

REVIEW



Obesity in the critically ill: a narrative review

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Abstract

The World Health Organization defines overweight and obesity as the condition where excess or abnormal fat accumulation increases risks to health. The prevalence of obesity is increasing worldwide and is around 20% in ICU patients. Adipose tissue is highly metabolically active, and especially visceral adipose tissue has a deleterious adipocyte secretory profile resulting in insulin resistance and a chronic low-grade inflammatory and procoagulant state. Obesity is strongly linked with chronic diseases such as type 2 diabetes, hypertension, cardiovascular diseases, dyslipidemia, non-alcoholic fatty liver disease, chronic kidney disease, obstructive sleep apnea and hypoventilation syndrome, mood disorders and physical disabilities. In hospitalized and ICU patients and in patients with chronic illnesses, a J-shaped relationship between BMI and mortality has been demonstrated, with overweight and moderate obesity being protective compared with a normal BMI or more severe obesity (the still debated and incompletely understood “obesity paradox”). Despite this protective effect regarding mortality, in the setting of critical illness morbidity is adversely affected with increased risk of respiratory and cardiovascular complications, requiring adapted management. Obesity is associated with increased risk of AKI and infection, may require adapted drug dosing and nutrition and is associated with diagnostic and logistic challenges. In addition, negative attitudes toward obese patients (the social stigma of obesity) affect both health care workers and patients.

Keywords: Obesity, Critically ill, Mortality paradox, Complications

Introduction

According to the World Health Organization (WHO), overweight and obesity are defined as the condition where excess or abnormal fat accumulation increases risks to health. Depending on the degree, duration and distribution of excess adipose tissue, these health risks are type 2 diabetes, hypertension, cardiovascular diseases, dyslipidemia, non-alcoholic fatty liver disease, chronic kidney disease, obstructive sleep apnea and hypoventilation syndrome, mood disorders and physical disabilities (Fig. 1). Some of these problems such as

obstructive sleep apnea or physical disabilities are a direct consequence of the increased fat mass (obesity per se) but the majority results from the obesity-associated metabolic phenomena.

The amount of body fat is generally estimated with the body mass index (BMI) [weight (kg)/height² (m)], which forms the basis for the WHO classification (Table 1). BMI, however, becomes a poor marker of excess body fat in patients with either increased or low muscle mass (sarcopenic obesity) [1]. More important, patients with similar BMI may have different obesity-related complications depending on the distribution of excess fat (visceral and ectopic versus subcutaneous fat) [2, 3]. Adipose tissue is highly metabolically active, and visceral adipose tissue has a more deleterious adipocyte secretory profile resulting in insulin resistance and a chronic low-grade inflammatory and procoagulant state. Subcutaneous fat in the lower body on the other hand may act as a metabolic sink for excess fat and protect other tissues/organs from

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lipotoxicity [2, 3]. Waist circumference (WC) and waist-hip ratio (WHR) are tools to assess fat distribution and contribute to risk stratification [3] (Table 1).

Since 1975, the prevalence of obesity has tripled worldwide and is still increasing. It is currently one of the biggest health issues that affects all age groups, populations and countries of all income levels. This is also reflected in the intensive care unit (ICU) population where recent studies report a prevalence around 20% [4, 5]. Although the impact of obesity on ICU mortality is debated, it seems to be associated with morbidity [4] and increased resource utilization [6]. In this narrative review, based on exploratory literature searches, invited international experts summarize recent developments in the management of obese ICU patients. It aims to discuss the impact of obesity on different organ systems with the intention to assist ICU physicians in the management of this vulnerable population. A discussion of specific issues related to bariatric surgery is beyond the scope of this article.

The obesity paradox

Large cohort studies in the general population have demonstrated an increased mortality risk in both overweight

Take-home message

Obesity in the critically ill appears to be associated with lower mortality but increases the risk of complications in several organ systems.

and obese individuals [7]. However, more recent data in hospitalized patients or patients with chronic illnesses showed a J-shaped relationship between BMI and mortality, with overweight and moderate obesity being associated with lower mortality compared with a normal BMI or more severe obesity. The phenomenon that obesity increases the risk of obesity-related disease but paradoxically is associated with increased survival in patients with these diagnoses is called “obesity paradox.” It has been observed in chronic diseases such as heart failure [8], coronary artery disease [9] and end-stage kidney disease [10], but also in acute conditions such as pneumonia [11], sepsis [12], acute respiratory distress syndrome (ARDS) [13] or critical illness in general [4, 14].

Whether the obesity survival paradox represents a real protective effect of adipose tissue has been challenged. First, admission of obese patients may be subjected

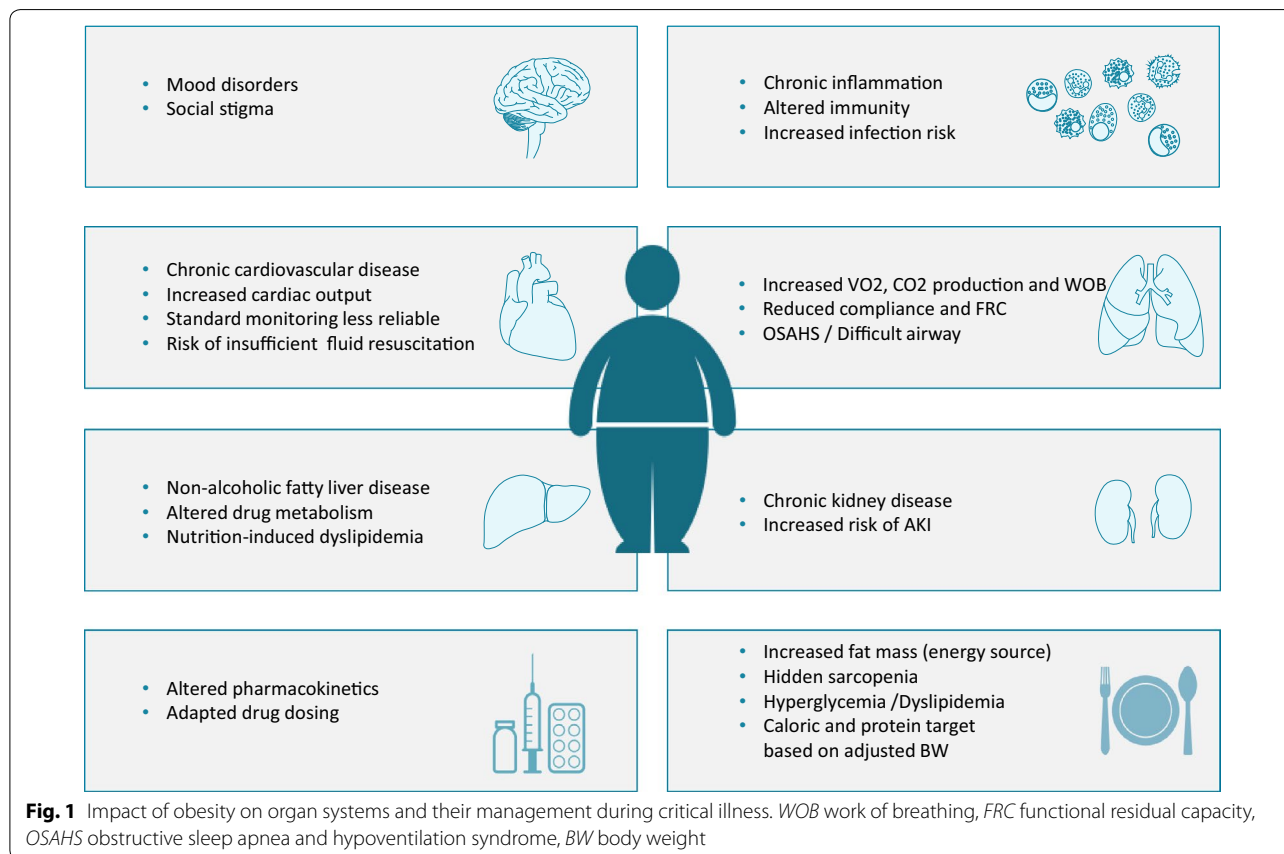


Table 1 WHO classification for obesity based on BMI

Diagnosis	BMI (kg/m ²)	Disease risk WC (cm) males ≤ 94 Females ≤ 80	Disease risk WC (cm) males > 94 Females > 80
Underweight	< 18.5		
Normal weight	18.5–24.9		
Overweight	25–29.9	Increased	High
Obesity class I (moderate obesity)	30–34.9	High	Very high
Obesity class II (severe obesity)	35–39.9	Very high	Very high
Obesity class III (very severe obesity)	≥ 40	Extremely high	Extremely high

Within each BMI category, disease risk may vary depending on fat distribution reflected in waist circumference (WC)

to selection bias because of a lower threshold for ICU admission of otherwise young and healthy obese patients only for surveillance to avoid possible complications. Second, the obesity survival paradox has also been related to therapeutic factors. Obese patients frequently receive a lower weight-based dosage of fluids and vasopressors, potentially attenuating side effects of these therapies [15]. Third, the obesity paradox is typically reported in observational trials and meta-analyses that are subject to confounding and reverse causation [4, 14]. Finally, the use of BMI as a measure of obesity has been criticized. Indeed, in some obese individuals the high BMI may be related to an increased muscle mass or they may have a more advantageous (subcutaneous) fat distribution that is not associated with metabolic comorbidities, referred to as “metabolically healthy obesity.” This condition has been linked to weaker adiposity-related inflammation and low risk of mortality. These phenotypes may confound the results of studies on the obesity paradox.

On the other hand, the hypothesis that “moderate obesity” in and by itself may be protective is increasingly adopted. Several mechanisms have been proposed. Being overweight or obese may be a marker of improved general health status (absence of illness-induced malnutrition) and better exposure to adequate health care. In addition, adipose tissue may also function as a fuel source and provide energy and lipid soluble nutrients during highly catabolic states [16]. A more likely explanation of the obesity survival paradox is that immunomodulatory substances secreted by fat cells (e.g., leptin, interleukin-10 and soluble TNF-alpha receptor) have immunomodulatory effects that might attenuate the inflammatory response and improve survival during severe illness [17, 18]. Most studies show higher leptin levels in ICU survivors [17]. Obese patients with acute lung injury have lower levels of proinflammatory cytokines (IL-6, IL-8) and surfactant protein D than non-obese patients [19]. Moreover, activated macrophages have been shown to infiltrate adipose tissue and switch from the proinflammatory M1 to the antiinflammatory M2 phenotype, with subsequent improvement in immune, antiinflammatory

and scavenging functions in critical illness [16]. Higher lipoprotein and cholesterol levels may neutralize circulating endotoxin and provide precursors for adrenal steroid synthesis [20]. Future studies should focus on identifying the pathophysiologic mechanisms related to the obesity survival paradox and should consider the underlying disease conditions, therapeutic interventions and possible phenotypes.

The respiratory system

One of the main objectives of the critical care management of obese patients is prevention of respiratory complications. Respiratory management of obese ICU patients may differ between patients with healthy lungs and those with ARDS at ICU admission [21].

Differences in respiratory system

Oxygen consumption, production of carbon dioxide, work of breathing and abdominal pressure are increased in obese patients, whereas compliance of the respiratory system and functional residual capacity are decreased [22]. Obesity is a major risk factor of obstructive apnea syndrome. These factors in obese patients may partly explain the higher incidence of difficult airway management, atelectasis and respiratory complications [23]. However, perioperative mortality is not higher in obese compared with non-obese patients [24].

One of the most life-threatening respiratory complications is ARDS. Incidence of ARDS is higher in obese patients, as suggested in a meta-analysis performed in 30,583 patients [25] [pooled OR 1.89 (95% CI 1.45–2.47)]. However, the prognosis of obese ARDS patients appears better compared with their non-obese counterparts (“obesity paradox”) [26].

Airway management

Obesity is a risk factor for difficult intubation and difficult mask ventilation [27]. Elevated Mallampati score, limited mouth opening, reduced cervical mobility, presence of an obstructive apnea syndrome, coma and severe hypoxemia are associated with difficult intubation in

(See figure on next page.)

Fig. 2 Schematic presentation of the effects of regional transpulmonary pressures at different airway pressures on aeration (light blue area), atelectasis (dark blue area) and regional perfusion (red bands) in mechanically ventilated obese patients in supine position. *PEEP* positive end-expiratory pressure, *P_{plat}* plateau pressure of the respiratory system, *P_{pl}* pleural pressure, *PL* transpulmonary pressure (distending pressure of the lung); **a** ventilation at low positive end-expiratory pressure (PEEP) levels is associated with minor regional stress and strain but increased shunt (higher red bands in dark blue areas). Low tidal volume (VT) at zero PEEP results in minimally increased transpulmonary pressure in the ventral while no changes in the dorsal regions at end-inspiration, minimizing overdistension and tidal recruitment. **b** Increased airway pressures promote alveolar recruitment, but increased regional stress and strain in the most dependent lung regions, with possible hemodynamic effects; high tidal volume at zero PEEP results in increased transpulmonary pressure in the ventral and dorsal regions at end-inspiration, increasing the ventilation/perfusion ratio (light blue and narrow red bands) with overdistension and tidal recruitment. On the other hand, increased shunt at end-expiration. **c** Ventilation at moderate PEEP levels optimizes the regional transpulmonary pressures as well as ventilation-perfusion. Low VT with moderate PEEP levels increases lung recruitment (light blue area) while still avoiding dynamic overdistension maximizing improvement in shunt (light blue and red bands). **d** Ventilation at excessively high PEEP levels increases regional transpulmonary pressures and stress and strain with vascular compression (light blue area and narrow red bands) resulting in negative hemodynamic effects and low tidal volume with higher PEEP levels, while ensuring maximal lung recruitment, causes ventral overdistension at end-inspiration and end-expiration

obese patients [27]. To limit desaturation during the intubation procedure, preoxygenation must be optimized. A preoxygenation of 5 min with noninvasive ventilation (NIV) in a sitting position, associating pressure support and positive end-expiratory pressure (PEEP) permits reaching an exhaled fraction in oxygen >90% more quickly than standard bag valve mask ventilation in obese patients [28]. The OPTINIV preoxygenation technique [associating a high-flow nasal cannula (HFNC) with NIV] was more effective at reducing oxygen desaturation compared with the reference method using NIV alone in a randomized controlled trial including obese and non-obese patients with severe acute respiratory failure [29].

Invasive mechanical ventilation

In non-ARDS

As in non-obese patients, protective ventilation should be applied in obese patients, using low tidal volume [set according to ideal body weight (IBW)], moderate-to-high PEEP and recruitment maneuvers [22]. The respiratory mechanics, alveolar recruitment and gas exchanges are significantly improved by application of $PEEP \geq 10$ cmH₂O (improvement of respiratory compliance and decrease of inspiratory resistance) [30]. However, commonly used PEEP by clinicians (11.6 ± 2.9 cmH₂O) was shown inadequate for minimizing atelectasis and “optimizing” ventilation in obese ICU patients [31]. A recruitment maneuver followed by PEEP titration significantly improved lung volumes, respiratory system elastance and oxygenation. Optimal PEEP levels were around 20 cmH₂O. More recently, a PEEP of 12 cmH₂O was found adequate to minimize atelectasis as monitored by electric impedance tomography in obese patients [32]. Another study [33] highlighted the positive impact of recruitment maneuvers on arterial oxygenation and available lung volume in the obese patient. However, higher PEEP levels have

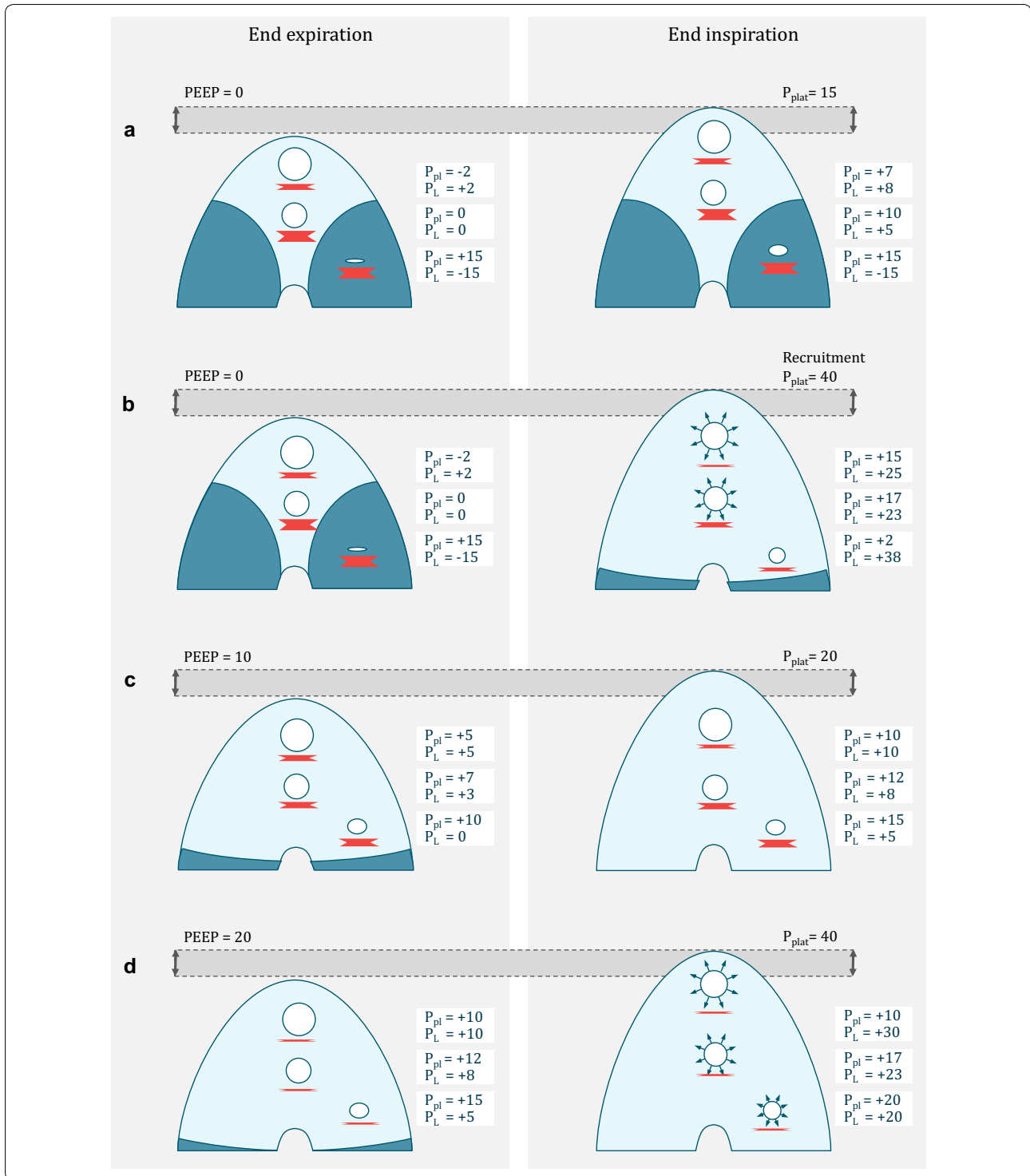
not been clearly shown to minimize clinically relevant complications and might impair hemodynamics, with additional need of vasoactive drugs, as well as fluid overload. Furthermore, recruitment performed by bag squeezing during surgery was associated with increased risk of postoperative pulmonary complications [34]. Thus, recruitment, when needed, should be performed under ventilator control [35].

In ARDS

High PEEP has been reported to be associated with better survival in obese patients with ARDS [36]. Contrary to non-obese patients, driving pressure might not be appropriated to assess the severity and prognosis of obese ARDS patients [37]. Of interest, low-to-negative values of transpulmonary pressure predict lung collapse and intratidal recruitment/derecruitment in obese patients [38]. These results further support the monitoring of transpulmonary pressure using esophageal pressure even if future studies are needed to demonstrate its safety and efficiency in obese patients with ARDS.

Figure 2 shows the effects of different airway pressures at end-inspiration and expiration on regional transpulmonary pressure as well as aeration and atelectasis.

Prone position is a therapy of choice in obese ARDS patients. The safety and efficiency of prone position in ARDS patients with a body mass index > 35 kg/m² were found similar to those of non-obese patients, the ratio of arterial oxygen pressure and the fraction of inspired oxygen (PaO₂/FiO₂) being even significantly more increased after prone position in obese patients compared with non-obese patients [39]. Reverse Trendelenburg position and optimal abdominal fat positioning can help to avoid complications of increased abdominal pressure as bowel ischemia. In case of severe ARDS



after failure or impossibility of using prone positioning and neuromuscular blockers, veno-venous extracorporeal membrane oxygenation (ECMO) can also be safely used in obese ARDS patients [40].

Noninvasive support therapy (oxygen therapy, noninvasive ventilation and CPAP)

NIV and continuous positive airway pressure (CPAP) have proven effective in small observational studies in preventing acute respiratory failure following extubation

Table 2 Ventilatory management in obese non-ARDS and ARDS patients

Intubation	CPAP (5–10 cm H ₂ O) or NIV (PS 10 cmH ₂ O and PEEP 5–10 cmH ₂ O) or HFNC (40–60 l/min) and NIV (FiO ₂ 0.8–1)	
Mechanical ventilation	No ARDS TV (6 ml/kg IBW) RR to achieve CO ₂ 35–45 mmHg or individual ETCO ₂ or pH > 7.25 PEEP 5–10 cmH ₂ O or individualized according to oxygenation, driving or transpulmonary pressure Pplat < 20 cmH ₂ O Driving pressure < 15 cmH ₂ O Recruitment maneuvers only if clinically needed FiO ₂ to keep SpO ₂ 92–95%	ARDS Tidal volume (6 ml/kg IBW) RR to achieve ETCO ₂ 35–45 mmHg or individual ETCO ₂ or pH > 7.25 PEEP 10–15 cmH ₂ O or individualized according to optimal oxygenation, driving or transpulmonary pressure Pplat < 30 cmH ₂ O Driving pressure < 15 cmH ₂ O Recruitment maneuvers (40–50 cmH ₂ O) only if clinically needed and/or before PEEP setting FiO ₂ to keep SpO ₂ 92–95% Prone position if PaO ₂ /FiO ₂ < 150 mmHg Use NIV cautiously and do not delay intubation
Extubation	CPAP or NIV after extubation, especially in patients with comorbidities and/or OSAHS with or without home ventilation and/or cardiac diseases; in patients with intraoperative oxygen desaturation and/or hemodynamic impairment and/or admission to high-dependency units	

ARDS acute respiratory distress syndrome, CPAP continuous positive airway pressure, NIV non-invasive ventilation, HFNC high-flow oxygen nasal cannula, IBW ideal body weight [for males: $0.9 \times (\text{height in cm} - 100)$, for females: $0.9 \times (\text{height in cm} - 106)$], RR respiratory rate, PaCO₂ arterial pressure of carbon dioxide (CO₂), ETCO₂ end tidal PCO₂, Pplat plateau pressure of the respiratory system, FiO₂ inspired oxygen fraction, SpO₂ peripheral capillary oxygen saturation, OSAHS obstructive sleep apnea and hypoventilation syndrome

of obese patients in the ICU and postoperative setting [21, 22]. A high-flow oxygen nasal cannula (HFNC) was not found superior to standard oxygen to prevent reintubation in 155 obese post-cardiac surgery patients [41]. In case of acute hypercapnic respiratory failure, curative NIV is as efficient in patients presenting an obesity hypoventilation syndrome as in chronic obstructive pulmonary disease patients [42]. In the absence of a large study in obese ARDS patients, NIV should be used with caution in this specific population. Among obese cardiothoracic surgery subjects with or without respiratory failure, the use of continuous HFNC compared with NIV did not result in more treatment failure [43]. Whatever the method of oxygenation chosen (NIV, CPAP, HFNC, standard oxygen), intubation should not be delayed in obese ARDS patients (Table 2).

Future agenda for ventilation

In non-ARDS

For airway management, preoxygenation using HFNC, and the use of video laryngoscopes in obese patients should be evaluated in large randomized controlled studies. The best mechanical ventilation settings to prevent ARDS occurrence must be determined. The benefit of post-extubation preventive NIV ± HFNC compared with standard oxygen therapy to prevent reintubation remains to be assessed in overall obese ICU patients.

In ARDS

The optimal mechanical ventilation settings in ARDS obese patients are still unknown, morbid obesity being often an exclusion criterion in the main ARDS studies. The best indications for using extracorporeal membrane oxygenation (ECMO) and extracorporeal carbon dioxide removal (ECCO₂R) in obese patients need to be investigated.

The cardiovascular system

Obese patients, depending on the degree, distribution and duration of obesity, are at increased risk of developing cardiovascular disease. Fat mass-related cardiovascular pathology includes increased blood volume and cardiac output, with secondary ventricular hypertrophy and diastolic dysfunction and finally ventricular dilation (obesity cardiomyopathy). Atrial fibrillation is a frequent complication of obesity, and pulmonary hypertension (secondary to elevated left atrial pressure, hypoxia from obstructive sleep apnea and hypoventilation syndrome or chronic thromboembolism) should also be suspected. In addition to fat mass-related problems, sick fat (visceral and ectopic fat) can affect the cardiovascular system through direct immune and endocrine effects or indirectly by the associated metabolic syndrome with hypertension (afterload), dyslipidemia and ischemic cardiopathy [2].

In view of the high cardiovascular risks, accurate hemodynamic monitoring is crucial, but may be challenging in obese patients. Oscillometric blood pressure

measurements are less accurate in obesity [44], and invasive blood pressure monitoring should be the standard in hemodynamically unstable patients. Transthoracic echocardiography often suffers from a poor acoustic window hampering accurate image acquisition. Evaluation of hemodynamic instability may require transesophageal echocardiography. Uncalibrated noninvasive cardiac output measurement based on pulse contour analysis has gained popularity in the ICU but appears inaccurate in obese patients [45]. This is not surprising since the transformation of the pressure wave form to a cardiac output relies on an algorithm including the dynamic characteristics of the vessel wall that may be significantly altered in obesity. If close monitoring of cardiac output is considered necessary, right heart catheterization or esophageal Doppler could be used. Limited data are available on the interpretation of hemodynamic parameters in obesity: a small study suggests it should not be different from non-obese patients on the condition that they are indexed to the body surface area.

Fluid resuscitation in the obese should account for both the increased blood volume and the risk of fluid overload and heart failure. Little guidance exists for initial fluid resuscitation in obese patients, and the surviving sepsis campaign does not mention this subgroup. Two studies in trauma [46] and septic shock [47] showed that obese patients received less fluid on a weight basis and had more persistent shock or needed more escalation of hemodynamic support and time to reach stability, potentially pointing to under-resuscitation. Similarly, using ideal body weight to guide fluid resuscitation prolonged metabolic acidosis in obese trauma patients [48]. A retrospective analysis of a large cohort of patients with suspected sepsis suggested that using an adjusted body weight to guide initial resuscitation may result in better outcomes than actual or ideal body weight [49]. There is no doubt that fluid resuscitation should be performed with even more cautiousness compared with non-obese patients. Regarding dosing of vasoactive drugs, a retrospective analysis in obese patients with septic shock showed a lower weight-based but similar absolute norepinephrine requirement [50] suggesting that vasoactive drugs should be titrated to their clinical effect rather than using weight-based dosing.

The kidney

In the past decade obesity was recognized as an important cause and cofactor in the development and progression of chronic kidney disease. This fact has been termed “obesity-related nephropathy” [51]. The association of BMI and renal injury is multifactorial. The increased renal perfusion and glomerular hyperfiltration augments intraglomerular pressure, sodium reabsorption

and metabolic demands leading to glomerulomegaly and focal or segmental sclerosis [52]. Moreover, obesity is associated with insulin resistance, type 2 diabetes and hypertension, all important risk factors for chronic kidney disease.

Obesity also appears to be an independent risk factor for acute kidney injury (AKI) [53]. There is a linear correlation between BMI and the incidence of AKI, with higher BMI being associated with higher incidence [53]. The underlying mechanisms have been incompletely established, but besides obesity-associated comorbidities (especially chronic kidney disease), endocrine effects of adipose tissue may play a role. Additional obesity-related AKI risk factors include increases in central venous pressure and intra-abdominal pressure [54].

Despite the increased risk of AKI in obese patients, once AKI occurs, the previously discussed “obesity paradox,” with lower mortality rates in obese patients, has also been described in AKI [53] and chronic kidney disease [10]. In renal dysfunction, this survival advantage may have some specific explanations, such as higher plasma concentrations of beneficial mediators, an improved hemodynamic stability during renal replacement therapy and potentially fat tissue serving as a “buffer” for uremic toxins [52]. The survival advantage in obese compared with non-obese AKI patients may, however, be offset by the increased incidence of AKI, which by itself is a risk factor for mortality.

Obesity may also represent a problem for the diagnosis and supportive therapy of AKI. Using actual body weight to apply the oliguria criteria may lead to a false-positive diagnosis of AKI. In addition, it is not clear whether dosing continuous renal replacement therapy on a ml/kg/h basis should use actual, adjusted or ideal body weight (formulas shown in Table 3 legend).

Immunity and infection

While it is increasingly recognized that adipose tissue is an active participant in the regulation of physiologic and pathologic processes, including immunity and inflammation, the evidence on how and to what extent obesity influences the immune response and the subsequent clinical outcome remains complex and conflicting. Clearly, there is cross-talk between immune cells and adipocytes, resulting in the (dys)regulation of both innate and adaptive immunity [18] (Fig. 3). Adipose tissue produces several hormones (adipokines) and classical inflammatory mediators resulting in chronic inflammation [18], as illustrated by elevated baseline CRP levels in obese patients [55]. Of interest, and illustrating the complex interplay of covariates present in obesity, it has become clear that this chronic inflammation plays a role in the development of

Table 3 Summary of the ESPEN and ASPEN guidelines for adjustment of nutritional therapy in critically ill obese patients

Suggestions	ESPEN guidelines	ASPEN guidelines
For calculating the energy target if measurement of REE is not possible		
In general	20–25 kcal/kg actual BW/day Below 70% of REE should be given during 'early' acute phase	25–30 kcal/kg actual BW/day
In obese	Same as above, but calculated according to adjusted BW ^a If REE measured, set target to 80–100% of REE after the early acute phase (within days 3–7)	11–14 kcal/kg actual BW/day if BMI 30–50 kg/m ² 22–25 kcal/kg ideal BW ^b /day if BMI > 50 kg/m ² If REE measured, set target to 65–70% of REE
For calculating protein target		
In general	1.3 g/kg actual BW/day	1.2–2.0 g/kg actual BW/day
In obese	Same as above, but calculated with adjusted BW ^a	2.0–2.5 g/kg ideal BW ^b /day
For adjustment of nutritional therapy according to serum markers ^c		
Glucose	Below 10 mmol/l (180 g/l) Consider lowering carbohydrate administration when > 6 U insulin/h is needed for > 24 h	Below 10 mmol/l (180 g/l)
Urea	Consider lowering protein administration if > 30 mmol/l: Probably only justified if protein administration > 1.5 g/kg BW/day	–
Triglycerids	Investigate and consider lowering fat administration if > 5.6 mmol/l	–
Examples for calculating energy and protein targets in obese ^d		
Example 1: male 120 kg, 185 cm ≥ BMI = 35.1 kg/m ² Ideal BW ^b = 77 kg ^b and adjusted BW ^a 86–88 kg		
Energy target	Calculated with adjusted BW ^a 25 kcal × 86–88 kg Target = 2150–2200 kcal/day	Calculated with actual BW 14 kcal × 120 kg Target = 1680 kcal/day
Protein target	Calculated with adjusted BW ^a 1.3 g × 92–96 kg ^a Target = 120–125 g/day	Calculated with ideal BW ^b 2.0–2.5 g × 77 kg ^b Target = 154–193 g/day
Example 2: female 140 kg, 165 cm ≥ BMI = 51.5 kg/m ² Ideal BW ^b : 53 kg and adjusted BW ^a = 70–75 kg		
Energy target	Calculated with adjusted BW ^a 25 kcal × 70–75 kg Target = 1750–1875 kcal/day	Calculated with ideal BW ^b 25 kcal × 53 kg Target = 1325 kcal/day
Protein target	Calculated with adjusted BW ^a 1.3 g × 70–75 kg Target = 91–98 g/day	Calculated with ideal BW ^b 2.0–2.5 g × 53 kg Target = 106–133 g/day

REE resting energy expenditure, BW body weight

^a Adjusted BW = ideal BW + 20–25% of difference between actual and ideal BW (actual BW – ideal BW)

^b Ideal BW: for males: $0.9 \times (\text{height in cm} - 100)$; for females: $0.9 \times (\text{height in cm} - 106)$ suggested in ESPEN guidelines, no specific suggestion for calculating ideal BW in ASPEN guidelines

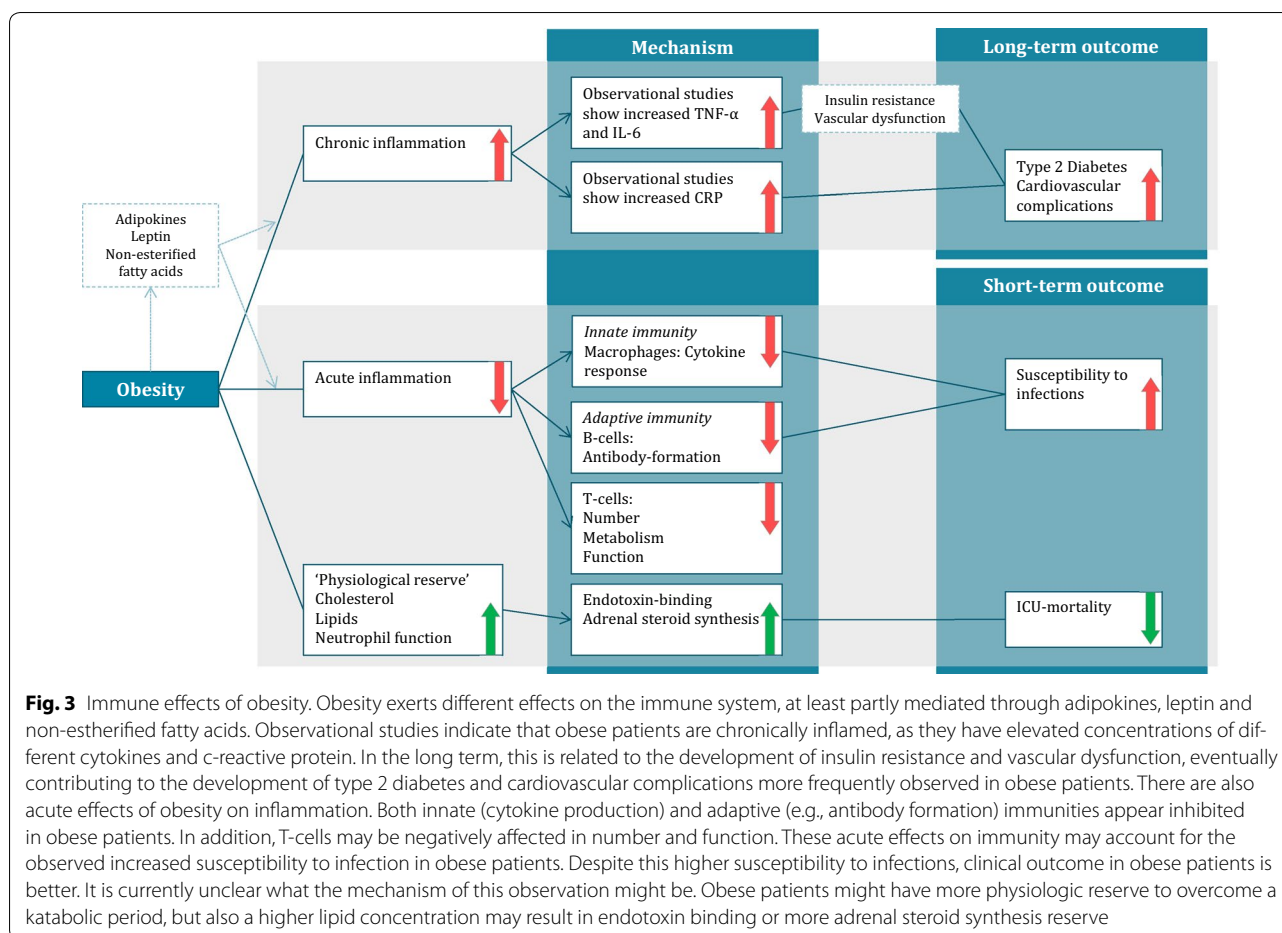
^c No difference in guideline targets regardless of whether applied to normal weight or obese individuals

^d The upper level of suggested energy targets in kcal/BW/day is taken as a basis for calculations

insulin resistance and the cardiovascular complications of obesity [7, 18].

While chronic inflammation is present, there are indications that the innate immune response is impaired in obese patients. The association between severe obesity and both community-acquired and nosocomial infection has indeed repeatedly been described [56], although it is not a universal finding [4]. Also, adaptive

immunity appears affected. Following influenza vaccination, the initial response (IgG levels) may be more pronounced, but 1 year later antibody titers were lower in obese patients compared with the non-obese [57]. In addition, obesity was found to be an independent risk factor for increased morbidity and mortality from pandemic H1N1 infection [58]. The effect of obesity on the adaptive immune response appears also to be mediated through perturbations in T cell numbers, metabolism



and functioning [59]. Nutrient, hormone and adipokine dysregulations in the obese patient may be implicated. Despite the evidence supporting the notion that obesity impairs immunologic responses and increases the susceptibility to become infected, the “obesity paradox” has also been described in patients with pneumonia and sepsis [11]. Overall, it appears that obesity is associated with chronic inflammation and several impairments in immunity, while the clinical outcome following an infection is not negatively influenced.

Other issues

Other obesity-related problems in ICU patients include increased risk of venous thromboembolism (VTE), both deep vein thrombosis and pulmonary embolism (PE) [60], abdominal compartment syndrome and skin problems. The mechanisms underlying the increased VTE risk are multiple. Obesity, especially visceral obesity, results in a proinflammatory, prothrombotic and hypofibrinogenic milieu [7]. Besides physical effects of body fat, limiting venous return and causing stasis, the underlying pathology leading to critical illness (e.g., sepsis), the

associated bed rest and possible subtherapeutic thrombosis prophylaxis may further increase the prothrombotic state. Diagnosis of VTE is difficult and particularly challenging in the obese. Clinical signs such as leg swelling may be obscured, and compression ultrasonography is hindered by the increased soft tissue thickness [60]. Other imaging procedures are often limited by the equipment (see below).

Optimal dosing of prophylactic anticoagulants in obesity is poorly documented [61]. Normal fixed doses of low-molecular-weight heparin (LMWH) may be inadequate. In small studies, inverse correlations have been found between body weight and anti-Xa levels [61]. For enoxaparin, both increased fixed doses (40 or 60 mg twice a day or 60 mg once a day) and weight-based dosing (0.5 mg/kg once or twice a day) have been suggested [61]. Most of these studies used the anti-Xa level (target peak 0.3–0.5 U/ml) as a surrogate outcome. However, monitoring of anti-Xa in clinical practice remains controversial [61]. The evidence for intermittent pneumatic compression is limited [61], and early ambulation remains important.

Immobilization of obese patients, especially the morbidly obese, increases the risk of skin breakdown and decubitus ulcers. Pressure-reducing devices and patient positioning are therefore of utmost importance but challenging.

Nutrition

Despite the clear role of nutrition in development of obesity, there is considerable uncertainty as to optimal nutritional therapy for obese people during critical illness. Obesity defined by BMI (Table 1) may occur with increased, normal or low muscle mass. Low muscle mass or sarcopenia occurs mainly with aging and may be substantial and not immediately obvious in the critically ill patient with a higher BMI [1]. Body composition (relative muscle and fat mass) has a stronger relationship with outcomes than BMI per se [62].

Independent of body constitution, the preferred route to provide nutritional therapy is enteral [63–65]. The estimation of caloric and protein requirements in critically ill obese patient may require an alternative approach to that used for ‘normal’ BMI patients. Furthermore, the underlying metabolic syndrome may require more intensified monitoring of hyperglycemia and hyperlipidemia.

Equations estimating resting energy expenditure (REE) are even more imprecise in obese patients. International guidelines therefore recommend measuring REE with indirect calorimetry in obese patients [64]. However, since indirect calorimetry is not always available, calculations remain a pragmatic estimate of energy expenditure, particularly in the acute phase of illness, to set target energy delivery (Table 2).

The American Society for Parenteral and Enteral Nutrition (ASPEN) guidelines recommend 65–70% of REE be delivered to obese patients and propose using either 11–14 kcal/kg actual body weight/day (for BMI 30–50 kg/m²) or 22–25 kcal/kg ideal body weight/day (for BMI > 50 kg/m²) to calculate this target [63]. The European Society for Clinical Nutrition and Metabolism (ESPEN) guidelines [64] recommend 20–25 kcal/kg adjusted body weight/day with no further adjustments below REE in obese patients after the early acute phase (Table 3).

The optimal amount of protein to deliver to obese critically ill patients is also contentious. ASPEN guidelines suggest a hypocaloric high protein diet with 2.0–2.5 g/kg ideal body weight/day [63], whereas the ESPEN guidelines do not support such high protein supply and recommend 1.3 g/kg of “adjusted body weight”/day (Table 3) [64]. Similar to starvation, in obese ICU patients the percentage of protein oxidation contributing to basal metabolic rate may be reduced with an increased utilization of ketone bodies [66]. Excess energy storage in fat tissue

may therefore attenuate muscle wasting during critical illness [66]. In older obese critically ill patients, a hypocaloric high-protein feeding increased blood urea concentrations [67], suggesting that high-protein feeding may not be appropriate for all patients. ESPEN guidelines suggest assessment of lean body mass and nitrogen balance in obese patients, whereas no specific recommendations for monitoring and management of glucose, urea or triglycerids are provided [64]. A more nuanced understanding of metabolism in the obese critically ill may inform future design of specialized nutritional therapies [68]. Until then, we support using a pragmatic approach as suggested in ESPEN guidelines [64].

Pharmacotherapy

Obesity can affect pharmacokinetics (relationship between drug dose and concentrations in the body) as well as pharmacodynamics (the pharmacologic effect resulting from a drug’s concentration). This is of potential major significance as most drugs elicit a strong concentration-effect relationship and dosing regimens are developed without consideration of the pathophysiologic effects of critical illness or obesity on pharmacokinetics.

Predicting the need for dose adjustment relies on an understanding of the physicochemistry of the drug (hydrophilic or lipophilic) and the effect that obesity itself, or critical illness, can have on altering pharmacokinetics and dosing requirements. Hydrophilic drugs mostly distribute into water-based sites in the body (e.g., interstitial fluid, muscle), whereas lipophilic drugs are more likely to distribute intracellularly and into adipose tissue. Where altered pharmacokinetics lead to an increased volume of distribution (Vd), this may lead to a requirement for greater doses, whereas changes to drug clearance can require a different dosing frequency.

Pharmacokinetic changes associated with obesity

The increased body weight associated with obesity is likely to affect the Vd of all drugs [69]. Hydrophilic drugs will have a small increase in Vd as a result of an increased blood volume and an increased lean muscle mass. Lipophilic drugs are more likely to distribute into the increased volume of adipose tissue, which will increase their Vd. For drugs with weight-based dosing, choosing the most relevant weight metric may be difficult. Almost always a lean body weight descriptor, including adjusted body weight, is likely to be more relevant than ideal body weight or actual body weight for both hydrophilic and lipophilic drugs alike [70]. It is rare that a dosing weight > 100 kg is ever required. It is important to note the importance that comorbidities may be present in patients with long-standing obesity (e.g., peripheral vascular disease,

chronic kidney disease, non-alcoholic fatty liver disease), as these may affect the pharmacokinetics as well.

Pharmacokinetic alterations caused by critical illness

There are many small studies describing the scenarios and drugs where critical illness may affect pharmacokinetics [71]. Principally, for renally cleared and hydrophilic drugs (low Vd), the effects vary from very high drug clearances (augmented renal clearance) to difficult-to-predict effects on drug clearance in the presence of renal replacement therapy and extracorporeal membrane oxygenation. Hypoalbuminemia may affect drug clearance for some drugs with high protein binding (e.g., ceftriaxone, phenytoin) as well as the Vd of hydrophilic drugs. However, the latter will be more affected by high-volume fluid resuscitation.

In summary, the magnitude of any altered pharmacokinetics in critically ill obese patients is generally driven by the presence of critical illness, with obesity itself being a lesser contributor. Because of the multiplicity of factors affecting drug levels, a one-size-fits-all approach is not justified, and maximal use of therapeutic drug monitoring is recommended.

Diagnostic and logistic challenges

Peripheral veins are often less accessible in obese patients. In addition, due to the absence of anatomical landmarks, establishing a central venous access may be particularly challenging. Ultrasound guidance should be the standard of care. Not unexpectedly, a femoral access has been shown to increase the risk of infection in this population [72]. Diagnostic investigations often yield reduced image quality due to the limited penetration of portable X-ray machines and ultrasound waves. Also, transporting the patient to the radiology department may be challenging. CT and MRI are limited by the aperture diameter of the equipment and the table weight limit. In addition, they require a supine position, which may cause respiratory difficulties.

Morbid obesity affects nursing workload and resource use. Indeed, caring for the obese will frequently require additional staff (some even suggest dedicated teams), staff training and specialized durable equipment (bariatric beds, mattresses, transfer devices, lifts, chairs, walkers) to enable safe care for both the patient and staff. Despite specialized equipment, early in-ICU mobilization and post-ICU rehabilitation of obese patients remain challenging.

Psychologic aspects

Health care workers are not immune to the social stigma of obesity, with negative attitudes and prejudices

towards obese patients. People living with obesity are often perceived as weak-willed, unmotivated, noncompliant, sloppy and accountable for their excess weight. In the ICU, nurses provide physically and emotionally demanding care to these patients with complex needs. A qualitative study among ICU nurses showed feelings of repulsion, disgust, anger, frustration, blame, discomfort and fear despite the intention to provide obese patients with the same level of care as normal weight patients [73]. These negative attitudes impact the obese patient both physically and psychologically with feelings of embarrassment, discrimination and distress [74]. Providing nurses with the appropriate equipment and infrastructure to alleviate the physical burden of caring for obese patients, increasing awareness of the impact of their attitudes and specific education, may help to systemically improve the health care experience for obese patients [74].

Conclusion

Obesity is associated with important health risks. It affects one-fifth of ICU patients. Although moderate obesity may paradoxically decrease mortality in ICU patients, increased adipose tissue has an impact on several organ systems, increases morbidity and requires an adapted ICU management, which is summarized in Fig. 1.

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Compliance with ethical standards

Conflicts of interest

Dr. Jaber reports receiving consulting fees from Drager, Fisher & Paykel and Xenios. P. Pickkers reports receiving travel reimbursements and consulting fees from AM-Pharma, Baxter, Adrenomed and EBI. J. Roberts has received

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