increased mortality in trauma hemorrhage and sepsis after ovariectomy [58]. In conclusion, estrogen seems to play a pivotal role in the immunocompetence of females following trauma hemorrhage. Therefore, the application of female sex hormones in ovariectomized or postmenopausal female patients could be beneficial for the therapy of immunodeficiency under such conditions.

Immune Modulation by Sex Steroids and Subsequent Therapeutic Properties

Immune-modulating properties of sex steroids have been shown in vitro: thymocytes from hemorrhaged animals incubated with DHT exhibited reduced expression of IL-3 but an increased apoptotic rate [36], while peritoneal macrophages showed an enhanced release of TNF when co-incubated with estradiol [63]. These immunomodulatory effects could be explained by the presence of estrogen receptors on immune cells, such as thymocytes, macrophages, and leukocytes [2, 64]. Recently another cell surface receptor, G protein-coupled receptor(GPR)-30, has been described to bind estrogen [65, 66].

Furthermore, altered synthesis of sex steroids is evident after trauma hemorrhage [67, 68]. Physiological plasma testosterone levels are associated with an impaired cellular immune defense [50–52, 55]. After trauma hemorrhage, however, an even enhanced DHT synthesis has been shown in male T lymphocytes. In contrast, in proestrus females an increased synthesis and a decreased catabolism of estradiol can be found in the T lymphocytes [67, 68].

Table 2. Experimental results on the effect of sex steroids and gender following trauma, shock, and sepsis

<table>
<thead>
<tr>
<th>Effect of trauma, shock, and sepsis</th>
<th>males</th>
<th>females (proestrus)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Immune response – gender</td>
<td>splenic/peritoneal Mφ cytokine release ↓</td>
<td>splenic/peritoneal Mφ cytokine release →</td>
</tr>
<tr>
<td></td>
<td>Mφ antigen presentation ↓</td>
<td>Mφ antigen presentation →</td>
</tr>
<tr>
<td></td>
<td>T cell proliferation and Th1 release ↓</td>
<td>T cell proliferation and Th1 release →</td>
</tr>
<tr>
<td></td>
<td>T cell Th2 (IL-10) ↑</td>
<td>T cell Th2 (IL-10) →</td>
</tr>
<tr>
<td>Immune response – sex steroids</td>
<td>immunosuppressive</td>
<td>immunoprotective</td>
</tr>
<tr>
<td>Mortality after sepsis</td>
<td>70% (in mice after CLP)</td>
<td>20% (in mice after CLP)</td>
</tr>
<tr>
<td>Effect of sex hormone depletion</td>
<td>castration: immunoprotective</td>
<td>ovariectomy: immunosuppressive</td>
</tr>
<tr>
<td>Treatment option</td>
<td>flutamide (testosterone receptor antagonist)</td>
<td>estrogen (in non-proestrus females)</td>
</tr>
<tr>
<td></td>
<td>estrogen</td>
<td>DHEA (via estrogen receptor)</td>
</tr>
</tbody>
</table>

CLP = Cecal ligation and puncture.
Endogenous testosterone impairs the immune response and organ function following trauma and blood loss. In this respect, the usage of the androgen receptor antagonist flutamide has been examined with regard to immune response in models of trauma and severe bleeding. Flutamide is frequently used for testicular cancer over prolonged periods, and no major adverse effects have been reported. Following trauma hemorrhage and resuscitation, the immune response was significantly increased as indicated by enhanced expression of cytokines from macrophages due to the administration of flutamide [69]. Moreover, in sepsis after trauma hemorrhage, the application of flutamide over 3 days also improved the release of cytokines from splenic macrophages, and significantly reduced mortality [70]. Therefore, the short-term use of this androgen receptor antagonist in male trauma patients could represent a new treatment for deleterious immunodeficiency as well as cardiovascular dysfunction in trauma and sepsis.

The steroid hormone dehydroepiandrosterone (DHEA) is a precursor for estrogen and testosterone, and is found in abundance in the plasma of both females and males. In males, DHEA and its metabolites develop mainly estrogenic effects [71]. On the basis of the beneficial effects of estrogen shown of DHEA and its metabolites, the authors state they have no conflict of interest to declare.

References


Disclosure Statement

The authors state they have no conflict of interest to declare.

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