Challenges in Achieving Collaboration in Clinical Practice: The Case of Norwegian Health Care

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Introduction:
This article summarizes and synthesizes the findings of four separate but inter-linked empirical projects which explored challenges of collaboration in the Norwegian health system from the perspectives of providers and patients. The results of the four projects are summarised in eight articles.

Methods:
The eight articles constituted our empirical material. Meta-ethnography was used as a method to integrate, translate, and synthesize the themes and concepts contained in the articles in order to understand how challenges related to collaboration impact on clinical work.

Results:
Providers’ collaboration across all contexts was hampered by organizational and individual factors, including, differences in professional power, knowledge bases, and professional culture. The lack of appropriate collaboration between providers impeded clinical work. Mental health service users experienced fragmented services leading to insecurity and frustration. The lack of collaboration resulted in inadequate rehabilitation services and lengthened the institutional stay for older patients.

Conclusion:
Focusing on the different perspectives and the inequality in power between patients and healthcare providers and between different providers might contribute to a better environment for achieving appropriate collaboration. Organizational systems need to be redesigned to better nurture collaborative relationships and information sharing and support integrated working between providers, health care professionals and patients.

Keywords:
integrated care; inter-professional collaboration; patient provider relationships; clinical work; Norway

Introduction
The health care system’s duty is to treat patients; i.e. clinical medical activity provided by various groups of healthcare professionals. In many countries, increasing numbers of older people and individuals living with chronic conditions in the community have created a need for better coordination and integration between different parts of the health system. In four separate but inter-linked empirical projects we explored the professional and organizational challenges to achieving collaboration in different health care settings in the Norwegian health system from the perspective of both providers and patients. In this article we explore the significance of these challenges for clinical work through a synthesis of the findings from across all four projects.

Clinical work
The basic activity of healthcare services is clinical work with individual patients and their next-of-kin. Clinical practice includes discretionary assessment and interpretation. A clinical conclusion is reached through logical reasoning based on, among other elements, clinical knowledge, i.e. the accumulated individual experience of work with patients. Communication is essential to delivering good quality medical care and a meaningful relationship between patient and provider is essential. Patients are all unique and individual patient needs are different. Providers need to ascertain an individual’s attitudes, values, and thoughts and give him or her the feeling of being understood. In addition to collaboration between patients and doctors, collaboration between providers is also required, especially when caring for older patients with complex problems and patients with chronic conditions,
for instance people with mental disorders and/or substance abusers.

**Collaboration**

Collaboration between providers can imply that providers from different specialities, disciplines or sectors work together. This includes a wide spectrum of activities — from simple electronically-conveyed messages or face-to-face encounters to comprehensive inter-professional, perhaps integrated, work. Integration implies a certain degree of collaboration among the parties who work together [1–3]. On a practical level it requires an effort to integrate and translate themes and schemes shared by different professional groups and the shared ownership of common goals, decision-making processes, and the integration of specialised professional knowledge and expertise. Barriers to overcome with regard to the successful integration between health care professionals include a blurring or misunderstanding of professional identities, roles and responsibilities [3–5]. A good knowledge of each other's work, a culture of mutual respect and recognition of each other's areas of expertise and competence, and the free and open exchange of information are also key elements. The significance of good relationships between providers and having sufficient time and resources for ongoing relationship-building has been well rehearsed in the literature [3]. Several empirical studies have also highlighted the key significance of effective organisational leadership, and appropriate funding arrangements for the achievement of successful collaboration and integration [3, 6, 7]. Singer et al. [8] recently addressed the concept of patient-centeredness in the provision of integrated care. In line with the clinical approach in the present study we also adopted a patient-centred perspective and included a focus on collaboration between patients and providers.

**The Norwegian healthcare system**

Health care in Norway is divided into two broad delivery systems: the specialist and the primary healthcare system, each of which is subject to different funding systems, laws, and central regulations. Four regional health enterprises owned by the state are responsible for the provision of hospital services. Hospitals are financed by a combination of block grants and activity-based financing, with hospital employees paid on the basis of a fixed salary. The 428 municipalities, which comprise the lowest governmental level, have responsibility for providing primary health care, long-term care services, home based care, and social care provision [9]. Health and social services are based on the classic Scandinavian Welfare model with financing and provision of universally accessible services to everyone. Nursing homes, home-based services and social care are public financed and mainly public provided but with an increasing part of private actors. As part of the Coordination Reform implemented in 2012 [10] specific responsibilities and resources are progressively being devolved from central to municipal government with the key aim of reducing hospital beds and widening access for health services within the municipality. Policies and programmes to achieve this aim are among other initiatives which established intermediate units between the two governmental levels, and the introduction of a daily penalty fee when municipalities are not able to receive those patients that are ready for discharge from hospital. Since 2001 all Norwegian citizens have been registered with a general practitioner (GP). GPs are in part paid by a capitation component depending on the number of patients on the list, and partly on the basis of fee-for-service.

As in other Western countries there have been a number of significant health care reforms in Norway over recent decades. The public sector modernisation, recommended by the Norwegian government, includes reorganisation inspired by core ideals in the New Public Management to achieve more cost effective solutions in care provision [11, 12]. These reforms involve among other factors a greater emphasis on measuring outcomes with the introduction of explicit standards, a range of performance metrics used to assess provider performance, and a greater degree of competition via the creation of quasi-market mechanisms. An example of this is the purchaser-provider split model which has a clear administrative distinction between those who assess the need for services and those who determine the scope of the services and provide the services. The contract contains detailed specifications from the purchaser and outcome control requires detailed reporting by the provider [11, 12]. The increasing professional differentiation and sub-division of healthcare services in Norway has been allied with attempts to improve collaboration and coordination between health care services. Since 2001 patients requiring coordinated services have had a statutory right to an individual care plan (IP). The latest initiative for improving the coordination of services is the Coordination Reform [10].

There is a vast literature on the range of challenges in organisational and professional collaboration, but there has however, been a paucity of empirical research on how these problems impact on clinical work. In this article we draw on a range of empirical work undertaken in four different contexts in the Norwegian health care system which explored patient and provider experience of collaboration in real world contexts.

**Aim**

The overall aim of this article is to explore how challenges of collaboration impact on clinical work through an examination of provider and patient experience and perceptions.

**Background and context**

Below we summarise the key findings from an empirical research project – “The challenges of collaboration in an integrated health care system” - which was undertaken between 2010 and 2012. The study comprised four distinct but inter-linked sub-projects. The key elements of the main project are presented in Box 1.
The main study was designed as a multi-site-research project and explored collaboration in four specific healthcare contexts:

i) an intermediate unit for older people
ii) mental health care services
iii) home-based rehabilitation services
iv) collaboration of GPs in the municipal care

Qualitative data were gathered through semi-structured interviews and participant observation as outlined in the Methods section (see Table 1). The findings from the study are summarised in eight articles, outlined below:

**Sub-project I** explored an intermediate unit recently established to improve the clinical pathway from hospital to home for patients aged 60 and over. The nurse-led unit had 15 beds, was located near a university hospital, and constituted collaboration between the hospital and four municipalities. The results of this study are summarised in three articles:

*Johannessen: Article I* explored the activities carried out and the conditions required to enable satisfactory work in the unit. The findings indicate that unfavourable environmental and adverse organisational factors exerted pressure on effective working and impeded patient clinical pathways [13].

*Johannessen: Article II* examined the unit’s role in a clinical pathway. Healthcare providers in the hospital, the intermediate unit and the municipalities had different opinions about who is a ‘suitable’ patient for the unit as well as the most appropriate time for hospital discharge. This resulted in lengthy negotiations between the hospital and the unit [14].

*Johannessen: Article III* explored the significance of professional roles in collaboration on patient transitions from hospital to home via the intermediate care unit. Collaboration within the unit and between the healthcare institutions was primarily viewed as ‘a nursing matter’. Apart from the physician, all the healthcare providers perceived the level of collaboration in the unit as being ‘uni-disciplinary’ rather than ‘inter-professional’ [15].

**Sub-project II** explored young adult mental health service users’ care pathways and focused on factors associated with continuity and disruption of care. The municipalities have the responsibility for primary care, including primary mental care services. The specialised mental health service is integrated with and run according to the same principals as other specialised health care services. The results are summarised in two articles:

*Ådnanes: Article I.* Key obstacles to continuity of care included the mental health system’s lack of access to treatment, poor integration between different specialist services, and inadequate tools for coordination [16].

*Ådnanes: Article II.* Users’ perceptions of services were influenced by fragmented care and a lack of user involvement. Concurrent problems were viewed in isolation rather than treated holistically. Some patients felt rejected when seeking to participate in key decisions regarding their medicine, diagnoses and treatment. Developing good relationships between providers and patients was considered crucial to effective working but in practice proved difficult to achieve [17].

**Sub-project III** explored how rehabilitation work was perceived and delivered by front-line services in two boroughs in Oslo. Norwegian municipalities are required to offer social, psychosocial, or medical rehabilitation to all inhabitants requiring such services and to establish a coordinating unit for rehabilitation. The results are summarised in two articles:

*Steinhaug: Article I.* Home-based rehabilitation received little attention in the boroughs, but participation in the project provided a broad discussion of rehabilitation. Starting with agreed policy guidelines and staff experience the researchers and borough staff jointly developed a model for organisation of and cooperation on rehabilitation [18].

*Steinhaug: Article II.* Results show that patients were rarely rehabilitated at home. The purchaser-provider organisation of home-based services, the rushed nature of service delivery, and limited resources were reported to impede effective rehabilitation work. There was a discrepancy between the high level of ambition of the health authorities and how these could be achieved by practitioners on the ground [19].
Subproject IV explored the various contextual barriers that served to attenuate effective collaboration between GPs and other health professionals working in managed primary healthcare services. The results are summarised in one article:

Paulsen: Playing a key role, GPs share their treatment of patients with many collaborative partners. GP’s collaborative patterns are dependent on individual priorities according to personal interests, considerations of importance, personal affinities, time schedules and practical barriers related to differences concerning branch-related organization and funding. The different branches of primary health care are organized according to an internal rationale. Conflicting principles of organization and funding between branches served to block effective collaboration. Cross-level professional axes between professionally interrelated branches of primary and specialist care complicated collaborative relations within primary care itself [20].

**BOX 1:** “The challenges of collaboration in an integrated health care system”.

**Material and methods**

**Material**
The eight articles arising from the project and outlined in **Box 1** constitute the empirical material underpinning this article.

**Methods**
We used the meta-ethnography method as originally developed by Noblit and Hare [21-23] as a guiding framework for integrating and synthesizing the findings across the eight articles. This method was developed for synthesizing published literature and involves taking relevant empirical studies to be synthesized, reading them repeatedly and noting down key concepts. The synthesis of these key concepts is achieved via a translation. The method comprises seven sequential phases as outlined in **Box 2**.

**Accomplishment**
Multidisciplinary teams are a useful approach to undertaking a meta-ethnography [22, 23]. In the present study, the analysis was conducted by four researchers (SS, A-KJ, MÅ, BP) each with different specialist professional backgrounds, namely: Medicine, nursing, psychology, and political science. In addition, four researchers with different disciplinary backgrounds, including health services research, economics, sociology and political science were involved in the wider research group. The whole group met at half-yearly workshops throughout the period of research and openly debated and discussed the analysis, emerging findings and their interpretation.

**Phase 1 – Getting started**
A formal systematic searching of the literature was not required as we were synthesizing results from our own eight empirical articles. In several workshops in the wider research group the authors discussed the content and findings of the eight articles in detail and agreed on their overall research quality based on, in addition to relevance, the following general criteria [22]:

- Are the objectives of the research clearly stated?
- Is the research design clearly specified and appropriate for the objectives?
- Do the researchers provide a clear account of the process by which their findings were produced?
- Do the researchers display enough data to support their interpretations and conclusions?
- Is the method of analysis appropriate and adequately explicated?

**Phase 2 – Deciding what is relevant to initial interest**
The participants of the wider research group jointly defined the focus of the study and developed the research question.

**Phase 3 – Reading the articles.**
Even though the authors knew all the sub-projects fairly well we read all the eight articles thoroughly at the start of the process of analysis. We first extracted information on the context, methods, and informants in all the projects (see **Table 1**). This step also included beginning the process of extracting themes in the articles. Each researcher extracted her/his own emerging themes using as far as possible the terms used in the original papers. We extracted eleven themes related to the challenges to collaboration in clinical work from across the eight articles.

**Phase 4 – Determining how the studies are related**
Preparation for comparison between studies requires listening and juxtaposing the themes and concepts used in each account. For comparing the themes and concepts in one article with themes and concepts in others we chose to use grids with the articles located along the X-axis and the themes located along the Y-axis. We compared studies, and our initial broad grouping of themes was gradually refined by merging, deleting, and establishing categories. Through group negotiation and discussion we eventually agreed on the three key themes: “collaboration with patients and between providers”; “different professional view”; “the significance of organisation for collaboration”.

**Phase 5 – Translating the studies into each another**
We explored each of the three key themes identified across all of the articles, see our final grid; **Table 2**. We chose an index study [14], characterised by high methodological
BOX 2: Noblit and Hare’s 7 phases for conducting a meta-ethnography.

| Sub-project | Article | Informants | Context | Recruiting informants | Data collection |
|-------------|---------|------------|---------|----------------------|----------------|
| 1           | Johannessen I | 16/8 | Intermediate unit | Strategic sample | Individual interviews Group interviews Observations in collaboration meetings |
|             | Johannessen II | 38/8 | Intermediate unit | Strategic sample Snowball sampling | Individual interviews Observations in collaboration meetings |
|             | Johannessen III | 38/0 | Intermediate unit | Strategic sample Snowball sampling | Individual interviews Observations in collaboration meetings |
| 2           | Ådnanes I | 0/9 | Mental health field | Recruited from municipal services, user organization, secondary school, and snowball sampling | Repeated individual interviews |
|             | Ådnanes II |         |         |                      |                 |
| 3           | Steihaug I | 24/0 | Home-based services | Strategic sample | Individual interviews Group interviews |
|             | Steihaug II |         |         |                      |                 |
| 4           | Paulsen | 10/0 | Primary health care | Strategic sample | Individual interviews |

Table 1: Articles, informants, context, and methods in the four sub-projects.

quality, a broad database, and systematic presentation, as a starting point. For each study we examined in detail the issues related to the given concept, for example “different professional view”. We related the core content issue of each paper to each other horizontally. The interpretations in the right hand column in Table 2 are results of our translating the studies into each other (the first order analyses).

Phase 6 – Synthesizing translation
For the synthesizing translation – the second order analysis – we used a method for systematic text condensation (STC). This comprises a four-step, cross-case method for thematic analysis suitable for developing descriptions of experiences within a field, in this case, how challenges of collaboration impact on clinical practice [24]. In the first
| Johannessen I | Johannessen II | Johannessen III | Ådnanes I | Ådnanes II | Steihaug I | Steihaug II | Paulsen | Our Translation |
|---------------|----------------|-----------------|-----------|------------|------------|------------|--------|----------------|
| The nurses in the intermediate unit collaborated appropriately and the patient felt well cared for. | The collaboration between different organisational levels was mainly a “nursing thing”. | Nurses had an inclusive collaborative culture which excluded other professional groups. Inter professional collaboration was poor. | Repeated disruptions occurred in the care pathway when the patient was transferred from one service unit or level to another. Individual plans were established, but did not work as intended. | A lack of appropriate cooperation was found between provider and patient and between providers. | In the relevant rehabilitation case the different services were established, but the providers did not collaborate appropriately. The patient was not involved in developing her rehabilitation plan. | Home-based rehabilitation was afforded little attention and seldom occurred in practice. The providers needed better inter-professional collaboration but framework condition were lacking. | Several providers wished to collaborate with GPs, but the GPs had to make priorities. GPs preferred to collaborate with their professional hospital colleagues. | Good professional collaboration was achieved, but the providers did not succeed achieving effective inter-professional collaboration. Collaboration between patients and providers in mental health was often poor. |
| A nursing perspective and nursing activities dominated in the unit. Physiotherapists and occupational therapists missed more rehabilitation. | There was disagreement between the collaborative partners about inclusion criteria and ‘suitable’ patients for the unit and about what were the unit’s role and tasks. | Different professionals had different opinions about inter-professional collaboration. The physician performed medical work, while the others wanted her to contribute to rehabilitation. There was medical dominance in inter-professional meetings. | Different views of “treatment” complicated the collaboration between the primary health care and the specialist services. | Patients and providers had different understanding of illnesses, diagnoses, treatment, and patient involvement. Providers in specialist services were regarded as “therapists”, while providers in primary services were “helpers”. | Different professional groups disagreed as to what rehabilitation is and had different foci in the rehabilitation process. Providers and managers disagreed on the aim of reducing the face-to-face communication in the home based services. | Different occupational groups worked separately with different methods against different goals in the relevant rehabilitation case. Providers in different positions and different levels disagreed about how best to prioritise rehabilitation. | Different professionals in the municipality had different views on inter-professional collaboration and different views on the GPs role as a collaborative partner. | Different professionals and different units had a range of conflicting perspectives and worked towards different goals. |
| The intermediate unit constituted collaboration between a hospital and four municipalities. This gives rise to a new collaboration interface in patients discharge. | The three organisational levels’ different aims and tasks hampered the collaboration. The hospital aims to discharge patients at once they were ready, the unit wanted patients needing rehabilitation, and the municipality wanted the patients to stay as long as possible. | Different symptoms were treated in different department in hospitals or in different hospitals. There were lacking collaboration structures between different department in specialist services and between the two levels. | Specialist services were responsible for “treatment” and the municipality for “following up” patients. The users found that their services were not coordinated appropriately between units in specialist services or between levels. Coordinating tools were found inadequate. | A coordinating units for rehabilitation was lacking in the boroughs. | Purchaser/provider organisation’s splitting up work into smaller, measurable units hampered inter-professional collaboration in rehabilitation work. | Each professional group had their own organisation without appropriate coordinating structure in place in the municipalities. Purchaser provider splits organisation hampered collaboration. | Organisation principles served as a barrier to appropriate collaboration between departments and between professionals. |

Table 2: Translating the studies into each other – first order analysis.
The mental health users reported both good and poor relationships with the provider [17]. They emphasised the importance of being seen, understood, and taken seriously and told powerful narratives about situations when this was not the case. Good patient-provider relationships required, according to the patients “good chemistry,” trust, and continuity.

Many mental health service users missed information and more influence on their services [17]. Several reported about how they “were admitted” and “were discharged” without being consulted.

The intermediate unit with different professionals working together to get patients “back on their feet” seemed to be a perfect place for inter-professional collaboration, but they were not successful in achieving this [15]. The employees reported that better collaboration could have prevented time-consuming discussions and that the patients’ care plans could have been more adjusted to the patients’ needs. The nurses described an inclusive and educational nursing collaboration in the unit, while the physiotherapists and the occupational therapists desired better inter-professional working. Providers in the municipal services also reported examples of poor collaboration [18–20]. The physiotherapists and the occupational therapists in the two boroughs reported limited inter-professional collaboration with their nursing colleagues and argued that the rehabilitation work suffered from lacking collaboration [15, 18]. Despite the fact that municipalities are required by law to provide rehabilitation, patients were rarely rehabilitated in their own home [18]. One rehabilitation case was found in one borough during one year, but the different professionals involved did not collaborate. Work with an IP was not initiated until several months had elapsed and the patient was not involved in decision made over her care needs. The providers emphasized that nobody had the overall responsibility for coordinating the work with an IP.

Different providers in the municipality reported wanting to collaborate with local GPs [18, 20], but GPs interviewed said that they were too busy to meet these demands and had to make priorities. Several professionals in the municipalities reported that they are more in touch with their professional colleagues in specialist services than with other professional groups on the same level [13, 14, 18, 20].

A case looks different from different perspectives
The differing perspectives on illness and treatment seemed to be one reason for a the existence of poor relationships between mental health service users and their providers. While professionals emphasised the need for providing an accurate diagnosis for the patient, the patients themselves were far more concerned about the cause of their mental health problems rather than being told the a name of their problems. Most of patients had received a diagnosis,
but individual patients reacted differently to their diagnosis. One patient diagnosed with a personality disorder, for example, disagreed with the providers’ diagnosis and did not find the diagnosis helpful at all. In fact she experienced not being taken seriously because her expressed opinion, for instance criticisms of the health care system or scepticism towards medication treatment, was interpreted as symptomatic of her personality disorder. She did not feel her needs were being “met”. Most of patients were medicated and some had felt pressured to take medicines. Several were sceptical about the use and impact of the medicines they were prescribed.

A recurring theme across the articles was the finding that different professional groups had different perspectives on illness and treatment and how they gave priority to different patients’ needs. The significance of profession and occupational position was clearly demonstrated across different contexts. Both in the intermediate unit [15] and the municipalities [18–20] physicians and physiotherapists revealed different understandings of their own and others’ roles and tasks. Rehabilitation is clearly defined, but nurses, physiotherapists, and occupational therapists disagreed about what rehabilitation is and ought to be [13, 18, 19]. In the intermediate unit the physician’s colleagues reported that she used to set the agenda in the inter-professional meetings and that medical issues often dominated at the expense of discussions concerning rehabilitation and recovery [15]. Health professionals at different levels in the hierarchy and in different positions also held different perspectives. In spite of the fact that the intermediate units’ collaboration with the hospital and the municipalities consisted primarily of collaboration between nurses, the providers had different opinions about what kinds of tasks the unit should perform [14]. The hospital and municipal informants maintained that the unit’s main task was rehabilitation while on the other hand employees in the unit reported that their role also included medical treatment, and that patient treatment should not be completed before discharge to the unit. Several providers reported that such disagreements often complicated and lengthened time of charge to the unit. Several providers told that the purchaser provider split model hampered the rehabilitation work because they missed being able to make their own assessments and to give priority to tasks other than those on the contract. As one physiotherapist complained [19]:

“You never know what happens next; I do not know what happens this summer, I do not know what happens next fall. Nothing! I only know that the providers I now have will quit their job. I do not know if I should continue treatment in department x or not.”

GPs and their collaborative partners in the municipalities, primary the home-based services, highlighted that the home-based services’ purchaser provider split organization was a hindrance to effective collaboration in practical work [18–20]. Historically, GPs could phone the home-based services and ask them to take blood-test. Now they had to make an application to the purchaser office [20]. “We cannot always wait for that”, a GP related. A number of providers told that the purchaser provider split model hampered the rehabilitation work because they missed being able to make their own assessments and to give priority to tasks other than those on the contract. As one physiotherapist complained [19]:

“It’s a problem that home nurses work according to decisions taken by others than the providers of the services, and they aren’t allowed to do anything other than what is stated in the decision. Needs change constantly during a rehabilitation process, and decisions have to be constantly altered – on application – by the service providers. We have to deal with a lot of red tape in order to get anything done. Collaboration has got so cumbersome since the purchaser-provider organisation was brought in.”

It was also reported out that no single authority in the municipality had the overall responsibility for promoting and ensuring effective collaboration.

**Discussion**

The overall aim of this article is to explore how challenges of collaboration impact on clinical work in healthcare services in Norway across four different healthcare contexts: (i) intermediate unit for older people, (ii) mental health care services (iii) home-based rehabilitation, and (iv) GPs’ collaboration in municipal care. The results from the meta-analysis of the eight articles highlight a lack of effective
collaboration between patients and providers and between health professionals and expose a range of barriers to collaboration.Providers described how poor collaboration made good clinical work difficult, and patients reported challenging meetings with providers and experienced a lack of holistic care. Both organizational and individual level factors were attributed to impede collaboration. Below, we discuss the challenges in collaboration arising from our analysis and detail how they impeded effective clinical work.

Insufficient collaboration hampers clinical work
Collaboration between patients and providers – the essence of health care
Relationships and collaboration between patient and provider are central to good quality clinical practice. The mental health service users articulated narratives relating to both good and poor collaboration with providers and highlighted how an unsatisfactory relationship sometimes resulted in them feeling rejected, insecure, and their needs not being adequately met. Patients and providers sometimes disagreed on the significance of the diagnosis and this disagreement complicated the collaboration. One patient reported that she felt rejected when her complaints were understood in light of her diagnosis. This aligns with the findings of a recent research project on user involvement in health services [25]. Mental health service users experienced that their complaint about the healthcare system were often attributed to mental health problems. Psychologists and psychiatrists aim to make diagnoses. Several patients in our study did not feel helped by a diagnosis, for instance ‘depression’; they wanted an explanation of the symptoms and an understanding of the reasons why the symptoms had developed; they needed to understand themselves. These different preferences may be attributed to different professional knowledge bases. As a medical specialty, the field of psychiatry is based on a medical logic in which mental disorders are classified into stable, universal categories - diagnoses where one diagnosis is associated with specific treatment methods with predicted effects [26, 27]. In clinical medicine, the point is not, however, to understand the illness but the ill person in a more holistic sense - to understand the person as a social and intentional human being that cannot be understood in isolation from his or her social environment. The patients’ ambivalence regarding use of medicine does perhaps reflect that the biomedical model of disease does not always fit with their experience of their illness.

Collaboration between professionals in a fragmented health care system
Providers in mental health services were not interviewed. Patients, however, reported issues related to fragmented services and a lack of collaboration between providers within the specialist mental health services and between specialist services and primary care. Ramsdal describes a schism between specialist and primary mental health services in Norway because the two have developed separately from one another based on different knowledge bases and different organization and management principles [28]. This description concurs with the perspective of that of Roger and Pilgrim [27]. In our study providers’ differing views on patients’ need for hospital admittance may be an example and this is presumed to complicate collaboration and prevent more integrated services. A small number of patients reported that services forming a whole despite serious mental problems and many providers involved. This indicates that some cases of successfully integrated services occurred. The professional groups in the intermediate unit were not successful in implementing appropriate inter-professional collaboration despite a favourable context. Different foci, cultures and ideologies across the different professional groups appeared to impede their daily work. The nurses emphasized the caring aspect, while the physiotherapist and occupational therapist tended to focus on the need for rehabilitation, and the physician was more focused on medical treatment. Insufficient inter-disciplinary collaboration seemed to hamper work with patients. In the municipality the nurses, physiotherapist and occupational therapist disagreed about what rehabilitation entailed, despite the fact that a national definition exists. This may be why home-based rehabilitation was rarely provided and only one patient who was rehabilitated at home was identified in the study. Aligning with other studies [3, 4, 5, 29, 30], these results indicate that different bases of professional knowledge and different understanding of professional demarcation roles and tasks were important barriers to effective collaboration. With a long patient list and a diversified patient perspective, GPs may consider it inappropriate to give priority to one particular patient group – irrespective of requests from other groups of health professionals. The nurses in primary care wanted more collaboration but GSs mentioned time pressure as one obstacle. Results of a Swedish study indicate that nurses are slightly more positive about collaboration than GPs and that a positive attitude to collaboration is part of nurses’ professional role to a larger extent than the GPs’ [31]. This may have influenced the GPs’ modest wish for inter-professional collaboration and preference to collaborate with providers in specialist services [32].

Disagreement also occurred within one professional group taking on different roles. In accordance with the findings of another Norwegian study [5] it appears that our informants’ perspectives were dependent on their position. Despite the fact that managers from the hospital and the municipalities had agreed on inclusion criteria for the intermediate unit, the nurses at different levels (hospital, unit, and municipality) had different opinions of “suitable” patients and what rehabilitation entailed. This could, according to the providers, postpone admissment to and discharge from the unit.

Differences in positional and economic power may also lead to significant barriers to inter-professional collaboration. The physician’s influence in the intermediate unit may well have a connection with the authority and power physicians have traditionally enjoyed through their use of scientific and diagnostic language, their monopoly on defining what constitutes disease and illness and what constitutes knowledge and expertise in clinical practice [33, 29]. More listening to the other professionals’ perspective might have made the physician a better collaborating partner, for instance in rehabilitation work. In collaboration between GPs and specialists in mental health, GPs
appear to predict clinical outcome [39]. Collaboration in clinical work – how and when?
The health care exists for the patients and the patient provider relationship is the core of clinical work. Fragmented healthcare services require effective collaboration. Strong support in both national reforms and legislation and various strategies and means have been used in Norway to improve collaboration in the healthcare system. Thus far, it has, however, had only a limited impact on for example the use of IP [35–37]. Holm [38] found a lack of time, muddled responsibility, and lack of rights and resources for the patient as reasons for the limited use of IP. Ahgren [35] argues that the modest effect of IP in practice is due to the fact that very little is done to change the existing fragmentation by means of organizing collaboration between care providers involved. Health systems integration requires policies and management practices that support relationship-building and information-sharing across organisational and professional boundaries [3]. Framing conditions and methods for developing collaboration is a responsibility for managers at different levels of the health system and presupposes managers’ thorough knowledge about and understanding of clinical work. The importance of including the clinical level in planning and implementing means for collaboration is emphasised because they are at the core of clinical work [35, 10, 18, 29]. On the other hand, if national policies are lacking, the collaboration arrangements in healthcare could develop in different ways with accordingly various effect for patients involved.

Collaboration is often represented as a “must”. We argue that the main questions ought to be what should be done for giving the patients the best possible services within available resources. The degree of collaboration itself does not appear to predict clinical outcome [39]. Collaboration is time- and resource consuming and a balance between time-consuming and time-saving is needed [31].

Methodological considerations
We used meta-ethnography as a method to synthesize the results across our eight articles focusing on how key barriers to collaboration can impact on clinical practice from the perspective of patients and providers. We found
the method well suited to this task. However, the analysis process was time-consuming; we had several discussions in developing the three main themes in the first order analysis and even more discussions in the text condensation process in second order analysis. There were, however, few problems in agreeing on the results. This may be due to the fact that we all knew all the sub-projects in advance though repeated discussions in the research group. A risk with knowing the projects so well could be that the researchers held common biases or took things for granted. The four participants in the half yearly workshops, however, asked questions, raised objections, and brought new perspectives. A strength of the study is that four different contexts were studied and a wide spectrum of professionals interviewed but a study limitation was that there were relatively few participants in some of the studies. In the mental health study a strength was that the patients were interviewed four times, a limitation was that providers were not included. Only the study of the intermediate unit was based on both observations and interviews with patients and providers. Observation as a method in the other studies might have brought valuable additional information.

The four researchers coming from different professions and perspectives performed the data analyses according to the meta-ethnography methods. Phase 5 in the meta-ethnography method – translating the studies into each other – revealed that numerous phenomena were found in all contexts and that new concepts and new relationships between the concepts could be established. In phase 6 – synthesising translation – we analysed the text of all the articles using the STC method. This allowed us to interpret concepts that encompassed more than one of the studies being synthesized. The significance for collaboration of power inequalities between different professionals was for instance clearly expressed in only one of the original projects [15] but through the synthesizing translation we found that power inequality occurred in all of the contexts: between providers in specialist and primary mental health care, between purchasers and providers in rehabilitation work, and between GPs and both their collaboration partners in the municipality and their colleagues in special health services.

The study’s new and original contribution to knowledge in this field is that our findings reveal how challenges to collaboration are intimately connected to a range of organizational factors across the four contexts under study. Patients’ and care providers’ experiences yielded bottom up perspective on how organizational conditions influence clinical work. Despite the study’s limitations we argue that our results have important lessons for other contexts and countries grappling with the design and operation of integrated care services in publicly funded and provided systems.

Conclusions
A range of both organizational and individual factors appear key to achieving collaboration in clinical practice. We have argued that challenges in collaboration are also due to differences, disagreement and conflicts, which are often implicit or even unconscious. Providers’ conflicting perspectives may indicate, as noted in the wider literature [3–5], problems with viewing issues from another party’s perspective as healthcare professionals are trained in one kind of logic and reasoning. If the collaborating parties can manage to surface differences and disagreements we argue that collaboration will be more easily accomplished. Focusing on inequalities and asymmetries in power and influence between the patient and the healthcare provider and between different providers might contribute towards the design of a better framework for communication and collaboration between the different actors. Organizational structures in the health system need to be redesigned so as to better nurture collaborative relationships which support integrated working and decision-making between providers, health care professionals and patients.

Ethics
Ethical approval was obtained from the Regional Committee for Medical and Health Research Ethics for sub-project II (ref.: 2010/1144). For the three other sub-projects, an ethics’ application was also submitted to the Committee, but the projects were not found to be part of the Committee’s mandate since it is not regarded as medical or healthcare research conducted with the purpose of generating knowledge about illness or health. Informed consent was obtained from all the patient participants before the interviews. Principles of confidentiality and anonymity have been applied in the conduct, reporting, and storage of data in accordance with the Act on Processing of Personal Information and the requirements of the Regional Ethics committee.

Acknowledgements
Thanks to the researchers Hilde Lurås, Anne Werner, and Jan Lippestad for valuable contributions in the half-yearly research workshops where the analysis and emerging findings were discussed and debated.

Reviewers
Dag Olaf Torjesen, Associate Professor, Ph.d., Dep. of Political Science and Management, University of Agder, Kristiansand, Norway. One anonymous reviewer.

Funding Statement
The study was funded by the Research Council of Norway.

Competing Interests
The authors declare that they have no competing interests.

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