Patients’ Willingness to Share Information in Online Patient Communities: Questionnaire Study

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Abstract

Background: Online patient communities provide new channels for users to access and share medical information. In-depth study of users’ willingness to share information in online patient communities is of great significance for improving the level of information sharing among the patient community and the long-term development of communities.

Objective: The aim of this study was to build a model of factors affecting patients’ willingness to share medical information from the perspective of both positive and negative utilities. Specifically, we aimed to determine the influence of online information support and privacy concerns, as well as the moderating effect of disease severity and information sensitivity of different patients on their willingness to share.

Methods: Data from 490 users with experience in online patient communities were collected through a questionnaire survey, and structural equations were applied to empirically verify the model hypotheses.

Results: Privacy concerns negatively affected the patients’ willingness to share information ($P<.001$), whereas online information support positively affected patients’ willingness to share information ($P<.001$), and information sensitivity negatively moderated the impact of online information support on sharing willingness ($P=.01$). Disease severity positively moderated the impact of privacy concerns on sharing willingness ($P=.05$). However, the hypotheses that information sensitivity is a negative moderator and disease severity is a positive moderator of the impact of privacy concerns on sharing willingness could not be supported.

Conclusions: To improve the level of user information sharing, the online patient community should design a safe user registration process, ensure the confidentiality of information, reduce the privacy concerns of users, and accurately identify the information needs of patients to provide personalized support services.

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KEYWORDS
online patient community; information sharing; willingness to share; questionnaire; structural equation

Introduction

Background

Social media is increasingly being used by health services to encourage the growth of social support networks among people with health-related issues [1,2]. Through participation on different platforms, patients are able to provide and receive social support in various ways, without the typical barriers and constraints experienced by this population [3]. In this way, problems of information asymmetry, as well as the shortages and uneven distribution of medical resources can be alleviated [4,5]. Jackson et al [6] studied the characteristics of patients who shared medical records with others, and found that participants who shared medical records were more likely to experience better health management. To obtain information support from other patients in the community, users need to share information on personal medical treatments, medication records, or medical diagnosis. However, disclosing such
information may also have serious adverse consequences [7].
In the face of risky decisions about online information demands
and privacy disclosure, patients may choose whether to share
such personal information according to their judgment on the
possible benefits and losses. Therefore, in-depth study of users’
information-sharing willingness and the decision-making
mechanisms of online patient communities is of great
significance to the improvement of information sharing among
patient communities and the long-term development of these
communities.

Most of the existing research on patient information sharing
focuses on the impacts of sharing willingness from the
perspective of the value brought by online communities [8-10].
However, users usually consider both the gains and risks
associated with information sharing when making decisions
about whether they should share information [11]. In this
context, we here establish a model of factors affecting patients’
willingship to share medical information combined with utility
theory, which considers both the influence of online information
support (OIS) and privacy concerns on patients’ willingness to
share information (WSI) from the perspectives of both the
positive and negative utilities brought by information sharing
to users. The information-sharing decisions of online community
users depend not only on the community environment (ie,
whether the community provides OIS and whether the
community environment can dispel users’ privacy concerns)
but also on the personal attributes of each community user (ie,
health status and sensitivity to information). If patients are not
sensitive to the positive utility of information sharing but are
sensitive to the negative utilities (ie, afraid of the risks), the
probability that they will choose not to share information is
high, whereas if patients are not sensitive to the risks of
information sharing but are very sensitive to the positive utilities,
the possibility of sharing information is high. Therefore, the
main contribution of this paper is to introduce patients’
information sensitivity and health status into the model as
moderating variables to explore the moderating effect of such
individual patient attributes on information-sharing intention.

**Literature Review and Hypotheses**

**Information-Sharing Behaviors in Online Patient
Communities**

Information sharing refers to the process during which
individuals share the information they own with other members
in the community in various forms and routes [12]. In most
cases, members of online patient communities are patients
troubled by different diseases [13]; therefore, they have
relatively strong empathy. Their similar situations enable them
to easily obtain both the emotional and information support they
need while relating to the feelings of other members of the
community during the process of communication [14].
Combined with the features of online patient communities,
shared information can be classified into general health
information and special health information. General health
information refers to common medical and health information,
which is usually open to the public and can be easily obtained
by the masses, such as information on doctors and hospitals,
prices of medicines, and features of a specific disease. Special
health information is usually concerned with privacy, which
refers to personal empirical information, including medical
records, treatment, and personal health conditions [15].

**Utility Theory**

Utility theory is often adopted when individuals make decisions
[16]. Utility is defined as the degree to which a user’s
psychological expectation of a certain factor is satisfied or
realized [17]. This judgment is completely controlled by
personal preferences, interests, or the subjective consciousness:
the higher the satisfaction degree, the larger the utility, and vice
versa [18]. The criterion is maximizing utility by weighing costs
and benefits [18], and is considered the ultimate expression of
reason and egotism, reflecting the decision maker’s attitude
toward risks. In the utility theory formula $U_p=f(B,C)$, $C$ refers
to the potential risk of privacy disclosure caused by information
sharing, which is a variable indicating negative utility to users;
$B$ refers to the positive utility obtained by users, which mainly
includes the OIS that patients receive; and $D$ refers to the
decision of users on whether or not to participate in information
sharing after judging the positive and negative utilities brought
by this behavior.

Information sharing creates value for users but also invites risks.
Different individuals have different sensitivities to personal
medical health information; such difference in sensitivity could
result in different concerns regarding information sharing,
thereby affecting their WSI. In addition, different patients have
different physical health conditions. If the disease of a patient
is severe, the patient may be urgently seeking OIS, which could
weaken relative concerns regarding privacy. In this case, the
patient is more likely to acquire the necessary information by
positively participating in information exchange in communities.
Based on the above analysis, we established an influence model
of OIS and privacy concerns for patients’ WSI (Figure 1). The
moderating functions of patients’ information sensitivity and
disease severity on the WSI were also considered. Specifically,
we consider two problems in this model structure: (1) the
influence of OIS and privacy concerns on the willingness to
share medical information, and (2) the functions of information
sensitivity and disease severity as moderating variables.

In this model, the information-sharing intention of patients is
the dependent variable, privacy concerns and OIS are
independent variables, and information sensitivity and disease
severity are moderator variables. That is, privacy concerns and
OIS are the main factors affecting patients’ intentions to share
medical information, the effects of which are regulated by
disease severity and information sensitivity. Moderator variables
with increasing negative effects (ie, negative moderator
influence factors) affect sharing intentions, whereas moderator
variables with decreasing negative effects (ie, positive moderator
influence factors) also affect sharing willingness. The control
variables include gender, age, and education level. Along with
the different social roles and characteristics between genders,
feelings and behaviors can differ according to gender regarding
the same issue [19]. As a result, gender may influence
information-sharing intentions. Age may also have effects on
information-sharing intentions since age influences physical
condition and life experience, which can result in quite different
opinions and feelings about the same issue [20]. With respect to education level, higher education levels are associated with stronger judgment and cognitive competence; thus, behavioral patterns may differ across this characteristic as well [21] to ultimately affect the information-sharing intentions of online health community users.

**Figure 1.** Model of information-sharing intentions of patients in online health communities.

**Research Hypotheses**

Based on the above theoretical model, six hypotheses are proposed. As shown in **Figure 2**, the hypotheses are composed of two parts: hypotheses of direct effects and hypotheses of moderation effects.

**Figure 2.** Research hypotheses on online information-sharing willingness.

**Hypotheses of Direct Effects**

**Relationship Between Online Information Support and Patients’ Willingness to Share Information**

In an online patient community, patients conduct information sharing to acquire medical assistance and personalized experience information [22]. In a study of two online communities, Hargreaves et al [23] found that patients share experiences, information, and emotions, and receive empathetic support in a supportive and warm community atmosphere. Based on the theory of planned behavior and the theory of technology acceptance, Zhou [24] found that social returns and perceived usefulness can determine the intentions to share personal health information. On the basis of a technology acceptance model, Zhang [25] found that users’ usefulness of acquiring perceptual information from virtual communities can affect their information-sharing behaviors. On the basis of social exchange theory, Gui and Hu [26] found that mutual benefits can significantly affect intentions to share health knowledge by conducting empirical analyses. Thus, research employing various theoretical perspectives has demonstrated that OIS has positive impacts on the information-sharing intentions of patients; in other words, support or help from other community members strengthens the intention to share information.

Thus, the following hypothesis is presented:

**H1: OIS has positive impacts on patients’ WSI**.
Relationship Between Privacy Concerns and Patients’ Willingness to Share

As electronic health records become ubiquitous in the health care industry, privacy breaches are increasing [27]. In an online patient community, patients can predict but not fully control these potential risks, thereby reducing their WSI [28]. Privacy concerns are one of the major factors affecting attitudes toward exchanging electronic health information [29,30]. The privacy of patients is prominent in health information exchange discussions, given that their potentially sensitive personal health information may be electronically shared for various health care purposes [31]. Based on a grounded theory, Holloway et al [32] found that issues of privacy and confidentiality were paramount to information sharing. In an empirical study, Zhang et al [33] found that privacy concerns have obvious negative impacts on the attitudes of online patient community users regarding information disclosure. The main objectives of different methods of health information exchange are to reduce privacy concerns [34]. Frost et al [35] studied the information-sharing preferences of cancer patients in the community and found that poor online privacy experience, age, and health status negatively affected the WSI.

Thus, the following hypothesis is presented:

H2: Privacy concerns have negative impacts on patients’ WSI.

Hypotheses of Moderation Effects

Information sensitivity, which is closely related to privacy concerns, refers to the extent of concern of individuals regarding information in a specific environment: the greater the extent of concern, the more sensitive the information [36]. Medical information also includes personal information that is not directly related to personal health, such as population statistics (ie, name and date of birth) and information used to identify a patient’s personal identity (ie, ID card number, family address, private phone number, financial information, and insurance status). In addition, such information includes details that may help in making a correct diagnosis and adopting proper treatment measures, such as the cause for seeking medical advice, symptoms, medical history of family members, drug allergy history, and medical history, as well as disease data arising from diagnosis and treatment, such as examination results, progress notes, operation consent, anesthesia consent, operation records, and anesthesia records. Since most of this information involves personal privacy, online sharing causes concern regarding privacy disclosure: the more sensitive the information, the greater the privacy concerns.

Many people believe that if sensitive information is shared or disclosed, it will have bad consequences or even cause damage [37]. Consumers’ intentions to disclose personal information are closely related to the sensitivity of such information [38]. Milne [39] found that the validity of information changes with the sensitivity of such information when collecting personal information from users. According to Okazaki et al [40], information sensitivity increases privacy concerns. Information sensitivity is frequently used as the prepositive variable of privacy concerns [41]. In a network environment, privacy concerns cause a reduction in the publishing of information by online users, whereas information sensitivity causes concerns regarding private information [42]. In this study, the information sensitivity of patients was used as a moderator variable to explain the behavioral intentions of patients regarding sharing information in online patient communities. The users’ privacy concerns during information sharing will be more serious with respect to more sensitive medical information, thereby affecting sharing behavioral decisions [43].

Thus, the following hypotheses are presented:

H3: Information sensitivity negatively moderates the impact of privacy concerns on patients’ WSI.

H4: Information sensitivity negatively moderates the impact of OIS on patients’ WSI.

Disease severity refers to the users’ perception of their own health status; that is, a subjective perception of whether one’s health is good or bad will affect their information-sharing intentions. Tinsado et al [44] studied the impacts of demographic characteristics and individual health on the concordance between medical records and health information sharing. Patients who considered their health to be poor were more sensitive about sharing their personal health information compared with others; these patients worried that they may face uncontrollable risks if their medical and health information were to be disclosed [45]. Health conditions as perceived by online patient community users translate into privacy concerns regarding their medical and health information, which ultimately affect their intentions to share this information [46]. In online patient communities, the priority of patients is to improve their health, interact with other community users, and seek medical assistance. The levels of participation and activeness in the community also vary according to diversified health conditions. When they are physically healthy, they need not consult doctors or tell others about their medical treatment experiences; considering the privacy of medical information, their intentions to share relevant medical and health information may be reduced. When they are physically unhealthy, they urgently need to acquire the information support and assistance provided by online communities to improve their health [47]. However, the precondition is that to receive this assistance, they will need to illustrate their health condition to relevant helpers. Therefore, they have to actively share medical information, and thus their intentions to share relevant information increase. Consequently, the disease severity perceived by online patient community users may affect their intentions in sharing personal medical information by regulating their privacy concerns and OIS demands. In this study, disease severity was used as a moderator variable to moderate the relationship between privacy concerns and OIS demand intensity when both sharing and intending to share health and medical information so as to verify the impact on sharing behavioral patterns.

Thus, the following hypotheses are presented:

H5: Disease severity positively moderates the effect of privacy concerns on patients’ WSI.

H6: Disease severity positively moderates the effect of OIS on patients’ WSI.
Methods

Questionnaire Design
The questionnaire was composed of three parts: Part 1 reflects the experience of the respondents in sharing medical information with any online health community (if such experience is available), Part 2 reflects the objective cognition of the respondents regarding online medical services, and Part 3 mainly concerns the respondents’ basic information and medical history.

The medical information sharing intentions, disease severity, and perceived health information sensitivity were measured according to the method of Bansal et al [48], and privacy concerns and OIS were measured according to the method of Liang et al [49]. A typical 7-point Likert scale was used to score opinions and conclusions, with dimensions of “quite disagree,” “disagree,” “basically disagree,” “neutral,” “basically agree,” “agree,” and “quite agree”, respectively. The specific scale is shown in Table 1.

| Component and measurement number | Scale |
|----------------------------------|-------|
| **Willingness to share (WSI)**   |       |
| WSI1                             | I am willing to provide my medical health information to online medical websites to get proper treatment. |
| WSI2                             | I agree to let online medical websites use my medical health information. |
| WSI3                             | I do not think it is improper if my medical health information is revealed to online medical websites. |
| WSI4                             | It is highly probable that I will publish my medical health information on some online medical websites. |
| **Privacy concerns (PC)**        |       |
| PC1                              | I worry that my medical health information posted online may be misused. |
| PC2                              | I worry that other people can see my medical health information online. |
| PC3                              | In my opinion, personal medical health information is more important than other information. |
| PC4                              | I worry that my medical health information posted online will be used by others in an unexpected manner. |
| **Disease severity (DS)**        |       |
| DS1                              | My body is seldom subject to long-term pain and discomfort. |
| DS2                              | I have no chronic disease. |
| DS3                              | My overall health is good. |
| **Online information support (OIS)** |       |
| OIS1                             | When I need assistance related to health issues, anyone from an online medical website can provide relevant advice. |
| OIS2                             | When I encounter any health problem, anyone from an online medical website can provide relevant information to help me overcome the problem. |
| OIS3                             | Anyone from an online medical website will help me find the cause and provide relevant suggestions for any health problem I might have. |
| **Information sensitivity (IS)** |       |
| IS1                              | Medicine taken |
| IS2                              | Current health condition |
| IS3                              | Medical history |
| IS4                              | Process of seeking medical advice |

Data Collection
To guarantee the validity of the questionnaire design and content, many experts and some respondents were invited to evaluate each question in the questionnaire. An internal test was conducted within a small range before issuing these questionnaires. After confirming that there were no errors, the questionnaires were issued through wjx.cn, an online platform for issuing questionnaires. Those who had experience sharing information in online health communities were invited to fill in these questionnaires. A total of 531 copies were collected, 41 of which were incomplete, unreasonable, or unqualified, leaving a total of 490 valid responses.
## Data Analysis

The statistical description of the correlated variables is shown in [Table 2](#).

### Table 2. Statistical description of variables.

| Variables                          | Sample, n (%) |
|------------------------------------|---------------|
| **Gender**                         |               |
| Men                                | 168 (34.3)    |
| Women                              | 322 (65.7)    |
| **Age (years)**                    |               |
| ≤20                                | 3 (0.6)       |
| 21-30                              | 216 (44.1)    |
| 31-40                              | 243 (49.6)    |
| 41-50                              | 28 (6)        |
| 51-60                              | 0 (0)         |
| >60                                | 0 (0)         |
| **Education level**                |               |
| Middle school and below            | 0 (0)         |
| High school                        | 10 (2)        |
| College                            | 43 (9)        |
| Bachelor                           | 357 (72.9)    |
| Master                             | 68 (14)       |
| Doctorate and above                | 12 (3)        |
| **Duration of using the internet (years)** |         |
| ≤1                                 | 0 (0)         |
| 2-4                                | 40 (8)        |
| 5-7                                | 123 (25.1)    |
| 8-10                               | 171 (34.9)    |
| >10                                | 156 (31.8)    |
| **Duration of using online medical websites (years)** |         |
| ≤1                                 | 93 (19)       |
| 2-3                                | 278 (56.7)    |
| 4-5                                | 101 (20.6)    |
| 6-7                                | 15 (3)        |
| >7                                 | 3 (0.6)       |
| **Annual frequency of getting ill** |           |
| ≤1                                 | 88 (18)       |
| 2-3                                | 291 (59.4)    |
| 4-6                                | 93 (19)       |
| 7-10                               | 15 (3)        |
| >10                                | 3 (0.6)       |
**Results**

**Measurement Model Testing**

**Reliability Test**

The inspection criteria for the reliability and validity tests were the load levels of the corresponding components, common factor loading, Cronbach alpha, composite reliability, and average variance extracted (AVE). The desired values for these indices [50] are shown in Table 3.

The measurement results for the data factor loading coefficients are shown in Table 4.

The factor loading measurements <0.7 in the questionnaire were deleted, with those remaining reaching the criteria. The results of the modified Smart PLS data analysis are shown in Table 5.

As shown in Table 5, Cronbach alpha for each component was larger than 0.77, indicating that the reliability for each question in the research design was good. The composite reliability coefficient of each component was larger than 0.78, the AVE for each component was greater than 0.57, and all components could be aggregated effectively.

| Evaluation index                         | Critical value |
|------------------------------------------|----------------|
| Factor loading coefficient               | >0.70          |
| Cronbach alpha                           | >0.70          |
| Composite reliability coefficient         | >0.70          |
| Average variance extracted                | >0.50          |

Table 3. Desired values for the reliability test.

| Component and scale | Factor loading coefficient |
|---------------------|---------------------------|
| Privacy concerns (PC) |                           |
| PC1                 | 0.934                     |
| PC2                 | 0.922                     |
| PC3                 | 0.767                     |
| PC4                 | 0.938                     |
| Online information support (OIS)       |                           |
| OIS1                | 0.836                     |
| OIS2                | 0.848                     |
| OIS3                | 0.808                     |
| Information sensitivity (IS)           |                           |
| IS1                 | 0.671                     |
| IS2                 | 0.951                     |
| IS3                 | 0.850                     |
| IS4                 | 0.756                     |
| Disease severity (DS)                  |                           |
| DS1                 | 0.717                     |
| DS2                 | 0.981                     |
| DS3                 | 0.776                     |
Table 5. Reliability of test results.

| Variable | AVE<sup>a</sup> | Composite reliability | $R^2$ | Cronbach alpha | Communality | Redundancy |
|----------|-----------------|-----------------------|------|---------------|-------------|------------|
| Age      | 1               | 1                     | N/A<sup>b</sup> | 1             | 1           | N/A        |
| DS<sup>c</sup> | 0.574         | 0.785                  | N/A  | 0.854         | 0.574       | N/A        |
| EDU<sup>d</sup> | 1             | 1                     | N/A  | 1             | 1           | N/A        |
| Gender   | 1               | 1                     | N/A  | 1             | 1           | N/A        |
| IS<sup>e</sup> | 0.631         | 0.869                  | N/A  | 0.846         | 0.631       | N/A        |
| OIS<sup>f</sup> | 0.690         | 0.870                  | N/A  | 0.775         | 0.690       | N/A        |
| PC<sup>g</sup> | 0.731         | 0.913                  | N/A  | 0.889         | 0.731       | N/A        |
| WSI<sup>h</sup> | 0.654         | 0.883                  | 0.331| 0.823         | 0.654       | 0.039      |

<sup>a</sup>AVE: average variance extracted.  
<sup>b</sup>N/A: not applicable. 
<sup>c</sup>DS: disease severity. 
<sup>d</sup>EDU: education level. 
<sup>e</sup>IS: information sensitivity. 
<sup>f</sup>OIS: online information support. 
<sup>g</sup>PC: privacy concerns. 
<sup>h</sup>WSI: willingness to share.

Validity Test

In general, two measurement indices are used for the validity test for each component: the discriminant validity test requires that the correlation coefficient between two components be less than its AVE value, and the convergent validity test requires that the load value at each proposed measured variable be larger than that loaded at any other measured variable in cross-factor loading. Table 6 shows the correlation coefficients used for measuring discriminant validity; diagonal elements contain the arithmetic square root of AVE, and nondiagonal elements contain the correlation coefficient for each corresponding component. Based on these correlations, the data have favorable discriminant validity by satisfying relevant requirements. Table 7 further shows that the factor loading loaded at the corresponding component was larger than that loaded at any other latent variable, demonstrating quite good convergent validity between two components.

Table 6. Correlation coefficients.

| Variable | Age | DS<sup>a</sup> | EDU<sup>b</sup> | Gender | IS<sup>c</sup> | OIS<sup>d</sup> | PC<sup>e</sup> | WSI<sup>f</sup> |
|----------|-----|---------------|----------------|--------|--------------|----------------|---------------|---------------|
| Age      | 1   | —             | —              | —      | —            | —              | —             | —             |
| DS       | −0.120 | 1           | —              | —      | —            | —              | —             | —             |
| EDU      | −0.204 | −0.085      | 1              | —      | —            | —              | —             | —             |
| Gender   | −0.044 | −0.021      | −0.100         | 1      | —            | —              | —             | —             |
| IS       | −0.158 | −0.038      | 0.093          | 0.0351 | 1            | —              | —             | —             |
| OIS      | 0.123  | 0.209       | −0.116         | 0.032  | 0.132        | 1              | —             | —             |
| PC       | −0.179 | 0.079       | 0.092          | −0.041 | 0.466        | 0.083          | 1             | —             |
| WSI      | 0.254  | 0.158       | −0.009         | 0.043  | −0.129       | 0.436          | −0.267        | 1             |

<sup>a</sup>DS: disease severity.  
<sup>b</sup>EDU: education level.  
<sup>c</sup>IS: information sensitivity.  
<sup>d</sup>OIS: online information support.  
<sup>e</sup>PC: privacy concerns.  
<sup>f</sup>WSI: willingness to share.  
<sup>g</sup>Not applicable.
### Table 7. Cross-factor loadings.

| Variable | Age | DS<sup>a</sup> | EDU<sup>b</sup> | Gender | IS<sup>c</sup> | OIS<sup>d</sup> | PC<sup>e</sup> | WSI<sup>f</sup> |
|----------|-----|----------------|----------------|--------|--------------|--------------|----------|------------|
| Age      | 1   | -0.120         | -0.204         | -0.044 | -0.158       | 0.123        | -0.179   | 0.254      |
| DS1      | -0.164 | 0.410          | -0.031         | 0.029  | 0.149        | 0.136        | 0.308    | -0.037     |
| DS2      | -0.137 | 0.980          | -0.089         | -0.009 | -0.009       | 0.214        | 0.123    | 0.144      |
| DS3      | -0.161 | 0.771          | -0.018         | -0.031 | 0.064        | 0.189        | 0.237    | 0.020      |
| EDU      | -0.204 | -0.085         | 1              | -0.100 | 0.093        | -0.116       | 0.092    | -0.009     |
| Gender   | -0.044 | -0.021         | -0.100         | 1      | 0.035        | 0.032        | -0.041   | 0.043      |
| IS1      | -0.090 | -0.071         | 0.080          | -0.017 | 0.570        | -0.008       | 0.243    | 0.009      |
| IS2      | -0.152 | -0.059         | 0.049          | -0.017 | 0.950        | 0.080        | 0.380    | -0.152     |
| IS3      | -0.138 | 0.015          | 0.153          | 0.117  | 0.851        | 0.217        | 0.487    | -0.063     |
| IS4      | -0.109 | -0.029         | 0.104          | 0.064  | 0.757        | 0.090        | 0.430    | -0.047     |
| OIS1     | 0.080  | 0.101          | -0.108         | 0.086  | 0.088        | 0.835        | 0.047    | 0.369      |
| OIS2     | 0.090  | 0.222          | -0.047         | 0.069  | 0.074        | 0.848        | 0.026    | 0.358      |
| OIS3     | 0.137  | 0.199          | -0.132         | -0.076 | 0.168        | 0.809        | 0.133    | 0.361      |
| PC1      | -0.100 | 0.056          | 0.094          | -0.054 | 0.447        | 0.102        | 0.934    | -0.221     |
| PC2      | -0.243 | 0.147          | 0.044          | -0.103 | 0.420        | 0.070        | 0.922    | -0.271     |
| PC3      | -0.125 | 0.092          | -0.005         | 0.015  | 0.285        | 0.202        | 0.570    | 0.003      |
| PC4      | -0.143 | 0.011          | 0.124          | 0.047  | 0.438        | 0.065        | 0.937    | -0.247     |
| WSI1     | 0.179  | 0.115          | -0.009         | 0.110  | 0.041        | 0.516        | -0.048   | 0.733      |
| WSI2     | 0.184  | 0.185          | -0.076         | 0.050  | -0.185       | 0.293        | -0.245   | 0.837      |
| WSI3     | 0.260  | 0.161          | 0.025          | -0.016 | -0.163       | 0.268        | -0.283   | 0.820      |
| WSI4     | 0.195  | 0.053          | 0.025          | -0.009 | -0.123       | 0.315        | -0.297   | 0.841      |

<sup>a</sup>DS: disease severity.
<sup>b</sup>EDU: education level.
<sup>c</sup>IS: information sensitivity.
<sup>d</sup>OIS: online information support.
<sup>e</sup>PC: privacy concerns.
<sup>f</sup>WSI: willingness to share

### Results of Structural Model Testing

After fully verifying the reliability and validity of the questionnaire, we next analyzed the structural model to test the various proposed hypotheses based on the following values: (1) the path coefficient between each variable, (2) the significance of the $t$ value, and (3) the degree of $R^2$ to which the dependent variable is interpreted. In line with the research model, significance verification was conducted in two steps: the first step assessed the significance of the influencing factors among the major components (namely, verification of the basic model), and the second step examined the effect of the regulated variable added to the model. The results are shown in Table 8.
by combining the features of users’ information sharing in online patient communities, this study adopted utility theory to probe users’ information-sharing decisions under the condition that both privacy concerns and information demands exist from the user’s perspective. In addition, based on the characteristics of shared information, we further analyzed the information sensitivity and the adjustment effect of the patients’ own perceived health status on their WSI. Under the combined effect of the above factors, assessment of users’ WSI in this manner expands the application of utility theory to studies on information-sharing behaviors, thereby laying the foundation for further development of utility theory.

This research also has practical significance. In view of the mechanism of users’ WSI, online service operators should establish a reasonable mechanism to ensure the security and confidentiality of users’ medical information through reasonable and safe user registration processes and appropriate protection measures, thereby minimizing users’ privacy concerns during the process. Moreover, the patients’ health status information should be obtained to accurately identify individual information requirements, so that the platform can offer personalized support services while providing appropriate guidance and incentives, thereby improving the information-sharing level of community users.

Limitations
As with all empirical studies, this study also has limitations. First, online patient communities comprise a large group of users distributed in various cities across the country. Although the data of this study were obtained from the Internet, the sample size is negligible compared with the enormous group of community users. In addition, the sample data were all obtained from China (without considering a potential Western cultural background), which may affect the conclusions of the study to a certain extent. Second, the subject of information sharing in online patient communities is necessarily “human,” and thus the sharing behavior strategy is inclined to be affected by multiple factors. Moreover, with constant changes in other users’ sharing behaviors, future studies could focus more on the dynamic adjustment of users’ sharing strategies, thereby providing a deeper understanding of the entire sharing process.

Discussion
Principal Findings
This study showed that privacy concerns have a negative impact on users’ WSI, and the OIS provided by online communities has a positive impact on the users’ WSI. The hypothesis that information sensitivity negatively regulates the impact of privacy concerns on WSI was not supported. The indirect impact of information sensitivity on WSI mainly occurs through negative regulation of the OIS offered by communities. This phenomenon may be caused by the platform’s privacy protection mechanism. In most cases, community platforms include certain user privacy protection measures to protect users’ personal data from being leaked. When patients participate in online communication, most of their personal information is blocked and invisible to others. Despite the involvement of relatively sensitive information, users can be free from concerns regarding disclosure of their personal identity, which greatly diminishes the effect of information sensitivity on privacy concerns. Although the privacy protection measures of the community platform dispel the inhibition effect of privacy concerns caused by patients’ information sensitivity on WSI to some degree, information sensitivity still negatively regulates the positive effect of OIS on WSI. If patients are in poor health and need to obtain information from other patients in the community, their privacy concerns will be reduced to a certain extent. The mental inhibition factors of information sharing are reduced, while the weakened privacy concerns further increase the willingness to participate in information sharing. In summary, these results suggest that a community platform should make all efforts to desensitize the information shared by users while providing effective information support for users so as to reduce the negative effects of information sharing, increase the benefits of information sharing, and improve the overall information-sharing level of community users.

Theoretical and Practical Significance
The information-sharing intention model established in this study enriches and improves theoretical knowledge in the following aspects. Primarily, utility theory was applied to study users’ WSI in online patient communities. Previous studies on users’ willingness to share have mainly focused on aspects of motivation, mutual benefits, social exchange, and the like. Yet,

Table 8. Verification of the basic model.

| Hypothesis                                                                 | $t$ value | $P$ value | Supported |
|---------------------------------------------------------------------------|-----------|-----------|-----------|
| H1: online information support has positive impacts on patients’ WSI      | 5.77      | <.001     | Yes       |
| H2: privacy concerns have negative impacts on patients’ WSI              | 3.41      | <.001     | Yes       |
| H3: information sensitivity negatively moderates the impact of privacy    | 0.96      | .34       | No        |
| concerns on patients’ WSI                                                |           |           |           |
| H4: information sensitivity negatively moderates the impact of online     | 2.58      | .01       | Yes       |
| information support on patients’ WSI                                     |           |           |           |
| H5: disease severity positively moderates the effect of privacy concerns  | 1.96      | .05       | Yes       |
| on patients’ WSI                                                         |           |           |           |
| H6: disease severity positively moderates the effect of online information| 0.68      | .50       | No        |
| support on patients’ WSI                                                 |           |           |           |
Conclusions
This study established a model to probe the factors influencing decision making on patients’ WSI from the perspectives of both positive and negative utilities. The results confirm the positive and negative effects of OIS and privacy concerns on users’ WSI. In addition, we showed that information sensitivity negatively moderates the impact of OIS on sharing willingness, whereas disease severity positively moderates the impact of privacy concerns on sharing willingness. Therefore, the management of the platform should focus on desensitizing the information shared by users, while providing effective information support for users to reduce the negative effects of information sharing, increase the benefits of information sharing, and improve the information-sharing level of community users. This research can help community platform operators to clarify the behavioral mechanism of users’ independent disclosure of medical information, provide theoretical guidance for operators to stipulate effective measures that encourage users to voluntarily disclose medical information, and offer a reference to improve the operators’ user management of online patient community services.

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Conflicts of Interest
None declared.

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Abbreviations

AVE: average variance extracted
DS: disease severity
EDU: education level
IS: information sensitivity
OIS: online information support
PC: privacy concerns
WSI: willingness to share information

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