By next year a likely scenario is this:

June 2007, the consultant sighs. The letter from the GP is tantalisingly brief: the reason for referral seems unclear and no test results are included. The 70-year-old patient, seeing the GP’s confusion, offers to show him her records. She moves closer to the screen, taps her pin and personal password into his computer, and uses a token to generate random numbers that will ensure she is securely identified. Seconds later, all her recent consultations appear on screen, followed by all the correspondence and all test results.

On the same day in A&E, a patient admitted with chest pain shows a junior doctor the reports of the last chest X-rays and ECGs ordered through his general practice. He displays this on his handheld organiser.

In Leeds, the pensioner, confused by the last consultation with her GP, logs on to her records. She sees what the GP was thinking and the results on which the conversation was based. She can also see, linked with the problem title of hyperlipidaemia, a short list of weblinks that offer further information. Record access has the potential for:

- improving care and safety,
- affecting the way care is delivered, and
- enhancing patients’ self-efficacy.

While the outcome of HealthSpace is awaited, in the UK and worldwide alternative systems exist that enable patients to access their full GP electronic record in the surgery. Similar access on the internet, which would enable the scenarios described above, is currently being piloted.

Different approaches to record access
In the UK, kiosks (see picture below), designed by independents outside the NHS Connecting for Health programme, offer access to the full GP electronic records with linked health-related information. Read codes in the records are linked to patient information leaflets which can be read on screen or printed. Access is authenticated by fingerprint and pin. A touch screen and reformatted records, designed with patient input, make content and navigation easier to understand. Patients can see their consultations, demographic details, letters, investigation results, and allergies. As of June 2006, there were 11 practices in England using the system, with a total of about 5000 patients registered with their fingerprints.

Some surgeries allow patients to take away their records on a memory stick or a CD. In others, patients can insert health data, such as blood pressure and peak flow, directly into their record (R Fitton, personal communication, 2006). In the US a number of health maintenance organisations offer record access.

A web-based service is also being piloted which is different from the NHS Connecting for Health, national ‘Spine’ database of information about patients’ health. The information is not placed on the Spine, but remains in the practices, being called up each time it is needed. As a result, the system:

- enables patients to keep control over who sees their record,
- keeps the record in the surgery, so its ownership is clear, and
- allows patients, if they wish, to share their data with clinicians anywhere in the world.

In all situations, security and legal protection are imperative.

What the law says
The Data Protection Act 1998 and Access to Medical Reports Act 1988 state that a patient or anyone authorised by the patient, parents of children under 16 years
of age, or a ‘Gillick competent’ child are entitled to access their medical records. Access can only be denied where the information may:

• cause serious harm to the physical health, mental health, or condition of the patient or any other person, or
• relate to or be provided by a third person who had not consented to the disclosure.

The Freedom of Information Act 2000 gives people a general right of access to information held by or on behalf of public authorities, unless that information would, or is likely to:

• endanger the physical or mental health of any individual (including the applicant, the supplier of the information, or anyone else), or
• endanger the safety of any individual.

The Copying Letters to Patients initiative, enabling patients to see all letters written about them, is advisory only. However, this initiative is being adopted across the NHS, with general acceptance by patients and clinicians.

What is known about record access?

Who looks at their notes? Of patients who have been registered to use the electronic kiosk access system, 75% have looked at their notes: the same proportion as those who used to look at their paper records. Some patients access their records repeatedly; most only do so occasionally for specific purposes. Patients with long-term conditions use it more than others.

Clinicians’ reactions. Clinicians are often initially sceptical and worried about the impact of record access. They fear that mistakes and confusion will be exposed and that litigation may increase. However, there is no evidence for this. On the contrary, evidence is clear that record access improves relationships between clinicians and patients. Experience with record access tends to convince clinicians that its benefits outnumber its potential problems.

Benefits of record access. The benefits of record access appear to be substantial. Patients describe improved trust and confidence in their clinicians, and they feel more informed and in control of their condition and its management. There is some evidence for improved health practices by patients. For example, improved compliance in heart failure and improved cigarette quit rates have been demonstrated.

In general, patients are keen on record access in principle and in practice. Record access can increase safety by alerting the practice to any recording errors. Furthermore, patients can save time for practices and themselves by looking in their records for information rather than asking reception.

Duration of consultations. Patients using record access do not appear to take more time with clinicians. This coincides with experience with the kiosks in general practice. Record access seems to help patients focus their medical agenda and, ultimately, record access may end up saving time and foster compliance (M Cauldwell et al, unpublished data, 2005).

Risks. There are potential risks if the process is not handled carefully. If, for example, patients see frightening information before it has been discussed with them or if they are able to access third party information. Simple precautions in letter- and result-handling, however, can virtually eliminate these risks.

An investigation revealed that 10% of patients were upset by what they read, although the vast majority of those were still pleased that they had seen their notes. Patients with cancer, in particular, were pleased to have read their full GP records. Evidence suggests that patients with psychiatric problems can find seeing their notes more distressing than those without mental health problems. However, the experience can be harnessed therapeutically.

Experience of record access so far has been with enthusiastic committed practices, and it may be that such safety precautions may not be so strictly adhered to when the majority of practices will be able to offer access.

Confidentiality. Confidentiality is well-maintained in the waiting room kiosk-based approach. The system uses both fingerprint and pin identification, and there are simple and effective ways of ensuring that it is only with the patient’s permission that the record may be seen by anyone else. A web-based approach may be more prone to abuse, despite secure authentication and access. It is possible that members of the family, for instance, will be able to see records, perhaps by glancing over patients’ shoulders or by coercing them. As with internet banking and other protected sites, once the record has been delivered, we can assume that the patient becomes fully responsible for its security.

Authentication. Ensuring that each patient sees his/her own records and no-one else’s

Box 1. Record access approach of the online EMIS/PAERS project

A key part of the system is user authentication. There are several steps that a user must follow to ensure that they are as secure as possible.

• An initial sign-up document has a once-only PIN number. Without this number the user cannot create an account and so no-one can set up an account in someone else’s name.
• The user needs to enter their practice ID number, to tell the system which practice they are registered with, and an access ID to identify them individually.
• Users set their own password, not known even to practice staff, and will soon be able to change it online if required.
• For medical record viewing they have to enter selected digits/letters from a second ‘pass phrase’, which is different each time. This ensures that no one can easily steal their passwords by filming them or installing key-logging software on their computer.

Although some patients initially find these awkward, almost all soon become adept at logging in and using the website. None of these stages can be removed without the risk of harming security. In a later phase of the project a token-based system will be offered and this is expected to provide a significantly higher level of protection for a user’s account login.
is essential. The kiosks’ fingerprinting is robust, but can be difficult for older people whose prints are less defined. Net-based access requires ‘two-factor authentication’: pin and password plus another approach such as a token. An example of the approach taken by the online EMIS/PAERS project is shown in Box 1.

The main risk to security in the current system is that an attacker could guess or steal a patient’s account login information and use it to login themselves. This risk is higher if patients use weak passwords, fail to keep their passwords secret, use public computers, or do not keep their home systems up-to-date and protected from viral and trojan software. Token-based two-factor authentication will mitigate much of this risk.

It will be important that, if different systems are developed for record access in the UK, access and authentication is not only of the highest standard, but should be similar in all systems to avoid confusion.

Standards and guidelines
To ensure high standards and avoid some of the risks listed above, standards and guidelines need to be written prior to the launch of record access on a national level. These standards and guidelines will best be created by national agencies, such as the Royal Colleges, patient groups and NHS Connecting for Health.

How do patients use record access?
Current research suggests that patients use record access to prepare for consultations and recap after them; for finding information that they would otherwise have to ask the practice for, such as immunisation history; it that they would otherwise have to ask the record access to prepare for consultations. (NHS Connecting for Health, unpublished data, 2006). Access to records can compensate for poor communication.

What impact could record access have on daily clinical life?
The shared record, if we get it right, offers a new dialogue in which clinician and patient can learn from each other. Thus, the task is not only to foster ease of use for the clinician, it is also to make the record useful for the patient. The electronic record could change from a repository of recorded clinical information for clinicians to a tool of communication and health management used by patients and health professionals. This could affect the structure, processes and outcomes of records in a number of ways.

Reading the record. GPs’ notes are full of abbreviations, technical terms, and other impediments to lay understanding. Patients tell us that they understand on average 60% of the record as it is currently written. If patients start relying on the record to make sense of their health and to participate in their own care, recording may have to change. For instance, the record might need to show ‘heart attack’, instead of the standard abbreviation ‘MI’, for the patient’s benefit which would be automatically generated. One system currently enables patients to see information about most of the Read codes by touching the screen to reveal a patient information leaflet on the topic. Web-based plans include automatic translation of technical terms.

Making sense of data. For patients to understand clinical data, record access will need to make sense of these data for them. For example, results could be displayed as charts with normal ranges instead of simple numerical values. These results can include clinicians’ comments on their significance as well as information about the tests.

Issues of equity and design. The design of record access will also need to ensure equality of care. Ethnographic studies suggest that people have different mental models of health which structure their health behaviour (C Poirier, unpublished data, 2000). Although design needs to be universal to ensure basic equity of access, it would also make sense to design the planned ‘Common User Interface’ so that it can be customised by patients and clinicians. Different groups of patients and clinicians will want data presented in different ways.

Patients correcting errors in the record. Experience has shown that record access can aid the correction of recording errors.13 Such errors often involve patient identification. They may also pertain to aspects of a consultation, for example, to a description of conditions or symptoms described by the patient, or outcomes. When patients use the PAERS system, for example, they sometimes identify when results are changing in an unexpected direction, when the wrong tests have been taken, or when test results are missing. Studies are being conducted to quantify error reduction outcomes in records provided by record access.

Patients entering their own data. A number of practices have experience of patients adding data directly to their record by email through a system called GePmail. Peak flow and blood pressure are the most common entries. The design may need to ensure that data are viewed by the clinician before being entered into the notes so that any relevant clinical response could be made.

Enhancing self-care. The system could be designed to recognise, for example, someone with diabetes whose blood pressure and sugar control are inadequately controlled. The patient would be reminded of this, with associated advice, when they looked at the records. They could be reminded of relevant appointments, automatically. HealthSpace is designing this for its future deployment.

Conclusion
Record access is increasingly being adopted around the world by clinicians and patients. We recognise the risks and potential pitfalls associated with this trend. Yet, we see record access as significantly improving shared care and improving not only trust in and respect of clinicians by patients, but also trust in and respect of patients by clinicians.

Record access will result in a closer relationship between GPs and their patients. Access enables patients to
participate in their own care, find out relevant information, and ask about their disease and health. It could enable GPs to improve patient safety. Patients can include their own recorded values and their own care management in the record, for the benefit of both clinician and patient. Record access also facilitates education as it can frame results and medical information directly from patients’ concerns. It appears safe when used with simple precautions, and is likely to form the basis of future choice and shared decision-making.

Brian Fisher, Richard Fitton, Charline Poirier and David Stables

Competing interests
David Stables is the medical director of EMIS which is a commercial limited company that has developed implemented clinical management systems for primary care. Richard Fitton is a director of Foldercare Health Systems Ltd which is owned by staff, patients and families. Its objectives are to develop patient-accessed records to support patient self-care. The

COMMENTARY
‘You can’t buy happiness, but thankfully you can sell it.’

Let’s explore the nirvana in the ‘scenarios that will be taking place next year’.

Despite an excellent computer system, we have an inadequate referral letter. The older patient, like so many who are ill, has welcomed Microsoft Vista™, memorised the web address, and learned to access the records remotely using a secure token and password. Having rubbed the cyber lamp, the information genie that was missing becomes apparent, even though the referral letter was generated automatically from the same consultation records. The patient is, of course, unconcerned that the orthopaedic surgeon now knows all about her recent dystrophic dyspareunia. They all had a giggle about the curious 1960 episode of the Taiwan conference, the sales director, and the unusual cigarette.

The patient admitted with chest pain interrupts the A&E doctor in his urgent duties with the display of the previous X-rays and ECGs. Intrigued, but realising these are currently as influential in management as the mp3 recording of the grandson’s trombone concert on the same PDA, our hapless F2 cracks on with acquiring the contemporaneous information upon which true emergency care is based.

Crisis resolved, the retrospective data may fine tune care. If NHS Connecting for Health hadn’t interfered we might now have had messaging systems for requesting and transferring relevant and necessary information between excellent small discrete and discreet databases using technology that has been established throughout the lifetime of your average Pentagon hacker.

In Leeds, another GP has confused another pensioner. This quack has failed to use his silver tongue, omitted the information sheet he might have printed — jargon minimised and reading age-adjusted — and missed the opportunity to print the cholesterol graph. And yet, this same GP has managed to document a few good words so fluent that his very thoughts are revealed as though from a mind map, erupting with all the relevant web links.

In Valencia, the tourist recovers despite the doctors being able to see his previous 24-hour ECG tape. Having similarly discovered that the tourist knew of his benign palpitations before booking the holiday, his insurers decline to fund his repatriation. In resolving that little misunderstanding, his wife discovered from the printout that, just before departure, our holiday marker told his GP that ‘………

Few people enjoy true autonomy outside the consultation. Patients with the greatest health needs, who have the greatest need of privacy, would be least able to protect their own records.

Many patients would be obliged by peer, social, familial, and financial pressures or intimidation to provide copies of, or access to, their internet record if the possession of a ‘HealthSpace’ record were to become standard. Parents of adolescents, employers of their staff, ‘friends’ of the elderly, and abusers in domestic violence will all want access. ‘HealthSpace’, as proposed, is a major threat to patient privacy. The creation of an internet record should require patient consent in the first instance and the HealthSpace record must be invisibly editable at the request of the patient by clinicians before it is placed on the internet, and at any subsequent stage.

Computer multinationals sold happiness to Tony Blair. What will the NHS end up buying?

Paul Thornton

REFERENCES
1. Caption from a cartoon distributed some years ago by Action on Smoking and Health featuring a tobacco salesman in front of his escalating sales chart. If anyone has a copy of the original we would be delighted to give full credit.
company is currently discussing conversion to a community interest company. Brian Fisher is a director of PAERS which has developed a system that enables patients to see their own records. He is paid both by PAERS and by Connecting for Health for consultancy. Brian Fisher is funded by Connecting for Health for research into patients’ experiences of record access.

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Mike Fitzpatrick

Blair’s legacy: a ‘patient-centred’ NHS

When Tony Blair visited the offices of the King’s Fund, Labour’s favourite health policy think tank, to give a speech commemorating a decade of NHS reforms, he received a generally positive response from the assembled ranks of health experts and professionals.1 King’s Fund chief executive, Niall Dickson, set the tone in his congratulations to the prime minister on the eve of his departure for providing ‘unprecedented levels of funding’ and ‘significant improvements in key areas’.

Speaking to the King’s Fund conference immediately before the prime minister, David Pink, chief executive of the Long Term Medical Conditions Alliance, a consortium of more than 100 patient organisations, welcomed the commitment of the government to a ‘patient-centred’ NHS. As Pink acknowledged, the presence on such a platform of somebody speaking from the perspective of patients was a potent symbol of the transformation of health policy under Tony Blair. He enthusiastically endorsed a number of initiatives that, while purporting to advance patient interests, reveal the destructive consequences of the government’s attempts to reform health care according to the rhetoric of choice and empowerment.

Pink welcomed the Quality and Outcomes Framework (QOF) as ‘a major national programme that has turned the NHS’s attention to helping people monitor and maintain their own health’. In practice, QOF operates as a financial lever to shift the direction of medical practice away from the diagnosis and treatment of disease towards intrusive and moralising interventions in patients’ personal lives, justified by the dogma that this improves health and prevents disease. Far from benefiting those with chronic conditions, this shift is depriving them of continuity of care, while doctors’ energies are consumed with medicalising the worried well. The provision of an incentive bonus to GPs for recording patients’ preferences about where they would like to die — a choice over which neither patient nor doctor is, in practice, able to exert much influence — aptly symbolises QOF’s contribution to patient empowerment.2

Another government health initiative approved by David Pink is the Expert Patient Programme (EPP), a series of formal training sessions through which people learn to manage their own chronic illnesses. Though the EPP has a commonsensical appeal, it is imbued with bad faith. It offers an illusory empowerment to patients with chronic illness and an illusory relief from the burden of caring for patients with chronic illness to doctors. If the EPP were widely taken up, it would affirm an identity as sufferer from chronic illness for a growing proportion of the population (17 million according to the Long Term Conditions Alliance) while imposing an increasing burden of responsibility for their own care on those with chronic disease.

While patronising patients, EPP implicitly degrades doctors, devaluing medical science and professional expertise.

David Pink is also a staunch advocate of ‘patient and public involvement’ in the NHS, another of the favoured slogans of the Labour government. This has led to the cultivation of the professional patient who purports to express the interests of patients in general. Of course, members of the public who are able and willing to assume these roles are inevitably unrepresentative of patients in general, and — unlike local councillors and MPs — are not subject to the episodic recall of the ballot.

In his enthusiasm for ‘patient and public involvement’ David Pink personifies the problems of these initiatives. While he speaks on behalf of people with chronic illnesses, he was not elected by people with chronic illnesses and he is in no way answerable or accountable to them. As the chief executive of a meta-quango, which strictly represents a number of organisations rather than individuals with chronic illnesses, he is as remote from such individuals as any politician (and vastly more remote than the average GP).

In fact, what emerges is that his status is conferred by government endorsement of his position rather than by any democratic mandate. It is thus perhaps not surprising to find that Pink provides the perfect warm-up man for Tony Blair on his tenth anniversary celebrations.

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