Inflammation and reduced endothelial function in the course of severe acute heart failure

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Systemic inflammation and elevated circulating levels of the endogenous nitric oxide inhibitor asymmetrical dimethylarginine (ADMA) have been associated with increased risk in cardiogenic shock (CS). In this prospective study, we assessed, over 4 consecutive days, the changes and possible associations between vascular function, markers of inflammation, and circulating ADMA levels in patients with CS (n = 12) and postcardiotomy heart failure (n = 12, PC-HF). Vasodilator function was measured as a reactive hyperemia index (RH-index) using a finger plethysmograph. Blood samples were analyzed for plasma ADMA, interleukine-6, interleukine-8, intracellular adhesion molecule-1, and vascular adhesion molecule-1. Baseline RH-index was significantly attenuated compared with healthy controls (2.28) for both CS and PC-HF (1.35 and 1.45, respectively, P < .001). Although vasodilator function improved in PC-HF patients, it remained attenuated in CS. Inflammatory markers were markedly elevated followed by a significant fall during the observation period in both groups. ADMA levels increased significantly during the observation period for PC-HF, whereas no pattern of change was observed for CS. No association was found between the longitudinal changes in RH-index, markers of inflammation, or ADMA in CS. However, an improved RH-index was associated with decreasing inflammatory markers in PC-HF. ADMA correlated to arterial lactate levels and the degree of organ dysfunction in CS. In conclusion, CS and PC-HF were characterized by a marked inflammatory activation accompanied by an attenuated vasodilator function. ADMA was related to organ dysfunction and degree of hypoperfusion during CS but showed no correlations to inflammation or hampered vasodilator function. The pathogenic significance of these responses needs clarification. (Translational Research 2011; - :1–11)

Abbreviations: ADMA = asymmetrical dimethylarginine; CABG = coronary artery bypass grafting; CCU = coronary care unit; CPR = cardiopulmonary resuscitation; CS = cardiogenic shock; DDAH = dimethylarginine dimethylaminohydrolase; HR = heart rate; IABP = intra-aortic balloon pump; ICAM-1 = intracellular adhesion molecule-1; ICU = intensive care unit; IL-6 = interleukine 6; IL-8 = interleukine 8; IQR = interquartile range; IS = inotropic score; MAP = mean arterial pressure; NO = nitric oxide; PC-HF = postcardiotomy heart failure; RH-index = reactive hyperemia index; SOFA = Sequential Organ Failure Assessment; TNFα = tumor necrosis factor α; vCAM-1 = vascular cell adhesion molecule-1

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AT A GLANCE COMMENTARY

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Background

Systemic inflammation and elevated levels of the endogenous NO-inhibitor asymmetrical dimethylarginine (ADMA) are associated with poor outcomes in acute severe heart failure. How these risk markers relate to each other and their significance in the pathophysiology and vascular dysfunction in the course of cardiogenic shock (CS) is, however, unknown.

Translational Significance

We observed a marked inflammatory response and attenuated vasodilator function in the acute phase of CS. ADMA correlated to the degree of hypoperfusion and organ dysfunction. A substantial degree of individual and time-related fluctuations in these parameters demonstrates the challenges in clarifying the pathophysiological mechanisms in the circulatory dysfunction during CS.

Recent clinical trials and registries on patients with cardiogenic shock (CS) have demonstrated the heterogeneity of this severely ill patient population with regard to etiology as well as clinical and hemodynamical presentation. The current understanding of the underlying pathophysiology that marks patients with an unfavorable course and lack of response to treatment is limited. This was evident in the recently published Triumph trial that addressed the possible adverse effects of a presumed unselective NO inhibitor in patients with persisting shock failed to reduce the duration of shock and mortality. However, observational studies have suggested that systemic inflammation and neurohormonal activation plays an important role in the pathogenesis of CS.

The naturally occurring NO-inhibitor asymmetrical dimethylarginine (ADMA) has been implicated in the pathogenesis of endothelial dysfunction and several chronic cardiovascular disorders. In contrast to its increasing recognition as a risk marker in cardiovascular disease, little is known about ADMA's role in the acute setting. Experimental studies, however, have demonstrated adverse hemodynamic effects after systemic administration of ADMA in humans. Furthermore, ADMA has been indicated to be an independent predictor for a poor outcome in critical illness and CS.

Elevated levels of ADMA also have been observed in patients with acute decompensated heart failure as compared with chronic heart failure. However, a recent study on acute heart failure failed to confirm these findings, and ADMA also did not seem to identify patients with increased risk for future cardiac events including episodes of decompensation. However, at this time, the biological significance and possible regulatory role of ADMA as a modulator of NO production and thus as a vascular function in acute heart failure is not known.

It is necessary to characterize and establish the different factors modulating the circulation in patients with severe pump failure. Ultimately, these observations hopefully will identify and tailor an optimal medical treatment for different subgroups of patients.

The main aim of this observational study therefore was to examine the changes and possible relations among the endogenous NO inhibitor ADMA, vascular vasodilator function and markers of inflammation, and endothelial activation during the course of severe acute heart failure (ie, CS and postcardiotomy heart failure).

In addition, we assessed their association with organ dysfunction and patient outcomes.

METHODS

Study population. The study population consisted of 24 prospectively included patients admitted to the intensive care unit (ICU) or coronary care unit (CCU) at the University Hospital of North Norway between 2007 and 2009 with either CS or postcardiotomy heart failure (PC-HF). Our institution is a university hospital serving a population of approximately 500,000. Informed written consent was obtained from each patient if possible or the nearest relative. The study protocol was approved by the regional ethics committee (REK-Nord 51/2004) and conforms with the ethical guidelines outlined in the Declaration of Helsinki.

CS was defined as cardiac dysfunction with persistent hypotension (systolic blood pressure <90 mmHg) or the need for vasopressor/inotropic drug support to keep systolic blood pressure greater than 90 mmHg in combination with signs and symptoms of tissue hypoperfusion despite a correction of preload. Signs of systemic hypoperfusion included altered mental state, cool skin and extremities, low urine output (<0.5 mL/kg/h), or elevated arterial lactate. PC-HF was defined as inadequate cardiac performance after open-heart surgery in need of inotropes/vasopressors and/or mechanical circulatory support for more than 2 h after surgery with or without signs of systemic hypoperfusion. The PC-HF group did not include patients where an intra-aortic balloon pump (IABP) was implanted preoperatively to improve...
Peripheral vasodilator function was assessed.

Invasive hemodynamic monitoring with pulmonary artery catheter was not obligatory in the study protocol and only applied on clinical indication. To give an estimate of the severity of circulatory compromise, the concomitant requirements of inotropic and/or vasopressor agents were presented as total vasopressor dose (epinephrine + norepinephrine) and inotropic score (IS). IS summarizes the total dosing equivalents of inotropes and vasopressors in μg/kg/min (dopamine + dobutamine + milrinone × 15 + epinephrine × 100 + norepinephrine × 100).20,21 The occurrence of organ dysfunction and failure, from admission through day 4, was assessed by the Sequential Organ Failure Assessment (SOFA) score.22 The maximal SOFA score was calculated based on the worst score for each organ component in this time period. The neurological subscore was not included because of the high number of patients under sedation both at onset of heart failure and during ICU/CCU treatment. A SOFA subscore ≥3 was defined as organ failure.

**Blood sampling and biochemical assays.** Blood samples were collected in ethylenediaminetetraacetic acid containing tubes at baseline (day 1) and for the next 3 consecutive days. Plasma samples were frozen immediately after centrifugation and stored at −70°C until subsequent analysis. To determine plasma ADMA concentration, samples were assayed using a commercially available competitive enzyme immunoassay kit (ADMA-ELISA, DLD Diagnostika GMBH, Hamburg, Germany). As a normal control, ADMA levels were determined in plasma from 12 healthy volunteers (mean age = 35 years). Markers of endothelial activation, (intracellular adhesion molecule-1 [iCAM-1] and vascular cell adhesion molecule-1 [vCAM-1]) and the inflammatory markers interleukine 6 (IL-6), interleukine 8 (IL-8) all were quantified in duplicates using a bead-based multiplex assay (Bioplex; Bio-Rad Laboratories, Hercules, Calif). Final plasma concentrations were calculated using the Bioplex software supplied by the manufacturer.

**Measurement of peripheral vasodilator function.** Peripheral vasodilator function was assessed with noninvasive digital pulse amplitude tonometry using the EndoPAT 2000 (Itamar Medical Ltd, Cae-

**RESULTS**

**Study population.** Patient characteristics including prior medical history, precipitating factors for acute heart failure, and in hospital procedures are presented in Table I. Most patients with CS presented with acute myocardial infarction and were treated with early percutaneous coronary interventions. Two patients in
this group had coronary bypass surgery during
hospitalization. Treatment with intra-aortic balloon
pumps and inotropes/vasopressors were applied in most
patients. The open-heart surgical procedures prior to
PC-HF included 4 isolated coronary artery bypass
grafting (CABG) procedures, 4 aortic valve
replacements (2 redo procedures), 1 mitral valve
replacement and CABG, 1 composite graft replacement
in the ascending aorta and CABG, 1 procedure on the
thoracic aorta, and 1 redo closure of left ventricular
rupture. No significant differences were found between
the 2 groups with regard to the need for inotropic/
vasopressor support. However, the duration of IABP
support tended to be longer for CS patients ($P = 0.057$).

Baseline hemodynamic variables and laboratory
variables are presented in Table II. Mean time from
the onset of acute heart failure to the initial blood
sampling and endothelial function measurement was,
respectively, 17 and 19 h. CS patients showed on aver-
age more severe derangements in their blood gas at
baseline suggesting a more profound hypoperfusion in
this group. This coincided with significantly lower
diuresis and systolic blood pressure.

**Markers of inflammation and endothelial
activation.** Circulating levels of IL-6, IL-8, iCAM-1,
and vCAM-1 were not different at baseline between the
groups (Table II). IL-6 levels decreased from baseline
through day 4 for both CS ($P = 0.013$) and PC-HF
($P < 0.001$) and was significantly lower on days 3 and 4
compared with baseline (Fig 3, A). Similar longitudinal
changes were observed for IL-8 levels with a significant
decrease for both CS ($P = 0.004$) and PC-HF
($P < 0.001$) (Fig 3, B). IL-6 was negatively correlated
with MAP in both CS ($r_s = -0.57, P < 0.001$) and
PC-HF ($r_s = -0.567, P < 0.001$). Increased IL-6 levels at
baseline also were associated with increased need for
vasopressors in CS patients ($r_s = 0.65, P = 0.02$).

Circulating levels of iCAM and vCAM did not
change from days 1 to 4 for any of the groups (Fig 3,
C and D). iCAM and vCAM were correlated in both
groups (CS $r_s = 0.66, P < 0.0001$, PC-HF $r_s = 0.73, P = 0.0001$).

**Digital vasodilator function.** Satisfactory measurements
were obtained in 86 out of 94 potential recordings. Poor
patient cooperation and failure to receive a readable dig-
tal pulse signal were the main reasons for missing data.
The measured RH-index at baseline (day 1) was
significantly attenuated compared with healthy controls
(RH-index, 2.28) for both CS (RH-index 1.35, $P = 0.001$) and PC-HF (RH-index 1.45, $P = 0.001$). The
baseline RH-index was not significantly different
between CS and PC-HF (Table II). Also, no significant
changes were noted in RH-index for days 1 through 4
for CS patients ($Fig 1, A, P = 0.521$). In patients with
PC-HF, the RH-index changed significantly through the
observation period ($P = 0.019$) and was increased at
days 3 and 4 compared with baseline ($Fig 1, A, P = 0.011$). The individual repeated measurements for both
groups are displayed in Figures 1, B and C and illustrate
the variability among patients both at baseline and
during the course of their illness. At baseline, RH-index
positively correlated with MAP in PC-HF patients ($r_s =
0.70, P = 0.017$). This association also was present
when all recordings were pooled ($r_s = 0.55, P < 0.0001$). RH-index also correlated to ADMA levels at
baseline ($r_s = -0.633, P = 0.036$) for CS patients but
not for PC-HF patients. No correlation was found
among the measured RH-index and systolic blood
pressure, IS, or vasopressor requirements at baseline.

A linear mixed effect model was used to examine for
associations among day-to-day changes in RH-index;
the concomitant changes in MAP; and plasma levels
of ADMA, IL-6, IL-8, iCAM-1, vCAM-1. In patients
with PC-HF, increasing MAP and decreasing levels of
IL-6 and IL-8 was associated with improvement in
RH-index. The only significant correlate for improved
RH-index in CS patients was decreasing levels of
vCAM-1. We could not detect a significant association
between the longitudinal changes in RH-index and
ADMA levels in either of the groups.

**ADMA.** The measured plasma levels of ADMA are shown in
Figure 2. Baseline levels of ADMA in CS were similar to those measured in healthy volunteers
(0.74 vs 0.79 $\mu$mol/L, $P = 0.67$) but significantly
elevated compared with PC-HF (0.55 $\mu$mol/L, $P = 0.04$). A group difference also was found in the course
of ADMA levels from baseline through day 4 (Fig 2).
ADMA levels increased in patients with PC-HF ($P =
0.001$) and was significantly elevated compared with
baseline at day 3 ($P = 0.017$) and at day 4 ($P = 0.001$). No significant longitudinal changes were
observed for CS patients ($P = 0.58$). Repeating
the analysis excluding patients receiving hemodialysis
or hemofiltration (2 patients in CS group) did not
difficult these relations. However, a significant positive
association was found between the longitudinal
changes in ADMA and creatinine ($P = 0.032$) through
the observation period. Patients who underwent
cardiopulmonary resuscitation prior to CS onset had
significantly higher baseline ADMA levels (1.05 vs
0.58 $\mu$mol/L, $P = 0.003$). In CS patients, baseline
ADMA levels correlated with both lactate measured at
baseline ($r_s = 0.85, P = 0.001$) and the maximal
lactate measured within 48 h ($r_s = 0.90, P < 0.001$).
This association was not present in PC-HF patients.
No correlations were found between baseline ADMA
and the need for vasopressors or IS. Baseline ADMA levels did correlate with baseline MAP for PC-HF patients ($r_s = .73, P = .0007$) but not for CS patients.

No association between day-to-day changes in ADMA and changes in MAP, IL-6, IL-8, iCAM-1, vCAM-1 could be observed in either group.

**In hospital outcome and organ dysfunction.** Hospital mortality rates in CS and PC-HF was, respectively, 42% and 17%. However, only 1 fatality occurred during the first 4 days of follow-up (CS group). No significant difference was found in RH-index and baseline levels of ADMA, IL-6, IL-8, iCAM-1, and vCAM-1 between hospital survivors and nonsurvivors. The maximal total SOFA score (excluding neurological subscore) for days 1 through 4 was not significantly different between CS and PC-HF patients (Table I).

**Table I.** Study population. Baseline characteristics, precipitating factors, and in hospital treatment and outcomes

<table>
<thead>
<tr>
<th></th>
<th>CS (n = 12)</th>
<th>PC-HF* (n = 12)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years (mean, range)</td>
<td>65.8 (49–85)</td>
<td>70.7 (58–79)</td>
</tr>
<tr>
<td>Male gender, n</td>
<td>8</td>
<td>10</td>
</tr>
<tr>
<td>Prior medical history, n</td>
<td>5</td>
<td>7</td>
</tr>
<tr>
<td>Coronary heart disease</td>
<td>5</td>
<td>4</td>
</tr>
<tr>
<td>Myocardial infarction</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Chronic heart failure</td>
<td>8</td>
<td>2*</td>
</tr>
<tr>
<td>Hypertension</td>
<td>0</td>
<td>6</td>
</tr>
<tr>
<td>Severe valvular disease</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>Diabetes</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Stroke/TIA</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Renal failure</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Chronic obstructive pulmonary disease</td>
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<td>0</td>
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<tr>
<td>Current smoking</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>Precipitating factors, n</td>
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</tr>
<tr>
<td>Acute coronary syndromes</td>
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</tr>
<tr>
<td>STEMI</td>
<td>7</td>
<td></td>
</tr>
<tr>
<td>Non-STEMI</td>
<td>4</td>
<td></td>
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<tr>
<td>Arrhythmias</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>CPR§</td>
<td>5</td>
<td>2</td>
</tr>
<tr>
<td>Treatment and procedures</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PCI, n</td>
<td>9</td>
<td></td>
</tr>
<tr>
<td>CABG, n</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Other heart surgery, n</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Mechanical ventilation, n</td>
<td>7</td>
<td>8†</td>
</tr>
<tr>
<td>Pulmonary artery catheter, n</td>
<td>6</td>
<td>3</td>
</tr>
<tr>
<td>Ventricular assist device, n</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>IABP, n</td>
<td>11</td>
<td>10</td>
</tr>
<tr>
<td>Duration of IABP support, h</td>
<td>167 (65–240)</td>
<td>55 (39–180)</td>
</tr>
<tr>
<td>Inotropes, n</td>
<td>11</td>
<td>11</td>
</tr>
<tr>
<td>Vasopressors, n</td>
<td>10</td>
<td>12</td>
</tr>
<tr>
<td>Duration of vasopressor/inotropic support, h</td>
<td>84 (55–148)</td>
<td>51 (24–94)</td>
</tr>
<tr>
<td>Maximal vasopressor dose$^{15}$</td>
<td>0.10 (0.01–0.28)</td>
<td>0.13 (0.07–0.21)</td>
</tr>
<tr>
<td>Maximal IS$^2$</td>
<td>18.5 (5.2–31.0)</td>
<td>17.2 (12.4–30.0)</td>
</tr>
<tr>
<td>Hemofiltration or hemodialysis, n</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Length of ICU/CCU stay, days</td>
<td>10 (5–17)</td>
<td>7 (3–14)</td>
</tr>
<tr>
<td>Maximal SOFA score$^*$</td>
<td>7.5 (6.0–14.8)</td>
<td>8.5 (6.3–11.5)</td>
</tr>
<tr>
<td>Organ failure$^{11}$ n</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>In hospital mortality, n</td>
<td>5</td>
<td>2</td>
</tr>
</tbody>
</table>

$^1$Continuous variables are presented as median with IQR if not stated otherwise.

$^2$PCI, percutaneous coronary intervention; TIA, transitory ischemic attack.

$^3$Surgical procedures are presented under methods.

$^4$Patients necessitating intubation exceeding the first postoperative day.

$^5$Given as the total dose of epinephrine and norepinephrine (ug/kg/min).

$^6$Cardiopulmonary resuscitation prior to inclusion.

$^7$Total sequential organ failure assessment score (SOFA) score except neurological SOFA subscore.

$^8$Failure of ≥1 organ other than cardiovascular failure.

$^9$P < 0.05 compared with CS.
In CS patients, the SOFA score was correlated to baseline RH-index ($r_s = -0.69$, $P = 0.014$), baseline ADMA levels ($r_s = -0.62$, $P = 0.003$), and IL-8 ($r_s = -0.79$, $P = 0.007$). Furthermore, baseline ADMA levels were strongly correlated to the hepatic ($r_s = 0.79$, $P = 0.003$) and respiratory ($r_s = 0.77$, $P = 0.004$) SOFA subscores, whereas no significant associations were observed for the other organs including the kidneys ($r_s = 0.55$, $P = 0.065$). ADMA levels increased in 7 patients and decreased in 5 patients through the observation period. A negative correlation existed between the absolute change in ADMA levels and the degree of hepatic and respiratory dysfunction ($r_s = -0.63$, $P = 0.024$ and $r_s = -0.71$, $P = 0.01$, respectively). CS patients with failure of an organ other than the central nervous system or cardiovascular system had lower baseline RH-index (median 1.22 vs 1.51, $P = 0.01$), elevated ADMA levels (median 0.58 vs 1.05 umol/L, $P = 0.005$), and increased IL-8 (median 27 vs 150 pg/mL) compared with patients without organ failure. A tendency was also noted toward increased IL-6 levels in CS patients with organ failure, but this finding was not statistically significant (132 vs 343 pg/mL, $P = 0.202$).

For patients with PC-HF, no association was found among SOFA scores and baseline RH-index, ADMA, or IL-6. IL-8 did, however, correlate to SOFA score ($r_s = 0.79$, $P = 0.002$). Accordingly, baseline RH-index, ADMA, IL-8, and IL-6 were not different in patients with organ failure compared with those without. Also, no correlation was found between the change in ADMA levels and any of the SOFA subscores. No association was evident among iCAM-1, vCAM-1, and SOFA scores in any group.

**DISCUSSION**

Our study shows that patients with the most severe forms of acute heart failure (ie, CS and PC-HF) are subject to a profound inflammatory insult evident by excessive circulating levels of IL-6 and IL-8, markedly higher than reported in acute decompensated heart failure and acute myocardial infarction. These levels are comparable with the levels measured in sepsis. This finding was accompanied by a sustained elevation in markers of endothelial activation. Although the systemic inflammatory insult and endothelial activation inflicted by open-heart surgery and use of extra corporeal life support were mitigated in both patient groups, patients with CS had a higher degree of organ failure, a greater inflammatory response, and more pronounced activation of the inflammatory-endothelial axis compared with patients with PC-HF.

**Table II. Baseline laboratory and hemodynamic variables**

<table>
<thead>
<tr>
<th>Variable</th>
<th>CS n = 12</th>
<th>PC-HF n = 12</th>
</tr>
</thead>
<tbody>
<tr>
<td>CK-mb (µg/L), maximal value</td>
<td>207 (47–493)</td>
<td>56 (25–133)</td>
</tr>
<tr>
<td>Plasma creatinine (µmol/L)</td>
<td>125 (99–183)</td>
<td>85 (81–121)</td>
</tr>
<tr>
<td>eGFR (mL/min)</td>
<td>57 (40–114)</td>
<td>84 (63–102)</td>
</tr>
<tr>
<td>Lactate (mmol/L), Baseline*</td>
<td>3.3 (1.8–7.9)</td>
<td>1.9 (1.5–6.2)</td>
</tr>
<tr>
<td>Highest value†</td>
<td>3.3 (1.8–8.4)</td>
<td>5.2 (2.1–7.0)</td>
</tr>
<tr>
<td>Arterial pH, Baseline*</td>
<td>7.26 (7.11–7.31)</td>
<td>7.36 (7.29–7.43)</td>
</tr>
<tr>
<td>Lowest value†</td>
<td>7.20 (7.05–7.30)</td>
<td>7.26 (7.13–7.34)</td>
</tr>
<tr>
<td>Base excess (mmol/L), Baseline*</td>
<td>-7.7 (–12.6––3.9)</td>
<td>-4.4 (–5.7––1.8)</td>
</tr>
<tr>
<td>Lowest value†</td>
<td>-8.1 (–17.8––3.9)</td>
<td>-6.2 (–12.5––2.7)</td>
</tr>
<tr>
<td>C-reactive protein (mg/L)</td>
<td>33 (9–135)</td>
<td>69 (10–121)</td>
</tr>
<tr>
<td>White blood cells (10^9/L)</td>
<td>17.2 (9.8–19.8)</td>
<td>9.5 (5.4–11.7)</td>
</tr>
<tr>
<td>ADMA (µmol/L)</td>
<td>0.62 (0.52–1.11)</td>
<td>0.51 (0.37–0.68)</td>
</tr>
<tr>
<td>IL-6 (pg/mL)</td>
<td>281 (131–459)</td>
<td>342 (197–691)</td>
</tr>
<tr>
<td>IL-8 (pg/mL)</td>
<td>121 (26–122)</td>
<td>57 (43–140)</td>
</tr>
<tr>
<td>iCAM-1 (ng/mL)</td>
<td>96 (64–152)</td>
<td>98 (88–269)</td>
</tr>
<tr>
<td>vCAM-1 (ng/mL)</td>
<td>349 (257–441)</td>
<td>367 (311–412)</td>
</tr>
<tr>
<td>RH-index</td>
<td>1.35 (1.27–1.56)</td>
<td>1.45 (1.00–1.86)</td>
</tr>
<tr>
<td>Mean arterial blood pressure (mmHg)</td>
<td>60 (56–67)</td>
<td>64 (57–68)</td>
</tr>
<tr>
<td>Systolic blood pressure (mmHg)</td>
<td>80 (75–85)</td>
<td>90 (80–99)</td>
</tr>
<tr>
<td>Diastolic blood pressure (mmHg)</td>
<td>53 (46–58)</td>
<td>50 (40–55)</td>
</tr>
<tr>
<td>Cardiac index (L/min/m^2)†</td>
<td>2.18 (1.51–2.47)</td>
<td>2.097 (1961–2576)</td>
</tr>
<tr>
<td>SVRI (dynes<em>sec/cm^5</em>m^2)‡</td>
<td>2097 (1961–2576)</td>
<td>2097 (1961–2576)</td>
</tr>
<tr>
<td>Diuresis, ml/hour</td>
<td>10 (0–29)</td>
<td>68 (25–91)</td>
</tr>
<tr>
<td>HR (beats per min)</td>
<td>110 (85–120)</td>
<td>88 (85–99)</td>
</tr>
</tbody>
</table>

Values are presented as median with IQR.
CK-mb, creatine kinase mb; eGFR, estimated glomerular filtration rate; SVRI, systemic vascular resistance index.
*First measurement after onset of CS or PC-HF.
†Measured within first 48 h.
‡n = 6. eGFR calculated using the modification of diet in renal disease formula.
§P < 0.05 compared with cardiogenic shock.
bypass is well described, the cause and consequence of this in CS is less well understood. However, the previously described association between IL-6 levels and vasopressor requirements to restore blood pressure in CS also was evident in our study as high IL-6 levels were both related to low MAP and increased vasopressor dose.6

The peripheral vasodilator function was assessed serially through the first 4 days after acute heart failure onset. The RH-index was impaired at baseline in both groups compared with healthy controls. This finding was, however, not uniform as several individuals had a preserved RH-index indicating a preserved vascular and endothelial function. A few prior studies have assessed vascular reactivity in shock and critical illness demonstrating impaired vascular response during reactive hyperemia. Kirschenbaum et al found an attenuated increase in forearm blood flow after reactive hyperemia in patients with cardiogenic and septic shock compared with healthy controls.31 A similar observation was made in patients with severe sepsis using peripheral arterial tonometry similar to the present study.59 The repeated assessments in our study also revealed a substantial individual day-to-day variation. This finding underlines that vascular function and probably endothelial function in acute disease are not static, which is an important consideration to make when assessing vascular function in acute versus chronic disease states.

The mechanisms behind the apparent attenuation in vasodilator function and the changes observed through the course of the disease are likely to be multifactorial and probably a composite of prior vascular/endothelial function and a multitude of superimposed acute factors.32-34 Also, fewer functional capillaries secondary to intravascular obstruction or edema could blunt the reactive hyperemic response in critical disease.55 The RH-index, as a measure of the peripheral circulations ability to respond to transitory local ischemia, is likely a result from all these factors. Being in part NO-dependent, the impaired RH-index
at baseline could suggest a reduced endothelial NO bioavailability. The negative relation observed between the endogenous NO-inhibitor ADMA and the RH-index at baseline in CS could suggest an inhibitory effect of ADMA on vasodilator function. However, no longitudinal association was found between the two to back up this hypothesis. Also, this relation was not observed in PC-HF. It is possible that attenuation of other important contributors to endothelium dependent dilation, such as prostacyclin or endothelium derived hyperpolarizing factor, play a bigger role. Interestingly, the observed improvement in vasodilator function in PC-HF was associated with a concomitant decrease in inflammatory cytokines. A transitory impairment of endothelial function after exposure to cytokines and transitory inflammation has been described in an experimental setting and possibly could explain the effects observed here. Furthermore, tumor necrosis factor α (TNFα) and IL-6 also have been shown to induce endothelium-dependent vasoconstriction in human arterial segments.

Role of ADMA in CS and PC-HF. Circulating ADMA levels were similar in CS and healthy controls. The normal values found in this study coincide well with a recently proposed normal reference value (mean 0.69, 95% confidence interval: 0.36–1.17 μmol/L) and other reports using a similar enzyme-linked immunosorbent assay technique. This is in contrast to previous reports measuring elevated levels of ADMA after acute decompensated heart failure and CS using high-performance liquid chromatography and mass spectrometry, respectively. A significant elevation was, however, evident in the most critically ill CS patients in our cohort with high arterial lactate and organ failure.

Baseline ADMA levels were suppressed in PC-HF, and although no pattern of change was apparent during the first 4 days in CS, ADMA tended to increase in PC-HF patients. This could possibly be caused by a hemodiluting effect of extra corporeal circulation, as initially suppressed postoperative ADMA levels previously have been demonstrated in patients undergoing open-heart surgery. This consideration is important to make when using ADMA as a risk marker in critically ill patients in whom surgery has been done. The relatively suppressed ADMA levels in PC-HF also could reflect that, in our population of patients, this condition is more benign and that most of these patients were in a state of recovery as compared with CS. However, no indication was given that the degree or course of inflammation and endothelial activation were related to ADMA levels. Except for the observed correlation with MAP at baseline in PC-HF, ADMA levels were

Fig 3. A, Circulating plasma levels of IL-6. B, IL-8. C, iCAM-1. D, vCAM-1. One patient in the CS group died at day 2. *P < 0.05 compared with day 1 for CS patients. *P < 0.05 compared with day 1 for PC-HF patients.
not related to blood pressure or vasopressor/inotrope requirements.

Elevated ADMA levels were associated with the overall degree of organ dysfunction (SOFA score) and elevated systemic lactate in CS. Accordingly, CS patients with organ failure had significantly higher ADMA levels. In particular, this elevation was related to the degree of respiratory and hepatic dysfunction. Although baseline ADMA levels were not significantly correlated with the renal SOFA score, the longitudinal positive association between ADMA and creatinine were consistent with prior reports that ADMA accumulates secondary to renal dysfunction. These observations are in line with findings presented in previous reports from patients with severe sepsis and in critically ill patients with organ failure. Accumulation of ADMA is thought to occur mainly through reduced metabolism secondary to dysregulation and inhibition of the dimethylarginine dimethylaminohydrolase (DDAH). This enzyme is highly expressed in both liver and kidneys, which are presumed to be the main routes for metabolic clearance of circulating ADMA. However, ADMA levels were highest in patients receiving cardiopulmonary resuscitation (CPR) either before or at the onset of shock. It is likely that these patients had the most profound systemic hypoperfusion. Unlike septic patients, the observed early elevated lactate levels in CS patients are likely a result of systemic hypoperfusion rather than liver failure. It is from the available data not possible to determine whether patients in shock already have accumulated levels of ADMA or if they develop this after shock onset secondary to acute reduction in hepatic or renal elimination. Increased ADMA levels in patients who had CPR and/or had elevated arterial lactate could indicate that ADMA as a marker of elevated risk in CS merely reflects the severity of hypoperfusion during shock. However, ADMA also could potentially play a causative role in hypoperfusion and development of organ failure through inhibiting the NO-mediated regulation of organ blood flow.

Although ADMA was increased in patients with failing organs compared with those without, these patients did not accumulate more ADMA during the observation period. On the contrary, ADMA decreased more in patients with the highest degree of hepatic dysfunction.

Our study did not support an association between the degree of systemic inflammation and circulating ADMA. In vitro studies on endothelial cells, however, have demonstrated that oxidative stress, stimulation with TNFα, and nitrosative stress resulting from an induction of inducible NO synthase cause an accumulation of ADMA secondary to reduced DDAH activity. However, increased DDAH activity secondary to stimulation with inflammatory cytokine have been reported in rat vascular smooth muscle cells, and a mechanistic knowledge of the important metabolic responses and the regulation of the NO-system and vasculature is still warranted.

LIMITATIONS

The vascular function measurements were performed in a clinical setting in the ICU/CCU on patients receiving several different pharmacological and mechanical treatments through the observation period. This lack of standardization could affect the quality of these measurements. Invasive hemodynamics were only measured in 1/4 of the patients, and thus, no evaluation of RH-index in relation to changing hemodynamics could be performed. Also, multiple statistical comparisons based on a limited number of patients inherits the risk of type I errors.

CONCLUSION

In this study, both CS and PC-HF were characterized by initially elevated levels of inflammatory cytokines suggestive of a profound inflammatory insult accompanied by an attenuated peripheral vasodilator function. The resolving inflammatory response was associated with improved RH-index in PC-HF patients, whereas no such relation was evident for CS patients. The circulating levels of the endogenous NO-inhibitor ADMA did not relate to the degree of inflammation or to the day-to-day changes in vasodilator function. However, in CS, ADMA, vasodilator function, and levels of inflammatory cytokines were correlated to the overall degree of organ dysfunction, and to hepatic dysfunction in particular. The increased ADMA levels in patients who had CPR and/or elevated arterial lactate suggest a relation between ADMA and the degree of hypoperfusion in CS.

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REFERENCES


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