Palliative Sedation for Children at End of Life: a Retrospective Cohort Study

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Research Article

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Abstract

Background: Palliative sedation is consciously reducing the patient's consciousness to alleviate the refractory symptoms. However, studies on palliative sedation for children are scarce. We aimed to survey the symptom control and risks for children with sedative therapy in end of life.

Method: This study was a single center retrospective cohort study. Children who died in the Department of Palliative Medicine were divided into palliative sedation (Group A) and non-palliative sedation group (Group B). The symptoms relief, survival time, and last hospitalization time were compared between two groups.

Results: From January 2012 to November 2019, 41 children died in department of palliative care. 24 children were sedated (Group A), meanwhile 17 children were not (Group B). The symptoms in Group A were more complex than Group B (p=0.013). Overall symptom relief in Group A was higher than that in Group B (24/24, 10/15 p=0.041). Pain relief rates (7/7, 20/21 p=0.714), maximum/pre-death opioid dose (30(20, 77.5), 18(9, 45) p=0.208) and pain intensity difference (5(4.6.5), 4(2.6) p=0.315) were not statistically significant difference in both group. After diagnosis, the survival time of the Group A was longer than the Group B (p=0.047). However, the length of hospitalization before death was similar in two groups (p=0.385).

Conclusion: Palliative sedation controls complicated, painful symptoms at the end of life and does not shorten the hospitalization time in children.

Background

The annual demand for palliative care for children in China ranges from 23 to 24 per 100,000 people [1], with approximately 3,037,950 cases of children have pediatric palliative care needs among 1.3 billion people [2]. In 2017, the Integrated National Mortality Surveillance System, which covers 24 percent of China's population, showed that 90,600 deceased children may benefit from palliative care [3].

Palliative care for children not only improves the physical and psychosocial symptoms of children with limited lives [4, 5], but also benefits those children themselves and their families [6–10]. There is now an increasing social emphasis on timely intervention in palliative care for children. Palliative care for children could control symptoms, reduce overtreatment and allow for end-of-life preparation of those children. These contribute to improve psychosocial outcomes during parental bereavement [11–13]. Various symptoms appear at children's end-of-life, which are intractable, unmanageable and un-relievable, and cause great suffering and distress to children, their families and caregivers [14, 15]. Palliative sedation, which aims to relieve end-of-life refractory symptoms by reducing patient's consciousness, is used in 12-64% of adults in palliative care [16–18] and in 48.4% of adults who die in hospital [19]. Although there has been demonstrated benefit of better symptom control in patients with a terminal illness, palliative sedation continues to gather some controversy [20, 21].

In the available studies, palliative sedation in children has been reported as an isolated case and experience sharing [22–25], and there is a lack of effect and safety evidence for palliative sedation in children. The aim of this study was to reveal the sedation procedure, symptom control and risks for children in end of life, in order to enrich the practical experience for children palliative sedation.

Methods

1.1 Patients

We retrospectively recorded pediatric patients admitted to the palliative care department of our hospital between January 2012 and November 2019. Inclusion criteria: 1. Age 0-18; 2. Received palliative care; 3. Died during hospitalization. Exclusion criteria: 1. Age >18; 2. No death during hospitalization; 3. Information is incomplete.

1.2 Methods

1.2.1 Group: Children were divided into palliative sedation group (Group A) and non-palliative sedation group (Group B) according to whether used sedative drugs (midazolam, chlorpromazine), comparing the demographic characteristics, principal diagnosis, types of symptoms and hydration of two groups at admission.

1.2.2 Observation: Information about symptoms relief, pain, dyspnea, survival time after diagnosis, and last hospitalization time was extracted from the electronic medical record. Symptoms mentioned in the medical records were recorded, such as pain, dyspnea, irritability, coma, convulsions, vomiting, and fever.

Description of pain was pain intensity difference (PID), pain relief rate, and pain relief (PAR) [26]. Pain intensity (PI) assessments used FLACC scale scores, Wang-Baker faces pain rating scales, or numerical rating scales (NRS), according to ages and ability of cognitive. Analgesic drugs (initial dose, maximum dose, pre-death dose and route), pain relief, and length of sleep were recorded. Dose of opioids every 24h converted to oral
morphine (conversion rate, oral morphine: intravenous morphine =3:1, oral morphine: subcutaneous morphine =2:1, oral morphine: fentanyl transdermal (25mcg/h patch)=60:25, oral morphine: oral oxycodone =2:1) [27–29]. Pain intensity difference (PID) meant the difference between the pain score at the beginning of the administration of analgesics and the pain score at each time point. Pain relief (PAR) grading: not alleviated; mild relief (pain relief about 1/4); moderate relief (pain relief about 1/2); significant relief (pain relief about 3/4); complete relief (pain disappearance).

Dyspnea: Medical records mentioned in breathing, shortness of breath, gasp, or dyspnea, were equal to dyspnea. Dyspnea relief: 1.Clearly documented alleviation after treatment; 2.If symptoms were not mentioned later, symptoms were deemed to be relieved.

1.2.3 Statistical Analysis: Statistical analyses were performed using SPSS 20.0. Descriptive statistics (e.g. mean, SD, median, interquartile range) was used for analysis, and frequencies were calculated for continuous and categorical data, respectively. For comparison in continuous data/ordinal data and categorical data, we used Wilcoxon rank sum tests and chi-square test. Spearman's rank correlation was used for the correlation test. P-values lower than 0.05 were considered statistically significant.

1.2.4 Ethics
The study was approved by the medical ethics committee of West China Fourth Hospital of Sichuan University (No. HXSY-EC-2021027).

Results
2.1 demographic characteristics of group A and B
Between January 2012 and November 2019, 80 children were admitted to the Department of Palliative Medicine, West China Fourth Hospital, of whom 41 died during their hospitalisation. There were 24 in the palliative sedation group (Group A) and 17 in the non-palliative sedation group (Group B). The demographic characteristics of the two groups are shown in Table 1. The differences between the two groups were not statistically significant.

| Demographic characteristics (N=41) | Group A (n=24) | Group B (n=17) | p     |
|-----------------------------------|---------------|---------------|-------|
| **Age**                           |               |               |       |
| 0-28 days                         | 0             | 3             | 0.228 |
| 29 days -1 year                   | 3             | 1             |       |
| 2-6 years                         | 11            | 6             |       |
| 7-18 years                        | 10            | 7             |       |
| **Gender**                        |               |               |       |
| Male                              | 15            | 13            | 0.40  |
| Female                            | 9             | 4             |       |
| **Type of disease**               |               |               |       |
| Blood tumor                       | 4             | 3             | 0.564 |
| Solid tumor                       | 16            | 8             |       |
| Congenital disease                | 2             | 3             |       |
| Diseases of the blood system      | 0             | 1             |       |
| Severe infection                  | 2             | 2             |       |
| **Hydration during hospitalization** | 21          | 14            | 0.679 |

1Fisher of the exact probability method.

2.2 Symptoms
The main symptoms were pain, dyspnea, irritability, fever, coma, vomiting, convulsions (see figure 1). Pain occurring in 28/41 (68%) children was the most common symptom affecting the children's quality of survival at the end of life. Four children in group A got four types of symptoms, ten children got three types, seven children got two types and three children with one symptom. None children got four symptoms in group B, four children got three symptoms, three got two symptoms and ten got one symptom. Distribution of symptoms in two groups see in Table 2. Types of symptoms differed between the two groups, \( p = 0.013 \). The symptoms in group A were more complex than those in group B. But overall symptom relief was higher in A Group which used sedative medication (\( p = 0.041 \)). Five patients in Group B whose symptoms remained unrelieved were fever, abdominal distension, and dyspnea. Three of five were non-tumor. Pain control rate was similarities between the two groups, 95.23% and 100% respectively.

### Table 2

| Type of symptom(s) | Symptom control | Pain control |
|-------------------|----------------|-------------|
|                   | 1 2 3 4 n      | No remission remission Obvious remission n No remission Mild remission Moderate remissions Obvious remissions |
| Group A           | 24 3 7 10 4 24 | 0 20 4 21\(^b\) | 1\(^c\) |
| Group B           | 17 10 3 4 0 15\(^a\) | 5 8 2 7\(^b\) | 0 2 4 1 |
| \( Z \)           |               | 2.045       | 0.367 |
| \( p \)           | 0.013\(^d\)   | 0.041\(^e\) | 0.714 \(^e\) |

a. 2 cases of coma excluded from 17, were not included in the assessment.
b. There were no analgesia in 3 cases of Group A and 10 cases of Group B.
c. too short hospital stay to evaluate.
d. the exact probability method.
e. rank sum test.

#### 2.3 Pain control

Twenty-one children in Group A used analgesic; besides three patients of Group A were not prescribed for analgesia because the main symptoms were convulsions and moans. In beginning, four children had been prescribed two types of analgesics at the same time, such as hydrocodone sustained-release tablets, acetaminophen and hydrocodone, morphine sulfate solution combined with continuous venous morphine by syringe driver, fentanyl transdermal combined with morphine sulfate solution. The other 16 patients received hydrochloride morphine continuous venous infusion and 1 patient used morphine sulfate oral solution alone. 7 children in group B used analgesic (7/17 cases who got fever, coma were not prescribed analgesic, and 3/17 cases who prescribed morphine only because of dyspnea were not included in pain assessment). In group B one child used hydrocodone sustained-release tablets, six used morphine continuous venous infusion. Pain scores on admission and after control were higher in group A than in group B (\( p = 0.014, 0.039 \)), but there was no significant difference in maximum opioid dosage and pre-death opioid dosage between the two groups. Pain control is shown in Table 3.

### Table 3

| N | Admission | Pain score (hours) | Sleep (hours) | Pain score | Pain duration (hours) | Sleep (hours) | Maximum opioid dose converted to oral morphine dosage/24 hours (mg) | Opioid dose before death converted to oral morphine dosage/24 hours (mg) | Pain intensity difference |
|---|-----------|--------------------|---------------|------------|-----------------------|---------------|------------------------------------------------------------------|------------------------------------------------------------------|--------------------------|
|   |           |                    |               |            |                       |               |                                                                  |                                                                  |                          |
| Group A | 21 | 8(7,10) | 24(20,24) | 6(4,7) | 3(3,4) | 4(2.5,7) | 8(6,10.5) | 30(20,77.5) | 30(20,60) | 5(4,6.5) |
| Group B | 7  | 7(6,7)  | 22(18,24) | 4(4.5,5) | 2(0,3) | 4(1.5,13) | 9(8,7) | 18(9,45)  | 18(9,45)  | 4(2,6)    |
| Z     | -2.501 | -0.959 | -1.481 | -2.068 | -0.065 | 0.615 | -1.358 | -1.278 | -1.004 |
| \( p \) | 0.014 | 0.294 | 0.139 | 0.039 | 0.948 | 0.538 | 0.175 | 0.208 | 0.315 |
Opioids were the first choice for moderate and severe pain in children at the end of life. In our study, before hospital admission 24 children were treated with morphine sulfate/hydrochloride, 2 with oxycodone hydrochloride, 1 with fentanyl and 1 with acetaminophen and hydrocodone mixture. The forms included sustained-release tablets, oral liquid, injection, and transdermal patches. Two children were treated with two types of opioids (morphine sulfate oral solution + morphine hydrochloride injection, fentanyl transdermal patch + morphine sulfate oral solution). Sixteen cases only used single opioid analgesia, 12 cases used 1-3 adjuvant analgesic drugs, including paracetamol, ketorolac trometamol, scopolamine butyrate, gabapentin capsule, valproate sodium, and ketamine.

2.4 Dyspnea

In the study, twelve children had dyspnea, aged 4 days to 14 years. Three patients had dyspnea as single symptom, and other nine combined with pain, irritability, fever, and other symptoms. There were six cases in Group A and six cases in Group B. Both groups had 83% remission of dyspnea. Both groups used morphine to relieve dyspnea. The survival time and the hospitalization time of the two groups were shown in Table 4 (The data are non-normal, expressed in quartile spacing). The sample size of children with dyspnea was too small to analyze.

|                | n  | Dyspnea relief rate | Survival time after diagnosis of diseasea | Hospitalization timea |
|----------------|----|---------------------|------------------------------------------|------------------------|
|                |    | % (n)               | (days)                                   | (days)                 |
| Group A        | 6  | 83% (5)             | 450 (253.5,1036.5)                       | 7.5 (12.1,5)           |
| Group B        | 6  | 83% (5)             | 63.5 (52.5,1585)                         | 12 (1.75,51.75)        |
| a. quartile spacing |    |                     |                                          |                        |

2.5 Duration of survival and hospitalization time after diagnosis in Group A and Group B.

The survival time after diagnosis in Group A was 20 days - 2190 days and hospitalization time was 1-85 days. The duration of survival in Group B was 5 days - 5475 days and the length of stay in hospital was 1 - 63 days (see Table 5). Longer survival time after disease diagnosis in group A compared to group B (p = 0.047). However, the hospitalization time before death was similar. Palliative sedation treatment seemed not shorten the hospitalization time of children at the end of life.

|                | Survival time after diagnosis of disease | Last hospitalization |
|----------------|------------------------------------------|----------------------|
| Group A        | 365 (112.5,730)                          | 9 (2.5,22,75)        |
| Group B        | 60 (30,795)                              | 3 (1.24)             |
| Z              | -1.987                                   | -0.869               |
| p              | 0.047                                    | 0.385                |

2.6. Sedation in Group A

In electronic records, children (age from 5 months to 14 years, 15 males, 4 blood tumors, 16 solid tumors, 2 Congenital diseases, 2 infection, were prescribed sedation drugs. There were one to four main symptoms types (pain, dyspnea, irritability, fever, coma, vomiting, convulsions). Palliative sedation was mainly caused by irritability in 13 cases, convulsion in 5 cases, pain in 7 cases, and dyspnea in 6 cases. Midazolam was used in 17 cases; chlorpromazine was used in 1 case. Six cases used midazolam combined with chlorpromazine. The hospital stays lasted from 1 hour to 85 days, and the use of sedatives lasted from 1 hour to 47 days. The purpose of sedation was to reduce consciousness to control painful symptoms. Before and after sedation, the average Ramsay score was 1(1, 1) and 2(2, 3). When death was coming, the deeper sedation score was observed 3(2, 4) (p < 0.01)(see Table 6). Beginning of the sedation, the initial dose of midazolam ranged between 0.5 - 5 mg/24 hours [mean 1.5(1, 2.4) mg/24h]. Maximum dose was 0.5-30 mg/24 hours [mean 3(1.6)mg/24h], and the pre-death dose was 0.5-25 mg/24 h [mean 3(1.5)mg/24h]. The maximum dosage of midazolam was similar to that before death (p = 0.066). In 4 cases, midazolam was down regulated before death for unknown reasons, but the degree of sedation was not reduced. Midazolam sedation duration did not appear to be associated with maximum dose and pre-death does.
### Table 6

| Ramsay score a | Pre- sedation score | Sedation score after symptom control | Pre-death sedation score | \( \chi^2 \) | \( p^b \) |
|----------------|---------------------|--------------------------------------|--------------------------|----------|---------|
|                | 1(1,1)              | 2(2,3)                               | 3(2,4)                   | 34.344   | <0.01   |

a. percentile spacing,  
b. Friedman Inspection

### Discussion

There are refractory pain symptoms in children at the end of life. These symptoms seriously affect the quality of life for children and their caregivers and bring a heavy mental and psychological burden to medical workers [14, 15]. Palliative sedation consists in consciously reducing the patient's consciousness to alleviate the symptoms. From January 2012 to November 2019, 58.5% (24/41) of children who died in the palliative care department were treated with palliative sedation, similar to the palliative sedation rate reported in previous studies [25, 30, 31].

End-of-life symptoms of children are complex, and pain is the most common symptom affecting children's quality of life [32, 33], followed by dyspnea, irritability, fever, and other symptoms. Symptom complexity in the palliative sedation group was higher than that in the non-sedation group.

During palliative sedation management, midazolam started from low dosage and was titrated until the refractory symptoms were relieved. Midazolam maximum dosage varies greatly between individuals, and the average Ramsay score after symptom control was 2 (percentile spacing 2-3). Most patients were sedative to 2-3 scores, only two children's Ramsay score was 5. All sedation were continued to death, sedation duration ranging from 11 hours to 25 days. There was no difference in hospitalization duration between the sedation group and the non-sedation group. Sedation until death should not short the last time of children. During sedation, children were all properly rehydrated and there was no serious damage due to rehydration. Whether it is necessary to change the level of sedation when patients could reach deep sedation? Four children's midazolam dosage were reduced before death, but the level of sedation did not improve.

Opioids were the first choice for moderate and severe pain in children at the end of life [34] (WHO, 2012). In our study, before hospital admission 24 children were treated with morphine sulfate/hydrochloride, 2 with oxycodone hydrochloride, 1 with fentanyl and 1 with acetaminophen and hydrocodone mixture. The forms included sustained-release tablets, oral liquid, injection, and transdermal patches. Two children were treated with two types of opioids (morphine sulfate oral solution + morphine hydrochloride intravenous infusion, fentanyl transdermal patch + morphine sulfate oral solution). Sixteen cases only used single opioid, 12 cases used 1-3 adjuvant analgesic drugs, including paracetamol, ketorolac trometamol, scopolamine butyrate, gabapentin capsule, valproate sodium, and ketamine. Sedation group had higher pain score at admission. Combination of sedative drugs and opioids avoided more opioids doses.

End-of-life dyspnea was treated with pathogenesis treatment, including anti-infection, bronchodilators, glucocorticoids, and other drugs to correct reversible factors. All children with dyspnea used morphine to relieve dyspnea, and six children chose palliative sedation. Both sedation and non-sedation groups achieved a high rate of symptom relief (both 83%).

This is the first study describing procedures for refractory symptoms for children in end of life. This study found that palliative sedation could relieve refractory symptoms for children in end of life, while could not hasten death. However symptoms of sedation group were more complex than those of non - sedation group, symptom relief rate was higher in sedation group. Titration with a sedative to alleviate suffering was safe management for children in palliative department. Sedation until death should not shorten the last time of children. The presence of multiple symptoms and refractory symptoms makes it easier for doctors and families to make decision to opt for palliative sedation. Midazolam is one of the commonly used drugs for palliative sedation in adults [35]. It is still the first choice for palliative sedation in children in our hospital because of its rapid effect and short duration of action. Severe adverse reactions such as central inhibition and respiratory inhibition were not observed after titration.

It was noted that palliative sedation decision may be associated with symptom complexity, degree of pain, refractory dyspnea, disease course, disease severity and expected survival time, as well as healthcare professionals' perceptions of palliative sedation and parental wishes. All cases were discussed before sedation by a team of palliative medicine experts to determine that the symptoms were refractory, and the disease
progression was irreversible and children's survival time was limited. Subsequently, a family meeting was organized, and an informed consent form was signed. None of these children attended the communication before sedation because these children were not the legal age to become their agents. All the palliative sedation informed consent forms were signed by their guardian agent.

However, palliative sedation for children seemed effective and safe in this study. Our study was a retrospective study and small sample size. There are fewer children in palliative care than adults or the elderly. Therefore, multi-center clinical studies would be needed for further confirmation.

Palliative sedation was debated by society and medical field. Whether to involve the child in the decision-making discussion about sedation and sedation until death prior to sedation remains an ethical discussion.

**Conclusion**

Palliative sedation controls complicated, painful symptoms at the end of life and does not shorten the hospitalization time in children.

**Declarations**

**Ethics approval and consent to participate**

The study was approved by the medical ethics committee of West China Fourth Hospital of Sichuan University (No. HXSY-EC-2021027).

All methods were carried out in accordance with relevant regulations. And all the palliative sedation informed consent forms were signed by guardian agent of patient.

**Consent for publication**

All author agree with submission to BMC Palliative Care.

**Availability of data and materials**

The data used and analysed during the current study are available from the corresponding author on reasonable request.

**Competing interests**

Not applicable

**Funding**

Not applicable

**Authors’ contributions**

M.D. Yang Chen helped in conceptualization, data collection, data analysis, writing, and editing. Jianjun Jiang and Wei Peng helped in data analysis, writing, and editing. Chuan Zhang helped in conceptualization, data collection, data analysis, and editing.

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**Figures**

**Figure 1**

The distribution of main symptoms in Group A and Group B