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One of the major concerns of the health care community and the public surrounding the SARS-CoV-2 pandemic is the availability and use of ventilators. Unprecedented surges of patients presented to intensive care units across the country, with older adults making up a large proportion of the patient population. This paper illustrates contemporary approaches to critical illness myopathy (CIM), critical illness polyneuropathy (CIP), or a combination of the two, critical illness polyneuromyopathy (CIPNM), particularly in older patients, can have immediate and lasting negative consequences if not detected and treated promptly. Weakness that develops while a patient is in the ICU complicates recovery and prolongs the duration of both mechanical ventilation (up to 13 days for patients displaying CIPNM versus 3 days for those who do not) and the length of hospital stay [1,2]. These conditions often go undiagnosed or are only considered when patients fail to wean from ventilatory support, despite evidence that up to 62% of patients who experience failure to wean from ventilators have some underlying form of neuromuscular weakness [3]. Accurate early diagnosis and treatment of these conditions can help decrease the intensive care patient burden caused by the surge of SARS-CoV-2 patients requiring ventilation.

A 2011–2012 survey indicated that 68.8% of respondents underestimated the incidence of critical illness acquired weakness [4]. Among critically ill patients, muscle wasting occurs early and rapidly within the first ten days of illness [5]. Improving awareness of CIM, CIP, and CIPNM incidence and interventions can facilitate shorter hospitalizations, decreased mortality, and help fewer patients suffer from chronic disability as a result of critical illness, including pain, motor weakness, pulmonary insufficiency, and psychological stress [3,6]. These dysfunctions are contributors to post-ICU complications.
intensive care syndrome (PCIS), which has its own set of complications [7]. While over 50% of patients will recover completely from CIM, CIP, or CIPNM, those suffering muscular weakness may continue to experience symptoms from 4 weeks to two years after ICU discharge [7,8]. The CDC indicates that during March 1–30, 2020, 74.5% of patients hospitalized for SARS-CoV-2 were ≥50 years old, with the highest rate among adults aged ≥65, so this is particularly an issue for older patients [9].

### 2. Incidence

Studies from 2015 and 2017 focused on determining the incidence of critical illness polyneuropathy, critical illness myopathy, and critical illness neuromyopathy as a group indicate that 40% of patients mechanically ventilated for seven or more days and 40.5% of patients who were ventilated for greater than 24 h developed ICU acquired weakness [10,11].

#### 3. Risk factors

Pre-existing conditions in addition to underlying disease etiology increase the risk for developing CIM/CIP/CIPNM. Studies identified age, hyperglycemia greater than three days, delirium, and mechanical ventilation greater than five days to be independent predictors of intensive care unit-acquired weakness [11]. A 2005 study involving sixty-one critically ill patients also indicated CIP was associated with the presence and duration of systemic inflammatory response syndrome and the severity of cranial, respiratory, and cardiovascular organ failures [2].

A 2018 meta-analysis also indicated a statistically significant association between corticosteroid use and ICU-acquired weakness. The study suggests that where possible, corticosteroids should be limited, or the administration time shortened to reduce the risk of ICU-acquired weakness [12].

#### 4. Proposed mechanisms

The pathophysiology of CIM and CIP is complex, multifactorial, and not completely understood. Both conditions are caused by some combination of critical insult to the body with cytokine overproduction leading to microvascular derangement, and metabolic and electrical (channel) alterations [13]. Although CIM and CIP can and do occur simultaneously in patients, their mechanisms are distinct and important to differentiate.

CIM is a combination of cachectic myopathy and acute necrotizing myopathy (ANM). Cachectic myopathy and ANM are characterized by loss of myosin [3,14]. Cachectic myopathy has been shown to trend with disuse and atrophy and is characterized specifically by loss of type 2 muscle fibers [15].

CIM differs from ANM as patients with CIM display normal creatinine kinase levels and CIM is due to loss of myosin along with further breakdown of the actin-myosin contractile bundle [15]. Decreased myosin heavy chain mRNA (as early as day five of ICU admission) and protein expression, along with increased expression of genes associated with atrophy in critically ill patients with myopathy have been demonstrated [16]. Studies have also found that high-dose corticosteroid exposure leads to selective thick filament loss [15,17–19]. Interestingly, animal models have shown both glucocorticoids and some sort of denervation (ex. non-depolarizing neuromuscular blockers) are necessary in the pathogenesis of CIM, further blurring the line between CIM and CIP [19]. This “two-hit” hypothesis is supported by a study which found dexemethasone only (no denervation) usage led to hypertrophy and increased levels of myosin heavy chain on newborn rat hearts [20]. Other stressors, both pharmaceutical and physiologic, have also been postulated to cause muscle wasting by driving an increased catabolic/anabolic ratio [6].

Interleukin-6 (IL-6) and serum amyloid A1 (SAA1) has been shown to be increased in patients with CIM, which fits with the apparent importance of acute phase response in pathogenesis of the disease. IL-6 is a proinflammatory cytokine that aids in the production of acute phase proteins and is also involved in host immunity. Langhans et al. noted statistically significant increased levels of IL-6 and SAA1 in the skeletal muscle of CIM patients as compared to non CIM patients. They also found SAA1 in CIM patients earlier in the hospital stay than in non-CIM patients [21]. This conclusion has been used to find novel treatment for the acute respiratory distress syndrome and long-term ventilator dependence seen in older patients with SARS-CoV-2 [22].

Membrane dysfunction has also been implicated [23–25]. Allen et al. demonstrated normal nerve conduction velocities in critically ill CIM patients, but slowed conduction within individual muscle fibers. This was reproduced in another study on CIM patients which used muscle electron microscopy to show a loss of thick filaments with normal nerve histology. Rich et al. showed, in animal models, that steroid denervated muscles failed to generate action potentials in response to physiological stimulation, bolstering the idea that excessive steroid usage in critically ill patients is a risk factor for CIM. Rich also found that steroid-denervated muscles had reduced and dysfunctional voltage-gated sodium channels, and the decreased concentration of these channels aligned with progressively decreased excitability [24].

The exact mechanism of injury in CIP is also unknown. One theory is that systemic inflammation and overproduction of cytokines, nitric oxide, and oxygen radicals causes hypoxic and anaerobic conditions leading to decreased circulation of local axonal survival factors and subsequent Wallerian degeneration [26].

Another theory is that the chronic inflammation in CIM patients increases vascular permeability and leads to vasogenic edema [27]. Feni et al. biopsied the superficial peroneal nerve of critically ill patients with CIP and found decreased myelin density and axonal degeneration. They also found increased levels of E-selecin in the walls of endoneurial blood vessels, suggesting edema has an important part in the pathogenesis of CIP [28].

One last theory suggests CIP is related to dysfunction of voltage gated sodium channels [29]. This shared mechanism with CIM may underlie why both conditions often occur together.

#### 5. Diagnosis methods

Difficulties in diagnosing CIM and CIP include, but not limited to intubation, sedation, delirium and cooperation. If sensation can be accurately assessed, CIM is distinguished by failure to wean off mechanical ventilation, flaccid limb weakness more common in proximal than distal extremities, rare extraocular muscle weakness, and facial muscle weakness more common than found in CIP. A reliable neurological examination for CIP will demonstrate atrophy and limb muscle weakness, decreased pinprick sensation peripherally, relative cranial nerve preservation, and absent or reduced tendon reflexes [30]. Shared symptoms of CIM and CIP include ventilatory muscle weakness, limb weakness with relative sparing of the extracranial and facial muscles, and reduced tendon reflexes. CIM begins within days, whereas CIP presents after two or more weeks [31].

It can be challenging to distinguish CIM from CIP. One way is by testing for muscle stretch reflexes: they will typically be present in CIM but will be absent in CIP. Also, sensation usually remains intact in CIM. This can be tested in the ICU by response to pinprick sensation if a patient is awake and alert [32]. As mentioned, creatine kinase and myoglobin levels may or may not be normal in...
CIM. It is also important to note that elevated lab values may be
due to other systemic illnesses.

Due to the high potential for altered sensorium in the older
population most susceptible to CIM/CIP, it is not always feasible
to conduct tests that involve patient cooperation or awareness.
As such, skeletal muscle index measurement using CT imaging-
based calculations can be used. At the time of admission to the
ICU, these scores have been shown to be a valid predictor for ICU
acquired weakness in septic patients [33]. A retrospective study
showed that CT within four days of ICU admission showing low
skeletal muscle area was a risk factor for death in critically ill
mechanically ventilated patients [34]. Since aging is known to be
characterized by sarcopenia and loss of skeletal muscle, the poorer
outcomes seen in older patients could be attributed in part to CIM/
CIP complications.

Nerve conduction studies and electromyography are also help-
ful diagnostic tools. CIM nerve conduction studies show reduced
amplitude compound motor action potentials with preserved sen-
sory response. Electromyography may show polyphasic motor unit
potentials with or without fibrillations. CIP nerve conduction stud-
ies will show decreased amplitude (or absence) of sensory nerve
action potentials. Electromyography would show axonal loss with-
out demyelinating features [35].

When nerve conduction studies cannot be obtained or relied on
due to patient cooperation, significant edema or lack of resources,
other methods for diagnosis exist. For example, Rich et al.
described a technique utilizing the fact that muscle is inexcitable
in CIM and remains excitable in CIP. Comparison of direct muscle
stimulation and nerve stimulation with measured amplitudes of
muscle action potentials allow for differentiation [36]. In addition,
the landmark Italian multicenter CRIMYNE-2 study published in
2014 concluded that the peroneal nerve test (PENT) to measure
compound muscle action potential amplitude showed 100% sensi-
tivity and 85% specificity in diagnosing CIM and CIP in critically ill
patients. This technique has gained favor because it is speedy,
readily reproducible and it does not require patient cooperation.
Note, however, that the CRIMYNE-2 study excluded diabetic patients
[37].

Ultrasound has not been demonstrated as a reliable diagnostic
tool, and therefore may continue to serve best in the setting of
research centers [38,39].

Finally, although rarely used due to its invasiveness and vari-
able findings, muscle and nerve biopsy are the only definitive ways
to diagnose CIM and CIP. CIM would show the characteristic atro-
phy of type 2 fibers with loss of myosin thick filaments, whereas
CIP would show axonal degeneration greater in distal as compared
to proximal segments [13].

5.1. Interventions

Modern interventions for CIM and CIP can be divided into pre-
vention and treatment of ICU-acquired weakness (ICUAW) with
the mainstay of prevention via reduction of the immobilized state.
These treatments are subdivided into pulmonary rehabilitation
and musculoskeletal mobilization.

5.2. Pulmonary preservation

Pulmonary rehabilitation is a multifactorial process focused on:
removing retained airway secretions, exercising both primary and
secondary respiratory muscles after being mechanically ventilated,
and increasing measurable pulmonary values.

The most conservative, familiar option is incentive spirometry
[40]. Another option is positioning the patient to increase ventila-
tion, ventilation/perfusion ratio, and oxygenation. Mobilization
and positioning the patient can increase respiratory volume, expi-
ratory flow rate, and residual functional capacity. On mechanically
ventilated patients, the ideal position for patients is seated at
approximately 30 degrees. This position typically results in effi-
cient minute ventilation, increased respiratory rate and inspiratory
flow rate due to the increased displacement of the ribs and allowing
gravity to facilitate increased expansion [40,41]. In patients with
SARS-CoV-2 on ventilation who have failed standard low tidal
volume ventilation, prone ventilation is used because it decreases
ventral alveolar distention and dorsal alveolar collapse [42]. Prone
positioning, especially in early stages, have shown mortality ben-
fits in patients with severe acute respiratory distress syndrome
(ARDS). Prone positioning has not been proven to prevent organ
system dysfunction, reduce length of stay in the ICU, or shorten
the length of need for mechanical ventilation [43].

5.3. Early mobilization

One of the most successful measures to reduce the incidence of
ICUAW is to begin mobilizing the patient as early as possible. A
2019 study showed starting mobilization as early as 5 days into
an ICU stay can lead to patients standing sooner, walking sooner,
having more ventilation free days, reducing the incidence of
ICUAW, and increasing the discharge to home rate significantly
[40,44]. Another study found even passive mechanical loading in
deply sedated and mechanically ventilated patients decreases
the effects of CIM [45].

In bed cycle ergometry for ICU patients can be utilized in those
who are awake and cooperative as well as those who are uncoop-
erative or unconscious (active vs passive settings). This activity
increases overall exercise capacity post discharge [40].

For patients who are unwilling or unable to begin early mobi-
лизation, basic positioning changes can be performed to signifi-
antly reduce nerve compression and increase blood flow to skin
and superficial musculature thus reducing the rate of decondition-
ing. Patients who are immobile may also benefit from passive
range of motion exercises and stretching. The muscle fiber archi-
tecture can be better preserved in this manner than without exer-
cise [40].

5.4. Mechanical ventilation

Mechanically ventilated patients with ARDS secondary to SARS-
CoV-2 are failing to wean from ventilators and as such, relying on
them for long periods of time. One proposed mechanism causing
long term dependence on mechanical ventilators is elevated IL-6
causing cytokine release syndrome and CIM/CIP [46]. This is based
on data from patients with severe SARS-CoV-2 as well as SARS and
MERS which showed increased IL-6 [46–49]. IL-6 inhibitors such as
Tocilizumab are in phase three trials in the United States as poten-
tial adjunct therapy for mechanically ventilated COVID-19
patients, and while anecdotal support exists for its efficacy, further
research is needed to ultimately determine its role in therapy.

6. Outcomes

Long term studies indicate CIM/CIP/CIPNM has worse outcomes
for older patients. A 5-year longitudinal study (average participant
was 61 years old at recruitment) of patients who were in the ICU
for at least 5 days demonstrated a 44% mortality. This study attrib-
uted the particularly high mortality rate to the increased baseline
age of its participants compared to other studies [50], further
strengthening evidence of increased risk for older populations with
CIM/CIP/CIPNM secondary to SARS-CoV-2. Another challenge to
long term outcome that disproportionately impacts older patients
is diaphragm dysfunction. A study of diaphragm dysfunction as an
independent variable revealed the 2-year survival was 64% in those without diaphragm dysfunction (average age 65) and 71% in those without diaphragm dysfunction (average age 47). Those who had ICU-acquired weakness in addition to diaphragm dysfunction had a survival rate of 36% compared to the survival of those with just ICU-acquired weakness which was 79% [51]. A separate prospective study showed diaphragm dysfunction had a negative effect on extubation success: there was 50% failure rate of weaning process and of those that failed, 50% had passed away [52].

7. Discussion

This review shows how and why CIM/CIP/CIPNM resulting from the SARS-CoV-2 pandemic can particularly impact older patients. We wish to emphasize the importance of early consideration of the potential impact of CIM/CIP/CIPNM, and the need for robust efforts to detect these syndromes and implement interventions for both improved individual patient outcomes and efficient utilization of mechanical ventilators in the current pandemic context. There are several potential limitations on the interventions discussed here, including the psychological consequences of critical illness, baseline cognitive function, sedation, and requirements of positioning needed to adequately treat SARS-CoV-2 patients. For example, elderly patients with decreased baseline cognitive function, ranging from mild impairment to frank dementia may have additional difficulty regaining consciousness in an unfamiliar setting. The deleterious psychological consequences inherent to critical illness may be exacerbated in the SARS-CoV-2 patient who recovers surrounded by unfamiliar providers dressed in personal protective equipment obscuring their faces and eyes, interfering with human connections that could aid recovery. Moreover, providers may be redeployed physicians or nurses with little familiarity working in critical care and their inexperience may contribute to difficulty in recognizing the need for, and initiating, proper treatments and rehabilitation in this challenging context. Due to the respiratory pathology of SARS-CoV-2, fully controlled mechanical ventilation and prone positioning necessitating heavy sedation may become necessary. Delirium resulting from sedation would delay further interventions to reduce the incidence of CIM/CIP/CIPNM. These and other complications may erode patient abilities, motivation, and cooperation essential for engaging with pulmonary preservation and early mobilization rehabilitation.

8. Ethical publication statement

We confirm that we have read the Journal’s position on issues involved in ethical publication and affirm that this manuscript is consistent with those guidelines.

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Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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