Advance care planning for adolescent patients with life-threatening neurological conditions: a survey of Japanese paediatric neurologists

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ABSTRACT

Objective To evaluate current attitudes and barriers to advance care planning for adolescent patients with life-threatening conditions among paediatric neurologists.

Design Cross-sectional study. A self-reported questionnaire was administered to assess the practice of advance care planning, advance directives and barriers to advance care planning for adolescent patients with life-threatening conditions. All board-certified paediatric neurologists in Japan were surveyed and those who had experience in taking care of adolescent patients with decision-making capacity were analysed. We compared the results with those of paediatric haematologists reported previously.

Results In total, 186 paediatric neurologists were analysed. If the patient’s prognosis was <3 months, only about 30% of paediatric neurologists reported having discussions with patients, such as ‘do not attempt resuscitation’ orders (28%) and ventilator use (32%), whereas more than 70% did discuss these topics with patients’ families. About half of the paediatric neurologists did not discuss advance directives at the end of life with their patients, whereas over 75% did discuss advance directives with patients’ families. Compared with paediatric haematologists, paediatric neurologists had more end-of-life discussions with patients, such as where treatment and care will take place, do not attempt resuscitation orders, and the use of a ventilator, if the patient’s prognosis was >1 year.

Conclusion About half or less of the paediatric neurologists discussed advance care planning and advance directives with their adolescent patients who had life-threatening conditions, even if the patient’s prognosis was <3 months. They tended to discuss advance care planning and advance directives more with families than with patients themselves.

INTRODUCTION

Advance care planning (ACP) is, at its most basic, a process of thinking ahead to treatment choices, goals of care and/or choosing another person (proxy) to speak for oneself at a point in the future.1 Paediatric ACP is internationally recommended and welcomed by healthcare providers.2,3 It can increase the capacity of children and parents to anticipate decisions, support family coping, provide peace of mind, increase sense of control and reduce suffering.4–6 One important result of an ACP discussion may be an advance directive (AD), a written order delineating explicit wishes regarding medical interventions.

ACP with adolescent patients is gaining increasing attention, but is a difficult area of practice. Reports show that adolescent patients have the desire and ability to share their values, beliefs and preferences for treatment at the end of life.7–10 A previous study found that 75% of adolescent patients with cancer believed it was appropriate to discuss
end-of-life decisions even before their condition worsened. However, some care providers exclude adolescent patients from ACP because they are deemed not legally competent to make decisions for themselves.\textsuperscript{3,5}

While palliative care emerged with the treatment of patients with terminal cancer, more recent developments in this field suggest that palliative care may be appropriate for any neurological patient living with advanced, progressive illness or multiple comorbidities.\textsuperscript{12} Neurological diseases are largely incurable and reduce life expectancy, so all neurologists should be knowledgeable about, and feel comfortable discussing, ACP with patients and families. However, in the adult setting, many patients have never heard of ACP, and according to foreign report, 30\% of patients with advance care plans do not share these with their physicians.\textsuperscript{13} There has been no research about the current ACP and AD practices among paediatric neurologists with regard to adolescent patients.

We reported previously that haematologists in Japan tended to discuss ACP and ADs more with the families than with the adolescent patients themselves. In fact, more than two-thirds of haematologists reported not having difficult discussions, such as do not attempt resuscitation (DNAR) orders, ventilator use, and prognosis, with their patients.\textsuperscript{14} To compare with haematologists, there is a possibility that it becomes clear whether the reason of the difficulty of ACP practice differs depending on the disease or depending on age.

The aim of the present study was to determine current practices regarding ACP and ADs among paediatric neurologists with regard to adolescent patients with life-threatening conditions. In addition, the study aimed to clarify differences between paediatric neurologists and paediatric haematologists in their attitudes and barriers to ACP and ADs for adolescent patients with life-threatening conditions.

METHODS
A self-reported questionnaire (see online supplementary file) was administered to paediatric neurologists to assess their practices regarding ACP and ADs, as well as barriers to ACP, for adolescent patients with life-threatening conditions.

Study population
All paediatric neurologists certified by the board of the Japanese Society of Child Neurology, the oldest and leading paediatric neurology academy in Japan, were surveyed. The total number of neurologists certified by the board was 1081. In order to compare the data of paediatric neurologists with those of paediatric haematologists, the data from paediatric haematologists regarding attitudes and barriers to ACP and ADs were extracted from a previously reported study that was conducted at the same time using the same questionnaire.\textsuperscript{14}

Data collection
Mailing information for all board-certified paediatric neurologists was obtained from the Japanese Society of Child Neurology’s web page.\textsuperscript{15} All eligible physicians were sent a letter explaining the study and containing a questionnaire, and requesting their participation. Paediatric neurologists for whom a mailing address was not available (n=37) were excluded from the study. A reminder letter was sent to all participants 4 months after the initial mail-out, except to those for whom the initial letters were returned because of incorrect addresses. Identifying information, such as names and addresses, was not linked to respondents’ answers. Data were collected from October 2015 to May 2016 and were analysed for paediatric neurologists who indicated that they had experience in taking care of adolescent patients with life-threatening conditions who were also capable of making decisions like a kind of muscular dystrophy.

Questionnaire
The questionnaire consisted of 82 items. Items were adapted from existing surveys\textsuperscript{16-17} after minor revision, following discussions with specialists, namely three palliative care physicians, two paediatricians, two nurses and two psychotherapists. The survey was pilot-tested by a convenience sample of 15 physicians, including both paediatric neurologists and haematologists, and revised according to feedback from cognitive debriefing. All questions were close-ended, requiring categorical responses or rating on a Likert-type scale.

Outcome measures
Practice of ACP
The survey items concerning the individual physician’s practices in relation to ACP were separated into two parts. The first part examined the physician’s practice of ACP with adolescent patients with life-threatening conditions who had decision-making capacity and included questions regarding how frequently the physician did each of the following if their patient’s prognosis was >1 year or <3 months:

- discuss the patient’s medical condition
- verify the patient’s understanding of his/her medical condition
- discuss the patient’s prognosis
- discuss the goals of treatment and care
- promote sharing the treatment and care goals between the patient and his/her family
- discuss where treatment and care are to take place
- discuss ‘do not attempt resuscitation’ (DNAR) orders with the patient
- discuss ventilator treatment if the patient’s condition worsens.

Physicians were required to answer these questions using a 5-point Likert-type scale: ‘always’, ‘often’, ‘sometimes’, ‘rarely’ or ‘never’. The second part of this section of the survey asked clinicians to answer the same questions, but with regard to discussions with the patients’ families instead.
Discussion about ADs at the end of life

Participants were asked about the discussions they have with patients with decision-making capacity about ADs at the end of life using the following binary question: ‘Do you usually discuss resuscitation and life-prolonging therapy with patients with decision-making capacity if their prognosis is less than 4 weeks?’ Participants were also asked about the discussions they have with patients with decision-making capacity regarding cardiopulmonary resuscitation and the use of ventilators, vasopressors, antibiotics, tube-feeding and intravenous fluids. The same questions were also used to investigate the participants’ discussions of ADs with the patients’ families.

Barriers to ACP

Using a 5-point Likert-type scale (always, often, sometimes, rarely or never), physicians were asked to rate how often 29 potential barriers were actual impediments to ACP. These questions were created on the basis of a previous study. Specific items assessed the physician’s perception of barriers related to patient/family behaviours and practices, including patient/family expectations, readiness to have the discussion, understanding the medical issues and prognosis, and conflict between the patient and family members. Potential barriers related to physician behaviours/practices included concern about taking away hope or losing trust, not knowing the right things to say, the lack of a relationship with the patient and/or family, not knowing the right time to hold the discussion, uncertainty about the prognosis, physician expectations, lack of readiness to have the discussion, lack of time, physicians not placing much importance on ACP, lack of training, ethical considerations, and lack of laws and/or guidelines.

Additional covariates

Sociodemographic characteristics

Survey respondents were asked to report their specialty, their experience (years practising), age, sex and the number of dying patients under 18 years of age they have cared for over their entire career. In addition, they were asked to specify the type of medical facility in which they were currently working from the following choices: university hospital, children’s hospital, other hospitals, rehabilitation centre, clinic or ‘other’.

Statistical analysis

Statistical analyses were conducted using SAS V.9.2 (SAS Institute). Demographic data were summarised using descriptive statistics. Group comparisons between paediatric neurologists and haematologists were made using logistic regression model adjusted by sex, postgraduate experience and the number of dying patients under 18 years of age they have cared for over their entire career. In the present study, questions regarding ACP practices rated using Likert-type scales were dichotomised as follows: 0, ‘sometimes’, ‘rarely’ or ‘never’; 1, ‘always’ or ‘often’. Similarly, responses regarding barriers to ACP were dichotomised as a barrier frequency variable as: 0, ‘sometimes’, ‘rarely’ or ‘never’; 1, ‘always’ or ‘often’. Although this system was used to dichotomise variables in the present study, there were no differences in results if we dichotomised variables as ‘rarely or never’ versus all other categories or ‘never’ alone versus all other categories.

RESULTS

Of the paediatric neurologists eligible to participate in the study, responses were obtained from 564 (response rate 54%). Finally, data were analysed for 186 paediatric neurologists who indicated that they had experience in taking care of adolescent patients with life-threatening conditions and decision-making abilities.

Sample characteristics

The characteristics of the study participants are given in table 1. Average year of postgraduate experience (SD) was 28.3 (10.7) and 70% of paediatric neurologists had experience in caring for more than 10 dying patients <18 years of age (table 1).

ACP practices of paediatric neurologists for adolescent patients with life-threatening conditions

The ACP practices of paediatric neurologists for adolescent patients with life-threatening conditions are summarised in table 2 and compared with paediatric haematologists. If the patient’s prognosis was >1 year, fewer than 50% of neurologists discussed with the patient his/her medical condition (48%), verified the patient’s understanding of his/her medical condition (43%), or
Table 2  ACP practices of paediatric neurologists compared with paediatric haematologists

| Number of paediatricians who answered ‘often’ or ‘always’ for each question | Neurologists (n=186) | Haematologists (n=227) | p Value |
|--------------------------------------------------------------------------|----------------------|------------------------|---------|
| **Adolescent patients**                                                  |                      |                        |         |
| Prognosis >1 year                                                        |                      |                        |         |
| Discuss medical condition                                                | 85 (48)              | 155 (69)               | <0.01   |
| Verify understanding of medical condition                                | 77 (43)              | 136 (61)               | 0.01    |
| Promote sharing of treatment and care goals between patients and families | 28 (16)              | 135 (61)               | 0.43    |
| Discuss goals of treatment and care                                      | 88 (49)              | 126 (57)               | 0.22    |
| Discuss where treatment and care will take place                         | 88 (50)              | 98 (44)                | 0.13    |
| Discuss prognosis                                                        | 64 (36)              | 52 (23)                | 0.37    |
| Discuss DNAR                                                             | 19 (11)              | 17 (8)                 | 0.18    |
| Discuss ventilator treatment if the patient’s condition worsens          | 35 (20)              | 17 (8)                 | <0.01   |
| Prognosis <3 months                                                      |                      |                        |         |
| Discuss medical condition                                                | 100 (56)             | 135 (59)               | 0.57    |
| Verify understanding of medical condition                                | 97 (55)              | 125 (55)               | 0.56    |
| Promote sharing of treatment and care goals between patients and families | 65 (37)              | 133 (59)               | 0.04    |
| Discuss goals of treatment and care                                      | 91 (51)              | 125 (55)               | 0.93    |
| Discuss where treatment and care will take place                         | 100 (57)             | 111 (50)               | 0.65    |
| Discuss prognosis                                                        | 84 (49)              | 67 (30)                | 0.64    |
| Discuss DNAR                                                             | 49 (28)              | 38 (17)                | <0.01   |
| Discuss ventilator treatment if the patient’s condition worsens          | 56 (32)              | 42 (19)                | <0.01   |
| **Family**                                                               |                      |                        |         |
| Prognosis >1 year                                                        |                      |                        |         |
| Discuss medical condition                                                | 151 (83)             | 201 (92)               | 0.05    |
| Verify understanding of medical condition                                | 151 (83)             | 200 (91)               | 0.04    |
| Promote sharing of treatment and care goals between patients and families | 143 (79)             | 192 (88)               | 0.58    |
| Discuss goals of treatment and care                                      | 144 (80)             | 188 (86)               | 0.13    |
| Discuss where treatment and care will take place                         | 135 (75)             | 181 (83)               | 0.11    |
| Discuss prognosis                                                        | 118 (67)             | 163 (75)               | 0.29    |
| Discuss DNAR                                                             | 84 (46)              | 96 (44)                | 0.78    |
| Discuss ventilator treatment if the patient’s condition worsens          | 94 (52)              | 95 (43)                | 0.18    |
| Prognosis <3 months                                                      |                      |                        |         |
| Discuss medical condition                                                | 158 (89)             | 213 (96)               | 0.03    |
| Verify understanding of medical condition                                | 157 (88)             | 212 (96)               | 0.10    |
| Promote sharing of treatment and care goals between patients and families | 156 (88)             | 207 (94)               | 0.23    |
| Discuss goals of treatment and care                                      | 153 (86)             | 207 (94)               | 0.18    |
| Discuss where treatment and care will take place                         | 152 (85)             | 202 (91)               | 0.30    |
| Discuss prognosis                                                        | 146 (82)             | 196 (89)               | 0.23    |
| Discuss DNAR                                                             | 124 (70)             | 162 (73)               | 0.68    |
| Discuss ventilator treatment if the patient’s condition worsens          | 136 (76)             | 166 (75)               | 0.30    |

Data are presented as n (%). Percentage was calculated based on the number of respondents for each item.

p Values were calculated by logistic regression adjusted for sex, postgraduate experience and number of dying patients <18 years of age cared for.

ACP, advance care planning; DNAR, do not attempt resuscitation.
Table 3  Discussions about ADs at the end of life

| Number of paediatricians who held discussions | Neurologists (n=186) | Haematologists (n=227) | p Value |
|----------------------------------------------|----------------------|------------------------|---------|
| **Adolescent patients**                      |                      |                        |         |
| Cardiopulmonary resuscitation                |                      |                        |         |
| Use of:                                      |                      |                        |         |
| Ventilator                                   | 99 (57)              | 70 (31)                | <0.01   |
| Vasopressor                                  | 73 (42)              | 54 (24)                | <0.01   |
| Antibiotics                                  | 62 (36)              | 47 (21)                | <0.01   |
| Tube-feeding                                 | 88 (51)              | 90 (40)                | <0.01   |
| Intravenous fluids                           | 74 (43)              | 74 (33)                | 0.03    |
| **Family**                                   |                      |                        |         |
| Cardiopulmonary resuscitation                | 162 (93)             | 215 (99)               | 0.10    |
| Use of:                                      |                      |                        |         |
| Ventilator                                   | 170 (97)             | 213 (98)               | 0.60    |
| Vasopressor                                  | 156 (89)             | 192 (88)               | 0.34    |
| Antibiotics                                  | 131 (76)             | 115 (53)               | <0.01   |
| Tube-feeding                                 | 145 (83)             | 160 (74)               | 0.10    |
| Intravenous fluids                           | 131 (75)             | 140 (65)               | 0.04    |

Data are presented as n (%). Percentage was calculated based on the number of respondents for each item. p Values were calculated by logistic regression adjusted for sex, postgraduate experience and number of dying patients <18 years of age cared for.

AD, advance directives.

Discussion about ADs at the end of life

Discussions between physicians and patients and their families regarding ADs at the end of life are summarised in table 3. More than half of the neurologists discussed the use of ventilators (57%) and tube-feeding (51%) with their patients. In addition, paediatric neurologists discussed all the AD topics more often with the patients’ families than with the patients themselves. Overall, neurologists were significantly more likely than haematologists to discuss all the AD topics with their patients (table 3).

Barriers to ACP

Table 4 lists the factors that physicians identified as often or always acting as barriers to ACP. More than 50% of neurologists reported that unrealistic family expectations (58%), differences in the understanding of the prognosis between the physician and family (54%), differences in the understanding of the prognosis between the physician and patient (53%), lack of training (53%) and insufficient laws and guidelines (50%) were barriers to ACP. In general, neurologists perceived more barriers than haematologists. There were significant differences between neurologists and haematologists regarding eight barriers to ACP, namely insufficient laws and guidelines, inability of family/patient to fully comprehend the issue, uncertainty about the prognosis, lack of physician time, lack of a relationship between the physician and the family/patient, and physicians not placing much importance on ACP (table 4).

DISCUSSION

To the best of our knowledge, the present study is the first study to investigate current ACP and AD practices among...
paediatric neurologists with regard to adolescent patients with life-threatening conditions.

The most important finding of the present study was that only about half or less of the paediatric neurologists discussed ACP and ADs with their adolescent patients who had life-threatening conditions, even if the patient’s prognosis was <3 months. They tended to discuss ACP and ADs more with the families than with their patients. In fact, if the patient’s prognosis was <3 months, only about 30% of paediatric neurologists reported having discussions, such as DNAR orders and ventilator use, with their patients, whereas over 70% did discuss these topics with the patients’ families. About half of the paediatric neurologists did not discuss ADs at the end of life with their patients, whereas over 75% did discuss ADs with the patients’ families. We reported previously

### Table 4 Barriers to ACP

| Number of paediatricians who answered ‘often’ or ‘always’ for each question | Neurologists (n=186) | Haematologists (n=227) | p Value |
|---|---|---|---|
| Unrealistic family expectations | | | |
| Differences in understanding the prognosis | | | |
| Between physician and family | 94 (54) | 94 (42) | 0.07 |
| Between physician and patient | 93 (53) | 103 (46) | 0.37 |
| Lack of training | 93 (53) | 112 (51) | 0.91 |
| Insufficient laws and guidelines | | | |
| Physician concerned about taking away patient’s hope | 85 (49) | 94 (43) | 0.75 |
| Family not ready to have the discussion | 86 (49) | 86 (39) | 0.97 |
| Unrealistic patient expectations | 87 (49) | 80 (36) | 0.28 |
| Ethical considerations | 85 (49) | 85 (38) | 0.24 |
| Family unable to fully comprehend the issue | 87 (49) | 69 (32) | <0.01 |
| Differences in understanding the prognosis between patients and family | 82 (47) | 87 (39) | 0.17 |
| Patient unable to fully comprehend the issue | | | |
| Patient not ready to have the discussion | 83 (47) | 93 (42) | 0.79 |
| Physician does not know the right time to address the issue | 76 (43) | 96 (43) | 0.33 |
| Physician concerned about taking away family’s hope | 67 (38) | 67 (31) | 0.70 |
| Physician uncertain about the prognosis | 63 (36) | 58 (26) | 0.04 |
| Physician does not know the things to say | 60 (34) | 59 (27) | 0.68 |
| Conflict between parents and other family members | 58 (33) | 58 (27) | 0.95 |
| Physician not ready to have the discussion | 56 (32) | 59 (27) | 0.92 |
| Conflict between patient and parents | 52 (30) | 54 (25) | 0.79 |
| Lack of physician time | 49 (28) | 40 (18) | 0.04 |
| No relationship between physician and family | 47 (27) | 24 (11) | <0.01 |
| No relationship between physician and patient | 42 (24) | 23 (11) | <0.01 |
| Physician concerned about losing the trust of the patient | 34 (19) | 33 (15) | 0.46029 |
| Physician concerned about losing the trust of the family | 33 (19) | 30 (14) | 0.57 |
| Physician considers ACP not that important | 27 (17) | 17 (8) | 0.01 |
| Unrealistic physician expectations | 27 (15) | 20 (9) | 0.49 |
| Physician concerned that the family will receive less attention from physicians | 27 (15) | 23 (10) | 0.40 |
| Physician concerned that the patient will receive less attention from physicians | 23 (13) | 22 (10) | 0.45 |

Data are presented as n (%). Percentage was calculated based on the number of respondents for each item. p Values were calculated by logistic regression adjusted for sex, postgraduate experience and number of dying patients <18 years of age cared for.

ACP, advance care planning.
that haematologists, both paediatricians and internists, tended to discuss ACP and ADs more with the families than with the adolescent patients themselves. These findings show that discussions about ACP and ADs with adolescent patients with life-threatening conditions are less frequent with all physicians, regardless of the type of clinical practice.

The second important finding of the present study was that paediatric neurologists were more likely than haematologists to have end-of-life discussions, such as prognosis and ADs with patients. In fact, if the patient's prognosis was <3 months, about 50% of paediatric neurologists reported they discussed prognosis with their adolescent patients, whereas only 30% of haematologists discussed it. A possible explanation for this is due to the difference in illness trajectories between neurological diseases and haematological malignancies. Patients with life-threatening neurological illnesses are more likely to experience sudden and critical changes in their disease condition due to respiratory failure, cognitive impairment and motor paralysis in the course of illness. Therefore, neurologists may be more accustomed to having end-of-life discussions than haematologists. Conversely, neurologists were less likely than haematologists to have more basic end-of-life discussions with patients, such as current medical condition and treatment and care goals. In contrast to cancer, neurological illnesses have a longer and more variable time course, and it is often hard to determine exactly when a patient is entering the terminal stages of life. Therefore, neurologists may feel less familiar having basic discussions with patients than haematologists.

The third important finding of the present study was the identification of barriers to ACP from the perspective of paediatric neurologists who had experience in taking care of adolescents with life-threatening conditions who had decision-making capacity. The ranking of barriers to ACP was almost the same between paediatric neurologists and haematologists, but paediatric neurologists felt more barriers to ACP than haematologists. Family factors were one of the most common barriers for paediatric neurologists as well as haematologists. In addition, patient factors were also identified as barriers to ACP. Neurologists felt more barriers than haematologists about patients being unable to comprehend the issues. This may reflect the fact that patients with neurological illness often have decreasing cognitive impairment during the course of disease progression. Similar to haematologists, a lack of training and the concern of taking away hope from patients were physician factors that acted as major barriers to ACP for neurologists. Providing opportunities for training about end-of-life communication may reduce barriers to ACP and improve for practice of ACP discussion.

The present study had several limitations. First, the response rate was not high. The motivation for either agreeing or refusing to participate in the study may be biased by previous experiences of the study topics, so a higher response rate is likely to make the proportion of neurologists reporting ACP and AD discussions smaller. Second, there would be the potential for recall bias. Third, the generalisation of the findings of the present study may be limited because the ACP process may be affected by cultural differences.

CONCLUSION
The findings of the present study indicate that about half or less of paediatric neurologists discussed ACP and ADs with their adolescent patients with life-threatening conditions, even if the patient's prognosis was <3 months. They tended to discuss ACP and ADs more with families than with the patients themselves. Compared with paediatric haematologists, paediatric neurologists were more likely to have end-of-life discussions with their patients.

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