Relationships between various attitudes towards self-determination in health care with special reference to an advance directive

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Abstract

Objectives—The subject of patient self-determination in health care has gained broad interest because of the increasing number of incompetent patients. In an attempt to solve the problems related to doctors’ decision making in such circumstances, advance directives have been developed. The purpose of this study was to examine relationships between public attitudes towards patient autonomy and advance directives.

Subjects and main outcome measures—A stratified random sample of 600 adults in northern Sweden was surveyed by a questionnaire with a response rate of 78.2%. The subjects were asked about their wish for control of their health care, their concerns about health care, their treatment preferences in a life-threatening situation (both reversible and irreversible), and their attitudes towards the application of advance directives.

Results—Numerous relationships between various aspects of self-determination in health care (desire for control, fears of over-treatment, and choice of treatment level) in general and advance directives, in particular, were found. Those who wanted to have a say in their health care (about 94%) also mainly supported the use of an advance directive.

Conclusions—The fact that almost 30% of the respondents were undecided concerning their personal use of advance directives points to a lack of knowledge and to the necessity of education of the public on these issues.

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Keywords: Self-determination in health care; general population survey; advance directive; patient education

Objectives

In industrialised Western countries the proportion of elderly people among the total population is constantly increasing. Simultaneously, the number of elderly among the patients in the health care system is growing. One of the concurrent problems of this development which health care professionals are encountering is the increasing proportion of incompetent patients, for example those who are not able to communicate their treatment preferences.4–3 Another group, who present similar problems, and who are also increasing in numbers, are patients suffering from AIDS.

Today, more than ever before, elderly people express fears of an overzealous use of life-sustaining procedures when they are severely ill; procedures which would just prolong their suffering and compromise their dignity and quality of life.4–5 These concerns have been manifested both in general population6–9 and patient surveys.10–11 These investigations revealed the public’s considerable and widespread interest in self-determination in health care.

One way of promoting patients’ self-determination is by the use of so-called advance directives, which allow individuals to express and document their treatment preferences at a time when they are competent, and to inform others (i.e., health care professionals) how they would like to be treated in case of incompetency.4–7, 9, 12–13 Such documents could reduce conflicts in the doctor’s decision-making process.5–14 Results from the above-mentioned population surveys clearly indicate the willingness of the public to use advance directives. A survey of 405 outpatients at the Massachusetts General Hospital in Boston, USA revealed that 93% desired an advance directive; similar results (92.3%) were obtained from a 909-subjects sample in Canada by Molloy and co-workers.9 Among our representative general population sample in northern Sweden almost 80% regarded it as extremely or very important to document in an advance directive the level of care they would like to get in the event of an acute life-threatening illness.6
These unequivocal results urge both health policy makers and health care professionals to prepare the way for increased patient self-determination and for the implementation of advance directives.

The aim of the present study was to analyze the relationships between various attitudes towards self-determination concerning medical treatment in general and towards advance directives, in particular. These might constitute a basis for identifying crucial points for the education of the public in these issues.

Subjects

GENERAL POPULATION SURVEY

A stratified random sample of 600 adults, 20 years of age or older was obtained from the registration office of the province of Västerbotten in northern Sweden. The subjects were representative according to age-decades and gender. During May 1996 they were mailed a questionnaire and a covering letter, in which the voluntariness of participation was emphasised. Those who did not return the questionnaire received a first reminder after three weeks. After another four weeks a second reminder was sent to the non-responders. Altogether 472 subjects returned the questionnaire, a response rate of 78.2%.

The project had previously been approved by the ethical committee at Umeå University.

Measurement

QUESTIONNAIRE

The questionnaire, originally developed by Molloy and co-workers from the Geriatric Research Group at McMaster University, Canada, was translated into Swedish according to established guidelines including appropriate use of independent back translations. In the first part of the questionnaire biographical data were ascertained (age, sex, civil status, occupational status, chronic disease). Subsequently, the subjects were asked about: a) their wish for control of their personal health care, b) attitudes towards current health care practice, c) their preferences for treatment in a life-threatening situation, both where it was reversible and where it was irreversible, and d) their attitudes towards the application of advance directives. An example of an advance directive form was presented at the end of the questionnaire. The respondents were asked to reply either on a five-point scale: extremely, very, somewhat, slightly, not important/concerned, or on a three-point scale: yes, no, undecided, respectively. Aims concerns treatment preferences in case of reversible or irreversible, acute life-threatening condition, four options were given: a) comfort measures only - pain relief, intravenous medications, no tests, b) limited efforts - pain relief, intravenous medications, tests, but no surgery, c) moderately aggressive treatment - no intensive care unit, no ventilator, but surgery, if indicated, and d) very aggressive treatment - intensive care unit and ventilator, if necessary.

The statistical analysis was performed by means of frequency tables and $\chi^2$-square tests. X-square was used as a test of statistical significance which is done in order to determine whether a systematic relationship exists between two variables.

Results

SAMPLE CHARACTERISTICS

The mean age of the subjects was 51.3±17.8 years and they were equally distributed within seven age decades, except the group of 50 to 59 years old, and the group of 80 to 89 years old (Table 1). The sexes were represented by 50% each. Further details are given elsewhere.6

IMPORTANCE OF CONTROL

The questionnaire comprised two questions regarding control of their own health care, one as a general nature and a second focusing on advance directives (level of care in the event of an acute life-threatening illness). The results showed that desire for control was significantly related to the following aspects: a) concerns about being subjected to tests and procedures without prior discussion and consent ($\chi^2 = 36.35; 65.02; p < 0.003; p < 0.001$); b) concerns about not being treated aggressively enough at hospital ($\chi^2 = 26.83; 46.95; p = 0.043; p < 0.001$); c) wish to discuss treatment options with the doctor ($\chi^2 = 57.01; 29.49; p < 0.001; p = 0.003$); d) wish that document desired level of care in the event of an acute life-threatening condition ($\chi^2 = 61.36; 96.83; p < 0.001; p < 0.001$); e) regarding that presented directive in the questionnaire as useful ($\chi^2 = 18.92; 42.05; p = 0.015; p < 0.001$); f) wish to use the presented directive ($\chi^2 = 14.34; 55.61$);

<table>
<thead>
<tr>
<th>years</th>
<th>20-29</th>
<th>30-39</th>
<th>40-49</th>
<th>50-59</th>
<th>60-69</th>
<th>70-79</th>
<th>80-89</th>
<th>total</th>
</tr>
</thead>
<tbody>
<tr>
<td>females</td>
<td>36/52.2</td>
<td>35/48.6</td>
<td>30/46.2</td>
<td>51/54.3</td>
<td>39/48.1</td>
<td>38/48.1</td>
<td>6/50.0</td>
<td>235/49.8</td>
</tr>
<tr>
<td>males</td>
<td>33/47.8</td>
<td>37/51.4</td>
<td>35/53.8</td>
<td>43/45.7</td>
<td>42/51.9</td>
<td>41/51.9</td>
<td>6/50.0</td>
<td>237/50.2</td>
</tr>
<tr>
<td>total</td>
<td>69/14.6</td>
<td>72/15.3</td>
<td>65/13.8</td>
<td>94/19.9</td>
<td>81/17.2</td>
<td>79/16.7</td>
<td>12/25.0</td>
<td>472</td>
</tr>
</tbody>
</table>

The questionnaire, developed by Molloy and colleagues from the Geriatric Research Group at McMaster University, Canada, was translated into Swedish according to established guidelines, including appropriate use of independent back translations. In the first part of the questionnaire, biographical data were ascertained (age, sex, civil status, occupational status, chronic disease). Subsequently, the subjects were asked about: a) their wish for control of their personal health care, b) attitudes towards current health care practice, c) their preferences for treatment in a life-threatening situation, both where it was reversible and where it was irreversible, and d) their attitudes towards the application of advance directives. An example of an advance directive form was presented at the end of the questionnaire.


Table 2 Evaluation of the usefulness of presented directive and the wishes for using it (in %)

<table>
<thead>
<tr>
<th>Question</th>
<th>yes</th>
<th>no</th>
<th>don't know</th>
</tr>
</thead>
<tbody>
<tr>
<td>Do you think this directive is useful?</td>
<td>49.5</td>
<td>5.9</td>
<td>44.5</td>
</tr>
<tr>
<td>Would you use this directive for directing the level of care provided to you in the event of reversible/irreversible life-threatening illness?</td>
<td>59.2</td>
<td>11.6</td>
<td>29.2</td>
</tr>
</tbody>
</table>

p = 0.073; p < 0.001), and g) the opinion of the respondents, as to which group of individuals the presented directive should be used by (χ² = 33.54; 50.31; p = 0.029; p < 0.001). With all of these associations the more important the subjects felt control over their own health care to be, the more pronounced were their wishes regarding self-determination, and the more frequently they expressed the opinion that such a directive should be used by everybody.

Furthermore, the perceived importance of desire for control by an advance directive was alone related with concerns about being treated too aggressively (χ² = 36.20; p = 0.003), indicating a specific association between fear of overzealous treatment and the importance of advance directives.

CONCERNS

All three sources of fear: of treatment and diagnosis without prior discussion and consent - χ² = 22.15, p = 0.005; of treatment that was not aggressive enough - χ² = 27.65, p < 0.001; and/or of treatment in hospital that was too aggressive - χ² = 17.62, p = 0.024, were related to the wish to document the desired level of care. In other words, the more fear the subjects expressed, the more they wanted to direct their level of care. Specifically, the subjects who were most concerned about overtreatment were the ones who tended to decide to use the presented directive (χ² = 14.51; p = 0.069).

TREATMENT LEVEL

The expressed wish for participation in the decision-making process (wish to discuss hospital treatment and investigation issues with the doctor and wish to document desired level of care) were related to regarding the directive as useful (χ² = 19.23; χ² = 82.56; p = 0.023; p < 0.001, respectively) and to the willingness to use this directive (χ² = 24.49; χ² = 92.29; p < 0.001; p < 0.001, respectively).

Those subjects who favoured a documentation of their desired level of care also suggested to a larger extent than other subjects that everyone should use this kind of advance directive (χ² = 57.35; p < 0.001).

The selected treatment level in the event of both an acute reversible and an irreversible life-threatening illness was associated with the perceived importance of having a say in one’s own health care decisions (χ² = 39.06; χ² = 36.05; p = 0.001; p = 0.003, respectively) and with the perceived importance of having an advance directive under such circumstances (χ² = 25.52; χ² = 32.91; p = 0.061; p = 0.008, respectively).

USE OF DIRECTIVE

About 50% of the respondents thought the presented directive could be useful, only 5.9% evaluated the directive as not useful, but 44.5% were unable to decide (table 2). Nearly two-thirds of the subjects wanted to use this directive, 11.6% did not want to use it, (table 2) and about two-thirds indicated that everyone should use this directive (table 3). Most of the respondents expressed the opinion that this directive ought to be reviewed and updated every 12 months (table 4).

Those individuals who considered the presented directive as useful would like to use it for themselves (χ² = 279.14; p < 0.001). Furthermore, both those who declared the directive to be useful and those who would use it advocated its general use (χ² = 125.15; χ² = 164.77; p < 0.001; p < 0.001, respectively) and the updating of an advance directive every 12 months (χ² = 28.04; χ² = 32.49; p < 0.001; p = 0.002, respectively).

On the other hand, those encountering difficulties in understanding this directive were less positive and more reluctant concerning its usefulness (χ² = 43.34; p < 0.001) and its personal use (χ² = 32.96; p < 0.001). Therefore, they were also more sceptical about a general dissemination of advance directives (χ² = 44.40; p = 0.001).

CONFOUNDING VARIABLES

When testing for confounding variables, for example, age, gender and presence of chronic ill-

Table 3 Who should use the presented directive? (in %)

<table>
<thead>
<tr>
<th>users</th>
<th>healthy</th>
<th>ill</th>
<th>chronically ill</th>
<th>acutely ill</th>
<th>everyone</th>
<th>nobody</th>
<th>others</th>
</tr>
</thead>
<tbody>
<tr>
<td>n in %</td>
<td>11.7</td>
<td>6.0</td>
<td>5.3</td>
<td>61.1</td>
<td>12.9</td>
<td>3.0</td>
<td></td>
</tr>
</tbody>
</table>

Table 4 Expressed opinion about the frequency of updating the advance directive

<table>
<thead>
<tr>
<th>months</th>
<th>3</th>
<th>6</th>
<th>9</th>
<th>12</th>
<th>not at all</th>
<th>others</th>
</tr>
</thead>
<tbody>
<tr>
<td>n in %</td>
<td>9.9</td>
<td>15.8</td>
<td>17</td>
<td>44.7</td>
<td>8.3</td>
<td>19.6</td>
</tr>
</tbody>
</table>
ness, the following few relationships occurred. Age was inversely correlated to choice of treatment level as regards reversible illness ($\chi^2 = 40.38; p < 0.010$) and positively correlated to concerns about being treated too aggressively ($\chi^2 = 51.85; p < 0.001$). Regarding gender differences, females attached more importance to having control in health care decisions ($\chi^2 = 22.46; p < 0.001$) and expressed generally more concerns ($\chi^2$ from 10.60 to 11.50; $p$ from 0.032 to 0.022) about their health care compared to their male counterparts.

Conclusions
The results confirm the correctness of our original assumption that it is necessary to study public attitudes towards self-determination in health care; and this is underscored by the high response rate obtained in our general population survey.

Numerous relationships between various aspects of self-determination with regard to medical treatment in general and advance directives in particular could be established. We found desire for control highly related to fears concerning: treatment and diagnosis without prior discussion; undertreatment, and overtreatment. Consequently, the wish for self-determination was associated with support for advance directives. These relationships became most pronounced with regards to the fear of being too aggressively treated. Those who wanted to have a say in their health care (about 94%) also predominantly expressed a wish to use the presented advance directive; only 11.6% of the subjects rejected it. The more important self-determination in health care was perceived to be, the more a lower level of treatment was selected. The great majority wish to use an advance directive to indicate their future choice of treatment level. Nevertheless, almost 30% of the respondents were undecided concerning their personal use of the presented directive. In addition, our findings of a relationship between difficulties among the respondents in understanding the directive and their rejection of it, and their general unfamiliarity with what an advance directive is, point to the necessity of educating the general public in the field of patient autonomy.

In ethics, autonomy implies acting with intention, with understanding and without controlling influences. All these characteristics underlining the individual desire for information. The finding of more concern about being treated too aggressively among the elderly age groups indicates a relation between older peoples' views on technology and the principle of beneficence in terms of ensuring their quality of life after treatment. Furthermore, the fear of overtreatment and possible prolonged suffering and pain during the dying process is related to the principle of non-maleficence, ie, the ethical requirement to do no harm.

Considering these findings, the following issues should be addressed as part of the process of educating the general public:

a) clarification of the treatment possibilities of modern medicine;

b) information about doctors' responsibilities concerning medical treatment according to law and medical ethics;

c) information about doctors' ethical and legal conflicts in medical decision making and their acceptance and support of advance directives;

d) information about patients' rights and the legal basis of self-determination in health care; and

e) information about advance directives, and education in completing such forms, covering consideration of the available options and the implications of patients' decisions.

In order to ensure the success of such large-scale educational measures mechanisms have to be found for their implementation. The family physician seems most suited to be the key person in this, which in turn requires that this group of health care professionals be prepared for this important and challenging task.

Promising steps have been taken in Canada by Molloy and his Geriatric Research Group at McMaster University in Hamilton, who have developed an advance health care directive and educational material which seem to be feasible, practical and well supported both by various patient groups and by health care professionals.

Martin Eisemann, PhD, is Associate Professor at the Unit of Medical Psychology, Department of Psychiatry and WHO Collaborating Centre, Umeå University, Umeå/Sweden and Jörg Richter, PhD, Associate Professor at the Psychiatric University Clinic, Rostock University, Rostock, Germany.

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News and notes

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Special attention will be paid to European traditions in health care ethics. All lectures and plenary sessions will be held in English.

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