The role of vasoactive agents in the resuscitation of microvascular perfusion and tissue oxygenation in critically ill patients
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Abstract  Purpose: The clinical use of vasoactive drugs is not only intended to improve systemic hemodynamic variables, but ultimately to attenuate derangements in organ perfusion and oxygenation during shock. This review aims (1) to discuss basic physiology with respect to manipulating vascular tone and its effect on the microcirculation, and (2) to provide an overview of available clinical data on the relation between vasoactive drugs and organ perfusion, with specific attention paid to recent developments that have enabled direct in vivo observation of the microcirculation and concepts that have originated from it. Methods: A MedLine search was conducted for clinical articles in the English language over the last 15 years pertaining to shock, sepsis, organ failure, or critically ill patients in combination with vasoactive drugs and specific variables of organ perfusion/oxygenation (e.g., tonometry, indocyanine clearance, laser Doppler, and sidestream dark field imaging). Results: Eighty original papers evaluating the specific relationship between organ perfusion/oxygenation and the use of vasoactive drugs were identified and are discussed in light of physiological theory of vasomotor tone. Conclusions: Solid clinical data in support of the idea that increasing blood pressure in shock improves microcirculatory perfusion/oxygenation seem to be lacking, and such a concept might not be in line with physiological theory of microcirculation as a low-pressure vascular compartment. In septic shock no beneficial effect on microcirculatory perfusion above a mean arterial pressure of 65 mmHg has been reported, but a wide range in interindividual effect seems to exist. Whether improvement of microcirculatory perfusion is associated with better patient outcome remains to be elucidated.

Keywords  Organ perfusion · Oxygenation · Microcirculation · Vasoactive · Shock · Vasomotor tone

Introduction

Although shock is defined in terms of a critically low blood pressure, a more generally accepted and physiology-based definition of shock is: the inability of the circulation to sustain the cellular respiration needed to maintain normal organ function [1]. This controversy in definition reflects an underlying dilemma with profound effects at bedside. Although intensivists are aware of the physiology-based mechanisms of shock, they have to rely on surrogate global upstream (e.g., blood pressure) and/or downstream (e.g., SvO2) hemodynamic variables to
diagnose ‘inadequate’ organ perfusion and guide therapy. Attempts to integrate regional indicators such as intestinal CO₂, lactate and indocyanine green (ICG) clearance, which have all been associated with prognosis, have had limited success.

Apart from these general limitations, it becomes clear that in distributive shock shunting in the microcirculation is an intrinsic complexity. Although theoretically the concept of shunting has been well known for many decades [2], new technologies have elucidated the role of ‘microcirculatory weak units.’ In a model of ischemia reperfusion, a speckled pattern of NADH fluorescence and pO₂ emerged after re-oxygenation, indicating an inhomogeneous distribution of cells more vulnerable to ischemic insults than others. Microsphere embolization determined these so-called ‘weak units’ at the capillary level [3]. Provided upstream oxygen delivery is adequate, shunted parts of the microcirculation remain hypoxic in the presence of well-maintained venous pO₂, explaining the persistence of elevated lactate and regional pCO₂ levels [4] not sensed by the downstream variables such as SvO₂ [5]. The net systemic result is an inability to increase tissue oxygen uptake, despite efforts to ‘(supra)normalize’ oxygen delivery, but whether this reflects a true extraction deficit as result of a preceding cellular metabolic shutdown or a potentially correctable state of inhomogeneous hypo perfusion remains to be established [6]. Because of the introduction of direct in vitro observation of the microcirculation [7], heterogeneity of blood flow has now become a key feature of distributive shock. It has not only been reported within the microcirculation [8], but also between different organs, within different regions of one organ, in response to therapeutic interventions and in a time-dependent way [4, 9], all challenging the simplicity of systemic hemodynamic monitoring under such conditions.

At first evaluation, these observations seem to add more data to the already dazzling complexity of treatment modalities in the critically ill. However, these new insights have established two main findings: (1) the independent perfusion behavior of the microcirculation in relation to classically measured systemic hemodynamic variables, albeit within certain absolute limits of minimal perfusion pressure [8], and (2) persistence of microcirculatory alterations are associated with morbidity and mortality under various conditions, irrespective of correction of systemic hemodynamics [10, 11]. The issue to be resolved is whether correction of microcirculatory distributive alterations will improve organ function and morbidity. In order to explore this exiting concept, it is not sufficient to ‘simply’ find novel therapeutic approaches that specifically aim to reopen the microcirculation directly. We also have to be prepared to (re)evaluate our current strategies at the bedside in the light of its potential effects from the perspective of the microcirculation and relate it to organ function and outcome.

Administration of vasoactive compounds and fluid therapy are the cornerstone of hemodynamic management of critically ill patients. In this paper we will review reported effects of vasoactive substances on microcirculatory perfusion and tissue oxygenation in shock. These compounds have been advocated to improve systemic perfusion pressure and oxygen delivery, with subsequent improvement of tissue perfusion and oxygen uptake [12]. But do they behave the way we expect them to do? Since shock is a syndrome rather than a well-defined pathophysiologic entity, disease itself as well as treatment might contribute to the clinical picture [13]. An appreciation of basic physiology with respect to manipulating vascular tone and its effect on the microcirculation is followed by a systematic review of the literature. We will focus our review on recent clinical research that includes microcirculatory effects of three classes of vasoactive substances: vasopressors, inotropes and vasodilators, and follow the suggested classification of shock etiology [2]. Preclinical studies will only be discussed in so far as they pertain to the clinical studies.

Materials and methods

In MedLine the following search strategy was performed. Article titles containing shock or sepsis or septic or organ failure or critically ill in combination with vasopressor or vasoactive or inotrope or inotropic or vasodilator or several specific vasoactive drugs [e.g., (nor)adrenaline/(nor)epinephrine, vasopressin, dopamine, dobutamine, nitroglycerin] were selected. The search was restricted to clinical studies in the English language, published over the last 15 years, where indicators of microcirculatory or regional perfusion were specifically mentioned, or in case the effect was evaluated with aid of specific microcirculatory or regional techniques [e.g., tonometry/capnography, ICG, orthogonal polarization spectral (OPS)/side-stream dark field (SDF) imaging, spectrophotometry (Table 1)].

Results

Physiology

Microcirculatory perfusion pressure is the net result of precapillary inflow pressure minus venular outflow pressure. Most of the pressure drop occurs upstream in small arterioles (resistance vessels) as the principal site of active diameter changes to achieve regulation of blood flow. Mean capillary pressure is therefore much closer to venous pressure than to arterial pressure, which is relevant for the maintenance of tissue fluid balance [14].
<table>
<thead>
<tr>
<th>Technique</th>
<th>Principle</th>
<th>Data expressed as</th>
<th>Area of interest</th>
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<tbody>
<tr>
<td>Tonometry</td>
<td>Balloon content (saline: tonometry, air: capnography) equilibrates with mucosal cellular fluid/gas. Content is quantified with classical bloodgas technology (pH, $pCO_2$, $HCO_3^-$)</td>
<td>Intestinal pH or difference between intestinal and arterial ($pCO_2_{g-a}$)</td>
<td>Mucosal capillary blood flow</td>
<td>Detects mucosal dysxia (imbalance between oxygen demand and supply) as a result of inadequate microvascular perfusion. Hypoxic hypoxia is not detected, serosal perfusion can influence measurements.</td>
</tr>
<tr>
<td>Indocyanine green clearance</td>
<td>Dye dilution method. The flow is related to the quantity of the injected indicator dye divided by the area under the curve of the downstream dye concentration, according to the Stewart-Hamilton equation</td>
<td>l/min</td>
<td>Hepatosplanchnic vascular compartment</td>
<td>Indocyanine green clearance is dependent on overall hepatosplanchnic blood flow and hepatic cellular blood flow function (altered in liver failure)</td>
</tr>
<tr>
<td>Laser-Doppler flowmetry</td>
<td>Shift in frequency between emitted and received signal related to (red) blood cell velocity (Doppler principle); flow = velocity $\times$ probe catchment area</td>
<td>m/s (velocity) or l/min (flow)</td>
<td>Catchment volume incorporates arteriolar, capillary and venular blood flow</td>
<td></td>
</tr>
<tr>
<td>Spectrophotometry</td>
<td>Light with a specific wavelength is emitted by a spectrometer. Given a specific distance, the solute concentration or color between the spectrometer and the receiving photometer determines the (change in) received light intensity, according to Beer’s law</td>
<td>Optical density units from which hemoglobin saturation (%) is calculated</td>
<td>Catchment area incorporates capillary and venular hemoglobin saturation</td>
<td>Hemoglobin saturation may be normal or elevated under conditions of microvascular heterogeneity in capillary blood flow due to shunting</td>
</tr>
<tr>
<td>Orthogonal polarization spectral imaging</td>
<td>Direct in vivo video microscopy. Absorbance of green light (540 nm) by hemoglobin projects red blood cells as black dots. Polarization prevents scattering of light</td>
<td>Semi-quantitative: microvascular blood flow index (convective transport) and perfused capillary density (diffusion distance) quantitative: functional capillary density (software analysis needed)</td>
<td>Images less than 1 mm$^2$ discriminates between capillaries and venules</td>
<td>Visualization of vessels depends on presence of red blood cells. Enables to visualize distributive changes in blood flow between individual capillaries</td>
</tr>
<tr>
<td>Sidestream dark field imaging</td>
<td>Direct in vivo video microscopy. Absorbance of green light (540 nm) by hemoglobin projects red blood cells as black dots. Ring of stroboscopic light emitting diodes around the camera prevents scattering of light</td>
<td>Semi-quantitative: microvascular blood flow index (convective transport) and perfused capillary density (diffusion distance). Quantitative: functional capillary density (software analysis needed)</td>
<td>Images less than 1 mm$^2$ discriminates between capillaries and venules</td>
<td>Visualization of vessels depends on presence of red blood cells. Enables to visualize distributive changes in blood flow between individual capillaries</td>
</tr>
</tbody>
</table>
Under normal conditions sympathetic vasoconstriction is reduced in case of venular hypoxia. This signal is forwarded upstream electrophysiologically via endothelial cells connected by gap junctions proximal to arterioles causing upstream dilatation. This process is regulated per microcirculatory unit arising from the respective arteriole feed vessels [15]. It is proposed that the ‘sensor’ for detecting changes in ambient $pO_2$ is located downstream and acts by release of nitric oxide (NO) in the venules [16]. In this respect additional exogenous precapillary and postcapillary vasoconstriction by means of a vasopressive drug results in a reduced net perfusion pressure over the microcirculation [17], despite increment of systemic blood pressure (Fig. 1). At the same time shunting may disrupt the normal venular signaling process for vasodilatation to prevent local hypoxia. Indeed, in ischemia/reperfusion local hypoxia persists in ‘microcirculatory weak units’ after restoration of oxygen delivery in a speckled pattern that appears to be determined at the capillary level, but not at the arteriolar level [3].

Apart from interference in the complex regulation of tissue perfusion, vasoactive drugs may also influence the homeostasis of tissue oxygenation. Tissue oxygenation is not only dependent on convective properties of red blood cells (flow), but is also determined by diffusion. Given the gas-specific characteristics, oxygen diffusion is related to the pressure gradient and inversely related to the distance between the capillary and the cell (Fig. 2). Closing

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\text{Fig. 1 Left panel arteriolar vasodilation increases the opening pressure of the microcirculation as result of a decrease in pressure drop prior to the microvascular compartment. Right panel combined arteriolar and venular increment of vascular tone reduces the net driving pressure over the microvascular compartment (from [17] by permission)}
\]

\[
\text{Fig. 2 Convective transport of oxygen through the capillaries depends on red blood cell velocity, capillary hematocrit and oxygen saturation. Oxygen transport from the capillary to the cell via diffusion is inversely related to the diffusion distance ($D_1$ and $D_2$) according to Fick's law}
\]

\[
\text{\[VO_2 = \frac{C \times A \times (cappO_2 - mitpO_2)}{D}\]
}
\]

\[
\text{VO_2 oxygen volume transported by diffusion}
\]
\[
\text{C diffusion constant}
\]
\[
\text{A capillary surface area}
\]
\[
\text{cappO_2 capillary pO_2}
\]
\[
\text{mitpO_2 mitochondrial pO_2}
\]
capillaries in order to maintain perfusion pressure increases diffusion distance and vice versa. Under resting conditions capillaries constantly perform changes in cal-iber in order to serve both purposes [15].

A third factor that influences microcirculatory oxygen delivery is capillary hematocrit. Mechanical interaction between red blood cells and vessel walls induces the formation of a plasma layer adjacent to the wall and increases hematocrit in the center. Since blood flow-velocity distribution has a parabolic shape, from zero next to the wall to a maximum at the center, the average red cell velocity is higher than overall blood velocity. As a consequence, red cell transit time is reduced. This results in dynamic lowering of the capillary hematocrit in comparison to the entering and discharge hematocrit, and is known as the Fahraeus effect [18]. Additional to this effect, capillary hematocrit is also determined by a phenomenon, originally described by Krogh as ‘plasma skimming’ [19]. At the diverging branches of the capillary network, distribution of red cells is related to the diameter of the daughter vessels (Fig. 3). During in vitro experiments at a systemic hematocrit of 50%, capillary hematocrit ranged from 6.8% during vasoconstriction to 38% under vasodilatation [20], with profound implications on capillary oxygen transport. From the perspective of microcirculation, the dilemma of the use of vasopressors to enhance organ perfusion and oxygenation cannot be better expressed than in the original observations by Krogh: ‘This plasma skimming is usually very pronounced when adrenaline is applied in small drops to muscle arteries of which all branches, even the smallest, react and show a contraction, which in a short time may become complete. The portion of the muscle supplied by the contracting artery becomes blanched and the capillaries often disappear completely from view, while application of adrenaline to capillaries and venules alone shows that these vessels do not react visibly to the sub-stance’ [19].

Despite abundant experimental data, little is known about human in vivo alterations of vasomotor tone during shock. Direct in vivo observations of the microcirculation in animal endotoxemia demonstrated loss of arterial vasomotion. Interestingly, not all vessels were affected at once, but areas with stagnant blood flow were observed next to areas with brisk flow [21]. In humans, topical endotoxin exposure produced acute venous hyporesponsiveness to NE or sympathetic nervous system activation. This was inhibited by glucocorticoids, but not mitigated by NO synthase- or cyclo-oxygenase inhibitors, suggesting other mediators than NO or prostanoids contribute to the overall vascular response to endotoxin [22]. Systemic LPS injection in healthy volunteers blunted the vascular baro-reflex response against nitroprusside-induced hypotension and caused complete uncoupling of the cardiac baroreflex from prevailing blood pressure [23]. At present, no direct observations of the microcirculation, other than in tongue or skin, are available from human endotoxemia models.

**Human studies**

**Vasopressors**

**Norepinephrine** (a and β-adrenergic, Table 2) is the most commonly advocated vasopressive drug in septic shock to maintain mean arterial pressure (MAP) at a minimum level of 65 mmHg [12]. Although suggested, improvement of outcome as a result of the use of NE has never been established [24]. Recent studies, which included direct in vivo observation of the microcirculation with Sidestream Dark Field (SDF) imaging, have established that in human sepsis further increment of MAP above 65 mmHg by means of NE does not improve sublingual microcirculatory perfusion, intestinal pCO₂ concentrations or arterial lactate levels [25, 26]. This confirmed previous reports in which stepwise increases of MAP with
cumulating NE doses did not change arterial lactate levels, skin capillary blood flow, arterial-to-gastric $pCO_2$ gradient ($pCO_2$ g-a), systemic oxygen uptake or renal function [27, 28]. However, substantial inter-individual differences were observed, related to baseline values of microvascular blood flow [25]. Furthermore, these effects may be time and situation dependent [29], and equal changes in MAP may have different drug-related effects on organ perfusion [30]. For renal function an optimal MAP of 75 mmHg in sepsis has been suggested, but direct in vivo observations of the renal microcirculation has not been performed [31]. These studies underline the concept that under conditions of a minimal systemic perfusion pressure, an additional rise in arterial pressure does not automatically improve microcirculatory perfusion in distributive shock, in accordance with physiological theory [17]. Clearly, despite important efforts to compare mortality, systemic and regional perfusion and cardiovascular side effects between (combinations with) NE and other vasopressors, these studies do not elucidate its individual effects on the microcirculation or establish a critical minimum MAP with respect to microcirculatory organ perfusion [32–35]. An observed relation between both hypotension and mortality, as well as vasopressor load and mortality, may be related to the event itself, but may also be a marker for severity of illness [36, 37].

In cardiogenic shock, inotropic agents, but not vasopressors, are advocated [38, 39]. Intra-aortic balloon counter pulsation instantly improved sublingual microcirculatory blood flow in cardiogenic shock, but additional NE

<table>
<thead>
<tr>
<th>Study [ref]</th>
<th>Vasopressor</th>
<th>Setting</th>
<th>Technique(s)</th>
<th>Effects</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>[11, 25]</td>
<td>Norepinephrine</td>
<td>Septic shock, $n = 20$, $n = 20$</td>
<td>SDF-imaging, tonometry, laser-Doppler</td>
<td>No change in sublingual microvascular blood flow, lactate or $pCO_2$ g-a</td>
<td>Stepwise increase MAP from 65 to 85 mmHg, individual effect baseline-dependent</td>
</tr>
<tr>
<td>[27]</td>
<td>Norepinephrine</td>
<td>Septic shock, $n = 10$</td>
<td>Laser-Doppler, tonometry</td>
<td>No difference in skin capillary blood flow, $pCO_2$ g-a, lactate or urine output</td>
<td>Stepwise increase MAP from 65 to 85 mmHg</td>
</tr>
<tr>
<td>[28]</td>
<td>Norepinephrine</td>
<td>Septic shock, $n = 28$</td>
<td>Creatinine clearance</td>
<td>No difference in lactate, urine output or creatinine clearance</td>
<td>Comparison between MAP 65 and 85 mmHg</td>
</tr>
<tr>
<td>[31]</td>
<td>Norepinephrine</td>
<td>Septic shock, $n = 11$</td>
<td>Doppler ultrasonography, creatinine clearance</td>
<td>Decrease in renal resistive index, increase in urine output, no change in creatinine clearance</td>
<td>Effect between MAP 65 and 75, no further change at 85 mmHg</td>
</tr>
<tr>
<td>[42]</td>
<td>Epinephrine</td>
<td>Septic shock, $n = 8$</td>
<td>Tonometry, hepatic vein lactate, ICG clearance</td>
<td>Decrease in mucosal pH and splanchnic blood flow, increase in hepatic vein lactate</td>
<td>Cross-over, in addition to norepinephrine plus dobutamine</td>
</tr>
<tr>
<td>[34]</td>
<td>Phenylephrine</td>
<td>Septic shock, $n = 32$</td>
<td>Tonometry, ICG clearance</td>
<td>No change in splanchnic blood flow, $pCO_2$ g-a, lactate, urine output or creatinine clearance</td>
<td>Increment MAP from 65 to 75 mmHg, comparison between norepinephrine and phenylephrine</td>
</tr>
<tr>
<td>[44]</td>
<td>Phenylephrine</td>
<td>Septic shock, $n = 15$</td>
<td>Tonometry, ICG clearance</td>
<td>Decrease in splanchnic blood flow, $pCO_2$ g-a, lactate or creatinine clearance</td>
<td>Switching from norepinephrine to phenylephrine under steady state MAP at 70 mmHg</td>
</tr>
<tr>
<td>[45]</td>
<td>Phenylephrine</td>
<td>Cardiopulmonary bypass, $n = 15$</td>
<td>SDF-imaging, laser-Doppler flowmetry, spectrophotometry</td>
<td>Decrease in sublingual capillary blood flow together with an increase in microcirculatory hemoglobin oxygen saturation and overall sublingual blood flow, suggesting shunt</td>
<td>Increment MAP from 47 to 68 mmHg under steady state cardiac output</td>
</tr>
<tr>
<td>[74]</td>
<td>Dopamine (high dose)</td>
<td>Septic shock, $n = 20$</td>
<td>Tonometry</td>
<td>Decrease in gastric mucosal pH</td>
<td>Increment MAP from $\leq 60$ to $\geq 75$ mmHg, under norepinephrine intestinal pH decreased</td>
</tr>
</tbody>
</table>

$SDF$ Sidestream dark field, $pCO_2$ g-a arterial-to-gastric $pCO_2$ gradient, MAP mean arterial pressure, ICG indocyanine green.
dosage was inversely correlated with microcirculatory blood flow [40].

**Epinephrine (α and β-adrenergic, Table 2)** in the treatment of septic shock has been subject to controversy. Hyperlactataemia during use of epinephrine in sepsis treatment may be caused by tissue hypoxia, but may also be explained by direct metabolic effects [32, 41]. Human data about direct effects of epinephrine on microcirculatory perfusion in sepsis are not available, but only in comparison to NE. In a crossover design, in comparison to NE plus dobutamine, epinephrine was associated with lower intestinal pH (pH3) and higher hepatic venous lactate levels under equal systemic hemodynamic conditions [42], but again interpretation of these data with respect to splanchnic perfusion is complicated by the intrinsic effect of epinephrine to cause hyperlactataemia and systemic acidosis [41, 43].

**Phenylephrine** (α-adrenergic, Table 2) has been used to increase MAP in human sepsis from 65 to 75 mmHg over a 12-h period [34]. Gastric-to-arterial pCO2 difference, indocyanine clearance, arterial lactate levels and urine output/creatinine clearance were found to be unaltered over time and in comparison to NE, indicating that intestinal microcirculatory organ perfusion neither improved nor deteriorated as a result of a phenylephrine-induced rise in MAP. However, switching from NE to phenylephrine in patients with septic shock while maintaining MAP at 70 mmHg at steady-state systemic hemodynamic variables revealed a rise in pCO2-g-a and a decrease in ICG clearance and urine output/creatinine clearance [44]. These indicators of impaired splanchnic and renal perfusion during phenylephrine infusion disappeared after switch-back to NE again.

In cardiopulmonary bypass (CPB)-induced hypotension, phenylephrine was used to increase MAP from 47 to 68 mmHg [45]. During hypotension, SDF-measured microcirculatory blood flow of sublingual capillaries was unchanged in comparison to pre-CPB hypotension, but after correction with phenylephrine microcirculatory blood flow was significantly reduced. At the same time, global tissue blood flow (laser Doppler) and spectrophotometric microvascular hemoglobin oxygen saturation (Hb(02)) of the tongue increased, underlining the distributive changes that take place as a result of phenylephrine administration, not sensed in the absence of direct observation of the microcirculation. Interestingly, in an animal model of CPB-induced hypotension, administration of phenylephrine also reduced tissue perfusion to all visceral organs despite correction of MAP from 40 to 65 mmHg [46]. However, correction of hypotension by increasing pump flow improved perfusion of the pancreas, colon and kidneys.

**Vasopressin** (Table 3) depletion is believed to play a causative role in sepsis-induced vasodilatory shock [47]. In a mixed population with catecholamine-resistant vasodilatory shock, additional low-dose arginine vasopressin (AVP) was associated with an increment of pCO2-g-a over time, but lower than NE alone [48]. Direct in vivo observation of the sublingual microcirculation with OPS imaging in a patient with non-septic distributive shock revealed marked microcirculatory alterations, which did not change after AVP injection [49]. In a randomized controlled open-label trial in patients with septic shock, additional use of AVP disclosed no changes in pCO2-g-a and a higher creatinine clearance in comparison to NE alone [50]. This was in contrast to other reports of patients with septic shock; both additional continuous infusion of low dose AVP as well as replacement of NE by high dose AVP resulted in a significant increase of pCO2-g-a [51, 52]. Equally conflicting data in animal studies are suggested to be related to volume status and cardiac performance [53]. AVP administration in sepsis has been associated with ischemic skin and tongue lesions in up to 30% [54], although in a recent trial the reported incidence was considerably lower and not different from norepinephrine [55].

In post-CPB cardiac surgery patients with high output failure, replacement of NE by AVP during steady-state MAP at 75 mmHg resulted in augmentation of pCO2-g-a and diminished laser Doppler-measured jejunal mucosal perfusion, but this was in conjunction with a significantly lower cardiac index [56]. In hemodynamically stable post-CPB patients, incremental doses of AVP were associated with a decline in renal blood flow and impairment of the renal oxygen demand/supply relationship, in the absence of changes in MAP [57].

**Terlipressin** (Table 3) not only acts as a long-lasting prodrug for AVP with a high V1a receptor affinity, but is also a fast-acting vasopressor peptide per se that evokes coronary vasoconstriction with reduction in cardiac output [58]. It was originally described as rescue therapy in catecholamine-resistant septic shock [59]. The dilemma of its use in septic shock is outlined in a case report in which a bolus of terlipressin produced significantly higher MAP and urine output while tapering NE doses, but was also associated with a complete shutdown of sublingual microcirculatory perfusion and progressive acidosis [60]. More human data of direct effects of terlipressin on microcirculatory perfusion are not available, but only in comparison to other (combinations of) vasopressors. In septic shock after volume resuscitation, there was no difference between a bolus terlipressin or NE with respect to lactate levels and creatinine clearance [61]. Additional terlipressin to NE in septic shock was accompanied by a rise in bilirubin and aminotransferases [62], but not in pCO2-g-a [63]. As with AVP, ischemic skin lesions are reported frequently [64].

With respect to splanchnic perfusion in relation to terlipressin, it is of note that it has become an established treatment modality in hepatorenal syndrome and gastrointestinal bleeding in liver cirrhosis. This condition is characterized by a marked vasconstrictive state in the liver in combination with systemic vasoplegia. As
opposed to septic shock, its potent ability to reduce splanchnic blood flow has become beneficial under these conditions, either to increase risk of bleeding or to redirect flow to the kidney [65].

Inotropes

**Dobutamine** (predominantly β2-adrenergic, Table 4) has both inotropic and mild vasodilatory effects. In an open-label setting addition of dobutamine in septic patients was associated with improved OPS-measured sublingual microcirculatory perfusion over time, irrespective of changes in systemic hemodynamic variables [66]. Interestingly, topical application of acetylcholine further improved microcirculatory perfusion, challenging the widespread concept of an irreversible endothelium in sepsis-induced hypotension. In a crossover setting, dobutamine attenuated intramucosal acidosis during sepsis, whereas dopamine did not change \( pCO_2 \) under equal effects in systemic hemodynamics [67]. Addition of dobutamine to NE in septic patients revealed a lower \( pCO_2 \), but no change in ICG plasma clearance [68, 69]; this could either be due to a direct effect on the microcirculation or a result of an increase in cardiac output. Whether an increase in general splanchnic perfusion indeed is followed by improved organ function is still a matter of debate [70].

**Dopamine** (β2-adrenergic and dopaminergic, Table 4; in high-dose also α-adrenergic, Table 2) was used in sepsis for many years in low concentrations for its potential beneficial effects on renal function as a result of afferent renal vessel dilatation. However, results of multiple studies have made such an effect of clinical relevance very unlikely [71]. Low-dose dopamine in addition to NE did not alter \( pH \) despite increased overall splanchnic oxygen consumption [72, 73]. High-dose dopamine decreased \( pH \) despite a rise in cardiac output, whereas NE increased \( pH \) at the same MAP [74].

**Dopexamine** (predominantly dopaminergic, also β2-adrenergic, Table 4) initially was introduced as an inotropic agent with specific pro splanchnic perfusion abilities [75]. However, though dopexamine infusion was associated with a partial correction of spectrophotometric gastric \( HBiq2 \), \( pCO_2 \) remained unaltered [76]. This might be compatible with persistence of microcirculatory weak units.

### Table 3: Vasopressin and analogues (summary of effects in human studies on microvascular perfusion, oxygenation and organ function)

<table>
<thead>
<tr>
<th>Study [ref]</th>
<th>Vasopressor</th>
<th>Setting</th>
<th>Technique(s)</th>
<th>Effects</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>[48]</td>
<td>AVP</td>
<td>Vasodilatory shock, ( n = 48 )</td>
<td>Tonometry</td>
<td>Increase in ( pCO_2 ) and bilirubin concentrations over time</td>
<td>Additional to 0.5 mcg/kg/min norepinephrine, MAP ≥70 mmHg; increase in ( pCO_2 ) lower in comparison to norepinephrine alone</td>
</tr>
<tr>
<td>[49]</td>
<td>AVP</td>
<td>Vasodilatory shock, ( n = 1 )</td>
<td>OPS-imaging</td>
<td>No difference in capillary perfusion</td>
<td>Baseline capillary perfusion markedly impaired</td>
</tr>
<tr>
<td>[50]</td>
<td>AVP</td>
<td>Vasodilatory shock, ( n = 23 )</td>
<td>Tonometry</td>
<td>No change in ( pCO_2 ) over time and in comparison to norepinephrine, no change in lactate, increase in creatinine clearance</td>
<td>In comparison to norepinephrine lower SOFA score, despite higher bilirubin</td>
</tr>
<tr>
<td>[51, 52]</td>
<td>AVP</td>
<td>Septic shock, ( n = 11, n = 12 )</td>
<td>Tonometry</td>
<td>Increase in ( pCO_2 )</td>
<td>In addition to norepinephrine or replacement of norepinephrine respectively</td>
</tr>
<tr>
<td>[56]</td>
<td>AVP</td>
<td>Vasodilatory shock, post cardiac surgery, ( n = 8 )</td>
<td>Tonometry, laser-Doppler flowmetry</td>
<td>Decrease in jeuna mucosal blood flow, increase in ( pCO_2 )</td>
<td>No change in MAP, decrease in cardiac output</td>
</tr>
<tr>
<td>[57]</td>
<td>AVP</td>
<td>Vasodilatory shock, post cardiac surgery, ( n = 8 )</td>
<td>Renal vein thermodilution, ( 5^1 )Cr-EDTA</td>
<td>Decrease in renal blood flow, increase in glomerular filtration rate, impairment of renal oxygen supply/demand relationship</td>
<td>No change in MAP</td>
</tr>
<tr>
<td>[60]</td>
<td>Terlipressin</td>
<td>Septic shock, ( n = 1 )</td>
<td>OPS-imaging</td>
<td>Complete shutdown sublingual microcirculatory blood flow, increase in urine output</td>
<td>Rise in MAP from 58 to 105 mmHg</td>
</tr>
<tr>
<td>[62, 63]</td>
<td>Terlipressin</td>
<td>Septic shock, ( n = 17, n = 15 )</td>
<td>Tonometry, laser-Doppler flowmetry</td>
<td>Increase of urine output and creatinine clearance, rise in bilirubin, no significant change in ( pCO_2 )</td>
<td></td>
</tr>
</tbody>
</table>

**AVP** Arginine vasopressin, **OPS** orthogonal polarization spectral, **\( pCO_2 \)** arterial-to-gastric \( pCO_2 \) gradient, **MAP** mean arterial pressure, **EDTA** ethylenediaminetetraacetic acid
in combination with shunting, not sensed by spectrophotometry due to incorporation of arteriolar and venular hemoglobin. The same dilemma was illustrated in another study. Dopexamine, in addition to volume loading and dobutamine, increased ICG-derived splanchnic blood flow, but with a concomitant decrease in pHi [77]. Apart from this heterogeneity in blood flow between regional circulation and microcirculation, experimental data also suggested regional heterogeneity in blood flow between the gastric and colon region during the use of dopexamine in sepsis [78].

In postoperative cardiac surgery no difference in pCO₂ g-a between low-dose dopexamine and placebo could be detected [79]. In another study pH did not rise despite an increment in ICG-measured splanchnic blood flow and cardiac output [80]. Apart from thermogenetic effects of dopexamine, this might also be explained by a distributive effect. To date, in postoperative non-cardiac surgery, the use of dopexamine remains a matter of debate [81, 82].

**Levosimendan** (calcium sensitizer, Table 4) improves cardiac contractility and has slight vasodilatory effects. In patients with sepsis-induced myocardial depression levosimendan in addition to NE plus dobutamine was superior to an incremental dose of dobutamine, with respect to pCO₂ g-a, laser Doppler-measured gastric mucosal perfusion, arterial lactate levels and creatinine clearance, either as a direct effect or as result of improved cardiac output [83]. After abdominal aortic aneurysm surgery pCO₂ g-a was lower in the levosimendan group in comparison to placebo, but despite a higher cardiac output this was not a result of better regional splanchnic perfusion [84]. In an experimental setting attenuation of sepsis-induced cellular hypoxia was observed, but simultaneous SDF-imaging failed to detect changes in microvascular blood flow [85].

**Nitroglycerin** (NTG, Table 5) as well as other organic nitrates undergo intracellular metabolism in order to produce nitric oxide (NO)-mediated vasodilatation [86].

<table>
<thead>
<tr>
<th>Study [ref]</th>
<th>Inotrope</th>
<th>Setting</th>
<th>Technique(s)</th>
<th>Effects</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>[66]</td>
<td>Dobutamine</td>
<td>Septic shock, n = 22</td>
<td>OPS-imaging</td>
<td>Increase in sublingual capillary perfusion over time</td>
<td>Microcirculatory effects irrespective of systemic hemodynamics</td>
</tr>
<tr>
<td>[67–69]</td>
<td>Dobutamine</td>
<td>Septic shock, n = 10, n = 14, n = 12</td>
<td>Tonometry, laser-Doppler flowmetry, ICG clearance</td>
<td>Decrease in pCO₂ g-a, increase in gastric mucosal perfusion, no change in overall hepatosplanchic perfusion</td>
<td>Substantial rise in cardiac output</td>
</tr>
<tr>
<td>[72, 73]</td>
<td>Dopamine (low dose)</td>
<td>Septic shock, n = 11, n = 16</td>
<td>Tonometry, ICG clearance</td>
<td>No change in intestinal pH despite increase in overall splanchnic blood flow</td>
<td>Effect on splanchic blood flow baseline-dependent</td>
</tr>
<tr>
<td>[76]</td>
<td>Dopexamine</td>
<td>Septic shock, n = 15</td>
<td>Tonometry, spectrophotometry</td>
<td>Increase in microvascular hemoglobin saturation, no change in intestinal pH</td>
<td>These combined observations suggest distributive failure</td>
</tr>
<tr>
<td>[77]</td>
<td>Dopexamine</td>
<td>Septic shock, n = 22</td>
<td>Laser-Doppler flowmetry</td>
<td>Increase in gastric mucosal blood flow</td>
<td>In addition to norepinephrine and in comparison to epinephrine</td>
</tr>
<tr>
<td>[79, 80]</td>
<td>Dopexamine</td>
<td>Cardiac surgery, n = 19, n = 14</td>
<td>Tonometry, ICG clearance</td>
<td>Increase in splanchnic blood flow and oxygen consumption, no change in intestinal pH</td>
<td>Significant rise in cardiac output</td>
</tr>
<tr>
<td>[83]</td>
<td>Levosimendan</td>
<td>Septic shock, n = 28</td>
<td>Tonometry, laser-Doppler flowmetry</td>
<td>Decrease in pCO₂ g-a and lactate, increase in gastric mucosal perfusion and urine output</td>
<td>In comparison to dobutamine</td>
</tr>
<tr>
<td>[84]</td>
<td>Levosimendan</td>
<td>Abdominal aneurysm repair, n = 20</td>
<td>Tonometry, ICG clearance</td>
<td>Overall splanchnic blood flow remained unaltered, pCO₂ g-a decreased</td>
<td>In comparison to placebo during aortic clamping, significantly higher cardiac output</td>
</tr>
</tbody>
</table>

OPS Orthogonal polarization spectral, pCO₂ g-a arterial-to-gastric pCO₂ gradient, ICG indocyanine green
Despite the fact that overwhelming NO production is believed to play an important role in sepsis-induced hypotension, it has also been suggested as therapeutic strategy to overcome heterogeneity in microcirculatory blood flow [5, 87]. This dilemma is illustrated by the fact that unselective blocking of NO-synthase indeed was associated with a substantial increase in blood pressure, but also with a higher mortality [88, 89]. The general idea that increasing MAP results in a higher net microcirculatary perfusion pressure is not in line with physiological theory of the microcirculation as a low-pressure vascular compartment [17]. For example, in the brain of healthy volunteers (with intact autoregulation), NE was unable to improve cerebral perfusion pressure despite a lower rise in MAP, whereas glyceryl trinitrate increased estimated cerebral perfusion pressure as a result of a lower zero flow pressure, even at a lower MAP [90].

In an uncontrolled setting, NTG was administered to septic patients who fulfilled static systemic hemodynamic resuscitation endpoints [91]. Despite fulfillment of these endpoints sublingual microcirculatory flow remained severely impaired. After a single bolus of NTG, instant improvement of microcirculatory blood flow was observed, indicating an NO-mediated endothelial ability for vasodilatation, which based on the analogy of topical acetylcholine challenged the general idea of endothelial hyporesponsiveness in sepsis-induced hypotension [66]. However, in a double-blind placebo-controlled setting, after fulfillment of dynamic systemic hemodynamic resuscitation endpoints, sublingual microcirculatory perfusion improved significantly over time, but no effect of NTG in comparison to placebo could be demonstrated [92]. In the uncontrolled setting, baseline values for microvascular blood flow in large vessels was considerably lower in comparison to the controlled setting, suggesting marked differences in systemic volume status.

In cardiogenic shock, impairment of sublingual microcirculation has also been observed and associated with morbidity and mortality [93]. Application of low-dose NTG in addition to standard care for cardiogenic shock was associated with a rise in sublingual perfused capillary density [94]. This was not accompanied by a change in cardiac output or MAP and disappeared after cessation of NTG administration. Interestingly, non-pulsatile systemic circulatory flow support by means of a percutaneous left ventricular assist device after acute myocardial infarction was also reflected in a better sublingual microcirculatory flow score [95].

In the specific setting of gastric tube reconstruction, topical application of NTG improved laser Doppler blood in the fundus without changes in local HbiO2 [96]. However, no effect was observed during intravenous NTG in the same setting, suggesting potential interference with bioactivation [97].

Ketanserin (serotonin 5-HT2 antagonist and weak α1-adrenergic blocker, Table 5) has been used in the treatment of hypertension with a reported beneficial effect on microhemodynamics and hemorheology due vasodilatation and inhibition of platelet aggregation [98]. Administration of ketanserin in hypertensive post-CPB cardiac surgery patients effectively lowered MAP, while sublingual capillary blood flow was preserved [99].
Prostacyclin (cyclic adenosine monophosphate activator, Table 5) was initially propagated as a drug for improvement of oxygen extraction deficits, suspected to cause tissue dysoxia [100]. Since then it has been studied extensively, especially in experimental settings, for its potential beneficial effect on organ perfusion, as it plays a crucial role in the physiological endothelial vasodilatory response to ischemia/reperfusion [101]. Administration of intravenous prostacyclin after conventional resuscitation was associated with higher pH and indicative for outcome [102]. Similar effects on pH were reported during aerosolized prostacyclin despite unaltered ICG clearance in patients with pulmonary hypertension [103].

During CPB, infusion of prostacyclin blunted jejunal vasomotion, resulting in a reduced ability to maintain laser Doppler-measured jejunal perfusion constant under variation of blood pressure (autoregulation) [104]. On the other hand, jejunal perfusion increased significantly in parallel to a drop in MAP, but unfortunately pCO2 g-a was not measured to establish the overall effect on jejunal microcirculatory perfusion.

Conclusions

Regardless of practical limitations, there is a growing perception that shock should be defined at the level of the microcirculation. Previous clinical research was limited to surrogate markers of organ microcirculatory blood perfusion such as tonometry, laser Doppler flowmetry and ICG clearance. Recent developments have brought direct in vivo observation of the microcirculation in critically ill patients within reach, albeit limited to accessible organs.

To date there are no data available in support of the idea that increasing MAP is beneficial from the perspective of microcirculatory perfusion and/or oxygenation, and such an idea is not in line with the physiological theory of the microcirculation as a low-pressure vascular compartment. In septic shock several reports have demonstrated an absence of effect on microcirculatory perfusion above a MAP of 65 mmHg, but a lower threshold has not been established. The role of vasodilators in sepsis as a therapeutic strategy to recruit ‘microcirculatory weak units’ is yet to be elucidated. In cardiogenic shock neither vasoressors nor inotropic agents have been proven beneficial in terms of organ perfusion, but increasing systemic perfusion by means of several assist devices are now associated with better microcirculatory flow.

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Conflict of interest ECB has no conflict of interest. CI holds a patent on SDF imaging, has stock in Microvision Medical and has received educational grants from Hutchinson Technology, Baxter, Novartis and Eli Lilly.

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