Implementing clinical governance: turning vision into reality

Aidan Halligan and Liam Donaldson

BMJ 2001;322:1413-1417
doi:10.1136/bmj.322.7299.1413

Updated information and services can be found at:
http://bmj.com/cgi/content/full/322/7299/1413

These include:

Data supplement
"Further details"
http://bmj.com/cgi/content/full/322/7299/1413/DC1

References
13 online articles that cite this article can be accessed at:
http://bmj.com/cgi/content/full/322/7299/1413#otherarticles

Rapid responses
8 rapid responses have been posted to this article, which you can access for free at:
http://bmj.com/cgi/content/full/322/7299/1413#responses

You can respond to this article at:
http://bmj.com/cgi/eletter-submit/322/7299/1413

Email alerting service
Receive free email alerts when new articles cite this article - sign up in the box at the top left of the article

Topic collections
Articles on similar topics can be found in the following collections

Quality improvement (including CQI and TQM) (598 articles)

Correction
A correction has been published for this article. The correction is available online at:
http://bmj.com/cgi/content/full/323/7314/685/b

Notes

To order reprints follow the “Request Permissions” link in the navigation box

To subscribe to BMJ go to:
http://resources.bmj.com/bmj/subscribers
Implementing clinical governance: turning vision into reality

Aidan Halligan, Liam Donaldson

Clinical governance was the centrepiece of an NHS white paper introduced soon after the Labour government came into office in the late 1990s. The white paper provides the framework to support local NHS organisations as they implement the statutory duty of quality, which was placed on them through the 1990 NHS act. Clinical governance provides the opportunity to understand and learn to develop the fundamental components required to facilitate the delivery of quality care—a no blame, questioning, learning culture, excellent leadership, and an ethos where staff are valued and supported as they form partnerships with patients. These elements have perhaps previously been regarded as too intangible to take seriously or attempt to improve. Clinical governance demands the re-examination of traditional roles and boundaries—between health professions, between doctor and patient, and between managers and clinicians—and provides the means to show the public that the NHS will not tolerate less than best practice.

In 1998 Scally and Donaldson set out the vision of clinical governance: “A framework through which NHS organisations are accountable for continually improving the quality of their services and safeguarding high standards of care by creating an environment in which excellence in clinical care will flourish.” In this paper we take the story forward. Two years on, how is clinical governance faring in the NHS, and, with the this paper we take the story forward. Two years on, how is clinical governance faring in the NHS, and, with the introduction of governance in the NHS, structures have been put in place to set standards and ensure that they are met.

New approaches are needed to leadership, strategic planning, patient involvement, and management of staff and processes. The NHS Clinical Governance Support Team is providing task based training for health professionals, who learn as they do.

Summary points

Clinical governance represents the systematic joining up of initiatives to improve quality Since the introduction of governance in the NHS, structures have been put in place to set standards and ensure that they are met.

Why clinical governance?

For most of its first 40 years the NHS worked with an implicit notion of quality, building on the philosophy that the provision of well trained staff, good facilities, and equipment was synonymous with high standards. The quality initiatives that followed, such as medical and clinical audit, took a more systematic approach. However, they were often criticised as professionally dominated and somewhat insular activities whose benefits were not readily apparent to the health service or to patients.

During the 1980s, managers and policymakers in many parts of the public sector, including health care, tried to apply the approaches of total quality management and continuous quality improvement. These approaches, which were developed in Japanese industry, were not widely accepted, perhaps because they were viewed as too management driven with no clearly identified role for clinical staff.

An internal market was introduced into the NHS in the early 1990s, but there was little evidence that opportunities were taken to embed quality improvements into the health service at a structural level. However, around the same time the NHS was given a national research and development function, and this forced it to re-examine the role of clinical decision making in improving quality. Adoption of the philosophy of evidence based medicine has resulted in more effective and consistent transfer of the lessons of research into routine practice. This has been carried forward as a core component of clinical governance.

Clinical governance was introduced at the end of a decade in which quality had been more explicitly addressed than ever before. It offers a means to integrate previously rather disparate and fragmented approaches to quality improvement—but there was another driver for change. The series of high profile failures in standards of NHS care in Britain over the past five years caused deep public and professional concern and threatened to undermine confidence in the NHS. Unwittingly, these events seem to have fulfilled a key criterion for achieving successful change in organisations—the need to establish a sense of urgency.
Key elements of the NHS quality strategy

Standards:
- National Institute for Clinical Excellence
- National service frameworks

Local duty of quality:
- Clinical governance
- Controls assurance
- Assuring quality of individual practice:
  - NHS performance procedures
  - Annual appraisal
  - Revalidation

Scrutiny:
- Commission for Health Improvement
- Educational inspection visits

Learning mechanisms:
- Adverse incident reporting
- Learning networks
- Continuing professional development

Patient empowerment:
- Better information
- New patient advocacy service
- Rights of redress
- Patients’ views sought
- Patients involved throughout the NHS

Underpinning strategies:
- Information and information technology
- Research and development
- Education and training

Framework to support quality improvement

Clinical governance is the central element of a framework that supports the delivery of quality. The box lists the national structures and mechanisms that help to develop and reinforce local clinical governance. The National Institute for Clinical Excellence and national service frameworks are important in setting quality standards. The National Institute for Clinical Excellence has a key role in appraising new technology (such as drugs and medical devices), providing guidance on the appropriate use of treatment interventions and procedures, and developing clinical guidelines for the management of specific diseases. The institute also produces clinical audit tools to support clinicians in local clinical governance activities. National service frameworks define evidence-based best practice for specific chronic diseases or patient groups. The standard setting mechanisms of these bodies are reinforced by the Commission for Health Improvement, which inspects clinical governance arrangements and provides feedback to local NHS organisations to inform development.

Policies to deal with poor practitioner performance and to learn effectively from adverse events and errors have been added to clinical governance structures to improve the safety of the clinical environment. The national system of rapid assessment to examine concerns about a doctor’s practice will enable poor performance to be recognised earlier and tackled through a range of flexible interventions. It will also be more effectively linked to a reformed system of professional regulation.

The NHS plan has strengthened ways in which patient and citizen participation can influence the quality of health services. A patient advocacy and liaison service will be established, and patient advocate teams (with access to chief executives and with their own executive powers) will be available for patients and their families. The plan commits the NHS to improving patient information, consent, and participation. There will be patients’ forums and more lay contribution through trust boards to the work of the National Institute of Clinical Excellence, Commission for Health Improvement, and professional regulatory bodies, as well as to the work of the new NHS Modernisation Board. A new NHS charter will formalise these commitments.

What might clinical governance look like on the ground?

From listening to NHS audiences across England over the past two years, we sense that healthcare professionals feel clinical governance is the right idea. Most want to work in an organisation with a strong positive culture of teamwork, and all want to find better ways to deliver quality care.

Delivery of clinical governance will include new approaches to leadership, strategic planning for quality, patient involvement, information and analysis, the management of staff, and process management. There is no one way to develop each of these areas, but certain underpinning organisational attributes are essential to successful implementation. Whatever their style, organisations need a clear understanding of what might be expected under each criterion.

Effective leadership

An organisation benefits from being clear about (and being able to describe) how it is led and how this leadership is followed through at every level in the organisation. A well-led organisation will know how the vision, values, and methods of clinical governance are being communicated effectively to all staff. Such communication gives staff a common and consistent purpose and clear expectations. Good leadership empowers teamwork, creates an open and questioning culture, and ensures that both the ethos and the day to day delivery of clinical governance remain an integral part of every clinical service.

Planning for quality

Clinical governance cannot be developed by doing what “seems right.” Health organisations need a plan to develop the quality of their clinical services. The plan should be based on an objective assessment of the needs and views of patients, assessed exposure to clinical risk, regulatory requirements, staff capabilities, unmet training needs, and a realistic appreciation of how present performance compares with that of similar services and best practice standards. It is also important to ensure that key underpinning strategies (such as information technology, education and training, and research and development) are serving the purposes of quality assurance and quality improvement. Ownership of the plans needs to be generated not just at board level but right down the organisation in individual teams.

Being truly patient centred

Health organisations must be clear how information and feedback from former and current patients is used to assess and improve the quality of services.
Empowering patients with information, and increasing their contribution to planning services, can greatly influence the development of clinical governance. Contributions from patients will affect not just the responsiveness and performance of services but the process through which quality improvement initiatives are identified and prioritised (box).

All staff need to be patient centred in their work—from the doctor discussing treatment options with a patient in the consulting room, to the primary care nurse ensuring that the elderly diabetic woman can get in contact for advice if she has worries, to the hospital manager spending time in wards and clinics to see the care patients receive and listen to their comments.

**Information, analysis, insight**

A health organisation establishing a culture of clinical governance must develop excellence in the selection, management, and effective use of information and data to support policy decisions and processes. For information and data to be useful they must be valid, up to date, and presented in a way that provides insight. Good data and information used to highlight, for example, differences in outcome, shortfalls in standards, comparisons with other services, and time trends, are essential. This information is vital to tell staff how they are doing and show where there is room to do even better (box).

**Ordinary people doing extraordinary things**

People who work in the NHS must be able to make the best possible contribution, individually and collectively, to improving health care. The ideal of a service that enables all staff to develop and use their full potential, which is aligned with the organisation’s objectives, is rarely met.

One step towards this goal is for education and training to support the organisation’s implementation of clinical governance so that knowledge and skills are reinforced in the workforce. However, developing a workforce that is fit for purpose goes much wider than this. At the most basic level it means ensuring that staff feel valued, that they share in the policy discussions about developing clinical governance, and that management is seen to be trying to tackle their problems and concerns as well as seeking their ideas for improvement and innovation.

An effective workforce also needs appropriate technical support—for example, access to valid best evidence to support clinical decisions. Finally, the creation of a culture that is free of blame and encourages an open examination of error and failure is a key feature of services dedicated to quality improvement and to learning.

**Good service design**

It is important to step back and examine how processes in the delivery of health care can be better designed. An organisation working towards implementing clinical governance could begin to describe how new, modified, and patient specific services are designed and implemented. It could include how changing patient requirements and changing technology are incorporated into healthcare service designs; how processes for delivering healthcare services are designed to meet patient, quality, and operational requirements (including best practice requirements); and how design and delivery processes are coordinated and tested to ensure trouble free and timely introduction and delivery of services. An integral part of process management includes examining how processes to design healthcare services are evaluated and improved to achieve better performance.

**Demonstrating success**

The ability to measure the quality of services is essential for successful implementation of a culture that supports clinical governance. Measures of effectiveness might include waiting times and turn around times, are essential. This information is vital to tell staff how they are doing and show where there is room to do even better (box).

---

**Case study: family centred care for children with complex needs**

**Traditional management (as described by professionals from the NHS Trust)**

- Children were referred to each therapist individually
- After referral the child and family attended clinics in various places at various times
- Reports were returned, at various intervals, to the referring doctor
- Reports were reviewed in isolation—without the benefit of collaboration between professionals
- The child and family attended several clinics and often became confused about objectives, possibilities, work required, etc.
- It could take 2 years for a child to reach the end of the evaluation process, during which time need had often changed

**Response**

Multiprofessional collaboration has facilitated the design of a family focused, effective, speedy package of care that is planned and delivered to suit the convenience and needs of the child and his or her family. A new service is currently being piloted.

**Solution**

- On first referral a child is visited at home by a member of the locality based team and the family’s health visitor
- Areas of need are identified and appropriate professionals arrange to assess the child and family at a time and place that suits the child and family
- All assessments are completed within 6 weeks and a single needs assessment report is produced in conjunction with the child and family
- Together, the child, family, and healthcare professionals agree the goals of healthcare intervention and formulate an action plan
- Progress is reviewed regularly

---

**Case study: sharing information to improve quality in trauma and orthopaedics**

**Traditional system (as described by professionals from the trust)**

- No system for benchmarking performance against national outcomes
- Clinical data were not shared
- No forum for discussion of clinical incidents or complaints
- No system to agree and implement new policy, guidelines, or protocols

**After multidisciplinary review and agreement of shared objectives**

- Weekly team meeting of all 7 surgeons, nurses, physiotherapists, managers, and junior doctors
- Care pathways and protocols have been agreed and are shared
- Agreed mechanism exists for implementing national recommendations and guidelines
- Mechanisms have been developed to review and deal with clinical incidents and complaints
- Clinical outcome data are shared and reviewed to allow modification of practice across the service
- Clinical outcome data are collected for benchmarking purposes
Education and debate

Urology: discharge summaries found to be of variable quality and value. Management protocols have been developed in an attempt to address these shortcomings. All professionals have agreed to implement the Edinburgh postnatal depression scale, training has been agreed and implemented, and interprofessional, evidence based assessment and management protocols have been developed.

Primary care groups where care of women with postnatal depression was found to be “hit and miss”—Early warning signs were often missed. Women with postnatal depression, and staff supporting them, felt that there was inadequate support. All professionals have agreed to implement the Edinburgh questionnaire, and the support for women with postnatal depression has improved. Examples of improvement initiatives undertaken by delegate teams are available at www.cgsupport.org.

Clinical governance development programme

The NHS Clinical Governance Support Team was established in 1999 to support the development and implementation of clinical governance. The team is now a part of the Modernisation Agency. Its aims are to promote the goals of clinical governance throughout the health service; to act as a focus of expertise, advice, and information; and to offer a training and development programme for clinical teams and NHS organisations.

The team runs a clinical governance development programme for multidisciplinary delegate teams drawn from organisations across the NHS. Delegate teams attend a series of five, task oriented workshops (learning days) punctuated by eight week action intervals spread over nine months. During this time delegates lead project teams in their organisations as they review, design, and deliver quality improvement initiatives. To date, 250 organisations have committed multidisciplinary teams to the five day programme.

The support team reinforces top down support for delegates by visiting health organisations and meeting the boards. The team helps boards to understand what staff have already achieved and plan support structures and dissemination strategies to spread clinical governance initiatives throughout the organisation. The visits help the board to develop an organisational culture that supports whole system, multilevel improvement initiatives and healthcare professionals who “learn as they do.”

The programme follows the RAID (review, agree, implement, demonstrate) model (figure) to initiate a project culture within their organisation. The first stage is a large scale review of current service. Delegates gather staff and patient views, come to understand and define the baseline existing service, and collect evidence about current best practice. The process encourages the examination of traditionally accepted unwritten rules and beliefs.

The agreement phase involves flagging up the route to initiate improvement. It ensures that all healthcare alliances and partners have been involved and are contributing to defining a vision for the service. This phase is about winning “hearts and minds.”

The implementation phase capitalises on the enthusiasm previously generated. Healthcare professionals are keen to measure, to know, and to prove that they are making an important difference for patients. They move naturally into the demonstration phase, where improvement activities are reflected in hard data that is then used to inform future development.

Each team of delegates works with a support team programme manager, who makes regular site visits. Delegates are helped to identify existing resources within their organisation and to secure more if necessary. Training, research, and educational materials are made available, and delegates have telephone and electronic access to the team and programme managers for advice and support. The box gives some examples of improvement initiatives that have been introduced by delegates. Further details of the programme are available on the BMJ’s website.

Conclusions

The first investigations into failing services carried out by the Commission for Health Improvement showed organisations that were poorly led. There were cliques and factions among groups of staff, management was ineffective, staff with concerns about standards of care were marginalised or worse, adequate systems were not in place, and the service was not seen through the patients’ eyes. The fact that these dysfunctional organisations were associated with such poor quality care will not surprise anyone who has read the succession of inquiry reports into NHS failings over the past 10 years.

The NHS has been late in realising that healthy organisations matter to patients. The challenge of
 clinical governance is to transform the culture and service delivery of NHS organisations throughout the United Kingdom. This revolution has begun.

We thank Laurence Wood, Ron Cullen, Eileen Smith, and Susanna Nicholls.
Competing interests: None declared.

10 Malcolm AJ. Enquiry into the bone tumour service based at the Royal Orthopaedic Hospital, Birmingham: Birmingham Health Authority, 1995.

Measuring quality of life
Who should measure quality of life?

Julia Addington-Hall, Lalit Kalra

One of the reasons behind the rapid development of quality of life measures in health care has been the growing recognition of the importance of understanding the impact of healthcare interventions on patients' lives rather than just on their bodies. This is particularly important for patients with chronic, disabling, or life threatening diseases who live without the expectation of cure and have conditions that are likely to have an impact on their physical, psychological, and social wellbeing.

Health professionals frequently make quality of life judgments when making decisions about the care of disabled patients, and the professional's view on expected quality of life is often the key factor in determining whether effective treatment for a life threatening condition will be given or withdrawn. Professionals' perceptions may, however, be at odds with those held by their patients. It is therefore important to ask patients to assess their own quality of life using one of a growing number of reliable and valid measures.

Choosing an appropriate measure and using it in clinical practice can be problematic. Deciding to use a measure, however, presupposes that patients are able to assess their own quality of life and complete a quality of life measure. Some patients—and in some conditions many patients—are unable to do this because of cognitive impairments, communication deficits, severe distress caused by their symptoms, or because the quality of life measure is too burdensome physically or emotionally. These may be precisely the patients for whom information on quality of life is most needed to inform decision making. Rather than lose all information on that patient, someone else (a family member or health professional) may be asked to act as a proxy or surrogate.

In this paper the use of proxies to measure quality of life is addressed. We consider the advantages and disadvantages of using proxies to rate quality of life, debate the reasons why a proxy's view and a patient's view may differ, and suggest directions for future research.

Can proxies provide useful information on quality of life?

Quality of life tools measure subjective experience. Completing a quality of life measure on behalf of someone else requires proxies to put themselves in another person's shoes, to imagine what it feels like to...