Clinical and electrophysiological studies reveal a high incidence of nervous system dysfunction in patients treated in major medical and surgical intensive care units (ICU’s). We evaluated autonomic nervous system (ANS) function in 29 ICU patients with various neurological disorders. **Methods:** Testing involved cardiac R-R variation (CRRV) as an index of parasympathetic function and the sympathetic skin response (SSR) for sympathetic assessment. **Results:** Only those 8 patients with sepsis-related neuropathy or encephalopathy had abnormal CRRV, while the SSR was absent in all but 2 patients. **Conclusions:** Our preliminary study revealed a high incidence of autonomic dysfunction in ICU patients with various neurological disorders.

**RÉSUMÉ:** La variation cardiaque R-R et la réponse cutanée sympathique chez des patients de l’unité de soins intensifs. **Contexte et objectif:** Le système nerveux central et le système nerveux périphérique sont souvent atteints chez les patients hospitalisés à l’unité de soins intensifs (USI), surtout chez ceux qui sont sous ventilation assistée prolongée et ceux qui sont atteints de septicémie ou d’un syndrome de réaction inflammatoire généralisée (SIRS). Cependant, le système nerveux autonome (SNA) a été peu étudié chez ces patients. Nous avons étudié la fonction du SNA chez 29 patients de l’USI atteints de différents problèmes neurologiques. **Méthodes:** Nous avons étudié la variation de l’intervalle R-R (VIRR) comme indice de la fonction parasympathique et la réponse cutanée sympathique (RCS) pour la fonction sympathique. **Résultats:** Seuls les 8 patients atteints de neuropathie ou d’encéphalopathie reliée à une septicémie avaient une VIRR anormale et la RCS était présente chez deux patients seulement. **Conclusions:** Cette étude préliminaire a révélé une incidence élevée de dysfonction autonome chez les patients atteints de différents troubles neurologiques qui sont hospitalisés à l’USI.

Since SIRS has widespread central and peripheral nervous system effects, we decided to examine the autonomic nervous system prospectively. The parasympathetic nervous system can be assessed by measuring the cardiac R-R variation (CRRV) and the sympathetic pathways are evaluated by using the QRS complex in the electrocardiogram is reduced in critically ill patients.

**ABSTRACT:** **Background and Purpose:** The central and peripheral nervous systems are often affected in intensive care unit (ICU) patients, especially those with prolonged assisted ventilation and sepsis or systemic inflammatory response syndrome (SIRS). The autonomic nervous system, however, has been under-investigated in such patients. We evaluated autonomic nervous system (ANS) function in 29 ICU patients with various neurological disorders. **Methods:** Testing involved cardiac R-R variation (CRRV) as an index of parasympathetic function and the sympathetic skin response (SSR) for sympathetic assessment. **Results:** Only those 8 patients with sepsis-related neuropathy or encephalopathy had abnormal CRRV, while the SSR was absent in all but 2 patients. **Conclusions:** Our preliminary study revealed a high incidence of autonomic dysfunction in ICU patients with various neurological disorders.
sympathetic skin response (SSR) that mediates cutaneous cholinergic (sudomotor) function via both central and peripheral pathways. The CRRV and SSR have been shown to be abnormal in patients outside the ICU who have a wide variety of disorders affecting these pathways.\textsuperscript{8,10}

**Methods**

Studies were performed on patients in a 30 bed medical-surgical ICU. Neurological and electrophysiological consultations were requested to investigate a variety of possible central and peripheral nervous system (CNS and PNS) disorders according to previously documented techniques.\textsuperscript{12} Limb temperature was monitored routinely. For CNS disorders, somatosensory evoked potentials were utilized. For PNS disorders motor and sensory nerve conduction and the needle electrode myography were used. There were 18 men and 15 women ranging in age from 19 to 83 years of age; 11 patients were over 60 years of age. The predominant neurological diagnoses were septic encephalopathy (2), anoxic- ischemic encephalopathy (1), Parkinson disease (1), drug induced extrapyramidal syndrome (1), traumatic encephalopathy (3), traumatic quadriplegia (2), critical illness polyneuropathy (14), and myopathy (5). At the time of testing patients were not receiving adrenergic blockers or drugs with anticholinergic properties that might affect the CRRV and SSR. Critically ill patients had widespread organ dysfunction, glucose intolerance and other manifestations of SIRS. They had a variety of reasons for ICU admission.

Both the CRRV and the SSR were measured according to the technique of Shahani et al.\textsuperscript{10} and were performed on an

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**Figure:** The normal result in healthy persons is shown in A for the CRRV and C for the SSR (upper trace is for the hand and the lower trace is the foot.) The abnormal response for the CRRV is shown in B (a fixed R-R interval) and for SSR in D (absent response from the hand and foot.) (Calibration A-200 ms and 200 µV/div; B-200 ms and 100 µV/div; C-1.0 sec and 500 µV/div; D- 1.0 sec and 200 µV/div.)
Advantage EMG system (Advantage Medical, London, Ontario, Canada) with special software designed for these procedures. Each patient was on a ventilator and testing was done while the ventilator was operating. The SSR was tested before any other stimulus was given to the patient to avoid habituation. The stimulus was applied to the left median nerve at the wrist randomly with duration of 0.2 milliseconds and 15-50 milliamps intensity. Five sets of stimuli were given at greater than 30 second intervals. Responses were recorded simultaneously with surface electrodes over the contralateral palm and sole of the foot. Sympathetic skin response was performed at filter settings between 0.5 and 100 Hz and gain between 200 and 500 µV/div. Sympathetic skin response was considered abnormal only if it was entirely absent at both the hand and the foot. Cardiac R-R variation was then tested, with filter settings between 0.5 and 100 Hz and gain from 100-500 µV/div, and expressed as a percentage of the average RR interval and was considered abnormal if the variation was less than either 12% or less than 5% for patients over 60 years of age.

Two patients, who had been studied while on ventilators in the ICU and were found to have absent SSR’s, were tested one month after discharge from the ICU while off the ventilator.

RESULTS

Eight of the 29 patients had an abnormal CRRV (Figure). Six of these had CIP (mild in 5 and moderate in 1) and 2 had mild septic encephalopathy.1 The SSR was absent (Figure) in all patients (see list in methods) except for a 73 year old female with anoxic-ischemic encephalopathy, and a 54 year old female who was on the ventilator briefly for a drug-induced extrapyramidal syndrome.

The median nerve sensory action potential (SNAP) recorded at the index finger from wrist stimulation was normal in 21, reduced in 1, absent in 1 and, for several technical reasons, not tested in 6 patients.

Two patients with moderate/severe CIP had absent SSRs while on the ventilator in the ICU and when tested one month later, while off the ventilator.

The typical results of the CRRV and the SSR for healthy persons and patients are shown in the Figure.

DISCUSSION

The study revealed a high incidence of dysfunction in critically ill patients involving both the parasympathetic (measured by the CRRV) and sympathetic (measured by the SSR) pathways, in the central and peripheral nervous systems. It is known that sepsis SIRS causes widespread dysfunction of the brain, peripheral nerve and muscle in the form of septic encephalopathy, critical illness polynuropathy and myopathy.13 It has been shown that cardiac muscle is also involved manifested as reduction in the amplitude of the QRS complex of the electrocardiogram.7 Examination of the sympathetic chain at autopsy in critically ill patients reveals axonal degeneration.12 However, studies have not been previously performed on the parasympathetic connections to the heart and it has not been previously determined whether unmyelinated fibres of the peripheral nervous system are involved. There is dysfunction of the parasympathetic and sympathetic systems in critically ill patients, which may relate to septic encephalopathy and CIP.

The absent SSR may explain the tendency to hypotension in the critically ill patients.8 Dysfunction of sympathetic pathways in the CNS probably explains the absent SSR in patients with anoxic-ischemic encephalopathy, traumatic encephalopathy, traumatic quadriplegia and Parkinson’s disease who did not have SIRS.

It seems unlikely that habituation from the repetitive action of the ventilator stimulating the lungs would cause the absent SSR since two of our patients with moderate to severe critical illness polynuropathy continued to have an absent SSR one month after discharge from the ICU and off the ventilator. The median nerve SNAP was normal in 21 patients, making it unlikely that dysfunction of the afferent limb of the SSR accounted for the absent SSR. Furthermore, none were on adrenergic blockers or drugs with anticholinergic properties that would block or blunt the SSR. This was a preliminary study; prospective studies should include repeat testing during and after the ICU stay in well defined patient groups. Recording of the quantitative sudomotor axon reflex8 would further establish the presence of sympathetic dysfunction and determine if it is proximal or distal to the sympathetic ganglia.

REFERENCES