Patients’ experiences of an open access follow up arrangement in managing inflammatory bowel disease

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Background: Improving access is a key policy issue in improving quality of care and extending patient choice and participation. People’s experience of changing from fixed outpatient appointments to more flexible direct access arrangements for chronic disease has been underexplored.

Objectives: To examine patients’ views on using an open system of access compared with fixed outpatient appointments as part of a guided self-management intervention for inflammatory bowel disease (IBD).

Design: Embedded qualitative study undertaken alongside a randomised controlled trial. Semi-structured interviews were undertaken to obtain an in depth understanding of patients’ experience of the change in access arrangements.

Participants: A purposive sample (n = 30) was drawn from the intervention group (n = 700) according to a range of responses to the trial baseline and follow up quantitative measures.

Results: 28 interviews were included in the analysis. Compared with the previous system of fixed appointments, preference for the new open access system was based on enhanced personal control in contacting services and the view that it fitted better with everyday routine management and the requirement for urgent medical contact when symptoms fail to respond to medication. Preference for retaining fixed appointments was based on a sense of security from gaining access which did not require the individual to initiate the request for medical help.

Conclusions: Open access may fit better with patients’ self-management of their condition and everyday routines, roles and responsibilities. Ensuring that outpatient organisational arrangements and personnel are responsive to patient initiated requests for appointments is likely to impact on the acceptability of this type of access arrangement. Some people may continue to prefer the fixed appointment system which should be retained if patient choice is to be respected.

Patients with chronic illness have an ongoing need for access to health services and have traditionally relied on hospital based services for treatment and advice which provide only limited support for day to day disease related problems between hospital visits. Previous studies have shown that, for people living with a chronic illness, their concern to continue to receive medical management and treatment is balanced against the competing need to emphasise normality and maintain a considerable degree of control over their lives.1-3 Many people with chronic illness therefore try to find a balance between asking for help from formal services and coping without. One way to support this self-management is to change access to outpatient clinics by allowing patients to self-refer when they judge advice is needed rather than forcing them to rely on appointments fixed by the hospital. Improving access accords with key policy issues in improving the quality of care, meeting demand for primary and secondary care, and patient participation initiatives.4

Reasons given by hospital doctors for long term follow up of people with chronic illness in outpatients departments include: a perceived need for a review of symptoms, medications and blood tests; unwillingness to discharge to primary care management; and a belief that patients expect hospital treatment.5 However, since most chronic diseases follow an unpredictable course of activity and remission, a fixed appointment system is unlikely to provide coincidence of attendance and relapse, which results in either many clinic non-attendances (approximately 12% across all specialties)6 or lack of access for those in need of urgent attention.

Inflammatory bowel disease (IBD) is used as an exemplar of a chronic disease to explore patient response to open access arrangements introduced as part of an intervention which was evaluated by a randomised controlled trial (RCT). Crohn’s disease and ulcerative colitis affect about 175 000 people in the UK.7 Current medical treatment is ameliorative rather than curative and many patients need maintenance drug treatment. Symptoms—which include bloody diarrhoea, abdominal pain and weight loss—follow a relapsing course with periods of remission. The RCT to which this study is linked is outlined in box 1.

Variations to traditional outpatient department access arrangements for IBD have been tested in two studies. A recent randomised trial explored the effectiveness of using the general practitioner (GP) as the first point of contact which reduced demand for outpatient attendance.8 Patients randomised to follow up through open access were asked initially to contact their GP about problems and were only encouraged to contact the hospital directly if they were unable or unwilling to see the GP first. However, this link with primary care may be unnecessary given, as the second study suggests, it is likely that secondary care services provide principally for the ongoing contact people have with a chronic illness such as IBD.9 Neither of these studies was able to identify and account for the processes underlying the fall in the number of outpatient appointments.

This study was undertaken to examine the experiences and views of patients with IBD on their use of a new system of open access to outpatient clinics compared with the traditional system of fixed appointments and to determine the rationale behind their preference.
Patients’ experiences of open access

Box 1 Inflammatory bowel disease RCT

The intervention was an evidence based package designed to bridge the gap between the requirement for continuity of clinical care and a patients’ own requirements for disease management (evidence suggests that patients with IBD feel insufficiently informed and want involvement in their treatment). The RCT had two arms and outcome measures were recorded at baseline and at 12 months. The intervention had four components:

**Improved information**
Provision of a patient guidebook containing information that is relevant, accessible and uses a combination of lay and traditional evidence based knowledge. Guidebooks for ulcerative colitis and Crohn’s disease were developed with patients before the study.

**Guided self-management**
A written guided self-management plan to which patients can refer when making decisions about treatment and the need for service contact.

**Changed professional response**
Promote flexibility in professional response through a patient centred approach provided by consultants trained in its methods.

**Changed access to services**
Provide direct access to services, allowing and enabling patients to self-refer based on their own evaluation of need.

The objective of our study was to determine whether this ‘whole systems approach’—which includes interventions at a number of levels (the patient, the professional method of working, and the organisation of services)—altered patients’ abilities to self-manage their condition, and whether they made more appropriate use of health service resources. The main results of the trial are reported elsewhere. Three outcome measures used in the trial provide the context for the qualitative analysis of the open access arrangements. These were: the number of appointments attended during the trial year; the number of made appointments not attended; and the percentage of patients who self-referred.

The number of appointments reduced by approximately one third in the intervention group (from 3.0 to 1.9) compared with the control group (from 3.1 to 3.0; difference \(-1.04, 95\% \text{ CI} \ -1.43 \text{ to} \ -0.65, p<0.001\)). The number of non-attendances during the trial was also lower for the intervention group than for the control group (difference \(-0.08, 95\% \text{ CI} \ -0.15 \text{ to} \ -0.01, p=0.034\)). More patients at intervention centres than at control centres self-referred for at least one appointment (43\% v 22\%, p<0.001).

At the end of the trial patients recorded whether they would prefer traditional fixed hospital appointments or the intervention system where they receive a personal self-management plan and can make their own appointments. Analysis was restricted to the intervention arm. Overall, 26\% of patients in this arm expressed a preference for the traditional system and 74\% expressed a preference for the open access arrangements. We concluded that changing access arrangements was generally acceptable to most patients with IBD and reduced demand for health service resources.

**RESULTS**

Of the 28 respondents, 17 had ulcerative colitis (UC), 11 had Crohn’s disease (CD), 16 were female, 17 were in employment, and their ages ranged from 22 to 68 years.

Comparison of direct access with fixed appointments

Judgement about the new access arrangements was formed against the backdrop of people’s previous experiences of a system of making and attending fixed regular appointments over a number of years. Previous attendances at outpatient appointments varied considerably across the patients’ accounts. At control centres patients mostly received appointments when they were invited by secretarial staff or receptionists in outpatient departments. Patients’ permitted attendance was based on previous attendance at appointments.

At intervention centres patients could make their own appointments. They could book appointments at any time in the future. The majority (79\%) were offered appointments for the entire duration of the study. Most patients made between one and four appointments during the study. However, 5\% of patients were offered appointments that they did not take.

**Comparison of direct access with fixed appointments**

When patients were offered open access but subsequently subsequently chose fixed appointments, they were more likely to have previously attended previous appointments (76\% vs 44\%, p=0.009). However, there was a significant difference in the number of appointments attended during the trial (mean 1.3 v 2.0, p=0.003) and the number of made appointments not attended (mean 0.7 v 1.4, p=0.003).

**Methods**

To extend and challenge the existing data and test the integrity of the findings from the main trial, we used aspects of grounded theorising to guide the sampling, collection, and analysis of qualitative data. Qualitative semi-structured interviews were undertaken to obtain an in depth understanding of patients’ experience of the intervention and to focus on the processes underlying the outcomes of the trial. Interviews were conducted by EN and took place between January 2001 and July 2001. All interviews were audiotaped, lasted 30–60 minutes, and were transcribed verbatim.

A purposive sample of 30 respondents was drawn from the intervention group according to a range of responses on the trial baseline and follow up quantitative measures to reflect the range of possible experiences underlying responses to the changing self-management and health service arrangements. A checklist of topics was initially generated by the research team and modified in an interactive and iterative fashion after initial interviews. Agreement around meaning was established through regular discussions and comment throughout the interviewing, collection, and analysis of interviews (EN, AK, AR). This ensured the key areas of having participated in the trial were comprehensively covered with each respondent while, at the same time, allowing issues which were important to patients to emerge.

The latter was consistent with a phenomenologically informed approach to thematic analysis aimed at exploring the interpretive meanings given to access arrangements. Interviewing ceased after 30 interviews as there were no new themes arising and repetition of accounts indicated saturation of the existing themes. Similar themes across interviews were identified and within or sub-themes scrutinised in depth. The NUDIST qualitative data analysis system was used to assist with the thematic analysis which involved examining the interviews thematically across the whole dataset as well as in the context of each patient’s account. The personal and social context of managing illness and coping strategies before the intervention and perceptions about patient experience of each component of the intervention (guidebook, patient consultation, and access arrangements) and overall impact on self-management and health behaviour.

Here we report only what was said or is relevant to the open access arrangements, which included the experience of negotiating the type of appointment system to be used, subsequent use of the arrangements compared with previous access arrangements, and perceptions and evaluation of intermediary processes—for example, contact with secretarial staff and receptionists in outpatient departments.

Twenty-eight of the 30 respondents in our sample discussed the open access arrangements for making appointments (two did not articulate an elaborated response when asked about this area of their experience). They included six respondents who were offered open access but subsequently renegotiated or reverted back to the fixed appointment system. The information gleaned from these later interviews together with the accounts given by all the respondents who had previously used the fixed appointment system allowed comparisons to be made between fixed appointments and open access.

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departments were frequently portrayed as personally inconvenient, wasteful of time, and ineffective organisationally.

“I think the waiting has got worse, it used to be quite quick really. But sometimes I wait ages and an hour and a half is a nuisance when you are busy and have young children.” (UC, female, age 36)

“The waiting time can be long, but you expect that. Generally you go with the knowledge that you can be a long time” (laugher) (UC, male, age 46)

Consideration of the specific difficulties faced by people with IBD in attending and the inconvenience relative to the perceived importance of what was undertaken during the appointment was also relevant.

“The clinic for Crohn’s is in the morning, now people with Crohn’s have very bad mornings getting up, dealing with toilets and getting down there. They have sent me appointments for 9 o’clock in the morning and I just found it stupid that all these people with bowel disorders are having to go to morning clinics. And when I’ve sort of said—oh the later the better—they say what about 10 o’clock and I can say you just pass it on a bit later—and they look at you as if you’re stupid so now I just tell them ‘I’ve got Crohn’s disease and I can’t get out of the bathroom for at least 3 hours… I hate to think of how the people manage who have to get there for 9 o’clock…” (CD, female, age 29)

“Oh it’s been a nightmare—honestly—especially when I took the children, because sometimes I’ve had to take both of them… They don’t do anything they just ask you how you’re going on and give you the medicine and the blood tests. I go every six months” (UC, female, age 53)

**Link between increased sense of control in managing IBD and increased satisfaction in using open access**

The qualitative data suggested that a preference for the open access system was based on more than a predilection for a less inconvenient system. The new access arrangements seemed to signify a transfer of control in using services from the hospital to the individual, which, in part, was also linked to people’s confidence in their ability to manage their own condition. The form of advice and support which replaced the traditional system was based on more than a predilection for a personal benefit of the open access arrangements. People were encouraged by the new approach.

“It was clear to me that I could ring to make an appointment or to chat. I thought that was a good idea because if you are managing your own life—like I’m used to anyway—a lot of the time you’re waiting time going back to the hospital.” (CD, female, age 60)

“It’s been good for me because when you are running your own business you don’t want to be nipping down to the hospital all the time—not if you don’t need to. This is definitely how I would like to carry on—definitely. I like the flexibility of this, it suits my life and I am happy to deal with the flare-ups. I’m fortunate, I don’t seem to go with the knowledge that you can be a long time.” (UC, male, age 43)

A sense of autonomy in deciding when and how to use hospital outpatient resources was reinforced by the knowledge that the service was there when and if it was needed. In addition, some people felt they were getting a better service even if contact with services was less, because the system was felt to map more closely onto people’s own needs rather than the person having to fit in with the organisational imperatives of the hospital system.

“I feel more in charge of my condition. I know that the safeguard (the phone) is there should I need it and I prefer this to having standard appointments. I share the decisions with him and I am prepared to take the responsibility for myself. Previously, as I said, the condition was managed for me and I was told what to do… And it then became not something that controlled me but something that I can live with and control so that it doesn’t affect my life.” (UC, male, age 46)

“I haven’t needed to use it. But I do have a vague recollection of him (the doctor) saying ring me and come in if you get any flare-ups like this again and I found that quite reassuring and [that] I wouldn’t [have to] lie in bed like I have before and suffer and do nothing—I wouldn’t hesitate to ring if I was ill again…” (UC, female, age 40)

Responses to the open access arrangement suggested that people saw it as fitting in with future long term management based on the other parts of the approach which made up the whole systems approach (see box 1). Increasing people’s confidence and ability to manage was consciously built into the design of the guided self-management information, with the idea of reinforcing this through evidence of introducing self-management and trying to reinforce it through the philosophy of patient centred consultations and greater freedom in using open access arrangements.

“I would like it to proceed exactly as it is … that if I need to see him then I want to be able to pick up the phone and say ‘things are not going well and I need a consultation’ and to be able to have one in quite a short space of time. Whether that would work I don’t know because I’ve never had to do it since the project started, but that’s how I would see it in an ideal world.” (UC, male, age 60)

“I prefer to make my own appointments. I thoroughly agree with this whole management system because, if you keep coming back every 6 months, you might have a flare-up in the middle and be fine when your appointment comes, so what’s the point? You need them to see you when you’ve got symptoms. I think it’s an excellent idea and it certainly works for me. I phoned on the Friday and I was in on a Thursday and I was quite happy with that. You need to have early access because you need to phone when you have a problem …” (CD, male, age 40)

The new arrangements were usually viewed as the intended outcome of negotiation and sense of trust and equality between doctor and patient which underlies the patient centred consultation training. Surprise was occasionally expressed when the new access arrangements were actually found to work:

“I did phone up once in the beginning when I had a couple of problems—I phoned his secretary and he actually did phone me back at work.” (UC, male, age 43)

The personal benefits of the open access arrangements were not seen in isolation. Rather, evaluations were made with reference to a collective responsibility towards the NHS. In the following quotation clear reference is made to being treated well in the context of overstretched services.

“I felt I get a lot of support from him, especially the fact that he gave me his secretary’s telephone number …. with the words ‘if at any time you want to get in touch phone my secretary and we’ll get you in’ and I was quite impressed with that, especially when you know the pressure that the NHS is under.” (UC, male, age 60)

**Reasons for retaining a preference for a fixed system of appointments**

Despite the fact that, as indicated by the quantitative findings, most expressed a preference for the new system, this was not always the case. While most people preferred the new system, a number actively expressed a wish for the old system and/or continued to use the old system of appointments. Some people who were initially introduced to the open access system subsequently requested to revert to the old system or were advised by the consultant that they could use both in tandem. In two instances the open access system had not been introduced as it should have been by the consultant.

“He normally made an appointment for me to go back and see him but he also said any time I wanted to make an appointment then phone his secretary and she would make me one. But I try to run between each appointment rather than do that—I try my best …” (CD, male, age 52)
Some individuals considered that the transfer of the onus onto the patient in making a decision to access care might erode the sense of security and certainty which came with the routine appointment system.

"I like to go about every 2 months just to see and talk to somebody. I go every couple of months and I prefer someone to keep an eye on me—I feel reassured." (UC, female, age 67)

"Well I think [fixed appointments] … it’s just a check up. You may go along one day and he says you don’t need to take that medication, or he may spot something—it’s nice to have that check.” (CD, male, age 41)

Some people also expressed a dislike of the assertiveness required to instigate an appointment or telephone assistance which accessed medical help directly and felt uncomfortable using the secretary as a mediator.

“Well, yes, I’ve got a number but I think there should be a nurse in attendance who we can talk to, you know—if you’re a bit worried. I mean his secretary is really nice but she’s only there for appointments and that, isn’t she …?" (UC, female, age 68)

The lack of confidence expressed in getting the new arrangements to work was sometimes confirmed by the system failing to respond to an individual’s request or to operate the open access arrangements as originally agreed. It was clear that, on occasion, some individuals lacked the confidence to negotiate directly with the system even when they had been able to identify the problem that they considered needed medical attention.

“And this lady on NHS Direct told me I must ring them and demand an urgent appointment and if not I must ring her back and tell her. Anyway, they did push me in at 12.20 the following day.” (UC, female, age 68)

“He would just give me an appointment, sometimes it would be cancelled more than once and at one time it was as long as 6 months. I do have his secretary’s number but when I rang she just said ‘no we haven’t got any bookings.’” (CD, female, age 29)

Finally, there was also some suggestion from the interviews that these individuals tended to feel less in control of their illness.

“I tend to suffer more at night which is partly my fault—I am trying to get myself into a routine now where I eat more during the day—every couple of hours I am supposed to have something to eat and one of my drinks to get the body used to eating and having food in it. But a lot of the time, because you feel so ‘shitty’ during the day—your stomach’s hurting, you feel sick, you’re on the toilet until dinner time or whatever—you do not physically feel you can eat anything because you feel you are going to bring it back up so sometimes I’m not eating till 6 o’clock at night. I know I’ve lost a lot of weight and I need to put some on so what do you do—not eat and lose more weight, or eat and suffer in the morning—you don’t know what to do for the best.” (CD, female, age 29)

**DISCUSSION**

The aim of this study was to explore the experiences of patients of modifying access arrangements as part of a new approach to managing illness. Preference for the new open access system was based on a sense of enhanced personal control in contacting services and the view that it fitted better with everyday routine management and the requirement for urgent medical contact when symptoms fail to respond to medication. Some people preferred to keep the fixed appointment system because of the sense of security this engendered. A limitation of the study is the extent to which it is possible to generalise from the experiences of access arrangements reported here to the use of health service access arrangements elsewhere and for different conditions. One fruitful avenue of future qualitative research would be to compare changing access arrangements across different service contexts and conditions.

The way in which the open access arrangements worked was, to a large degree, dependent on patients feeling able to self-refer based on their own evaluation of need. Compared with the previous system of fixed appointments, the findings suggested that many people felt able to do this once this option had been explained to them by their consultant. Satisfaction with the open access arrangements was based on the achievement of a closer fit between the ways in which people wished to exercise greater control in their everyday management of IBD and the decision to seek medical advice and input as a safety net system when self-management options had been exhausted. These findings suggest that the benefits of open access arrangements identified by respondents are likely to fit with the philosophy and principles of contemporary patient participation initiatives. The Expert Patients Programme which includes a self-management education training initiative is designed to improve people’s confidence and ability to self-manage and engage in collaborative shared decision making with health professionals.

However, despite the fact that most of the patients preferred the new system, there was a preference among some patients for retaining the fixed appointment system. This was based on the need for a predictable and reliable system which was independent of the requirement to articulate the need for assistance on the part of the individual. In some instances this was a result of direct experience of a failure of organisational arrangements to respond to an individual’s requests for assistance made under the open access system.

The underlying reasons for people’s preferences need to be seen in the broader context of the way in which outpatient departments have traditionally determined arrangements for appointments, people’s orientations towards health services, and the style with which they manage their own illness. Previous research suggests that appointment making is a complex social process where outcomes are negotiated. In this study the use or otherwise of the open access system was not simply a matter of patients choosing one system over another. The consultants were able to influence the ability of patients to use open access arrangements by the extent to which they decided that such a system was appropriate for particular individuals and the extent to which they communicated and set up arrangements with other parties working within the outpatient department. In this respect, open access represents a paradox given that the move towards a more independent and patient driven system came initially from a relatively dependent doctor-patient relationship in which the consultants granted patients “permission” to use the new system. The relevance of this relationship is evident in the decision by a small minority of consultants to ignore the request to introduce these new arrangements or to offer both in tandem.

Intermediaries such as secretaries also assume a discretionary role in determining access to medical staff. At times it seems that the “gate keeping” activities of this group may have resulted in the failure to facilitate requests made through the open access system. Given the dependence of people on outpatient departments to administer and facilitate patient initiated requests for appointments, it is understandable that some people felt more able than others to assert their needs for an appointment and to negotiate this with hospital personnel. An implication of this is that, when the introduction of open access arrangements is being considered, there is a need to address the way in which access and appointment systems in outpatients have traditionally been centred on hospital and health staff routines (including those of secretarial and receptionist staff) and the ways these can be changed in a manner which is likely to
Key messages

- There is an interaction between people’s experience of health care and their subsequent use of services and illness management strategies.
- People make judgements about new access arrangements based on their comparison with existing arrangements for outpatients’ appointments, and the extent to which people find that the new arrangements work in practice.
- An open access system is preferred by patients because of its opportunities for autonomy in making a decision to consult, the closer fit it has with illness trajectories, everyday routines and coping strategies for managing symptoms.
- An expressed preference for retaining fixed appointments is predicated on the perceived availability of planned and certain access which does not rely on patient initiation.

inspire and encourage confidence in patients to use the new system. Even when this is achieved, it is likely that there will be a role for retaining some access to fixed appointments alongside open access arrangements in order to meet expressed patient choice and to be able to respond to the different ways in which people wish to manage their condition and use the health service.

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