Vitamin C is an essential water-soluble vitamin that is involved in many biosynthetic and metabolic processes. In healthy fasting humans, circulating levels of ascorbate, the redox form of vitamin C, are typically in the range of 50–70 \( \mu \text{mol/L} \) (1). Levels <25 \( \mu \text{mol/L} \) reflect marginal deficiency (hypovitaminosis C). Severe (scurry-like) vitamin C deficiency is present when levels fall below 10 \( \mu \text{mol/L} \) (2).

In the critical care setting, vitamin C is particularly notorious for its strong immunomodulating and antioxidant activity. As such, vitamin C is promoted as adjuvant therapy in conditions characterized by excessive oxidative stress or crippled immunity such as ischemia-reperfusion disorders, trauma, and various inflammatory disease processes (3).

Controlled studies on the effect of vitamin C in critically ill patients are scarce. Moreover, study outcomes are difficult to interpret because heterogeneous patient populations are studied, different and mostly low doses of vitamin C are used, oral and intravenous vitamin C regimens are mixed, and other antioxidative agents (e.g., vitamin E and selenium) are often associated (3). Most relevant experience with adjuvant vitamin C has been obtained in severe burn injury. Adding a high-dose vitamin C to standard fluid resuscitation significantly reduced fluid requirements and net fluid balance in a sheep burn model (4). Two small studies in burn patients assessed the effect of high-dose vitamin C (66 mg/kg/hour) given within the first 24 hours after thermal injury. One prospective study randomized patients to receive fluid resuscitation with or without adjuvant vitamin C. Vitamin C treatment reduced resuscitation volume and resulted in better gas exchange and less days on mechanical ventilation (5).

A more recent retrospective study confirmed reduced fluid requirements and also reported an increased urinary output in vitamin C recipients (6). Both studies could not demonstrate a mortality benefit.

Evidence is emerging that parenteral administration of high-dose vitamin C may be a beneficial adjuvant therapy of severe sepsis and septic shock. An excessive inflammatory response indeed enhances metabolic turnover of vitamin C. As a result, patients with severe sepsis often have very low plasma vitamin C levels that sometimes enter the “scurvy” zone (7). In animal models of sepsis, intravenous ascorbate rapidly and persistently improved capillary and microcirculatory blood flow, decreased microvascular permeability, and attenuated inflammation. Vitamin C also restored endothelial barrier function, prevented apoptosis, and exerted antibacterial effects (3,8). Additionally, vitamin C acts as a cofactor to optimize activity of the enzymes dopamine \( \beta \)-hydroxylase and peptidylglycine \( \alpha \)-amidating monooxygenase which synthesise respectively norepinephrine and vasopressin (9,10). This opens perspectives to use vitamin C for diminishing exogenous vasopressor need in clinical septic shock (11).

Despite these robust experimental benefits, few studies have evaluated the effect of vitamin C supplementation in human sepsis. Fowler et al. studied intravenous infusion of 50 or 200 mg/kg/day ascorbic acid every 6 hours for 4 days in patients with severe sepsis. As compared with placebo, patients receiving ascorbic acid exhibited less inflammation and prompt reduction of organ failure (12). Zabet et al. compared infusion of 25 mg/kg ascorbic acid every 6 hours for 72 hours with placebo in a small cohort of patients with septic shock who required norepinephrine treatment. During the study period, mean dose of norepinephrine and duration of norepinephrine infusion were significantly
lower in the ascorbic acid than in the placebo group. Moreover, subjects who received ascorbic acid had lower 28-day mortality (13). In both studies, ascorbic acid infusion was well-tolerated and devoid of adverse effects.

Many important issues must be addressed before vitamin C can be incorporated in sepsis treatment protocols. The most effective dose and the best time for administration remain to be determined. Under normal physiological conditions, 100 to 300 mg vitamin C per day is sufficient to reach adequate plasma concentrations (14). In contrast, up to 3 g daily is needed to restore normal plasma concentrations in critically ill patients (15). High-dose vitamin C infusion resulted in a 20 to 500-fold (!) increase in ascorbic acid plasma concentrations (5,12), yet a pharmacological dose (concentration)/effect has not been identified. Also, it is not known whether septic patients with normal or slightly decreased vitamin C levels may benefit from supraphysiologic supplementation. Hyperoxaluria and formation of calcium-oxalate stones in the kidneys is a potential adverse effect associated with high-dose vitamin C administration (16). However, a brief course of high-dose ascorbate is unlikely to elevate the risk of oxalate stone formation, except in patients with uncontrolled severely impaired renal function. Septic shock patients are increasingly initiated on continuous renal replacement therapy (CRRT). Vitamin C is substantially cleared by dialysis (17) and at least 50 % of ascorbate may be lost during CRRT (18). Current guidelines recommend a high, albeit still within physiological range, vitamin C dose during CRRT (17) which probably by no means guarantees to cover the needs in critically ill patients. Moreover, the daily intravenous dose in subjects with vasopressor-dependent septic shock treated with CRRT should probably even be higher. A temporary increase of up to 12 g per day has been suggested (18). “Stress” doses of steroids are commonly administered in patients with vasopressor-dependent or refractory septic shock. Steroids increase cellular vitamin C uptake (19) whereas vitamin C may restore glucocorticoid receptor function (20). This potentially relevant synergistic effect should be elaborated in future trials. Toxicity of high-dose vitamin C has not been reported in published clinical trials. However, theoretical concerns exist that a high vitamin C intake might cause pro-oxidant effects, excess iron absorption, vitamin B12 deficiency, or allergic reactions. It is wise to avoid large doses of ascorbate in patients with a history of oxalate nephrolithiasis or in documented glucose-6-phosphate dehydrogenase deficiency, paroxysmal nocturnal hemoglobinuria, and hereditary hemochromatosis.

Awaiting the results of ongoing clinical trials investigating vitamin C supplementation in vasopressor-dependent septic shock, following recommendations can be made. A baseline vitamin C concentration must be obtained in all patients. Vitamin C should be given if levels are below 25 μmol/L. In patients not treated with CRRT, 3 to 6 g vitamin C daily should be supplemented as long as vasopressor treatment is required. If CRRT is running, this dose can be temporarily increased to 12 g per day.

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Footnote
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