Organ Dysfunction in Critically Ill Patients
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Abstract

Intensive care unit (ICU) patients are the most common victims of a clinical condition known as the multiple organ dysfunction syndrome (MODS), multiple organ failure, multiple systems organ failure, or through some of its more prominent manifestations, as the acute respiratory distress syndrome (ARDS) or disseminated intravascular coagulation (DIC) (DIC). The goal of this study is to identify early signs of organ malfunction in critically sick patients and so reduce their risk of death and morbidity. The metabolic and nutritional characteristics of many critical conditions, particularly those originating from trauma or severe sepsis, are similar. In many critically ill patients, whether or not they are infected, there is an array of systemic inflammatory responses that impair immune function and tissue healing, increasing the risk of organ failure (the so-called multiple organ failure syndrome) as well as prolonged hospitalisation and high mortality.

Key words: Organ Dysfunction, Critically Ill Patients.

1. Introduction

Patients with critical illness face a life-threatening, multisystem process that may result in substantial morbidity or fatality. Evidence shows that early signals of deterioration in the patient's physical health are often ignored in patients with serious disease. To provide a successful "chain of reaction" that includes accurate recording and reporting of aberrant results, patient evaluation, and appropriate intervention, all clinical personnel have a vital role to play (1).

Identifying patients at risk of deterioration and significant adverse events may be done via early warning systems. All clinicians should be able to recognise a critically sick patient and begin beginning care of the patient's condition, since good results are dependent on quick recognition, diagnosis, and final treatment (1).

Death rates grow as the number of malfunctioning organs and the length of this failure increase, as seen by severity-scoring systems for illnesses. Because they were designed to standardise criteria of organ malfunction, morbidity (rather than death) could be compared. Consequently, the Multiple Organ Dysfunction Score (MODS), which measures the degree of organ failure, has been demonstrated to represent the development of organ dysfunction when evaluated sequentially and to be highly correlated with the eventual probability of ICU and hospital death (2).

The European Society of Intensive Care Medicine's (ESICM) Sequential Organ Failure Score (SOFA) is likewise well-validated, straightforward, and dependable. Logistic Organ Regression Score (LORS) is a third approach that was built utilising data from a large group of ICU patients. Even if mortality prediction isn't better than other scores, it was designed with that goal in mind (2).

Medications (such as antibiotics for infection source control, circulatory and respiratory support, organ-specific drugs, correction of abnormalities of coagulation, electrolyte, acid-base, metabolism) and specific organ support systems are required for critically-ill patients with multiple organ dysfunction syndrome. A worldwide plan for supporting individual organs and managing the combined effects of multiple organ cross-talk should include all of these therapies (3).

In addition to oxygenation and ventilation, multi-organ support treatment comprises veno-venous and invasive mechanical ventilation. Mechanical circulatory support (intra-aortic balloon pump, venous and arterial ECMO, percutaneous and surgical ventricular assist devices [VADS], and total artificial heart), renal replacement therapy (RRT), and extracorporeal liver support (ECLS) are all examples of ECMO, ECCOR, and ECCo2R, respectively ( molecular and sorbet recirculating system, plasmapheresis and sorbet therapies). There is a lack of knowledge regarding how these approaches interact with native organs and other artificial organ support systems in the intensive care unit, where they are presently employed (3).

The goal of this study is to identify early signs of organ malfunction in critically sick patients and so reduce their mortality and morbidity.

1. Critical Illness

An illness, disease, or corrective measure such as Cancer, Kidney Failure, Coronary Artery (Bypass) Surgery (or bypass), Heart Attack (Myocardial Infarction), Heart Valve Surgery, Major Organ Transplantation (or major organ donation), Primary Pulmonary Arterial Hypertension (PPAH), Paralysis, and Coma are all examples of "critical illness."

In the context of this policy, "disease/illness" refers to an ailment that first reveals itself during the Policy Period and requires medical attention from a Medical Practitioner (4).

A method for dealing with severely unwell people

Pain, anxiety, psychosis, weakness, and sleep deprivation are only some of the painful symptoms experienced by critically sick patients throughout their hospitalisation. ICU patients are typically kept sedated, immobilised, and socially isolated to manage these symptoms because of the difficulty of caring for them. Individuals in the medical field have traditionally operated in walled systems, with daily objectives of care based on individual organ failure rather than a more holistic, integrated approach. A growing body of evidence demonstrates that symptoms and the ICU
team's treatment of them might have significant unfavourable prognostic effects (5).

It is becoming more difficult for society to deal with an ever-growing public health concern associated to inefficient symptom management because of the ageing population, rising healthcare expenses, and the increasing number of people who survive serious disease (often referred to as Post Intensive Care Syndrome, PICS). Early mobility and exercise; family involvement and empowerment; assessment of delirium; prevention and treatment of pain; as well as the use of analgesia and sedation during spontaneous waking and breathing trials are all part of the ABCDEF bundle. (6)

Other evidence-based multicomponent ICU therapies vary in key ways from the ABCDEF bundle. Because it applies to all ICU patients on a daily basis, regardless of whether or not they are on mechanical ventilation or have been admitted with a diagnosed illness, As it focuses on the evaluation, prevention and treatment of symptoms rather than disease processes, it is especially useful early in the course of critical illness and may be used in combination with other life-sustaining therapies. Additionally, the ABCDEF bundle's ultimate purpose is to generate patients who are more alert, intellectually engaged, and physically active, all of which aid in their autonomy and their capacity to communicate unmet physical, emotional, and spiritual demands (7).

For the ICU Liberation Project, the outcomes of the quality improvement (QI) project (intended as adherence to the bundle and its components) will be presented in a separate article. The major goal of this study was to investigate the relationship between ABCDEF bundle performance and patient, symptom, and healthcare system outcomes. Specifically, we expected that the ABCDEF bundle would be linked with better clinical outcomes if it was completed and dose-related (i.e., proportionate) (8).

The three primary types of care

A patient's care may be broken down into three basic categories: supportive care, primary critical disease therapy, and avoidance of secondary insults at this point.

Syndrome of Organ Dysfunction (MOD) ICU admission may be caused by multiple organ dysfunction syndrome, which is described as "the development of potentially irreversible physiological derangement that involves two or more organ systems not implicated in the illness which resulted in ICU admission.” Moderate-to-severe encephalomyelitis (MODS) is divided into two categories: early (primary) and late (secondary) (9).

A set of organ-specific criteria for sepsis dysfunction was developed in 2005 at the International Pediatric Sepsis Consensus Conference. Several grading schemes have been tailored specifically for use in paediatrics. Adults used the scoring systems to evaluate their own health and the effectiveness of treatment. Organ dysfunction scores are often employed in adult research, including the LODS, SOFA, and multiple organ dysfunction score (MOD) (MODS). Extrapolating from some of these systems, paediatric scoring systems have been developed (10).

Mortality Rates and Incidence

Trauma, sepsis, or shock-related multiple organ failure syndrome is the most prevalent cause of death and post-ICU morbidity in adult patients. Shock, sepsis, and tissue hypoperfusion are the most prominent risk factors for MODS, however there are numerous more risk factors. MODS affects 11% to 40% of adult ICU patients, while 14% to 56% of paediatric ICU patients suffer from this condition as a result of their hospitalisation. Because of demographic variance, geography (i.e., availability to healthcare and nutrition), and discrepancies in the definition of MODS, such a large range might be expected. There is a larger frequency of MODS in juvenile patients with sepsis than in adults, with a range of 30-73%. Hypoxic–ischemic encephalopathy and hematopoietic stem cell transplant are further risk factors for MODS in severely unwell children (11).

The severity of MODS (Scoring Systems)

Numerous grading systems have been devised to quantify MODS severity and predict its fate based on clinical data. There is no reliable way to predict the long-term fate of MODS, its reaction to inflammation, or its ability to recover, even if these clinical grading systems are used as an initial evaluation of MODS severity in everyday practise. Table 1 lists the most widely used scoring systems, along with a brief description of each one's key features. MODS severity evaluation methods based on molecular mediators have recently been proposed as an alternative. When innate immune cells are activated, matrix metalloproteinase (MMPs) are secreted as a critical mediator of inflammation. A possible biomarker for organ damage severity evaluation might be MMP plasma levels, which were shown to favourably correlate with tissue levels (12).

System classification for grading

The scoring methods used in critically sick patients are not classified in an agreed-upon manner. Both one-time and recurring scoring may be used for different data sets. Among the options are:

- Anatomical scoring. These depend on the anatomical area involved. Anatomical scoring systems are mainly used for trauma patients [e.g. abbreviated injury score (AIS) and injury severity score (ISS)].
- Therapeutic weighted scores. These are based on the assumption that very ill patients require a greater number of interventions and procedures that are more complex than patients who are less ill. Examples include the therapeutic intervention scoring system (TISS).
- Organ-specific scoring. This is similar to therapeutic scoring; the underlying premise is the sicker a patient the more organ systems will be involved, ranging from organ dysfunction to failure [e.g. sepsis-related organ failure assessment (SOFA)].
- Physiological assessment. It is based on the degree of derangement of routinely measured physiological variables [e.g. acute physiology and chronic health
evaluation (APACHE) and simplified acute physiology score (SAPS)].

- Simple scales. It is based on clinical judgement (e.g. survive or die).
- Disease specific [e.g. Ranson's criteria for acute pancreatitis, subarachnoid haemorrhage assessment using the World Federation of Neurosurgeons score, and liver failure assessment using Child-Pugh or model for end-stage liver disease (MELD) scoring]. (13).

**Clinical Course**

The clinical course and evolution of MODS is dependent on a combination of acquired and genetic factors. It is well known that certain genetic polymorphisms predispose individuals to earlier and more severe MODS. As previously mentioned, lungs are usually the first organ involved in both adult and pediatric patients. Specifically in pediatric patients, myocardial involvement follows a close second, with neurologic involvement as the third most common system involved. Pediatric patients tend to have earlier development of MODS and faster multi-organ involvement, as opposed to the sequential involvement described in adults. Children typically develop organ failure in the first or second day after admission and median maximal duration of organ failure varies from 2 to 3 days (14).

Multiple laboratory derangements are often seen in MODS. Hyperglycemia is a common finding due to increased hepatic glucose production and peripheral insulin resistance. Thrombocytopenia and normocytic anemia are also frequently seen. There is also hyperlipidemia, most frequently in the form of hypertriglyceridemia, because of increased lipolysis and fatty acid recycling. Hyperlactinemia may occur because of tissue dysoxia; however, it may also result from a stimulatory effect of epinephrine on muscle lactate production and cytokine-induced increase in lactate production, among other factors. In addition, patients with sepsis and MODS enter a catabolic state, with skeletal muscle breakdown and negative nitrogen balance, resulting in a decrease of lean body mass (15).

**Cellular Biology**

The development of MODS and its result may be linked to changes in intracellular pathways, gene expression regulation, and the interplay between external stimuli and intracellular pathways.

Genes. Lung, liver, and kidney tissue in a mouse model of MODS were examined for differential gene expression (DEGs). There are several different types of cytokines that may activate the Janus kinase signal transducer and activator of transcription, which is triggered by many of these cytokines. 13 In individuals with acute pancreatitis worsened by the development of MODS, STAT levels were found to be changed (16).

These serine/threonine-based protein kinases, known as mitogen-activated protein kinases (MAPK), are involved in a variety of cellular activities, including cell growth and proliferation, as well as apoptosis and stress responses. In the pathophysiology of MODS formation in hemorrhagic shock and ventilation-induced lung damage, MAPK pathways have been linked to MODS specifically. Attenuation of MODS development, reduction of polymorphonucleocyte infiltration of the lungs and gut, reduction of pro-inflammatory cytokine production, and reduction of intracellular adhesion molecule-1 and P-selectin expression were all seen in mice after MAPK pathway suppression. Apoptosis and tissue damage were both decreased as a result of these alterations (17).

Molecular Mechanisms Underlying Dysfunction in Individual Organs

Septic shock has an inflammatory immunological environment, although organs may differ significantly in their ability to fight against infection.

Because of the activation of several transcriptional processes in multiple organs during sepsis, this event happens for the first time, researchers have studied the temporal sequence of sepsis-induced gene expression patterns in several organs and tissues. Depending on the organ, researchers discovered that sepsis-induced gene expression was either exclusive to that organ, detected in several organs, or sometimes divergent in select organs. Sepsis-induced gene activation in the brain proved to be less severe than in the other organs studied, and genes that increased inflammation were not infrequently counterbalanced by genes that decreased inflammation. Both of these findings are important: (1) different organs may have different immune responses to sepsis; and (2) the pathophysiology of organ failure during MODS may not be consistent. Mechanistic ideas that explain specific organ dysfunction during MODS are summarised in the next section (18).

**Examinations in the laboratory**

- **Systems in MODS for Assessing Organ Failure**
  - The primary purpose of organ failure ratings was to aid in the comparison of diverse ICU patients’ conditions. Both APACHE and SAPS, as well as clinical and laboratory markers that may be used to stage the degree of organ failure, are used in scoring systems. As a result, a variety of methods have been developed in which an overall score is calculated depending on the level of dysfunction in specific organs. “MultipleOrgan Dysfunction Score” (MODS), “Sequential Organ Failure Assessment Score” (SOFA), and “Logistic Organ Dysfunction System” are the most often utilised scoring methods (LODS). All three methods create a total score based on data from six different organ systems (respiratory, hematologic, hepatic, cardiovascular, central nervous system, and renal). All 3 systems have somewhat different variables. (19).

- **Acid-base**
  - A multifactorial metabolic acidosis typically occurs in patients with MODS.

  - Although acute renal failure or hyperlactatemia are common causes of an anion gap increase, in many individuals the anion gap increase cannot be fully explained. A significant percentage of the aniongap
increase may be attributed to the presence of "unidentified anions" in critically unwell individuals. These anions may represent Krebs cycle intermediates, according to new research. During critical illness, a rise in unmeasured anions seems to be connected with an increased risk of death. As a result of a reduction in the strong-ion difference, patients with circulatory shock who are resuscitated with significant volumes of normal saline may develop hyperchloremic metabolic acidosis. Twenty-one.

Total Number of Red and White Blood Cells

Multiple factors are involved in the development of normocytic anaemia in patients with MODS. Gastrointestinal bleeding, recurrent phlebotomies, and hematopoietic dysfunction are all factors that contribute to the condition. However, the white blood cell count may be abnormally low. The frequency of thrombocytopenia has been estimated at 50% in critically sick patients, according to certain research. In trauma or sepsis patients, individuals with a thrombocytopenia and signs of DIC have an excellent sensitivity and specificity for MODS. DIC, thrombotic thrombocytopenic purpura, and secondary thrombotic microangiopathy have all been linked to a thrombocytopenia-associated multiple organ failure syndrome (21).

Preliminary liver tests

Shock, sepsis, mechanical ventilation with positive end-expiratory pressure, and major surgery are all significant risks for liver impairment during critical illness. This is true even if other organs are functioning normally.

Standard liver function tests, which are notoriously imprecise, make early detection of hepatic disease challenging. Serum bilirubin and/or transaminases are used in most organ failure score systems to determine the degree of hepatic dysfunction. Cholestasis is the most common cause of significant direct hyperbilirubinemia. Acalculus Cholecystitis and Hemolytic Reaction should also be taken into account as possible causes of hyperbilirubinemia that may be treated (22).

Albunin in the bloodstream

The aetiology of hypoalbuminemia during critical illness is usually due to other factors including cytokine-induced suppression of albumin synthesis (e.g. reprioritization of protein synthesis), albumin catabolism, dilutional hypoalbuminemia, and third-space losses, even though hypoalbuminemia is often interpreted as an indication of protein-calorie malnutrition. Hospital mortality may be predicted by hypoalbuminemia, however it has a limited sensitivity and specificity (23).

Procalcitonin Is a Plasma Marker for Sepsis

A misdiagnosis of an illness may lead to inappropriate antibiotic treatment, which can have a negative impact on the patient's health. It is possible to employ plasma indicators of infection such as procalcitonin (PCT) for this purpose. A lack of specificity limits PCT's therapeutic relevance as a marker of severe systemic inflammation, infection and sepsis. PCT levels may also be increased in many noninfectious disorders, such as inhalational damage, pulmonary aspiration, pancreatitis, trauma, and burns, and postoperatively, as well as in patients who have had surgery. To a significant extent, the lack of specificity is related to the low sensitivity of existing tests (eg, LUMITest). PCT's diagnostic value may be enhanced by newer tests (e.g., Kryptor) that have greater sensitivity (24).

DELIRIUM (critically sick patients): an acute state of cognitive impairment.

Alzheimer's disease is a severe danger to the quality of life of many elderly individuals, even those who are physically well, who succumb to acute sickness; for those whose physical activities were already restricted, cognitive decline is a big additional concern (25).

It is becoming more important for both the general public and healthcare professionals to focus on the preservation of cognitive capacities, the avoidance of functional deterioration, and the quality of life for people who survive severe illness. Only recently has a growing body of research in the fields of ageing and critical care come to agree on the need of examining cognitive outcomes in those who have survived ICU stays. An international panel of specialists convened in Brussels in 2002 for a roundtable conference titled "Surviving Intensive Care" recommended strongly that more research into neurocognitive impairments in patients who had survived intensive care (26).

Sedation and delirium detection in the intensive care unit

In a nutshell: Delirium is a condition in which a person's mental state rapidly changes, with inattention and altered awareness. The Society of Critical Care Medicine’s (SCCM’s) 2002 recommendations for sedation and analgesia propose that all critically sick patients be evaluated for sedation level and delirium concurrently. The degree of arousal and the content of consciousness must be monitored by ICU nurses and doctors using data derived from well-validated, reliable, objective, and quick evaluation methods. A two-step strategy to sedation and delirium may simplify this neurologic monitoring in the ICU. Twenty-seven.

Objective sedation evaluation is the initial stage in assessing a patient's degree of consciousness/sedation in the ICU. In order to minimise oversedation and hasten the patient's exit from mechanical breathing, it is important to employ objective evaluation scales. To assist the multidisciplinary team communicate about patients' objectives and treatments, sedation scales are useful. It has been decades since the Ramsay Scale has been the most extensively utilised diagnostic tool in clinical practise and research. The Richmond Agitation Sedation Scale and the Sedation Agitation Scale have recently been validated in critically sick patients and distinguish sedation degrees via verbal and physical stimuli. These are commonly used in ICUs around the globe (28).

Pathology and origin

The brain's natural inflammatory response to systemic infections and injuries involves the creation of
cytokines, cell infiltration, and tissue destruction. Delirium may be caused by a local inflammatory reaction that alters neural activity patterns. The stimulation of the central nervous system's immune response is accompanied by the synthesis of substantial quantities of peripherally generated tumour necrosis factor-, interleukin-1, and interferon-. It is proposed that the brain may have a role in both the development and resolution of multiple organ dysfunction syndrome, or in both occurrences (29).

It's probable that the processes causing ICU delirium and long-term cognitive damage are related. Delirium is hypothesised to be linked to abnormalities in the synthesis, release, and inactivation of neurotransmitters that modulate the regulation of cognitive function, behaviour, and mood from the neuroscience viewpoint. Dopamine, -aminobutyric acid (GABA), and acetylcholine are three neurotransmitter systems implicated in the pathophysiology of delirium. GABA and acetylcholine, on the other hand, reduce neuronal excitability. Neuronal instability and erratic neurotransmission may occur from a deficiency in one or more of these neurotransmitters. A lack of acetylcholine and an excess of dopamine are two physiologic issues that have been linked to delirium. Serotonin imbalance, endorphin hyperfunction, and enhanced central noradrenergic activity are all hypothesised to play a role in the onset of delirium, as are other neurotransmitter systems (30).

Sepsis is characterised by multiple organ dysfunction. Patients who have a severe infection and sepsis are likely to have both systemic inflammation and extensive tissue damage. There is a severity spectrum ranging from sepsis through septic shock to MODS. Infection, which may lead to sepsis and organ failure, is frequently the first step in the clinical process. The American College of Chest Physicians and the Society of Critical Care Medicine collaborated to produce descriptions of the different phases of this procedure

The inflammatory reaction to the presence of microorganisms or the invasion of normally sterile host tissue by these organisms is diagnostic of infection. Viral infections, on the other hand, might manifest in a manner that is almost identical to illnesses caused by bacteria (31).

Organ dysfunction and injury mechanisms
In sepsis, it is still unclear exactly how cell damage and organ failure are caused. The following four mechanisms have been postulated to explain part of the endothelial and parenchymal cell damage associated with MODS.

Hypoxic hypoxia is a condition in which there is inadequate oxygen in the

The septic circulatory lesion interferes with tissue oxygenation, changes metabolic control of tissue oxygen supply, and leads to organ failure. The septic microcirculatory deficit in sepsis is caused in part by microvascular and endothelial abnormalities. Anxieties to the microcirculation are further exacerbated by erythrocytes' incapacity to travel through it because of

the reactive oxygen species (ROS), lytic enzymes, and vasoactive chemicals (e.g. NO and endothelial growth factors). (32).

The cytotoxic effect of direct contact
TNF-, NO, and endotoxin all have the potential to disrupt mitochondrial electron transport, resulting in an imbalance in energy production. The inability to use oxygen even when it is there is known as cytopathic or histotoxic anoxia (33).

Apoptosis
Disabled cells are often destroyed by apoptosis (programmed cell death). Other tissues (such as the gut epithelium) may experience accelerated apoptosis as a result of the proinflammatory cytokines delaying apoptosis in activated macrophages and neutrophils. As a result, sepsis tissue damage may be attributed to apoptotic derangement. (11)

Immunosuppression
Inflammatory and anti-inflammatory mediators may interact in a way that causes an imbalance. Both an inflammatory response and an immunodeficiency may be present. (34).

Dysfunction of the diaphragm
Eventration, weakness, and diaphragmatic paralysis all fall under the umbrella phrase "diaphragmatic dysfunction."

The hemidiaphragm is permanently elevated as a result of thinning. Paralysis, on the other hand, indicates that the diaphragm has completely lost its ability to create the essential pressure for effective breathing. Depending on the aetiology, this disease may be either unilateral or bilateral, short-term or long-term. A hernia occurs when an organ or tissue from the abdomen protrudes through a hole in the diaphragm. (35).

The patient's physical appearance
Unilateral diaphragmatic dysfunction may be asymptomatic, which is why it is commonly discovered on a chest X-ray when an abnormality is seen in the hemidiaphragm. Obese people and those with underlying heart or pulmonary disease are more likely to have more severe symptoms. Dyspnea on exercise is the most common symptom; however, nocturnal hypoventilation and reflux may also be present.

Decreased respiratory sounds in the lower hemithorax and probable dullness to percussion are the only physical findings that may be used to diagnose this condition. Occasionally, the thoraco-abdominal area moves in a strange way during sleep. These patients, according to some research, tend to sleep with their healthy hemidiaphragm in the lower region of their chest. (36).

Variables that can predict the future.
Based on this method, potential predictor variables were identified and categorised a priori as part of the PIRO system. Both baseline (time-fixed) and daily (time-varying) variables were included in the study, including factors such as age, gender and other predisposing factors (i.e. age, gender, immunodeficiency, cardiovascular disease and respiratory and renal insufficiency), infection characteristics (i.e. time of
acquisition, site of infection, and causative pathogen), and response characteristics (i.e. C-reactive protein and white blood cell count). Compound disease severity markers such as the Simplified Acute Physiology Score (SAPS) or the Acute Physiology and Chronic Health Evaluation (APA-CHE) score were excluded because they were only defined for a (first) 24-hour observation window in the ICU and were therefore deemed less suitable for "real-time" bedside prognostication. (37).

Patients with Covid-19 SARS who are critically ill Coronavirus-2 (SARS-CoV-2) is a new coronavirus that causes coronavirus sickness in 2019. It was initially discovered in Wuhan, China (Covid-19). As of January 19, 2020, there have been more than 400,000 verified cases of Covid-19 globally, including the first recorded cases in the United States. (38)

More than 3700 verified cases of Covid-19 and 175 fatalities have been reported in Washington as of March 27, 2020. There have been 1760 Covid-19 cases and 125 fatalities in King County, which covers Seattle and the surrounding suburbs. Researchers have shown that the spread of SARS-CoV-2 was a consequence of local transmission, which means that the source of infection cannot be traced back to a known exposure. Nursing homes and other long-term care institutions in the Seattle region have also seen evidence of transfer from one individual to another. (39).

China and Italy's first studies indicate a high death rate and an overloaded intensive care unit (ICU). In the United States, reports of Covid-19-infected patients in the ICU are few. Covid-19 infection in critically sick patients needs to be better understood to help guide decisions about critical care capacity and resource allocation. (40)

Intensive Care for Patients with COVID-19 Factors Associated with Needing ICU Care

Both preparing for an influx of new patients and determining how best to treat those who have been infected are critical benefits of understanding the normal clinical signs and illness course of the disease itself. A majority of the critically ill patients we’ve treated were over 60 years old, with comorbid illnesses including diabetes and heart disease accounting for 40% of the patients we’ve treated. Perinatal exposure, however, has been linked to a higher risk of severe disease in children than in adults. Pregnant women who have been infected with influenza A(H1N1)pdm2009 have so far experienced a mild illness, but the small number of cases means that it is difficult to make accurate predictions about the course of the disease. Most patients' conditions worsened over the course of 9 to 10 days before they were admitted to the intensive care unit. 4 Two-thirds of patients in intensive care have fulfilled criteria for acute respiratory distress syndrome (ARDS), making it the most well-documented cause for their need for breathing assistance (41).

Distinguishing It From Similar Illnesses

Reverse transcriptase–polymerase chain reaction and bacterial cultures are used to distinguish COVID-19 from other respiratory viruses, notably influenza, which are prevalent in the environment at the time of the study.

Radiographic alterations such as ground-glass opacities on computed tomography are suggestive but nonspecific. As a public health and clinical priority, timely access to diagnostic testing data is essential for effective patient triage and infection control measures to be implemented (42).

Patients and healthcare workers should be safeguarded.

It is vital to reduce the danger of a nosocomial epidemic spreading to additional patients and healthcare staff by preventing viral transmission. It is critical to keep patients with suspected or confirmed COVID-19 at a distance of at least 2 metres apart, to think about the use of medical masks for those who are experiencing symptoms, and to admit those who are at risk to private rooms. As a hospital administrator, you must ensure that all hospital workers are properly educated in infection control procedures, as well as the usage of appropriate personal protective equipment. Endotracheal intubation and diagnostic bronchoscopy both produce aerosols, thus doctors doing these operations should wear N95 respirators or comparable face masks and eye shields or goggles for their patients' safety. (33)

In critically ill patients, nutrition and fluid balance are critical.

Today’s critical care facilities include nutrition care as a standard part of their services. As a result of the work of numerous professional organisations, including ASPEN, the Canadian Critical Care Group, and ESPEN, recommendations and guidelines have been developed. Specific nutrients such as glutamine and other all-in-one combination components are also a major concern, as is whether or not to use EN as opposed to PN for certain indications. A lack of practical guidance on how to maintain homeostasis during nutrition treatment or clinical problem management in the event of issues is evident in the recommendations that are available. Although certain nutritional advice has been called into doubt, it has been included into this information. Due to the lack of hunger and thirst signals in critical illness, nutrition care in the ICU presents a number of complications. A person's intake is controlled externally, yet nutrients may interact with numerous organ systems in a complicated manner (44).

An evaluation of a person’s diet.

The Malnutrition Universal Screening Tool, a validated screening tool, is recommended by the UK’s National Institute for Health and Care Excellence (NICE) for screening patients upon admission to the hospital and frequent monitoring of adult nutritional intake (MUST).

Must is a five-step technique that detects individuals who are overweight, malnourished, or at risk for nutritional deficiency. Using BMI, weight loss, and an acute illness impact score, MUST calculates an overall risk of malnutrition. A dietician or nutrition team should conduct a formal nutritional risk assessment if there is a high risk (score 2) and insufficient oral intake is anticipated, according to the American Society of Parenteral and Enteral Nutrition (ASPEN) and Society of Critical Care Medicine (SCCM) guidelines in 2016.
Patients are classified as low (score 0-4) or high (scoring ≥5) risk by 3NUTRIC, depending on comorbidities and clinical condition, in order to better tailor nutrition to present conditions and illness status.3 (45).

I.V. (intravenous) administration of food

EN should begin after 48 hours following ICU admission, ideally once hemodynamic stabilisation has been attained, according to ESPEN and ASPEN.

According to the studies we’ve looked at, “early” refers to initiating feed as soon as possible, up to 48 hours.

Full feeding (growing from 10 ml/h to 20 ml/h) or trophic (increasing from 10 to 20 ml/h) It has been determined to be safe, with less gastrointestinal issues, and is suggested for up to six days after admission in the ICU. 3 Allowable Underfeeding or Standard Enteral Feeding in a pet This study, known as PERMIT, randomly assigned 894 critically ill patients to either permissive underfeeding (40-60 percent of their daily calorie needs) or normal feeding (70-100 percent of their daily calorie needs), with equal protein intake goals in both groups (1.2-1.5 g kg⁻¹ day⁻¹). Regardless of dietary risk, there was no difference in 90-day death or other outcomes. 6 1000 mostly medical patients with acute lung injury (ALI) were randomised to receive either trophic or full enteral feeding (EDEN) in the initial phase of the trial. However, there was a reduction in gastrointestinal intolerance with the use of trophic EN for up to six days without improving ventilator-free days, 60-day mortality, or infectious complications. 10 An anasogastric tube should be used to administer EN to the stomach as a first-line treatment. (46) Parental nutrition is provided.

It is still unclear whether the best time is to begin PN therapy in critically sick patients. Patients who are unable to take EN should begin intravenous nutrition (PN) within 3 to 7 days, according to ESPEN. There is no evidence that early PN improves mortality or critical care outcomes. ESPEN advises that before contemplating additional PN, all other EN techniques should be tried first. ICU admission without nutritional support for 14 days is related with higher mortality (21 percent vs 2 percent, P<0.05) and a longer hospital stay (36.3% vs 23.4 days, P<0.05) compared to the use of nutritional support. Patients at risk of malnutrition were part of the Early vs Late Parenteral Nutrition in Critically Ill Adults (EPA NIC) experiment, which examined the effects of starting intravenous nutrition (PN) sooner rather than later. An earlier beginning of mechanical ventilation was related with increased ICU survival, shorter mechanical ventilation duration, and reduced RRT use. EPA NIC studies demonstrated no benefit in patients who could obtain EN through supplementary PN. Infection rates and healthcare expenses were greater in the early PN group. There was no difference in 60-day mortality or ICU infection rates between the early PN trial and a regular EN, PN, or no early feeding regimen for patients with relative contraindications to early EN. (47) PN-related complications

Although the CALORIES and NUTRIREA-2 studies found that PN problems with appropriate CVC care may be less serious than previously thought, patients who received PN had a greater rate of CVC-associated infection (bacterial/fungal) than those who did not. When cleanliness is inadequate and CVCs are inserted in an emergency, the risk of infection increases and the length of CVC usage increases. For example, hyperglycemia, electrolyte abnormalities, Wer-encephalopathy, nickie’s nutritional excess or deficiency, liver dysfunction, and the refeeding syndrome are all possible side effects of pre-diabetes. Routine monitoring of blood glucose, hydration intake, and electrolyte levels is necessary. (48).

In the Critical Care Setting, Fluid Management

Patients who are seriously unwell or having major surgery need life-saving intravenous fluid treatment. General anaesthesia, stress, dehydration, age, and arterial pressure all affect the pace at which injected fluid is distributed and eliminated. For example, under general anaesthesia and surgery, only 5-10% of the infused volume is eliminated within 2 hours, but in an awake condition, 75-90% is excreted within 2 hours. An abrupt haemorrhage or induction of anaesthesia may cause an increase in efficiency (the percentage of infused fluid retained in the circulation) of 100 percent when arterial pressure is reduced by 20-30% or more. Infusions of crystalloids may now be better regulated thanks to kinetic analysis and outcome-oriented research. Compared to what was previously assumed, infusing crystalloids into patients having surgery while under general anaesthesia had a far better short-term impact on plasma volume expansion (49).

A critical care medicine practitioner’s handbook for effective strategic planning.

Strategic planning is seen as a chance for a company to fulfil its objectives. Employee and stakeholder morale, contentment and efficiency may be improved by strategic planning. This might lead to better patient care and lower healthcare expenditures if implemented in a healthcare setting. The practise of critical care medicine is resource-intensive. Intensive one-on-one care, high-tech equipment, and periodic biochemical and radiological imaging all contribute to the high cost of treatment. Costs associated with intensive care units (ICUs) are estimated to be 1% of GDP. As a percentage of overall hospital costs, it’s about 13%. That these expenses will rise in the future is an acceptable assumption. The world’s population is ageing, but we’re becoming better and better at sustaining it. As an example, during a 15-year period, the death rate for patients with severe sepsis and septic shock dropped from 45 percent to around 20 percent. (50).

Find out what the organization’s goals are.

Understanding the organization’s official and informal mandates can help you grasp the requirements, expectations, and limits that may confront your strategic plan. The feeling of being confined may be avoided by defining the requirements. Knowing the rules makes it less likely that you’ll break them. Having the department head in charge of strategic planning is a good idea for this reason alone. In many cases, they have been a part of
the hospital for a long time and are aware of the rules and regulations that govern it (51).

Revisit the company's goal and values

In order to better understand an organization's purpose, a mission statement might be helpful. However, this is an important element in strategic planning and may be laborious. It may serve as a roadmap for the organization's long-term goals. A good starting point for crafting a mission statement is figuring out what the organization's objectives are. Care quality, budget allocation, and public demands are often discussed in critical-care settings. (52)

Critically Ill Patients' Mortality Reduction
Efforts to Decrease Mortality
Ventilation That Isn't Invasive

A positive pressure supply to the airways and lungs without the use of an intratracheal tube or an extraglottic device is known as noninvasive ventilation (NIV). Noninvasive inspiratory positive-pressure ventilation (NIPPV), in which an expiratory positive airway pressure is nearly always present, and continuous positive airway pressure (CPAP) are both included in the term "NIV." National Institutes of Health (NIH) research shows that NIV can help prevent or treat acute respiratory failure (ARF) by preserving or increasing lung volume, reducing work of breathing, and avoiding or reducing complications of intubation. NIV can also be used outside the ICU or in patients who are not suitable for intubation. While NIV might be beneficial in certain cases, it can also be harmful if a patient cannot manage their own secretions or if they need to protect their airway. NIV's popularity has grown steadily over the previous two decades. In acute care settings, several studies have assessed its effectiveness and limitations. (53)

Acute Respiratory Distress Syndrome Mortality and Lung-Protective Ventilation

Acute respiratory distress syndrome (ARDS) is still a major issue in critical care nearly 50 years after its first description. It is relatively common in the ICU population, affecting about 5% of hospitalised mechanically ventilated patients, and its current mortality is greater than 40%, with high long-term morbidity. A number of its treatment aspects are also controversial or not yet clearly defined. First, the risk of ventilator-induced lung injury (VILI) is increased in ARDS patients due to the disruption of lung architecture, which results in poorly compliant and heterogeneously aerated lungs; and, secondly because mechanical ventilation itself may act as a second "hit" that causes ARDS (according to the so-called multiple hit theorem). ARDS pathophysiology and clinical management are therefore linked in both directions to the mechanisms of VILI. This practise is widely accepted as the standard in ARDS patients due to the disruption of lung architecture, which results in poorly compliant and heterogeneously aerated lungs; and, secondly because mechanical ventilation itself may act as a second "hit" that causes ARDS (according to the so-called multiple hit theorem). ARDS pathophysiology and clinical management are therefore linked in both directions to the mechanisms of VILI. This practise is widely accepted as the standard in ARDS patients without ARDS. Low tidal volumes (VT), moderate to high positive end-expiratory pressure (PEEP), and recruitment manoeuvres are all part of the treatment (i.e., a transitory increase in transpulmonary pressure aimed at opening atelectatic alveoli). (55)

The Use of Tranexamic Acid in Patients with Trauma

As a major public health problem, traumatised individuals face significant personal and societal consequences. Hospital trauma fatalities account for a third of all multiorgan failure deaths, and haemorrhage plays a significant role in both of these outcomes After a traumatic or surgically-induced severe vascular damage, the hemostatic system aids in the maintenance of circulation. Both major surgery and trauma elicit comparable hemostatic reactions, and excessive blood loss puts the coagulation system under high stress, potentially leading to pathological stimulation of clot disintegration (fi brinolysis). In severely damaged trauma patients, hyperfi brinolysis has been shown to contribute to an early coagulopathy that is linked with higher mortality. Patients undergoing surgery who have normal or excessive fi brinolytic reactions had less blood loss with antifi brinolytic drugs than those who have an increased risk of postoperative problems. Early injection of tranexamic acid (TXA), an inhibitor of fi brinolysis, was shown to have a good impact on survival in a large multicenter placebo-controlled experiment (CRASH-2). This validated the use of TXA in trauma patients. Strict Glycemic Stability

Acutely unwell and postoperative patients are more likely to develop stress-induced hyperglycemia, with a 50% and 13% prevalence, respectively. Homeostasis, or the capacity of the body to maintain a physio-logic equilibrium, is affected by critical disease. When this equilibrium is upset by external or internal factors, the body enters a condition of "allostasis," which aims to bring all systems, including metabolism, to a new stable state. This reaction is viewed as adaptive in acute critical illness, but maladaptive in prolonged/chronic critical illness. Upon infection, macrophages release tumour necrosis factor (TNF), which crosses the blood-brain barrier and stimulates the hypothalamic-pituitary-adrenal (HPA) axis, resulting in increased cortisol production and hepatic glycogenolysis/gluconeogenesis. Acutely Ill Patients' Use of Hydroxethyl Starch

In many critically sick patients, hypovolemia may lead to a bad prognosis because it reduces cardiac output and organ perfusion. The resuscitation of these individuals relies heavily on fluid treatment. In critically sick patients who are hypovolemic, the colloid hydroxyethyl starch (HES) has been regularly utilised as a resuscitation fluid for decades. It is thought that the big starch molecules in colloid would raise intravascular osmotic pressure, leading to better hemodynamics with less fluid utilisation. This is the justification for using HES instead of crystalloids." Due to safety concerns, such as tissue deposition and renal and hemostatic damage, the early generations of HES with their large molecular weight and substitution ratio were, however, refined. These starches were touted as having an overall favourable impact by the producers because of their decreased molecular weight and substitution ratio. A considerable percentage of the data supporting HES has been withdrawn owing to scientific misconduct, and the evidence that supports it is restricted to lower-quality studies on HES (small sample size, short follow-up
period, and high bias risk). Patients in critical condition may now be informed by data from big randomised clinical trials and meta-analyses on the best option of fluid treatment (58).

4. Conclusion

Metabolic and nutritional characteristics are common to many critical illnesses, especially those that are brought on by trauma or sepsis. In many critically ill patients, whether or not they are infected, there is an array of systemic inflammatory responses that impair immune function and tissue healing, increasing the risk of organ failure (the so-called ‘multiple organ failure syndrome’) as well as prolonged hospitalisation and high mortality.

References


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