Short report

Long-term outcome in patients with critical illness myopathy or neuropathy: the Italian multicentre CRIMYNE study

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ABSTRACT

Background: Critical illness myopathy (CIM) and polyneuropathy (CIP), alone or in combination (CIP/CIM), are frequent complications in patients in the intensive care unit (ICU). There is no evidence that differentiating between CIP and CIM has any impact on patient prognosis.

Methods: 1-year prospective cohort study of patients developing CIP, CIM or combined CIP and CIM during ICU stay.

Results: 28 out of 92 (30.4%) patients developed electrophysiological signs of CIP and/or CIM during their ICU stay, which persisted in 18 patients at ICU discharge. At hospital discharge, diagnoses in the 15 survivors were CIM in six cases, CIP in four, combined CIP and CIM in three and undetermined in two uncooperative patients. During the 1-year follow-up of six patients with CIM, one patient died and five recovered completely within 3 (three patients) to 6 (two patients) months. Of three patients with CIP/CIM, one died, one recovered and one with residual CIP remained tetraplegic. Of four patients with CIP, one recovered, two had persisting muscle weakness and one remained tetraparetic.

Conclusion: CIM has a better prognosis than CIP.

Differential diagnosis is important to predict long-term outcome in ICU patients.

Acquired neuromuscular disorders are common in patients in the intensive care unit (ICU); incidence in patients with sepsis, multiple organ failure or prolonged ICU stay is 46% (95% confidence interval 43% to 49%).

Muscle wasting and paralysis are common clinical features, which are ascribed to critical illness polyneuropathy (CIP), critical illness myopathy (CIM) or a combination of CIP and CIM (CIP/CIM).

Differential diagnosis between CIP and CIM is often not possible in patients in the ICU because they cannot cooperate for accurate sensory and (in some cases) motor testing, and for conventional needle electromyography (EMG) evaluation, which can help distinguish between the two entities but only with fully cooperative patients. Recent research has shown that diagnosis of CIM in patients in the ICU can be established by a combination of needle EMG, direct muscle stimulation and plasma creatine kinase. However, there is no evidence that differentiating between CIP and CIM has any impact on patient prognosis.

We conducted a multicentre prospective study in 92 critically ill adult patients admitted to nine Italian ICUs, and followed them with serial clinical and neurophysiological investigations during their ICU stay and for 1 year after acute care hospital discharge. Results on the acute phase have been reported previously. We now report the long-term follow-up of diseased patients.

METHODS

This multicentre prospective cohort study was performed between January 1998 and March 2001 in nine Italian ICUs belonging to the Gruppo Italiano per la Valutazione degli Interventi in Terapia Intensiva (GIvITI). Patients over 15 years of age were included. Written consent was obtained from the patient whenever possible; otherwise, written information was given to their next of kin. Written consent was obtained from all surviving patients as soon as they regained mental competency.

The acronym CRIMYNE (critical illness myopathy and/or neuropathy) identified the current study among the participating centres; this acronym was coined to emphasise that differential diagnosis between CIP and CIM was not a study target during the ICU period.

Twenty-eight patients developed electrophysiological signs of CIP and/or CIM during ICU stay, which persisted in 18 (64.5%) patients at ICU discharge. At hospital discharge, diagnoses in the 15 survivors were CIM in six cases, CIP in four, combined CIP and CIM in three and undetermined in two uncooperative patients.

Follow-up

Three, six and 12 months after hospital discharge, patients underwent a four part evaluation to define their degree of clinical recovery: (i) complete neurological examination, (ii) muscle strength evaluation, (iii) global motor performance and (iv) complete electrophysiological investigations of limbs. Investigators who performed the follow-up visits were blind to the diagnosis at hospital discharge.

Muscle strength was evaluated according to the Medical Research Council (MRC) scale. A test of global motor performance consisted of standing up from a chair, and walking unaided for at least 50 m with no evidence of ataxia. Clinical recovery was complete if patients had an MRC grade of 4/5 in all muscles, and successfully performed the global motor performance test. In all other cases, clinical recovery was judged as incomplete, and patients were defined as having muscle weakness; patients with an MRC score of 0–1 in all four limbs were defined as having tetraplegia or tetraparesis.

Electrophysiological investigations consisted of conventional motor (median and common peroneal nerves) and sensory nerve (median and sural nerves) conduction studies, and needle EMG in the...
RESULTS
Of the 15 patients followed-up, seven were critically ill non-surgical patients (pneumonia, n = 2; pulmonary oedema, pancreatitis, intracerebral haemorrhage, metabolic encephalopathy, post-anoxic encephalopathy, n = 1 each), five had multiple trauma and three had head trauma. Twelve patients were male (80%) and mean age was 44.7 (SD 14.9) years. Mean Simplified Acute Physiology Score II scores in patients with CIM, CIP or CIP/CIM were 41.7 (SD 8.1), 43.5 (7.9) and 46.0 (9.6), respectively. Median Sequential Organ Failure Assessment scores were 9 (range 6–14), 7 (5.0–12.0) and 11.0 (11–12), respectively. Duration of ICU and hospital stay were 24.1 (13.4) days and 53.7 (68.6) days, respectively.

Patients had variable degrees of muscle impairment (table 1). Incomplete recovery and persisting disability with tetraparesis and tetraplegia were common. Muscle weakness was asymmetrical but recovery of muscle strength was asymmetric in two patients who eventually had unilateral foot drop. Muscle atrophy, reduced or absent deep tendon reflexes and distal sensory loss was observed in the lower limbs of all patients with CIP or CIP/CIM. One patient refused the follow-up visit and was classified as lost to follow-up; on telephone interview, the patient (a medical doctor) referred he had regained normal muscle strength and had returned to work.

Severity of muscle weakness was not correlated with the clinical and electrophysiological diagnosis but the rapidity and completeness of recovery were. At 3 months, three of the five survivors with CIM had complete clinical and electrophysiological recovery and two patients recovered within 6 months. Conversely, patients with CIP or CIP/CIM had a slower recovery, or did not recover at all.

DISCUSSION
We found that 15 of 28 (53.6%) patients developing CIP, CIM or CIP/CIM during their ICU stay had persisting neurological and electrophysiological signs of neuromuscular abnormalities on discharge from the acute care hospital. Of these 15 patients, five (33.3%) recovered normal muscle strength and global motor performance within 3 months after hospital discharge whereas 10 (66.7%) experienced prolonged and severely disabling muscle weakness and paralysis. Patients with a definite diagnosis of CIM recovered earlier and better than those with CIP, the majority of whom remained severely disabled 1 year after hospital discharge.

In a systematic review on long-term outcome, 28.1% of patients with CIM or CIP had severe disability; mean duration of follow-up was 3–6 months, but the range was extremely variable (2 days to 8 years). Data were insufficient to judge whether different electrophysiological diagnoses were associated with different outcomes. Therefore, this study is the first, to our knowledge, indicating that differential diagnosis of CIP and CIM based on accurate clinical evaluation and complete electrophysiological examination is important to define the risk of prolonged disability after ICU discharge. Clinical evaluation alone is not helpful in this respect as both CIP and CIM may cause profound muscle weakness and paralysis.

Our results are different from those of Lacomas et al who, in a retrospective series, found similar functional outcomes in patients with CIM and CIP.11 A possible explanation is that patients where followed-up for only 4 months, when CIM is still largely prevalent. Our data, which need to be confirmed in larger series, suggest that the longer the interval between hospital discharge and electrophysiological examination, the higher the probability that CIM has resolved. This might explain the results of Fletcher et al who found electrophysiological findings consistent with previous CIP in a series of 22 critically ill patients studied at a median of 43 months (range 12–57) after ICU discharge.

We conclude that CIM has a better prognosis than CIP. Differential diagnosis is important to predict long-term outcome in patients in the ICU. Therefore, comprehensive clinical and electrophysiological investigations should be performed to precisely define the pathological diagnosis of patients discharged from the ICU.

Ethics approval: Ethics approval was obtained.

REFERENCES
Richard Bright and epilepsy

Before the work of Todd, Hughlings Jackson, and Herpin, Richard Bright in the 1830s was largely responsible for the notion of a cortical basis for epilepsy, which opposed the established view that epilepsy originated in the medulla. Bright described several partial and complex partial seizure patterns. He also located epileptogenic lesions in the grey (cineritious) matter whence they might be transferred to produce disordered and involuntary motions.

When Richard Bright began clinical practice, the prevailing idea was that epilepsy was essentially a rapidly reversible disorder of consciousness, the genesis of which lay in the medulla. After the primitive notions of divine, diabolical and mystical causes had waned, epilepsy was considered an idiopathic disease without discernible cause. The concept of cortical epileptogenesis was largely initiated by Bright (1789–1858), in 1831, and was later developed by Robert Bentley Todd (1809–1858), in 1831, and was later developed by Robert Bentley Todd. When an underlying cause was suspected, attacks were labelled epileptiform (now called symptomatic epilepsy). As a result of minute bedside observations, Bright corrected many erroneous notions of his day and distinguished several clinical patterns of epilepsy, notably some 50 years before Herpin’s (1799–1865) and Hughlings Jackson’s (1834–1911) invaluable papers.

Bright’s 1831 text, Diseases of the brain and nervous system (vol 2), contains 25 elegant coloured plates, and neuropathology detailed for more than 200 patients. It gives a clear account of what we now call absences and temporal or frontotemporal complex partial seizures:

“...it is simply a momentary absence of mind, the eye is fixed as in thought, yet gazing vacantly, no convulsion, no sound, the occupations of the hand ceases, while the mind for a moment is annihilated, the cloud passes off, the intellect returns, and often, unconscious that its operation has been suspended, the patient resumes the occupation in which he was engaged. At other times,

APPENDIX

The CRIMYNE study

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