BACKGROUND

Coordinating health care across different care levels poses a challenge for health systems around the world. This is particularly true for health systems in which primary health care acts as a gate-keeper and coordinator of patient care throughout the health care continuum. Rapid technological advances, increasing specialization and new ways of organising services mean that a growing number of professionals and services are involved in the health care of patients, thus...
jeopardizing its coordination; a problem which particularly affects patients with chronic conditions and multi-morbidities (1, 2). Faced with this scenario, health systems should adopt models of care provision that foster collaboration across different levels of care in order to improve care continuity, efficiency, and particularly the quality of care and health of patients (2, 3). In the public health system of Catalonia, one of the Spanish regions, as in any system based on primary care, key factors for operational effectiveness include: exchanging information; communicating fluidly across different levels of care; and making agreements between the professionals involved in the clinical management of patients, including their follow-up and appropriate access to services across different health care levels. Despite the extraordinary increase in the number of publications on care coordination in the last decade, the lack of consensus on definitions among disciplines, such as primary care, mental health, and disease management, still remains (4, 5). Many of them are limited to particular patient populations, settings, transitions or types of coordination. The broad conceptual framework adopted in this study (6) defines clinical coordination, according to Longest and Young (7), as the harmonious connection of different health services needed to provide care to a patient throughout the care continuum in order to achieve a common objective without conflicts. Following Reid et al (8), two different interrelated types of clinical coordination are distinguished (9): firstly, the coordination of clinical information, which refers to the exchange of patients’ clinical information to harmonize care activities between providers, consists of the transfer of clinical information and its use; and secondly, the coordination of clinical management, which refers to the provision of care in a sequential and complementary manner by the different services and levels of care involved, consisting of the coherence of care, patient follow-up, and accessibility across different levels of care. Care coordination refers to health care services and can be analysed through service-based indicators or by taking into account the views of health personnel using qualitative methods, such as in-depth interviews, or quantitative methods, such as surveys (10). In contrast, continuity of care refers to how patients experience the coordination of services received, which can be analysed only from the users’ perspective (8).

Despite the relevance attributed to care coordination across different levels of care, few studies adopt a comprehensive approach in order to include the different types and dimensions of coordination, different transitions between care levels and the general patient population (11). Most studies that measured care coordination from the perspective of health professionals focused on health care coordination for a particular type of patient or care coordination (12–14), the evaluation of a specific care coordination mechanism (15–17), or a level of care, mainly primary health care (18). In Catalonia, previous research has analysed clinical coordination in health care networks by exploring the patients’ perceptions of continuity across different care levels (19, 20) and measuring the degree of clinical coordination using service-based indicators (9, 19). However, factors influencing clinical coordination or the experiences of doctors were rarely analysed (21–23).

To the best of our knowledge (11, 24), the only comprehensive instrument to measure clinical coordination across care levels from the perspective of doctors is the COORDENA questionnaire, which was first developed and applied in six Latin American countries in 2015 (25), based on the same theoretical framework of Vázquez et al (6). It consists of three main parts: a) doctors’ experiences of clinical information and clinical management coordination across care levels and the general perception of doctors of the degree of coordination in their health care network; b) doctors’ knowledge and use of clinical coordination mechanisms across levels of care; and c) the factors that potentially influence care coordination (available on: www.equity-la.eu). The objective of this study is to adapt and validate an online version of the COORDENA questionnaire and its method of application for use in the public health system of Catalonia.

METHODS

The COORDENA questionnaire was adapted to the context of the public health system in Catalonia in two stages. In the first stage, the language and contents were revised and updated on the basis of a literature review, previous qualitative research results, meetings with experts and two pre-tests. In the second stage, the pre-tested online version was piloted (Fig. 1).

STAGE 1: ADAPTATION OF THE CONTENTS OF THE QUESTIONNAIRE

In order to adapt the questionnaire to the context of the public health system in Catalonia1, two steps were taken: a revision and update of contents (face or content validity) and language; and two pre-tests.

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1 Spain has a decentralised national health system. The health competences were devolved to the 17 regions (autonomous communities). Therefore, they may differ in the way they organize and deliver health services. Adapting the questionnaire to the context in Catalonia meant to consider what specific coordination mechanisms do exist in the health services of Catalonia and which organizational factors could be of relevance.
REVISION OF CONTENTS (FACE OR CONTENT VALIDITY) AND LANGUAGE
A literature review was conducted to identify new studies that analysed clinical coordination across levels of care and associated factors – such as organisational, interactional, and work-related attitudes – as well as instruments that were available to measure care coordination. The results of qualitative studies on care coordination conducted in Catalonia and elsewhere (21–23) were also taken into consideration. The COORDENA questionnaire was translated into Catalan, terms were culturally adapted to the context of the health system in Catalonia, and it was then translated back into Spanish. New questions were included related to coordination mechanisms available in the health care networks of the Catalan public health system as well as additional factors potentially associated with coordination that were relevant to the context, as identified in the literature review. Some questions relating to the influencing factors were also removed. The preliminary draft of the questionnaire was discussed in two sessions with an expert group to assess face validity and a first draft was developed for pre-testing. The expert group consisted of: members of the Health Care Integration Evaluation Group, or GAIA, which consists of health services professionals who are involved in health services research or quality evaluation processes and thus have a good knowledge of the subject and the context; and researchers who created the COORDENA questionnaire.

PRE-TESTS
Two pre-tests were conducted in order to evaluate: firstly, comprehensiveness, understanding, acceptability, sequence of themes and questions, and the length of the questionnaire; and secondly, the functioning of the online version. The first pre-test was carried out through face-to-face cognitive interviews with primary care and secondary acute and long-term care doctors. The selection of doctors was based on the following survey inclusion criteria: doctors had worked for at least one year in the health care organization, doctors provided direct care to patients, and whose daily practice involved contact with doctors from other care levels through, for example, the patient referral process. In the first pre-test, eight doctors participated: three from primary care, two from acute secondary care and three from long-term secondary care.

Based on the results of the first pre-test, changes were made to the questionnaire, followed by the development of the online version in both Catalan and Spanish. Its functioning was tested first by seven members of the research team and then by six doctors, three from primary care and three from secondary care, in the three health care areas of Baix Empordà, Osona and Alt Empordà. A number of improvements were subsequently made to the online version of the instrument.

STAGE 2: PILOT STUDY
A pilot study was conducted in order to test the newly adapted version, COORDENA-CAT, and the feasibility of the online survey under real conditions.

STUDY AREA
The study area was the network of health services within the Catalan public health system located in the Southern Metropolitan Area of Barcelona and comprised: 19 primary care teams of the Servei d’atenció primària Delta del Llobregat; one acute hospital, Hospital de Viladecans; and one long-term care hospital, Hestia Duran i Reynals. The primary care teams and the acute hospital were managed by the same public entity, the Institut Català de la Salut, and the long-term care hospital was managed by a private entity, Hestia Alliance.

STUDY POPULATION
The study population consisted of primary care and secondary acute and long-term care doctors that had worked for at least one year in a centre of the network, provided direct care to patients, and whose daily practice involved contact with doctors from other care levels through, for example, the patient referral process.

DATA COLLECTION PROCESS
Data collection was programmed to take place over two weeks. On day one, each health care provider sent, to all doctors working in their respective institution, an email containing a personal invitation to participate and a link to access the online questionnaire. Each link was unique and randomly generated, allowing doctors to respond anonymously. Doctors could access the questionnaire at different times at their convenience and continue answering at the point where they had left off, as the previously filled in answers were automatically saved. All answers were automatically registered in an Excel database to which only the coordinator had access. After a week, a second email was sent to all invited doctors to encourage them to participate or to thank them for their participation if they had already answered.
STRATEGIES FOR ENCOURAGING PARTICIPATION

A number of strategies were used to boost participation levels both before and during the survey:

- **Information sessions** for managers of participating health care centres, to publicize the project, explain the methodology and encourage participation, commenced two weeks before the beginning of the pilot survey.

- **Posters**. A poster was designed to explain the survey’s objective and online procedure, emphasizing how important it was for doctors to express their opinions in order to develop a realistic picture of care coordination in...
their organization and identify elements for improvement. The posters were displayed one week before the beginning of the survey in spaces commonly used by doctors, such as meeting rooms, libraries, and canteens, in all participating centres.

- **News published on the intranet.** A short article explaining the project was published on the corporate intranet of each participating centre one week before the beginning of the survey.

- **Participation follow-up.** Over the survey’s two-week period, the coordinator monitored the response rate of each centre. In those primary care centres with a low rate, specific actions to encourage participation were taken, such as extra reminders by email.

**DATA ANALYSIS**

In order to evaluate the way in which the survey was conducted, the following variables were analysed, both globally and for each health care level: number of responses per day, rate of access to the questionnaire, global response rate, and proportion of doctors who fully completed the questionnaire. To assess the performance of the COORDENA-CAT questionnaire, a descriptive univariate analysis was first conducted, in order to identify questions with a high no-response rate or low response variability. Secondly, a correlation analysis of questions on the same construct was performed, to identify questions that provided little added value. Finally, the open-ended question on difficulties in answering the questionnaire was analysed. All the analyses were performed using Excel.

**ETHICAL CONSIDERATIONS**

Approval for the study was granted by the ethics committee of Parc de Salut Mar and Bellvitge Hospital. Participation in the study was voluntary. All participants read and granted informed consent before gaining access to the questionnaire and were permitted to withdraw at any moment. The researchers had no access to any personal data of participating doctors. Anonymity was guaranteed by randomly assigning a code to each participant, not collecting names, and having an aggregated analysis of the data.

**RESULTS**

**ADAPTATION OF THE QUESTIONNAIRE**

Regarding the contents of the questionnaire, following the literature review and expert group discussion and consensus, the most important change was the addition of questions addressing the existing clinical coordination mechanisms in the health service networks of the Catalan public health system. Furthermore, according to their contextual relevance, certain questions referring to potentially associated factors were removed, others were reworded to make more sense in the context, and some new ones were added (Fig. 1). Regarding face or content validity, the expert group found that both types of clinical coordination across levels of care – information and clinical management coordination, and their dimensions and attributes – were represented in the questionnaire. In addition, all existing mechanisms for care coordination across care levels and potentially influencing factors were included in the questionnaire.

The first pre-test showed a relatively good understanding of the questions, an adequate sequence of themes and questions, and an adequate length of approximately 15 minutes. However, a need was identified to make some modifications, including: rewording some questions to make them easier to understand, such as those regarding the shared clinical history in the area; simplifying by fusing two questions into one; adding response categories; and refining the instructions for some sections. The second pre-test, of the adapted online version, confirmed a better understanding of the revised questions and identified a few more elements requiring refinement, such as: the information given in the consent form; the layout; and problems in its online functioning, such as the lack of filters and a progress indicator, and spelling mistakes (Fig. 1).

The final version of the COORDENA-CAT questionnaire consists of seven sections (Box 1) and is very similar to the original questionnaire. Changes in contents were introduced in sections three, four, five and seven. In section three, one question was added on doctors establishing a patient care plan together. The fourth and fifth sections refer to the knowledge of doctors and their use of clinical coordination mechanisms across different health care levels. The fourth section now has an additional question regarding the perceived usefulness of the mechanisms and adds four mechanisms to the original questionnaire: shared clinical history of Catalonia, shared clinical history of the network, virtual consultations through the clinical history, and case managers. The fifth section adds two mechanisms: shared clinical records and virtual consultations through the clinical record. The seventh section now groups all questions related to factors that potentially influence clinical coordination, in contrast to their having been in different sections in the original questionnaire. Furthermore, some questions were added or removed in this section including: for organizational factors, two added and two removed; for interactional factors, two added and
one removed; for job-related attitudes, two removed; and for employment factors, one removed. Questions on demographic characteristics did not change.

**Box 1. Contents of the Coorden-a-Cat questionnaire**

1. Informed consent
2. General information: experience in the health care network
   - Level and type of care
3. Experience of coordination between levels of care (16 items)
   - Coordination of clinical information (exchange, use and needed information)
   - Coordination of clinical management
     - Care coherence: related to treatment, diagnostic tests and shared care plans
     - Follow-up across levels of care: (back) referrals, recommendations, and consultations
     - Accessibility across levels of care: waiting times when (back) referred
   - Perception of coordination across levels of care
4. Coordination mechanisms between levels of care in your centre
   - Knowledge, frequency of use, and opinion on usefulness
     - shared clinical history of Catalonia-HC
     - shared clinical history in the centre, joint clinical sessions, virtual consultations through the clinical history, e-mail, telephone, referral report, discharge report, shared protocols/clinical guidelines, case managers/liaison nurses
5. Characteristics of use of coordination mechanisms between levels of care
   - Available information, difficulties, reasons of use: shared clinical history, joint clinical sessions, virtual consultations through the clinical history, e-mail, telephone
6. Suggestions for the improvement of clinical coordination across care levels
7. Factors related to coordination across levels of care
   - Organizational, job related attitudes, interactional, employment conditions, demographic

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**EVALUATION OF THE METHOD AND QUESTIONNAIRE**

**EVALUATION OF THE METHOD**

Participation was irregular over the survey period, although this increased after having sent reminders to doctors. In the first three days, 36.4% of responding doctors accessed the questionnaire, with increases to 71.6% and 88.6%, respectively, after having sent the first and second reminders. A similar pattern was observed for both primary care and secondary care doctors, with higher levels for primary care doctors. With regard to response rate, all doctors of the participating centres were invited to participate in the survey, and of these, 36.9% accessed the questionnaire and 33.8% agreed to participate (Table 1). Only two doctors provided a reason for not participating, in that they did not fulfil the inclusion criteria. From those doctors who agreed to participate, 83.9% fully completed the questionnaire. Differences in participation were observed according to health care levels: while the proportion of invited primary care doctors who participated, 40.7%, was much higher than that for the invited secondary care doctors, 19.0%, the percentage of those who completed the questionnaire was high in both groups, 81.1% and 96.6%, respectively (Table 1).

**EVALUATION OF THE QUESTIONNAIRE**

**Sample characteristics**

Most (70.4%) of the participants were women. Almost half of the sample (45.4%) were between the ages of 41 and 54 and most (88.0%) were born in Spain. Most were primary care doctors (82.0%); a majority (63.8%) had over 16 years of experience working in the same organization; and most (71.9%) had a permanent contract (Table 2).

**Descriptive analysis of the questions**

The descriptive analysis showed, firstly, that none of the questions presented a high no-response rate, low response variability or unexpected responses (Table 3).

Moreover, answers were generally in line with what was theoretically expected. With regard to clinical information coordination across different levels, most doctors reported that: they usually shared information on the patients they have in common (64.0%); that shared information is necessary for the care of these patients (66.9%); and that they use it (81.5%).

With respect to clinical management coordination across levels and care consistency, most doctors reported that they usually
### TABLE 1. DOCTORS’ PARTICIPATION IN THE SURVEY ACCORDING TO HEALTH CARE LEVEL

<table>
<thead>
<tr>
<th></th>
<th>Invited doctors</th>
<th>Doctors who accessed the questionnaire*</th>
<th>Doctors who agreed to participate*</th>
<th>Doctors who fully completed the questionnaire**</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>N (%)</td>
<td>N (%)</td>
<td>N (%)</td>
</tr>
<tr>
<td>Primary care</td>
<td>324</td>
<td>140 (43.2%)</td>
<td>132 (40.7%)</td>
<td>107 (81.1%)</td>
</tr>
<tr>
<td>Secondary care</td>
<td>153</td>
<td>36 (23.5%)</td>
<td>29 (19.0%)</td>
<td>28 (96.6%)</td>
</tr>
<tr>
<td>Acute hospital</td>
<td>145</td>
<td>29 (20.0%)</td>
<td>23 (15.9%)</td>
<td>22 (95.7%)</td>
</tr>
<tr>
<td>Long-term care hospital</td>
<td>8</td>
<td>7 (87.5%)</td>
<td>6 (75%)</td>
<td>6 (100%)</td>
</tr>
<tr>
<td>Total</td>
<td>477</td>
<td>176 (36.9%)</td>
<td>161 (33.8%)</td>
<td>135 (83.9%)</td>
</tr>
</tbody>
</table>

*Calculated for the number of doctors invited
**Calculated for the number of doctors who accepted to participate

### TABLE 2. SAMPLE CHARACTERISTICS

<table>
<thead>
<tr>
<th>Variable</th>
<th>n</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sex</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(n=161)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Women</td>
<td>95</td>
<td>59.0</td>
</tr>
<tr>
<td>Men</td>
<td>40</td>
<td>24.8</td>
</tr>
<tr>
<td>Missing</td>
<td>26</td>
<td>16.1</td>
</tr>
<tr>
<td><strong>Age</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(n=161)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>30–40 years</td>
<td>27</td>
<td>16.8</td>
</tr>
<tr>
<td>41–54 years</td>
<td>59</td>
<td>36.6</td>
</tr>
<tr>
<td>54–65 years</td>
<td>44</td>
<td>27.3</td>
</tr>
<tr>
<td>Missing</td>
<td>31</td>
<td>19.3</td>
</tr>
<tr>
<td><strong>Country of birth</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(n=161)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spain</td>
<td>117</td>
<td>72.7</td>
</tr>
<tr>
<td>Other</td>
<td>16</td>
<td>9.9</td>
</tr>
<tr>
<td>Missing</td>
<td>28</td>
<td>17.4</td>
</tr>
<tr>
<td><strong>Health care level</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(n=161)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Primary care</td>
<td>132</td>
<td>82.0</td>
</tr>
<tr>
<td>Secondary care (SC)</td>
<td>29</td>
<td>18.0</td>
</tr>
<tr>
<td>- SC acute hospital</td>
<td>23</td>
<td>14.3</td>
</tr>
<tr>
<td>- SC long-term care hospital</td>
<td>6</td>
<td>3.7</td>
</tr>
<tr>
<td><strong>Experience in the organization</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(n=161)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 6 years</td>
<td>13</td>
<td>8.1</td>
</tr>
<tr>
<td>6–15 years</td>
<td>34</td>
<td>21.1</td>
</tr>
<tr>
<td>16–25 years</td>
<td>41</td>
<td>25.5</td>
</tr>
<tr>
<td>&gt; 25 years</td>
<td>42</td>
<td>26.1</td>
</tr>
<tr>
<td>Missing</td>
<td>31</td>
<td>19.3</td>
</tr>
<tr>
<td><strong>Type of contract</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(n=161)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Permanent</td>
<td>130</td>
<td>80.7</td>
</tr>
<tr>
<td>Temporary</td>
<td>5</td>
<td>3.1</td>
</tr>
<tr>
<td>Missing</td>
<td>26</td>
<td>16.1</td>
</tr>
</tbody>
</table>
agree with the treatments prescribed by doctors from another care level (72.8%); and considered that contraindications and/or duplication in the treatments prescribed are rare (59.3%); as is the repetition of tests already carried out at the other level of care (66.7%). However, most doctors (86.0%) reported that they rarely establish a treatment plan for patients together with other doctors, when needed. With regards to the follow-up of patients across levels of care, most doctors (90.8%) found that the referrals of primary care doctors to secondary care doctors were generally appropriate, as were the back referrals of secondary doctors (79.1%). However, some differences according to health care level were observed regarding patient follow-up and accessibility across levels of care (Table 3). In terms of the general perception of clinical coordination across care levels, most doctors (77.6%) found that patient care was not coordinated in their area, with small differences between primary and secondary care doctors (Table 3).

Analysis of correlations

The correlation analysis performed on questions addressing the same construct, or dimension, did not identify any strongly correlated questions. Hence, all questions provided added value and were therefore considered relevant for the independent analysis.

Difficulties in use of the questionnaire

The analysis of the open question on any difficulties encountered in answering the questionnaire revealed no relevant difficulty. Out of 96 doctors who expressed their opinion, half of them (51.0%) did not encounter any problems. Among those doctors who mentioned some kind of difficulty, the most frequently cited problems were the lack of time and the length of the questionnaire (12.4%), followed by the need to further qualify some answers but having no space in which to do so (5.0%).

DISCUSSION

Despite the fact that health care coordination is a health policy priority of health systems worldwide, to the best of our knowledge, the COORDENA questionnaire is the first tool to comprehensively evaluate health care coordination across different levels of care, taking into account the different types and dimensions of clinical coordination, different transitions, and the perspective of both primary and secondary care doctors, including a variety of specialties. It provides the perspective of one of the main actors, the doctors, and will be useful to complement other sources of information, such as indicators or the perspectives of patients. The online version was developed and piloted in the public health system of Catalonia following a systematic process, and in accordance with the conceptual framework, which guided all phases of the study. It has proven to be a valid instrument and method to evaluate clinical coordination across health care levels. Used periodically in the health system, it should allow us to identify and address problems of health care coordination across levels and their influencing factors in a particular area; serve as a benchmark across areas; and hence help to identify interventions to improve them. Moreover, changes in its contents are minimal compared to the original version (25) and these refer mainly to the inclusion of questions on the clinical coordination mechanisms existing in the health system and on influencing factors. This means that cross-country comparisons will be possible in order to analyse levels of achievement, and to identify contextual factors that might explain different results and require appropriate interventions.

Online surveys are easier to apply, faster, and less expensive than face-to-face surveys; however, they have a lower response rate, especially among doctors (26). Although the response rates of the COORDENA questionnaire, when applied by means of face-to-face interviews in Latin American health services networks, were significantly higher (approximately 90%) (10), the response rate achieved here (33.8%) was similar to another online survey of primary care doctors in Madrid (39.4%) (26), and higher than others (27). There were significant differences in the response rate between primary care doctors (40.7%) and secondary acute care doctors (15.9%) which is probably due to the different levels of involvement of their management teams. The primary care management team was actively involved and sent additional specific emails encouraging the participation of centres with low response rates. However, in the hospital, only reminders were sent with no further actions taken. With the aim of boosting participation, two suggestions emerged from the discussion around the pilot results with the primary care and hospital management teams: firstly, health managers from all levels could be more actively involved; and secondly, more face-to-face meetings could be programmed at all levels with the organisations participating in the survey.

Regarding the contents, the results on the doctors’ experiences with clinical coordination are generally in line with what was expected. For example, the relatively high level of information exchange can be attributed to measures taken to implement information coordination mechanisms, such as shared electronic medical records or virtual consultations (28). With respect to clinical management coordination across levels of care, doctors generally reported experiences of frequent
TABLE 3. EXPERIENCE OF THE DIFFERENT TYPES OF CLINICAL COORDINATION AND GENERAL PERCEPTION

<table>
<thead>
<tr>
<th>Health care level</th>
<th>Primary care</th>
<th>Secondary care</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>N=132</td>
<td>N=29</td>
<td>N=161</td>
<td></td>
</tr>
<tr>
<td>α (%)</td>
<td>α (%)</td>
<td>α (%)</td>
<td></td>
</tr>
</tbody>
</table>

### Experience of clinical coordination

#### Transfer and use of clinical information between levels
- **Primary and secondary care doctors share information on the care of patients we have in common (diagnosis, complementary tests, treatments) (n=136)**
  - Frequently: 71 (65.1)
  - Rarely: 38 (34.9)
  - Do not know/Do not answer: 0 (0)
- **The information we share is as required for the care of these patients (n=136)**
  - Frequently: 72 (66.1)
  - Rarely: 37 (33.9)
  - Do not know/Do not answer: 0 (0)
- **Primary and secondary care doctors use the information that we share (n=135)**
  - Frequently: 88 (71.5)
  - Rarely: 18 (14.7)
  - Do not know/Do not answer: 2 (1.5)

#### Health care consistency between levels
- **We agree with the treatments prescribed or directions given to the patients by doctors of the other level (n=136)**
  - Frequently: 79 (72.5)
  - Rarely: 27 (24.8)
  - Do not know/Do not answer: 3 (2.4)
- **There are contraindications and/or duplications in the treatments prescribed by primary and secondary care doctors (n=135)**
  - Frequently: 43 (39.8)
  - Rarely: 63 (58.3)
  - Do not know/Do not answer: 2 (1.9)
- **Primary and secondary care doctors establish a treatment plan together for patients that require this (n=136)**
  - Frequently: 13 (11.9)
  - Rarely: 96 (81.9)
  - Do not know/Do not answer: 2 (1.9)
- **We repeat the tests that doctors have already carried out at the other level (analysis, imaging) (n=135)**
  - Frequently: 33 (30.6)
  - Rarely: 74 (68.5)
  - Do not know/Do not answer: 1 (0.9)

#### Adequate health care follow-up between levels
- **Primary care doctors refer the patients to secondary care when appropriate (n=152)**
  - Frequently: 120 (97.6)
  - Rarely: 1 (0.8)
  - Do not know/Do not answer: 1 (0.8)
- **Secondary care doctors send the patients back to primary care for follow-up when appropriate (n=153)**
  - Frequently: 101 (81.5)
  - Rarely: 23 (18.5)
  - Do not know/Do not answer: 0 (0)
- **Secondary care doctors make recommendations to the primary care doctor on the follow-up of patients (diagnosis, treatment, other guidelines) (n=153)**
  - Frequently: 51 (41.1)
  - Rarely: 73 (58.9)
  - Do not know/Do not answer: 0 (0)
- **Primary care doctors clarify any doubts on the follow-up of patients with the secondary care doctors (n=153)**
  - Frequently: 61 (49.2)
  - Rarely: 61 (49.2)
  - Do not know/Do not answer: 2 (1.6)
- **Primary care doctors are informed when their patients are discharged from the hospital (n=150)**
  - Frequently: 72 (59.9)
  - Rarely: 47 (38.5)
  - Do not know/Do not answer: 3 (2.5)

#### Health care accessibility between levels
- **On being referred in the normal way to secondary care, the patient has to wait a long time to be seen (n=152)**
  - Frequently: 124 (100)
  - Rarely: 0 (0)
  - Do not know/Do not answer: 0 (0)
- **On being referred urgently to secondary care, the patient has to wait a long time to be seen (n=152)**
  - Frequently: 106 (85.5)
  - Rarely: 18 (14.5)
  - Do not know/Do not answer: 0 (0)
- **On being sent back to primary care, the patient has to wait a long time to be seen (n=150)**
  - Frequently: 42 (34.1)
  - Rarely: 81 (65.9)
  - Do not know/Do not answer: 0 (0)

### General perception of care coordination in the area
- **I think that in this area patient care is coordinated between primary and secondary care doctors (n=152)**
  - Frequently: 25 (20.2)
  - Rarely: 98 (79.7)
  - Do not know/Do not answer: 2 (1.5)

*Frequently: Always/Very often

**Rarely: Rarely/Never
coordination, with two exceptions: the joint establishment of patient care plans and accessibility across levels, both of which are consistent with current practice and available indicators (29). However, these are descriptive results and further analyses of experiences and opinion are needed that take into account potential influencing factors, such as the level of care.

One of the most relevant findings emerging from the results was the contrast between doctors’ generally positive experience of most attributes of clinical information and clinical management coordination across levels, and their general perception of limited coordination across care levels in their health care areas. While this gave rise to a number of potential explanations, its most important consequence was the modification of the questionnaire to include the additional open-ended question “Why?” following the item on perception in the final version of the questionnaire.

CONCLUSION

In conclusion, the adapted COORDENA-CAT questionnaire has proved to be a valid instrument for comprehensively measuring health care coordination across different health care levels in Catalonia, from the viewpoint of both primary and secondary care doctors. It can be applied by health providers and authorities to: identify coordination problems across levels of health care in a specific area of the health system; be used as a benchmark tool across areas; and, periodically, to monitor the performance of health services regarding clinical coordination across levels of care in order to address any emerging problems. The results can complement other sources, such as indicators, or perspectives, such as those of patients. By having retained most of the contents of the original questionnaire, it can also be used for comparisons across different health systems and countries. While its adaptation to different contexts is relatively easy, certain recommendations can be inferred from the results for its application in other contexts. Firstly, an appropriate adaptation of the language and contents of the questionnaire is required, for which a preliminary analysis of the existing mechanisms of coordination between levels of care in the networks/health system is useful. Secondly, to achieve a greater response rate, it is advisable to involve the management teams of participating centres, and to implement specific face-to-face actions in order to motivate doctors, especially those working in acute care hospitals, to participate. Lastly, the survey results should be used to give feedback to health professionals in order to involve them in the process of identifying problems and, more importantly, solutions.

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3 All references were accessed on 8 December 2018.


