Management of Delirium in Adult Critically ill Patients: An Overview

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ABSTRACT - Delirium is a common manifestation of acute, reversible, brain dysfunction in critically ill patients. It is associated with increased morbidity and mortality in the intensive care unit (ICU). Detection and prevention of risk factors for delirium is critical. Daily assessment for delirium should be part of the treatment strategies. Although, non-pharmacologic treatment have been successful, often, pharmacologic intervention is necessary. Currently, there are no approved medications indicated for the treatment of ICU delirium. The objective of this review article is to provide a comprehensive overview of non-pharmacologic and pharmacologic options for the treatment of ICU delirium.

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BACKGROUND

Delirium is defined as a disturbance of consciousness characterized by an acute onset of impaired cognitive function (1). Its development is associated with the underlying medical conditions, substance abuse, adverse effects of medications, substance withdrawal, or a combination of these risk factors (2). Agitation, a manifestation of delirium, is defined as frequent movements of the head, arms, or legs, and/or ventilator – patient dysynchrony despite efforts to calm the patient (3). Patients may also experience delirium as a form of mental disturbance that is less obvious and referred to as quiet agitation (4). Delirium are classified as either hypoactive (not agitated), hyperactive (agitated), or mixed (5).

The pathophysiology of delirium in intensive care unit (ICU) is poorly understood. It is hypothesized that there is an imbalance in the synthesis, release and inactivation of neurotransmitters (6). Cholinergic deficiency and excessive dopamine are thought to contribute to delirium. Dopamine increases and acetylcholine decreases result in neuronal excitability (11). Other neurotransmitters such as GABA and serotonin have been implicated in delirium (9, 12).

Although delirium is considered to be common in ICU, there are few studies that have evaluated its incidence, risks, and outcomes (3, 4). Risk factors are categorized into two groups: predisposing risk factors and precipitating factors (4, 13, 14). (Table 1). ICU delirium is associated with increased morbidity, and mortality and increased cost to the healthcare system (15, 16).

In addition to the uncertainty of the incidence of ICU delirium, there is a lack of information about the effects that certain pharmacological treatments have on delirious patients. For example, do certain medications exacerbate/alleviate delirium, increase/decrease its incidence, prolong/reduce delirious states, increase/decrease ICU length of stay, and/or increase/decrease morbidity and mortality?

ASSESSMENT

There are numerous assessment tools available to identify, diagnose and classify the severity of ICU delirium. The two most widely validated assessment tools used are the confusion assessment method-ICU (CAMICU), and intensive care delirium screening checklist (ICDSC) (17, 18). The ICDSC is a screening checklist of eight items based on diagnostic and statistical manual of mental disorders (DSM) criteria and features of delirium including inattention, disorientation, hallucination-
Table 1. Risk factors for ICU delirium

| PREDISPOSING FACTORS | PRECIPITATING FACTORS |
|----------------------|-----------------------|
| Age greater than 65  | Prolonged pain         |
| Respiratory disease  | Psychoactive drugs such as benzodiazepines |
| Cognitive impairment | Sleep deprivation      |
| Depression           | Severity of illness    |
| Alcoholism           | Severe sepsis         |
| Smoking              | Hypoxemia             |

delusion psychosis, psychomotor agitation or retardation, inappropriate speech or mood, sleep/wake cycle disturbances, and symptom fluctuation according to a total score system of 0 to 8 points. A patient with a score of more than 4 points is defined as delirium positive. It is designed to be easy to use by the clinicians at the bedside and requires minimal time for the assessment. There is evidence to suggest that the ICDSC enhances the ability to accurately detect ICU delirium (18).

NON-PHARMACOLOGIC PREVENTION

Non-pharmacologic interventions are targeted to prevent or reverse potential contributors to the development of delirium. Recent studies have shown that delirium may be an independent risk factor for increased mortality in an ICU setting (19). Because delirium is a multi-factorial syndrome, it usually treated with a multi-component protocol which focuses on six delirium factors: 1) cognitive impairment, 2) sleep deprivation, 3) dehydration, 4) vision/hearing impairment, 5) early mobilization 6) psychoactive medications (20). It has been shown that multi-factorial interventions are effective and decreases the duration of delirium (21-23). The limitation for these studies is, they were conducted in non-ICU patients. However, the risk factors addressed in the studies are commonly seen in ICU patients.

Daily awakening trials

Often ICU patients receive analgesics and sedatives for pain control, comfort, and minimize anxiety while receiving mechanical ventilation (5). Numerous medications used in the ICU can have deleterious effects and could increase the risk for delirium. Evidence suggests there is an independent dose related association between lorazepam and progression to delirium (1, 4, 16). At present, there is little evidence to show whether other benzodiazepines have the same effect on the incidence and duration of delirium. Evidence of delirium associated with morphine is unclear (1). However, patients received morphine for hip fracture have been shown as a risk factor for Delirium (24). Daily waking trials involve discontinuing sedation and analgesia to allow patients to awaken and responsive. Once patients reached certain targeted responsiveness and awareness, the sedation and analgesia are to be re-started at 50% of the dose and titrate to effect. This process will ensure the lowest dose of sedation and analgesics used and avoid drug accumulation in patients who may require long ICU stay. The daily interruption of sedation and analgesia provides an opportunity for the medical staff to assess patients’ neurologic status, and pain level (25-27). Studies showed that daily interruption of sedation result in improved psychological functions, shorter duration of mechanism ventilation requirement, decreased length of stay in ICU and reduced need for diagnostic tests without an increase in complications (27-29).

Environmental factors

There are no data on primary prevention of ICU delirium by controlling environmental factors. However, there are studies that focused in non-ICU settings aimed on minimized these risk factors. It has been recognized that the environmental factors such as hospital, contribute to delirium, and making changes to the environment surrounding the patients can prevent delirium (30, 31). Non-pharmacologic strategies include repeated re-orientation of patients to day/night cycle, maintain regular activities for cognitive stimulation, providing and using eyeglasses, hearing aids, and reduce or minimize noise level (32, 33). Because ICU delirium prevention data is not currently available, it may be helpful to extrapolate the evidence from non-ICU delirium until ICU delirium studies are conducted.
Noise Reduction
Several enhancements and improvements can be implemented to promote a healing environment for patients in ICU. In a survey to evaluate experiences of critically ill patients in the ICU, 54% of respondents were concerned with the noise level in the ward (34). It has been shown that ICU with all the alarms and multiple monitoring devices is one of the noisiest units in the hospital (35). National agencies such as the World Health Organization, has set standards for maximum noise levels in the hospital (36). The noise recommendation from the World Health Organization for hospital settings are 35 dB(A) during the day and 30 dB(A) at night. In many institutions attempts are being made to reduce the level of noise by discontinuing the use of unnecessary monitors or equipment, minimize telephone use and conversation around patient bedside, adjust alarm volume to the lowest safe level, and use of ear plugs for patients having difficulties with noise (5, 37).

PHARMACOLOGIC TREATMENT
Currently, no drugs have received approval from Health Canada and US Food Drug Administration for the treatment of ICU delirium. Nevertheless, antipsychotic agents are routinely used based on outcome evidence. These medications may be helpful in treating the symptoms of delirium by exerting a stable balance by antagonizing the neurotransmitters such as dopamine and serotonin (38, 39).

Typical antipsychotic
Haloperidol
Haloperidol is a butyrophenone antipsychotic. It is a D2 receptor antagonist. The exact mechanism of action in ICU delirium is unclear. It is theorized that blockade of D2 receptor results in amelioration of hallucination, delusions and unconstructed thought patterns (40, 41). It is recommended as the drug of choice to treat ICU delirium by the Society of Critical Care Medicine and American Psychiatry Association.(42-44). Intravenous haloperidol, although off-label, is currently used to treat ICU delirium (5). The effective dose for ICU delirium has not been established. Depending on the severity of the delirium the common dose used is 2 – 10 mg intravenously every six hours(43). Serious side effects include extrapyramidal symptoms such as acute dystonic reactions, subactue parkinsonism, and akathisia, dose-dependent QTc prolongation, and neuroleptic malignant syndrome (44). Continuous infusions of haloperidol have been used to treat severe ICU delirium (45, 46). Close cardiac monitoring is recommended as QTc prolongations were reported by the authors. The use of continuous infusions should be limited to patients who required rapid titration of haloperidol to control acute symptoms, and large intermittent doses (47). Recent evidence indicated that haloperidol in high doses may cause a paradoxical effect by prolonging the duration of delirium (5, 48-50).

Chlorpromazine
Chlorpromazine is a phenothiazine antipsychotic. It is a potent D1, and D2 antagonist. Inhibition of these receptors accounts for ameliorates different antipsychotic symptoms. Currently no study has been conducted in ICU patients with chlorpromazine. It is equally effective compared to haloperidol in terminal AIDS patients with delirium but chlorpromazine has a less favourable side effect profile (51). Undesirable side effects caused by chlorpromazine include anticholinergic activities, hypotension, tachycardia, cardiac arrhythmias, and EPS which limits its use. In addition, chlorpromazine has been shown to lower seizure threshold (52). Due to the serious side effects, chlorpromazine is generally not recommended for ICU delirium.

Atypical antipsychotic
Atypical antipsychotics such as olanzapine, risperidone, quetiapine, and ziprasidone, began to gain popularity in the treatment of ICU delirium in early 2000. It is believed to be as effective but have fewer side effects such as extrapyramidal symptoms than haloperidol. Because atypical antipsychotics have a relatively high affinity for 5-HT2A receptor than D2 receptor, it is thought to be associated with less extrapyramidal symptoms (53-55) (Tables 2 – 3). Currently, there is limited evidence to support the use of atypical antipsychotics in ICU delirium.
Table 2. Adverse effects of receptor blockade

| RECEPTOR | ADVERSE EFFECTS |
|----------|-----------------|
| D₂       | Extrapyramidal symptoms |
| α₁       | Hypotension |
| H₁       | Sedation and weight gain |
| 5-HT₂    | Increased appetite and weight gain |
| M₁       | Cognitive deficits, dry mouth, constipation, increased heart rate, urinary retention, blurred vision |

D, dopamine; α, alpha, H, histamine; 5-HT, serotonin; M, muscarinic; adopted from 11, 56, 57.

Table 3. Relative receptor affinity for atypical antipsychotic agents

| RELATIVE RECEPTOR AFFINITY | Olanzapine | Quetiapine | Risperidone | Ziprasidone |
|---------------------------|------------|------------|-------------|-------------|
| Dopamine (2)              | +++        | ++         | +++         | +++         |
| Serotonin                 | +++        | +          | +++         | +++         |
| α₁                        | ++         | +++        | +++         | ++          |
| Histamine                 | +++        | +++        | +           | +           |
| Muscarinic                | +          | -          | -           | -           |
| ADVERSE EFFECTS           |            |            |             |             |
| Sedation                  | +++        | +++        | +           | ++          |
| Extrapyramidal symptoms   | +++*       | +          | +++*        | +           |
| Anticholinergic            | +++        | +          | +           | +           |
| Hypotension               | +++        | +++        | +++         | ++          |
| Hyperglycemia             | +++        | +          | +           | +           |
| QTc prolongation          | ?          | ?          | +           | +++*        |

+, very low; ++, low; ++++, moderate; ++++, high; +++++, very high; ?, uncertain, -, minimal or no activity, *, dose dependent; adopted from 56, 58-66.

**Olanzapine**

A prospective randomized controlled trial compared olanzapine with haloperidol in the treatment of ICU delirium (67). The study showed the once daily 5 mg olanzapine doses were as effective as haloperidol 2.5 mg – 5 mg enterally three times a day. The duration of the study was for five days. There was no difference in the reduction of delirium index between two groups. A significantly more rescued intravenous haloperidol doses were required in the haloperidol group. However, the dose and route of administration of haloperidol used in the study is not standard of practice for ICU delirium. The standard of practice is to administer haloperidol intravenously for ICU delirium. In the study, it was given enterally. Because the bioavailability of oral/enteral formulation is different than intravenous formulation, haloperidol given IV maybe more superior to oral olanzapine. Also, the doses in the study were not consistent with the standard haloperidol doses used in the ICU. Extrapyramidal symptoms were reported in six patients in haloperidol group, and none in olanzapine group. However, the investigators did not assess for QTc prolongation. The authors concluded that olanzapine is a safe alternative to haloperidol.

**Quetiapine**

A double-blind, placebo controlled, multicenter study evaluated the efficacy and safety of quetiapine (68). Although this is the first study to compare the drug with placebo, rescue intravenous haloperidol were allowed to be used in both groups. The results showed quetiapine associated with quicker resolution of delirium, reduced time of delirium and agitation, and reduced haloperidol requirement compared to placebo. There was no difference in the duration of ICU stay, length of hospitalization, and hospital mortality. More adverse events such as somnolence, and hypotension were reported in quetiapine group than...
placebo but not statistically significant. There was no difference in the incidence of QTc prolongation in both groups. In the study, patients must tolerate enteral nutrition, and without a complicating neurologic condition such as hypoactive delirium. Gastroparesis is a well-known complication developed in ICU patients and often some patients could not tolerate enteral feeding despite the administration of a GI motility agent. Also, by excluding hypoactive delirium, there is a strong potential for premature assumptions about resolution of symptoms. Due to the strict exclusion criteria, patients in the study do not truly represent commonly seen patients in the ICU. In general, quetiapine was well tolerated. The authors suggest further randomized, controlled trials are needed to evaluate the role quetiapine in ICU delirium.

**Risperidone**
Currently, there is only one published study on risperidone in the treatment of delirium. The subjects in this study are mixed ICU and non-ICU patients (69). There were no statistical differences in the mean delirium rating score, time to effect, and frequency of response between the groups. Because of the small sample size (24 patients), and the lack of consensus as to optimal dosage of haloperidol or risperidone, the authors could not offer conclusive evidence as to superiority of one drug over another. There was no report of clinically significant side effects. One patient in the haloperidol group show mild symptoms of akathisia but did not require medical treatment. Nevertheless, there have been reports of increased risk of delirium and stroke with risperidone, particularly in elderly patients (70, 71). Risperidone should be used with caution especially in elderly. Regulatory agencies such as FDA and Health Canada issued a black box warning regarding elderly patients with dementia treated with risperidone are at an increased risk of cerebrovascular adverse events such as stroke and transient ischemic attacks, and death compared to placebo. The authors suggest larger randomized, controlled studies are necessary to confirm the results.

**Ziprasidone**
Ziprasidone is a newer atypical antipsychotic agent. It is potent D$_2$ and 5-HT$_2A$ receptor antagonist (72). A multicentre, randomized, double-blind, placebo-controlled trial was conducted in ICU delirium with ziprasidone, haloperidol and placebo (73). All three groups were given additional intravenous haloperidol on an as needed basis. The duration of the study was 14 days regardless of the patient clinical status. A total of 101 patients distributed evenly in all three groups were included in the study. The results showed treatment with antipsychotics did not improve the number of days alive without delirium or coma nor it increased adverse events when compared with placebo. However, more patients in the placebo group required additional haloperidol compared to the treatment groups. Reported incidences of akathisia were similar and not clinically significant in both antipsychotic groups. Fewer patients in the ziprasidone group experienced extrapyramidal symptoms than the haloperidol and placebo groups. However, this difference did not reach statistically significance. There was no difference in the incidents of QTc prolongation among the three groups. The authors concluded that the use of antipsychotics in ICU patients is feasible but larger studies are needed to determine the appropriateness of these agents in this setting.

**ALPHA-RECEPTOR ANTAGONISTS**

**Dexmedetomidine**
Dexmedetomidine, a potent selective $\alpha_2$ agonist indicated for sedation during procedures and in ICU for long term (> 24 hour). It produces sedation and analgesia without causing respiratory depression (74). Common side effects are hypotension, and bradycardia. Clinical trial showed dexmedetomidine had a lower incidence and duration of delirium compared to midazolam (45). A double blinded, randomized controlled trial using dexmedetomidine compared with lorazepam for sedation to determine the effects of on the duration of delirium and coma mechanically ventilated ICU patients (75). The treatment group experienced significantly more median days alive without delirium or coma (75). Dexmedetomadine was associated with less incidence of delirium and shorter duration of mechanical ventilation requirement. Patients in the treatment group had a higher incidence of sinusbradycardia, and there was no difference in self-extubations between two groups. There was a trend, although insignificant toward increased in atrial fibrillation in dexmedetomidine group. Economic assessments of
the use of dexmedetomidine in sedation demonstrated a better clinical consequences and economic impact when compared with midazolam (75, 76). However, there are conflicting data indicating dexmedetomidine increases the incidence of delirium (77). Although studies showed promising results, more data are required to determine the efficacy and safety of dexmedetomidine in the prevention and treatment of ICU delirium.

Clonidine
Clonidine is an α₂ agonist indicated for hypertension. It is also has analgesic, sedative and anxiolytic properties (78). It has been used as an adjunct in the treatment of narcotic, ethanol, and tobacco withdrawal (79-81). In a study with intravenous clonidine, the investigators concluded clonidine reduces the severity of delirium, improves the respiratory function, decrease the weaning duration and the ICU length of stay after surgery (82). A limitation in using clonidine in Canada is, currently, intravenous clonidine is only available through Health Canada Special Assess Programme.

OTHER AGENT

Rivastigmine
Evidence suggests that impaired cholinergic neurotransmission plays an important role in the development of delirium (8, 10, 12). It has been shown that there is a strong correlation between increased anticholinergic activity and the development of delirium in particular with elderly patients (83). Acetylcholinesterase inhibitors inhibit the cholinesterase enzyme preventing the destruction of acetylcholine. This results in increased level duration of acetylcholine. Rivastigmine is an acetylcholinesterase and butyrylcholinesterase inhibitor indicated for Alzheimer’s disease, Parkinson’s disease dementia, and Lewy-body dementia. Cholinesterase inhibitors such as rivastigmine have been successfully used to treat delirium in elderly patients (84-87). A recent study investigates the use of rivastigmine as an adjunct to haloperidol in ICU patients with delirium (88). Unfortunately, the study was terminated prematurely due to increased mortality in the rivastigmine group. The investigators concluded that rivastigmine did not reduce the duration of delirium, hence, the use of rivastigmine in the treatment of ICU delirium was not recommended.

COMPLEMENTARY AND ALTERNATIVE MEDICINE

Complementary and alternative medicine (CAM) is a broad category of therapies, treatments, and use of products that are not considered mainstream conventional (Western) medicine (89, 90). CAM is classified into five categories:(89)

1. Medical systems which includes ayurvedic medicine, Chinese medicine, and homeopathy/naturopathic medicine.
2. Mind-body medicine includes animal-assisted therapy, art therapy, guided imagery, meditation, music therapy, and prayer.
3. Biologically based practices include dietary supplements, and herbal products.
4. Manipulative/body based practices involves acupressure, chiropractic/manipulative therapy and massage.
5. Energy medicine consists of Qi gong, reiki, therapeutic touch and bioelectromagnetic-based therapies.

It has been suggested that CAM may provide additional modalities for managing patients with delirium in ICU. Most critical care nurses view the use of CAM positively and are receptive to their use (91-95). The common CAMs used in ICU are diet, counseling/psychology, relaxation techniques, massage, prayer, and music therapy.

Music Therapy
Music intervention consists of the use of preferred music to achieve a therapeutic goal (92). Studies in mechanically ventilated ICU patients listening to preferred music, and relaxing music, through a headphone showed a reduction in anxiety, and physiological stress response from stress (96-99).

Promotion of Sleep
Most ICU patients develop impaired sleep pattern due to the use of opioids and benzodiazepines as well as other environmental factors which affect the rapid eye movement sleep (13, 14, 100). One of the consequences from impaired sleep is delirium (101). There are strong association between sleep deprivation and delirium in elderly, postoperative patients, and ICU patients (13, 14, 101).
Quiet time protocol
In an effort to promote sleep in the ICU, a quiet time protocol was developed to reduce noise and light levels at specific time twice a day. The study showed a significant increased of observed sleep after the protocol was implemented (37, 102).

Relaxation
Progressive muscle relaxation can decrease muscle tension, reduce stress, and help in pain control (92). This will help patients achieve the state of relaxation which will promote better sleep.

Massage
Massage is commonly utilized by nurses in the ICU to prepare patients for sleep (103, 104). Massage does not require special equipment and basic techniques can be easily learned and remembered by caregivers.

Melatonin
Melatonin, a hormone, plays an important role in the regulation of the sleep-wake cycle (105). Disruption of the sleep-wake cycle, with fragmented sleep during the night and sleepiness during the day, is frequently observed during delirium (106). Studies indicate an association between delirium and interrupted melatonin secretion (107-111). Despite evidence suggest that exogenous melatonin supplementation would regulate sleep cycle, results from studies did not demonstrate better sleep (112-114). The limitations of these studies include a lack of consistent sleep assessment, and control of environmental factors. Due to lack of randomized controlled studies to show safety and efficacy in the treatment or prevention of ICU delirium, melatonin is not recommended.

It is evident that critical care nurses are using CAM in the ICU. However, they acknowledge that there are many barriers in their ability to implement the practice. The barriers included a lack of knowledge and training in CAM, a lack of allowed time and physician reluctance to use CAM (95). Also, CAM is not innocuous. For example, relaxation is typically considered a positive response for patients who are under stress. However, relaxation tends to reduce blood pressure, heart rate and respiratory rate (115), which a critically ill patient may not be able to tolerate. In addition, patients may be hypersensitive to touch and may experience negative response such as anxiety. Before incorporating CAM into the plan of care for ICU patients, a careful evaluation of ICU confounders such as openness to use of CAM by providers, and resources available is needed.

CONCLUSION
Delirium is a serious complication in critically ill patients associated with increased morbidity and mortality. Patients should be assessed for delirium daily. Currently, there is a lack of randomized control trials with pharmacological treatments in ICU delirium. The complexity of medical condition exists in ICU patients limits the feasibility in recruiting homogenous study subjects for large randomized, controlled trials. Non-pharmacologic approach in the treatment of non-ICU delirium has been implemented in the ICU with success. However, caution is warranted when extrapolating results from pharmacologic treatments in non-ICU patients to ICU patients. In the case of rivastigmine, it was shown to be effective non-ICU delirium but increased mortality in ICU patients without benefit. The recommended drug of choice by both the Intensive Care Society and the American College of Critical Care Medicine is haloperidol. The strategies in managing ICU delirium should include prevention, identify risk factors, diagnosis and treatment with non-pharmacologic and pharmacologic modalities.

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