A guide to clinical governance reviews

Acute trusts
The Commission for Health Improvement (CHI) is a non departmental public body. It was established under the 1999 Health Act as part of the government’s reforms to help improve patient care. It has statutory powers and is accountable to government for its work, but operates independently. It has been apparent for some time that the standard of care offered by the NHS varies greatly. It can vary between hospitals, between departments in the same hospital and between general practices.

There is not always an obvious reason for this variation. CHI’s purpose is to assist the NHS in England and Wales to address unacceptable variations in patient care and to assure, monitor and improve the quality of clinical care. It collaborates closely with the NHS as well as with other bodies such as the royal colleges, professional organisations, regulatory and voluntary bodies.

Its main functions are:

- to provide independent scrutiny of local clinical governance arrangements to support, promote and deliver high quality services. CHI is carrying out a rolling programme of reviews of clinical governance arrangements in every NHS organisation
- to conduct or assist with investigations into serious service failures. CHI has the capacity for rapid investigation
- to carry out studies that monitor and review the implementation of national service frameworks, National Institute of Clinical Excellence (NICE) guidance and other key NHS policy priorities
- to provide national leadership to develop and disseminate clinical governance principles and to identify and share good practice

CHI has adopted six key principles that underpin all its work:

- the patient’s experience is at the heart of CHI’s work
- CHI will be independent, rigorous and fair
- CHI’s approach is developmental and will support the NHS in continuous improvement
- CHI’s work is based on the best available evidence and focuses on improvement
- CHI will be open and accessible
- CHI will apply the same standards of continuous improvement to itself that it expects of others
PREFACE

CHI has written this guide mainly for those involved in the clinical governance review process, including managers, clinical and non-clinical staff. It may be of interest to government departments, other regulatory and audit agencies in the NHS, academics and NHS commentators, patient representative bodies and members of the public. It is available on CHI’s website www.chi.nhs.uk. (See Appendix A for contact details).

The clinical governance review process explained in this edition of the guide has recently been improved. It is more efficient, reducing the time taken to conduct a review to 17 weeks. We have done this by making existing tools more effective, introducing new methods and removing those components that have not worked so well.

This guide is part of CHI’s commitment to being open and accessible in all its work. It describes what is involved in the clinical governance review process and explains CHI’s methods. It sets out the review framework and process, the data and documents required and the rationale for each stage of the process.

Trusts will benefit most from reviews if they are well prepared in advance. We hope that this guide will be a useful contribution to that preparation and assist trusts in their progress in implementing clinical governance. In addition, it includes appendices that show in detail the issues CHI will be focusing on during a review.

The new review approach was tested at two trusts in the last quarter of 2001 (Appendix B). CHI will continue to refine its review methods, meaning that aspects of the methodology will develop as CHI carries out more clinical governance reviews.

CHI would like to thank all those who helped to develop clinical governance reviews, including those trusts where the review methods were piloted. A key feature of the reviews is the involvement of NHS professionals as review team members and we are grateful both to them and to their seconding organisations. Each team also includes a lay member, who can take the viewpoint of patients and the public, and we are fortunate to have their involvement. They all have the opportunity to encourage good practice throughout the NHS and thus be a force for change and continuous improvement in patient care.

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1. THE CLINICAL GOVERNANCE REVIEW FRAMEWORK

What is clinical governance?

The government’s white paper, A First Class Service, and the Welsh Office paper, Quality Care and Clinical Excellence, defined clinical governance as ‘a framework through which NHS organisations are accountable for continuously improving the quality of their services and safeguarding high standards of care by creating an environment in which excellence in clinical care will flourish.’

The purpose of clinical governance is to ensure that patients receive the highest quality of NHS care possible. It covers the organisation’s systems and processes for monitoring and improving services, including:

- patient and public involvement
- risk management
- clinical audit
- clinical effectiveness programmes
- staffing and staff management
- education, training and continuing personal and professional development
- use of information to support clinical governance and health care delivery

Effective clinical governance should therefore ensure:

- continuous improvement of patient services and care
- a patient centred approach that includes treating patients courteously, involving them in decisions about their care and keeping them informed
- a commitment to quality, which ensures that health professionals are up to date in their practices and properly supervised where necessary
- a reduction of the risk from clinical errors and adverse events as well as a commitment to learn from mistakes and share that learning with others

Aims of clinical governance reviews

CHI’s rolling programme of clinical governance reviews will cover every NHS organisation in England and Wales. The reviews look at the effectiveness of organisations’ clinical governance arrangements and have five principal aims:

- to provide the public and people using NHS services with objective and fair assessments of NHS organisations’ progress towards introducing effective clinical governance
- to help the NHS achieve evident and continuous improvements in the quality of patient care
- to help the NHS reduce unacceptable variations in the quality of clinical services
- to identify and disseminate good practice in clinical governance

to increase understanding of clinical governance and the factors that determine its effectiveness

Guiding principles

CHI’s reviews of clinical governance incorporate the six key principles that guide all its work.

The patient’s experience is the central focus. The inclusion of a lay member in every review team reinforces this focus. Reviews capture information about the direct experience of NHS patients across the services they use in a trust. They also look at how the trust perceives the experiences of the patients it treats. CHI is particularly interested in waiting times, how care is organised, whether patients are treated with privacy, dignity and respect, environmental issues such as cleanliness and clinical effectiveness and outcomes.

CHI has designed the review process to be independent, rigorous and fair. CHI and its review teams collect, analyse and assess evidence according to a consistent framework (see ‘Assessing clinical governance’ in this chapter). The health care professionals in the review team work within the NHS and understand the overall context of acute trusts but do not work in the area in which the trust under review is located. All review team members undergo a rigorous selection process and are chosen for their ability to take an objective and independent standpoint.

The review process is about development and support for continuous improvement. CHI helps trusts to plan and prepare for the review, using existing information wherever possible. This process helps the trust to look carefully at its own performance. CHI also shares notable practice identified during reviews.

CHI’s work is evidence based and focused on achieving improvement. Review findings are based on robust evidence collected before and during the review. Review reports do not contain specific recommendations for change. Instead they highlight areas for action. The trust also receives a separate document containing a structured evidence summary to help them work out the most appropriate means of achieving change in its specific context. The trust then produces an action plan, in response to the review.

CHI is publishing this guide as part of its commitment to being open and accessible about every aspect of the review process and its development. In addition, all review reports are published in hard copy and on the internet, once the trusts have agreed their factual accuracy.

CHI applies the principle of continuous improvement to itself and its review methods. It recognises that there is much to learn from other review and inspection bodies and seeks to combine their best practice with its own. Using the lessons learned since the review programme began in October 2000, CHI has redesigned its clinical governance review process, drawing on the experience of both CHI staff and NHS organisations. This new process was introduced in January 2002.
Assessing clinical governance

CHI has developed a systematic framework for assessing clinical governance in trusts so that judgements made in reports of reviews are reliable, fair and consistent. The assessment framework was developed with the NHS Clinical Governance Support Team in England and the Clinical Effectiveness Support Unit (CESU) in Wales to ensure that consistent messages are given to trusts about clinical governance. (See Appendix A for contact details).

CHI’s model for clinical governance (Figure 1) illustrates its belief that effective clinical governance depends upon a culture of continuous learning, innovation and development and will improve patient experience of care and treatment in the NHS. CHI uses the information it accumulates from reviews to help determine which aspects of clinical governance are the most important for improving patients’ experience and outcomes.

Figure 1: CHI’s model for clinical governance

Work is in progress to identify the dimensions of the patient experience and outcomes under the ‘Results’ part of the model so that CHI can assess the information it collects about what it is like to be a patient and interpret information about clinical processes and care outcomes. CHI looks specifically at the environment, privacy and dignity, clinical effectiveness and outcomes, access and organisation of care.

CHI evaluates clinical governance by exploring three key, interlinked areas identified in the model:

- strategic capacity: how far does the trust’s leadership set a clear overall direction that focuses on patients? How well is it integrated throughout the organisation?
- resources and processes: how robust are its processes for achieving quality improvement, such as patient and public involvement and clinical audit?
How effective are the trust’s arrangements for staff management and development?

- **use of information:** what information is available on patients’ experience, outcomes, processes and resources, and how does the trust use it strategically and at the level of patient care?

Each of these areas comprises a number of components that CHI examines in every trust. CHI has so far identified seven components of ‘RESOURCES AND PROCESSES’ and ‘USE OF INFORMATION’ (Figure 2). Work is being carried out to identify the components of ‘STRATEGIC CAPACITY’.

**Figure 2: Components of clinical governance - resources and processes and use of information**

<table>
<thead>
<tr>
<th>COMPONENT</th>
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<tr>
<td>Resources and processes</td>
<td>Patient and public involvement</td>
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<td>(i) processes for quality improvement</td>
<td>Clinical audit</td>
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<td></td>
<td>Risk management</td>
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<td>Clinical effectiveness programmes</td>
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<td>(ii) staff focus</td>
<td>Staffing and staff management</td>
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<td></td>
<td>Education, training and continuing personal and professional development</td>
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<tr>
<td>Use of information</td>
<td>Use of information to support clinical governance and health care delivery</td>
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CHI’s review teams assess how well clinical governance is working throughout the trust by making enquiries about each of these seven components at corporate and directorate levels and in clinical teams. This involves collecting information systematically about review issues that have been defined for each component. To help with analysis and reporting, the review issues are grouped into themes:

- accountability and structures
- strategies and plans
- application of policies, strategies and plans
- quality improvements and learning
- resources and training for staff

The full set of review issues for each component and their grouping into themes is included in Appendix C.

CHI will introduce similar methods to assess information collected about components of ‘STRATEGIC CAPACITY’ in future rounds of reviews. Dimensions may include: partnership working; leadership; direction and planning; and patient involvement.
On the basis of the evidence collected, CHI’s reviewers assess each component of clinical governance against a four-point scale:

- **I** = little or no progress at strategic and planning level, or at operational level
- **II** = worthwhile progress and development at strategic and planning levels but not at operational level or:
  - worthwhile progress and development at operational level but not at strategic and planning levels
  or:
  - worthwhile progress and development at strategic and planning levels and at operational level but not across the whole organisation
- **III** = good strategic grasp and substantial implementation. Alignment across the strategic and planning levels, and the operational level of the trust
- **IV** = excellence – coordinated activity and development across the organisation and with partner organisations in the local health economy that is demonstrably leading to improvement. Clarity about the next stage of clinical governance

There is wide variation within trusts in progress made developing the component parts of clinical governance. At this stage of development, CHI believes it is most useful to trusts to assess each component separately to help them prioritise their development of clinical governance. It will not make judgements to produce an overall rating for an organisation.

**Review phases**

Reviews take approximately 17 weeks to complete from the start of the review to the final preparation of the report. Prior to this the trust has three months in which to collect data and information requested by CHI. This timescale is long enough to collect and rigorously analyse data, but intensive enough to mean that the evidence on which the review findings are based is current and useful.

There are five key phases in a review: pre review; pre visit preparation; visit week; reporting; and action planning. Each review follows the same timetable:

- **Pre review (weeks -12 to 0)**
  During this phase, CHI sends the trust an information manual detailing the data and information required and collects the data and information requested.

- **Pre visit preparation (weeks 1-7)**
  Between weeks one and seven, CHI holds an initial meeting with the trust, collects information from stakeholders, analyses data, identifies the areas for detailed review during the visit week and gives the trust preliminary feedback.
- **Review week (week 8)**

A CHI review team visits the trust to interview staff, observe practice, verify information already obtained and gather further information. Feedback on the issues identified during the visit is given to the trust on the final day of the visit.

- **Reporting (weeks 9 - 17)**

With the information gained from the review, the review manager writes a report on CHI’s findings. The report is quality assured internally, sent to the trust for comments on factual accuracy, finalised and published in week 17. The report is publicly available in hard copy and on CHI’s website.

- **Action planning**

Following the review, the trust develops an action plan in response to the key areas for action outlined in CHI’s report.

Each of these phases is described in more detail in Chapters 3 – 7.
2. THE REVIEW PROGRAMME

Selecting organisations for review

CHI’s rolling programme of clinical governance reviews started with acute trusts but is now expanding into all NHS organisations across England and Wales. CHI will review all primary care trusts, NHS trusts, NHS Direct sites and, in future, local health boards in Wales (subject to legislation) on a similar basis.

CHI selects trusts to review in two ways:

- on a random basis using a sampling technique that ensures that the number of trusts selected is spread proportionately across a regional area
- where appropriate, by grouping acute, primary care trusts, ambulance and mental health organisations geographically

Reviews are not normally triggered by special concerns. However, CHI has the capacity to ‘fast track’ clinical governance reviews of certain organisations and bring these forward in the programme. ‘Fast track’ clinical governance reviews may be triggered by a request from an individual or organisation, a recommendation made as a result of a CHI investigation or where a request is made for an investigation but a review is more appropriate. All requests for ‘fast track’ reviews and investigations are assessed against a set of guiding principles. If a decision is made to ‘fast track’ a review, the trust is informed.

In addition, CHI liaises regularly with each strategic health authority, the regional directorates of health and social care and the Welsh Assembly Government to look back at completed reviews and to discuss the future review programme. This enables CHI to identify organisations where the review should be delayed, for example because of an impending merger or change of management.

Approximately six months before the start of the 17 week review process, CHI informs the strategic health authority, the regional directorate of health and social care and the Welsh Assembly Government of the proposed review schedule. Four months before the start of the review the trust is informed of its selection as a clinical governance review site. CHI also gives the trust the opportunity to highlight any reasons why it thinks the review should be delayed.

CHI teams

Three different teams work with the trust during the review process: the CHI pre review team, the CHI internal team and the CHI review team.

The pre review team deals with the information and data request period before the review begins and is the trust’s first point of contact for help and advice until the review begins.

The internal team comprises a review manager, a review coordinator, analysts and a communications officer. Their roles are as follows:
• the review manager works with the trust throughout the review, ensures that all relevant evidence is collected and analysed, leads and supports the review team during the visit week and writes the report

• the review coordinator ensures the efficient running of a review, liaising closely with the trust and the CHI teams

• the analysts analyse data provided by the trust and other national data. They also support the review by undertaking ad hoc analyses as requested and by reviewing the trust’s information systems

• the communications officer advises the internal team and, where applicable, the trust on internal and external communications issues. They also publicise the review and help to identify stakeholders

The review team, led by the review manager, carries out clinical governance reviews and is multidisciplinary. The review team includes a nurse, a doctor, an NHS manager, a lay member and another clinical professional who is not a doctor or a nurse, for example a physiotherapist. CHI informs the trust of the membership of the team in advance of the visit week.

CHI recruits reviewers through national advertising. It requires high standards of its reviewers and operates a rigorous competency based selection process. Potential reviewers attend a one day selection centre.

Once selected, all reviewers attend an intensive two and a half day training course simulating the clinical governance review process. It provides reviewers with a thorough grounding in CHI’s review methods and helps develop the skills needed for reviewing, such as interviewing and listening, note taking and analysis. CHI requires all its reviewers to comply with its code of conduct. They also sign a confidentiality agreement that continues after they have finished working for CHI as a reviewer, as well as a declaration of interests.

Reviewers are on short term secondments. They spend around 10 days on each review and normally carry out one or two reviews per year while remaining in their current job within the NHS. This means that they are up to date with current practice and understand the context within which trusts work. They can also help spread identified good practice within their own organisations. In addition, the lay reviewers bring the patient’s and the public’s perspective to the review.

CHI welcomes applications from those interested in becoming a reviewer. Please contact the human resources team at CHI or consult CHI’s website, which contains details of vacancies.
3. THE REVIEW PROCESS - THE PRE REVIEW PHASE  
(WEEKS - 12 TO 0 )

During this phase, CHI collects data and information, both from the trust and from national and other sources. The framework for assessing clinical governance determines the data to be collected. CHI analyses the data to build up a picture of the trust and to identify areas to focus on during the visit week. These include areas of notable practice and areas for further development.

CHI aims to minimise the work for the trust during this phase, ensuring that there is sufficient time for data collection (a copy of the data request is available on the CHI website). The trust is sent a comprehensive information manual explaining CHI’s data and information requirements and can gain further advice and support from the pre review team. Over time, CHI will reduce the data it requires by careful targeting so that the supply and analysis of data is more economical and effective. The redesigned clinical governance review process has already started this process, reducing and focusing the number of documents required.

Key to the review’s success is visible commitment from the trust’s senior management and communication to staff. CHI’s communications team has prepared a handbook for trusts to help them with the internal and external communications aspects of the review. CHI also offers assistance to trusts in media handling and other communications issues when reports are published.

Contacting the trust

Three months prior to the start of the review, the organisation is formally informed of its selection and of the timing of the review team visit to the trust. CHI sends an information manual to the trust signalling the beginning of the pre review phase. The manual contains information to support the trust through the review including guidance, a trust questionnaire, a data and information request, a communications handbook and timetabling information.

CHI asks the trust to nominate a key contact – the trust coordinator – to collect information and arrange local meetings. This is a crucial role (figure 3) in ensuring that the review runs smoothly and review teams value the trust coordinator’s assistance highly. The time commitment required for this role will vary from trust to trust, but on average it will involve around 30–40 days of the coordinator’s time from start to finish. The trust coordinator should have an understanding of how the organisation as a whole operates and be senior enough to have influence at board level. Appendix D describes the role of the trust coordinator in more detail.
Figure 3: Role of the trust coordinator

The trust coordinator:

- acts as a link between CHI and the trust
- communicates information throughout the trust about CHI and the review process
- works with the CHI review manager to make sure that the review process runs as smoothly as possible
- coordinates the return of the trust’s data and information
- organises the briefing and preliminary feedback meeting
- coordinates the response to the CHI presentation at the preliminary feedback meeting
- arranges the timetable for the review team’s week long visit and schedules appointments
- is available throughout the visit week
- coordinates the return of comments on the factual accuracy of the report to CHI
- arranges the action planning workshop

Collecting data and information

To inform the review, CHI collects and uses national data sets such as clinical indicators. It also asks the trust to provide:

- **documentation from internal sources**. This includes information on the trust’s profile, strategies and business plans and information about the individual components of clinical governance
- **documentation from external sources**. This includes reports of other external organisations that visit the trust, for example, external auditors and Investors in People
- an extract of data from the trust’s **patient administration system (PAS)** over the previous four years

The request for data and information is comprehensive in its coverage of clinical governance and there should not normally be a need for the trust to provide documentation in addition to that which CHI has requested. Looking at this data and these documents helps to focus the visit week and avoid duplicating work already carried out by the trust or other external reviewers.

CHI asks the trust to return the data and information in electronic form before the start of the pre visit preparation phase. CHI analysts begin to produce a summary of evidence that will assist in identifying areas to look at during the visit week.

CHI also asks the trust to complete a questionnaire about its progress towards implementing clinical governance, including public and patient involvement. This tool provides further important background information to the review team and helps to inform the planning of the visit week. It is also an opportunity for the trust to highlight areas of good practice as well as those that may require further development.
Staff survey

The staff survey is an important component of the clinical governance review and enables CHI to gain a comprehensive picture of a number of clinical governance issues through the beliefs, attitudes and perceptions of staff working in the trust. The survey is carried out by means of a questionnaire sent to a random sample of staff. The survey is confidential and the responses are anonymous.

To enable CHI (or a nominated organisation carrying out the work on CHI’s behalf) to carry out the staff survey, CHI asks the trust to provide a database of all staff currently employed by the trust. CHI uses the staff database to draw the sample of staff and send out the questionnaires. The completed questionnaires are then analysed and the results considered alongside all the other sources of information gained during the pre review phase.
4. THE REVIEW PROCESS - PRE VISIT PREPARATION
(WEEKS 1 TO 7)

Initial meeting with the trust

Once the pre review phase is complete and the review has started, the review manager, analyst, review coordinator and CHI communications officer meet with the trust to discuss the review. The meeting normally includes the trust’s chief executive, relevant board members, the trust coordinator for the review and the trust’s clinical governance lead. The aim is to explain the process, the trust coordinator’s role, the preparation needed and the support that CHI’s communications team will provide to the trust, as well as answer any queries.

Meetings with stakeholders

Achieving an understanding of the trust’s local context and external perspectives on the trust’s clinical governance arrangements, is a significant feature of the review. CHI therefore spends around three days meeting with local people and non statutory organisations with an interest in the trust. They include members of the public, voluntary and not for profit organisations, staff associations and trade unions. Staff from the trust may also attend. CHI holds the meetings at a local venue with disabled access and conducts them privately. Information may also be received by letter, telephone or email.

CHI spends a further day conducting formal meetings with the Welsh Assembly Government, the strategic health authority, the community health council, primary care trusts, local authorities with social services responsibilities and the external auditors. In addition to providing context and helping to focus the review, these meetings help to raise local awareness of the review and action planning processes.

CHI is currently researching and developing a strategy for reaching patients who might have difficulty in making their views heard. This might be because they do not speak or read and write English or because they have a sensory or physical disability. The review process in the future will include methods for seeking these people’s views.

Summary of evidence and selection of clinical teams

CHI analyses the internal and external documents provided by the trust, the staff survey, the trust questionnaire and the results from stakeholders and summarises the evidence into a table. Its purpose is to provide background information to the review team and to help identify key issues and areas of good practice to follow up through the visit week. It provides initial findings on patient outcomes and experience, strategic capacity, resources and processes and use of information (see ‘assessing clinical governance’ in Chapter 1).

The summary of evidence is a working document that is constantly updated throughout the review process and shared with the trust prior to the visit week. It informs the visit week, supports CHI’s eventual findings and is used as the basis for
the final report. It collates information in a way that the trust can use to assess its own performance.

CHI uses the summary of evidence to select three clinical teams in the trust for the review team to look at in more depth. The aim is to test whether clinical governance arrangements are working at grass roots level, not to carry out service reviews. The areas chosen are therefore not intended to be representative across the whole trust but to provide evidence of clinical governance effectiveness. CHI also gives the trust the opportunity to nominate three to five teams which it thinks represent good practice worthy of sharing elsewhere. One of these is included in the team selection.

A clinical team comprises the staff who care for specified groups of patients. It is based around staff who work on a particular ward, outpatient clinic, theatre suite or other location in a hospital but also extends to other staff who care for the patient. A clinical team caring for patients who have fractured a neck of femur for example, might be based around the staff on an orthopaedic ward. Other staff who work in the accident and emergency department, theatres, rehabilitation, pharmacy, diagnostic units and the outpatients department have an input into the care of the patient and may also be included in the clinical team.

CHI informs the trust which clinical teams it has chosen at the end of week three so that the trust coordinator can start scheduling interview appointments with staff.

**Briefing the CHI review team**

In week five, to help the review team understand both the trust’s context and how it is approaching clinical governance, CHI sends each reviewer information about the trust and a summary of findings to date.

**Preliminary feedback**

In week six, the review manager, analyst, review coordinator and the CHI review team visit the trust. The purpose of this visit is to present the data, information and emerging issues identified so far in a preliminary feedback meeting. The trust can also use this meeting to present their progress on clinical governance and update CHI on relevant information.
5. THE REVIEW PROCESS – VISIT WEEK (WEEK 8)

The purpose of the visit week is to test out areas selected in the earlier stages of the review, to validate information already collected and to gather further evidence about the trust’s progress with clinical governance. Visiting the trust also allows the review team to assess the ‘softer’ issues, such as communication between and within teams and relationships within the trust, for example between managers, clinicians and other staff.

The review manager, review coordinator, analyst and the review team spend an intensive five days on site, including at least one visit to the trust at night. An example timetable is shown at Appendix E.

For the first three to four days of the week long visit, reviewers interview staff, observe what happens in areas of the trust, and fill in any data or information gaps. The review team works in a collaborative and non confrontational way.

Every lunchtime and evening during the visit week, the reviewers meet with the review manager to discuss their findings. This allows team members to exchange notes and highlight any issues to follow up in the next round of interviews. It also enables them to identify any further evidence needed to complete the assessment.

The final part of the visit is spent conducting any necessary additional interviews and preparing to give the trust feedback. This feedback is presented on the last day of the visit.

The review manager plays an important role in coordinating the review team’s work during the visit. He or she may also participate in the interview sessions.

The review manager’s main tasks are to:

- act as the formal link between CHI, the review team and the trust
- manage the process and make sure it runs as smoothly as possible
- provide leadership and support to the review team
- quality control the process and the reviewers’ activities
- facilitate review team discussions and run debriefing and planning sessions with the review team throughout the week
- work with the analyst and reviewers to analyse information on site, allowing more immediate and structured feedback to the trust at the end of the week

**Interviewing staff**

The review team interviews a cross section of staff of all grades and professions, including non clinical staff, in scheduled interviews lasting between half an hour and an hour. Most interviews are carried out by reviewers in pairs, allowing one person to ask questions while the other takes notes. Some interviews are group interviews, but the majority are with an individual member of staff.

The structured interviews cover the main components of clinical governance. For example, as part of assessing patient and public involvement, interviews with team members seek to find out whether:
- staff are aware of patients’ views of the service and whether action is taken as a result
- staff are committed to keeping patients and carers informed of progress
- patient privacy and dignity are respected
- patients are involved in the planning and delivery of their care
- care is organised around patients’ needs
- lessons are learned from patient complaints and changes made as a result

At the end of each interview, the reviewer completes a site visit recording form.

Information provided by individual members of staff is non-attributable in CHI’s final report. However, if a member of staff raises serious issues such as allegations of professional misconduct, CHI has a responsibility to act and it may not be possible to guarantee that person’s anonymity. CHI provides training and consistent guidance to all reviewers on the procedure to follow in such situations.

**Observation**

In addition to interviewing staff, the review team carries out scheduled observation sessions in a variety of areas for example, wards, the accident and emergency department, waiting areas, the x-ray department and the cafeteria. The review team does not observe within consulting rooms, operating theatres or treatment areas.

These sessions involve talking to staff who have not been included formally on the visit timetable, visiting operational areas and watching what happens in these areas. They enable the review team to capture information about privacy, dignity and respect for patients, patient confidentiality, communication between professionals, management of environmental risks and how facilities meet patients’ needs.

The review team only talks to patients after consultation with, and with the agreement of, the member of staff in charge. Reviewers do not talk to patients without the patient’s verbal consent.
6. THE REVIEW PROCESS - REPORTING (WEEKS 9 TO 17)

Report writing

After the visit week, the review manager drafts a report. The audiences for the report are the health and social community and its users, and it is made publicly available. It is therefore written in a clear, accessible and jargon free style.

The purpose of the report is:

- to provide a picture of where clinical governance is working well and where the trust needs to take action
- to highlight areas of good practice
- to provide information for the trust to use in identifying priorities for improvement

CHI considers the evidence gathered throughout the review process (from the data and information request, the trust questionnaire, the staff survey, stakeholders, interviews and observation during the visit week). In reaching conclusions, CHI weighs carefully the robustness of the evidence. This ensures that judgements made in reports are supported by information from a number of sources (Figure 4).

Figure 4: Evaluating supporting evidence

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<tr>
<th>Degree of confidence</th>
<th>Amount of evidence and sources</th>
<th>Reporting back to the trust</th>
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<td>✓ = include in written report or verbal feedback</td>
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<td>? = may appear in written report or verbal feedback</td>
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<tr>
<td>Very confident</td>
<td>A number of sources: data; documents; interviews; observation</td>
<td>✓ Report</td>
</tr>
<tr>
<td></td>
<td></td>
<td>✓ Verbal feedback</td>
</tr>
<tr>
<td>Confident</td>
<td>Several items of information from the same source type (e.g. interviews) from different areas or organisations</td>
<td>✓ Report</td>
</tr>
<tr>
<td></td>
<td>One interview or observation confirmed by an independent source</td>
<td>✓ Verbal feedback</td>
</tr>
<tr>
<td>Some confidence</td>
<td>Several items of information from the same source type (e.g. interviews) from the same area or organisation</td>
<td>? Report</td>
</tr>
<tr>
<td></td>
<td></td>
<td>✓ Verbal feedback</td>
</tr>
<tr>
<td>Little confidence</td>
<td>One interview or observation only</td>
<td>× Report</td>
</tr>
<tr>
<td></td>
<td></td>
<td>? Verbal feedback</td>
</tr>
</tbody>
</table>

The report contains key findings and action points, as well as examples of good practice. It does not contain recommendations. This enables the trust to consider the best way of achieving change in its specific context and circumstances.
The trust receives the full summary of evidence table to assist them in action planning but this does not appear in the report.

**Quality assurance**

The report is subjected to extensive quality assurance by CHI staff before it is sent to the trust to comment on for factual accuracy. These staff include assistant directors, the communications team, members of the review team and analysts. The report is then sent to the Department of Health or Welsh Assembly Government before being sent to print.

**Publishing**

The report is published in week 17 of the review.

CHI’s communications team liaises with the trust on publication dates and, where necessary, helps with media handling. They also publicise the publication of the report.
7. THE REVIEW PROCESS - ACTION PLANNING

The trust is required to generate an action plan in response to the key areas for action outlined in the CHI report.

Action plans should have:

- clarity about the action required and accountability for the action agreed
- clear and measurable outcomes which focus on continuous improvement to the patient experience
- clarity about the system for monitoring the action plan

To help develop the action plan, CHI asks the trust to run a workshop with trust staff and with other partners in the health community. It is the responsibility of the trust to organise the workshop.

The workshop takes place in either:

- week 13 of the review when the chief executive of the trust and the review manager are confident that the discussion of key themes emerging can be inclusive and encompass all those in the health community
- or within three weeks of the CHI report being published

Once the action plan has been finalised, it is placed on the CHI website alongside the report.
GLOSSARY

**Audit:** A review that establishes how well a service meets pre determined standards or criteria

**Clinical audit:** The continuous evaluation and measurement by health professionals of how far they are meeting standards that have been set for their service (standards can be set by health professionals, themselves, or others). Successful clinical audit also involves changing practice to meet the standards

**Clinical effectiveness:** For individuals, this means the degree to which a treatment achieves the health improvement for a patient that it is designed to achieve. For whole organisations, it means the degree to which the organisation is ensuring that ‘best practice’ is used wherever possible

**Clinical governance:** A First Class Service (DoH) and Quality Care and Clinical Excellence (Welsh Office) define clinical governance as “a framework through which NHS organisations are accountable for continuously improving the quality of their services and safeguarding high standards of care by creating an environment in which excellence in clinical care will flourish”. It is about the systems the organisation has for ensuring high quality care.

**Clinical governance review:** A systematic review of the arrangements an organisation has put in place to implement clinical governance

**Clinical governance review report:** CHI’s published findings of each clinical governance review which are available to the public. The purpose of the report is to identify areas for improvement and to encourage the spread of good ideas.

**Clinical indicators:** Selected measurements of clinical care which help NHS staff to judge how well they are doing.

**Clinical information:** Any information about treatments or services which can be used by patients or health professionals to help take decisions about patient care

**Clinical risk management:** The systematic use of information and expertise of individuals within the organisation to identify and reduce clinical risks to patients

**Community health council (CHC):** CHCs represent the public interest in the NHS and have a right to be consulted on health service changes in their area

**District auditors:** The external auditors for all NHS trusts, local authorities and other bodies

**Health economy:** The collection of organisations that plan and provide health services in an area including health authorities, NHS trusts, primary care groups and voluntary health organisations

**Investors in People:** Investors in People is a national quality standard which sets a level of good practice for improving an organisation’s performance through its people

**Lay member:** A person from outside the NHS who brings an independent voice to CHI’s work

**National data sets:** A standard set of data items (statistical evidence), concepts and definitions to enable the production of national and nationally comparable data

**National indicators:** Statistics recorded by the Department of Health (DoH) on a range of specific treatments to allow comparison and measurement of NHS organisations

**NHS trust:** A self governing body in the NHS which provides health care services. They employ a full range of health care professionals including doctors, nurses, dieticians, physiotherapists etc. An acute trust –provides medical and surgical services usually in hospital

**Orthopaedics:** A branch of surgery interested in disorders and treatment of the spine and repair of joints and bones
**Performance indicators:** Nationally agreed standards and measures to indicate how well an organisation is performing

**Postgraduate deaneries:** Postgraduate deans commission, manage and develop postgraduate medical and dental education. They are responsible for the training of all medical and dental trainees within their region or part of a region (deanery)

**Primary care trust (PCT):** Primary care trusts are self governing bodies that will evolve from primary care groups. They will have the same functions as primary care groups but will also commission some hospital based health care services for their population and directly provide community health services

**Qualitative:** Data that can not be expressed using numbers e.g. interview statements, diagrams, documents

**Quantitative:** Data which can be measured in terms of numbers

**Sampling technique:** A way of selecting a small group that is representative of a bigger group or the total population

**Welsh Assembly Government:** The devolved tier of government in Wales (previously known as the National Assembly for Wales)
APPENDIX A

CONTACT DETAILS

Commission for Health Improvement

For further details about clinical governance reviews, please contact one of the following:

John Dennis – Assistant Director
Andrea Groom – Assistant Director
Jane Farleigh – Assistant Director

At:
Commission for Health Improvement
Finsbury Tower
103-105 Bunhill Row
London EC1Y 8TG
Telephone: 020 7448 9200
Fax: 020 7448 9222
Minicom: 020 7448 9292

Or:
E-mail CHI at the following address:
information@chi.nhs.uk

You can also consult CHI’s website, which includes a full description and a complete range of documents relating to the clinical governance review process, and other information about CHI’s work at:

www.chi.nhs.uk

The NHS Clinical Governance Support Team (CGST)

CGST runs a series of unique programmes to support the implementation of clinical governance ‘on the ground’. Clinical governance is the framework which helps NHS organisations provide safe and high-quality care. Fundamental to making this happen is creating and enabling a cultural, change within the NHS. Through its innovative programmes, the support team enables a wide variety of NHS organisations to involve staff and patients in improving services and to continue to do so. Clinical governance is about changing the way people work, demonstrating that leadership, teamwork and communication is as important to high-quality care as risk management and clinical effectiveness.
For further information about its work in England please contact:

**Clinical Governance Support Team**
2nd Floor, St John’s House  
East Street  
Leicester LE1 5ZW  
Telephone: 0116 295 2000  
Or access its website at: [www.cgsupport.org](http://www.cgsupport.org)

**The Clinical Effectiveness Support Unit (CESU)**

CESU closed at the end of March 2001. An Assembly-based NHS Wales Clinical Governance Support Unit has now being established as part of the NHS Quality Division. For further information about its work in Wales please contact:

NHS Quality Division  
Welsh Assembly Government  
Cathays Park  
Cardiff  
CF1 3NQ  
Telephone: 02920 826206  
Or access its website at: [www.cesu.wales.nhs.uk](http://www.cesu.wales.nhs.uk)
APPENDIX B

PILOT SITES

CHI piloted its initial review process in full at four pilot sites, starting in April 2000. They were:

Southampton University Hospitals NHS Trust
City Hospitals Sunderland NHS Trust
North West Wales NHS Trust (Bangor)
Chesterfield and North Derbyshire Royal Hospital NHS Trust

The redesigned review process was tested at two pilot sites, starting in October and November 2001 respectively: They were:

Northumberland Healthcare NHS Trust
Tameside and Glossop Acute Services NHS Trust

We are very grateful to the management and staff of all trusts for their help in developing the reviews and in our evaluation of the review methods.
APPENDIX C

REVIEW ISSUES

Patient and public involvement

Accountabilities and structures
- committee responsibilities for patient/service user/carer and public involvement
- staff responsibilities for patient/service user/carer and public involvement work
- reporting and monitoring – to/by management teams, committees and the board e.g. of trends in patient/service user-initiated areas of concern and complaints

Strategies and plans
- strategy and implementation plans for patient/service user and public involvement work
- connection of the strategy with wider clinical governance and quality improvement programmes
- involvement of patient/service users/carers, or their representative organisations, in policy and planning of services, e.g. through: public participation groups; citizen juries; stakeholder conferences; lay/citizen and patient/service user/carer representation on board and clinical governance committees
- involvement of partners in strategy development and implementation plans
- resources (staff and budget) to support the implementation of the strategy for patient/service user and public involvement

Application of policies, strategies and plans
- information to the wider public about what the organisation is doing e.g. communications work, reporting on involvement work
- information to the wider public about how well the organisation is doing e.g. performance information
- ‘customer’ care practice to ensure patient/service users’ privacy, dignity and confidentiality about themselves and their treatment e.g. codes of conduct; attitudes and behaviours of staff
- availability and quality of written or other information for patient/service users about treatments, services and facilities
- involvement of patient/service users or carers in treatment choices, including processes for patient/service users to consent to treatment
- arrangements to meet patients/service users’ particular needs e.g. cultural, dietary
- support to patients and carers in self-management of their care and treatment e.g. ‘expert patient’
- access by patient/service users to information about their care e.g. shared care plans patient held records; copies of correspondence between health professionals
- arrangements for patient/service user/carers to voice concerns, issues and compliments about services e.g. comment cards; suggestion boxes
- systems for individual patient/service user and carers to seek redress e.g. complaints system; PALS; Independent Complaints Advisory Service

**Quality improvements and learning**
- involvement of patient/service users, carers and the public in monitoring the quality of care e.g. research into patient/service user/carers’ views; monitoring and evaluation of services from patient/service user/carers’ perspectives
- analysis of all feedback (including complaints) from individual patient/service users on their experience of the organisation
- improvements to the quality of service outcomes (performance) and to the quality of decision making (governance) as a result of patient/service user involvement work
- dissemination of lessons learnt from consultation and patient/service user involvement activities

**Resources and training for staff**
- training for staff in patient/service user (customer) care; communication skills; obtaining patient/service users’ consent to treatment; confidentiality issues; complaints handling
- support for individual patient/service users e.g. patients’ advocates; support for carers; interpreters; translation services; signers; link workers
Clinical audit

Accountabilities and structures
- committee structure for clinical audit
- staff responsibilities for clinical audit
- reporting and monitoring – to/by management teams, committees and the board

Strategies and plans
- strategy for clinical audit – including priority given to participation in national, regional and local audits – and programmes
- integration of clinical audit with quality improvement programmes e.g. to audit compliance with evidence-based practice protocols, guidelines and care pathways etc
- involvement of patient/service users and carers in clinical audit strategy and programme development
- involvement of partners in cross-organisational clinical audit
- support and resources for clinical audit including:
  - central clinical audit unit to support audit design, data collection and analysis
  - budgets for clinical audits

Application of policies, strategies and plans
- clinical audits carried out including:
  - connections with other clinical governance activities
  - staff awareness and involvement
- participation in national confidential enquiries

Quality improvements and learning
- processes to consider the results of clinical audits
- compliance with evidence based practice shown by audits
- quality improvements as a result of clinical audits
- dissemination of lessons learnt from clinical audit

Resources and training for staff
- training and development for staff in audit skills
Risk management

**Accountabilities and structures**
- committee structure for clinical risk management
- staff responsibilities for risk management
- reporting and monitoring – to/by management teams, committees and the board
- strategy and implementation plans for risk management

**Strategies and plans**
- integration of all risk management activities (clinical, non-clinical, health and safety)
- integration of risk management with audit and quality improvement programmes
- consideration of risk in decision making processes
- involvement of patients, service users in carers in risk management
- involvement of partners in developing risk management strategies where risk is to patients/service users who are cared for by more than one organisation e.g. other health organisations, social services, police
- resources for risk management including:
  - budgets for risk management activities
  - specialist teams and support e.g. for infection control and pressure sore control & tissue viability

**Application of policies, strategies and plans**
- risk assessment, including:
  - the collation of information from all sources about risks and monitoring of incidents and trends
  - inclusion of information from patient/service users e.g. from complaints
  - involvement of partners e.g. at discharge meetings for users at risk
- incident and near miss reporting and investigation
- risk management, including use of trigger events, protocols
- prevention and control of specific risks e.g. for: misuse of drugs; use of medical devices; lone workers; infections; pressure sores; violence/self harm

**Quality improvements and learning**
- analysis of individual risks and events and trends
- quality improvements as a result of risk management activities
- dissemination of lessons learnt from risk management activities

**Resources and training for staff**
- training and education for staff in risk prevention and management
Education, training and continuing personal and professional development

Accountabilities and structures
- committee structure for education, training and CPD issues
- staff responsibilities education, training and CPD
- reporting and monitoring – to/by management teams, committees and the board
- strategy and plans for education, training and CPD

Strategies and plans
- links between training and CPD programmes and wider quality improvement programmes, and with individuals personal development plans
- partnerships with educational establishments; joint training programmes with partners e.g. other health organisations, social services, police
- budget for professional development, education and training (excluding SIFT and MADEL)

Application of policies, strategies and plans
- personal development planning
- mandatory training including CPR; manual handling
- work based training schemes
- CPD programmes
- schemes for obtaining relevant professional, or further, qualifications

Quality improvements and learning
- improvements to services and facilities following external assessments (e.g. by Royal Colleges) and internal evaluations of training and education programmes
- dissemination of knowledge of effective education, training and CPD methods

Resources and training for staff
- time, financial and other support for staff undergoing formal education and for individuals’ CPD activities
Clinical effectiveness programmes

Accountabilities and structures
- committee responsibilities for clinical effectiveness programmes
- staff responsibilities for clinical effectiveness programmes
- reporting and monitoring of implementation of, and compliance with, evidence based practice– to/by management teams, committees and the board

Strategies and plans
- strategy and programmes for clinical effectiveness work, including research to identify effective clinical practice
- coordination of clinical effectiveness strategy and programmes with the wider clinical governance and quality improvement programmes
- involvement of partners in clinical effectiveness strategy development and programmes
- involvement of patient/service users and carers in clinical effectiveness strategy development and programmes
- resources (staff and budget) to support research, development and implementation of the effective clinical practice

Application of policies, strategies and plans
- collection and distribution of evidence based practice to the relevant teams and staff, including:
  - results of the organisation’s own research
  - published evidence of effective practice, including NSF and guidance issued by NICE
- research projects to identify effective clinical practice
- implementation and application of effective clinical practice e.g. integrated care pathways; evidence based guidelines for disease management
- monitoring the effectiveness and application of evidence-based practice e.g. cycle of data collection; use of performance indicators; clinical audit; team discussion; guideline amendment

Quality improvements and learning
- improvements to the patient/service user experience as a result of the implementation of evidence based practice
- dissemination of learning from the implementation of evidence-based practice

Resources and training for staff
- accessibility of research results and evidence of effective practice e.g. libraries; Internet; journals; intranet (or other local electronic library)
- training for staff e.g. in critical appraisal skills; literature, database and internet search skills
Staffing and staff management

Accountabilities and structures
- committee structure for staffing issues
- staff responsibilities for staffing
- reporting and monitoring – to/by management teams, committees and the board

Strategies and plans
- strategy and workforce planning for staffing including:
  - delivery of national priorities including targets in Working Together and Improving Working Lives
  - links to service plans
  - current and future number requirements; skill requirements
- joint approaches to staffing with partner organisations, including compatible systems e.g. with social services

Application of policies, strategies and plans
- HR employment processes e.g.:
  - equality of opportunity
  - good race relations
  - checking qualifications and registration
  - disciplinary and grievance procedures
- workplace induction
- individuals’ performance appraisal
- clinical supervision and mentoring schemes
- systems for dealing with cases of poor performance (including procedures for whistle blowing)
- deployment of appropriate staffing and skills e.g.
  - minimum ’safe’ numbers and mix
  - schemes of delegation and supervision
  - protocols for staff working in extended roles (e.g. nurse prescribing)
- compliance with working time directives
- assessment and management of risk to staff e.g. violence to staff; workplace health and safety

Quality improvements and learning
- systems for staff feedback e.g. from staff attitude surveys; staff appraisal processes; exit interviews
- consideration of feedback from staff and improvements to the patient/service user experience as a result
- dissemination of lessons learnt from staff feedback

Resources and training for staff
- employee support services e.g.
  - occupational health services
  - independent confidential advice services
  - support against bullying and harassment
Use of information to support clinical governance and health care delivery

Accountabilities and structures
- committee responsibilities for IM&T
- staff responsibilities for IM&T
- reporting and monitoring of performance targets and achievements– to/by management teams, committees and the board

Strategies and plans
- strategy and plans for IM&T
- priority given to IM&T in strategic plans for clinical governance and to the needs of clinical governance in strategic plans for IM&T
- identification of the clinical and other information needed by the board, executive team, management teams, and clinical teams to support clinical governance and health care delivery
- involvement of patients in identifying information needs
- involvement of partner organisations in IM&T (e.g. LIS strategy)
- resources (staff and budget) to support the implementation of the IM&T strategy

Application of policies, strategies and plans
- information used:
  - to monitor performance and outcomes
  - to support performance review and improvement
  - to inform clinical governance activities
  - to support implementation of policies and guidelines e.g. Mental Health Act
- information management systems (including links to enable sharing of information with staff from other organisations)
- health care records systems, including electronic patient/service user records (including communication of patient information with staff from other organisations)
- processes to ensure confidentiality of information about patient/service users e.g. Caldicott guardianship; application of Data Protection Act
- process and systems for assuring data quality

Quality improvements and learning
- use of information to review and improve clinical practice e.g. clinical indicators
- dissemination of methods of effective use of information

Resources and training for staff
- training and support for staff in the interpretation and use of clinical information
- analytic support to users of information
APPENDIX D
THE ROLE OF THE TRUST COORDINATOR

What is a trust coordinator?

The trust coordinator plays an important role in the success of the CHI clinical governance review. He or she acts as the main point of contact between CHI’s pre review, internal and review teams and your trust, assisting with collecting information, planning the review, and helping ensure the visit week runs smoothly. Their help is much valued by the review team.

Who should be a trust coordinator?

You should identify your trust coordinator. The trust coordinator needs to have an understanding of how your trust operates as a whole and knowledge of clinical governance in the trust would also be helpful. The trust coordinator also needs good organisational skills and be senior enough to influence at board level. The CHI pre review team and review manager will provide support to the trust coordinator during the pre review phase and during the review process.

We estimate that a trust coordinator will spend around 30 – 40 days helping with the review from start to finish. They will need to help with the preparation of the review, be available during the visit week to assist with any queries, and be involved during the reporting and action planning stages.

What does a trust coordinator do?

Your trust coordinator will assist in all phases of a review – pre review, pre visit preparation, visit week, reporting and action planning.

During the preparatory phases, we will ask the trust coordinator to publicise the review to staff and patients. CHI will work closely with your communications staff to supply methods of publicising the review. We will also ask the trust coordinator to send all of the information requested by CHI to the pre review team on time. The trust coordinator will need to help with the practical arrangements for the review – such as scheduling, and arranging work areas and catering for the review team whilst they are at your trust.

During collection of the data and information requested by CHI during the pre review phase, the trust co-ordinator will have a key role. We will give the trust coordinator a checklist to help them through the data collection process, which they will be required to complete and return. This checklist will log and identify every document the trust needs to return to CHI. Full instructions for completion of this checklist are supplied within the ‘pre visit request for existing data and information’. The trust coordinator will also help with the administration of the staff survey.

During the visit week, the trust coordinator will assist in any logistical arrangements, such as accompanying the review team members from appointment to appointment and re-arranging interviews if necessary.
After the review visit, the trust coordinator will coordinate comments on the factual accuracy of CHI’s draft report, and assist with the action planning for your organisation.

**What information is available for the trust coordinator?**

A comprehensive information manual to support the review process will be sent to the trust at the start of the pre review phase. This will include a ‘mock time-table’ to use as a basis for scheduling meetings and visits during the visit week.

During the pre review phase, all communication will be via the pre review team – they will be happy to provide help and advice. Contact details will be provided at the start of the pre review phase.

At the initial meeting marking the start of the review, the CHI review manager will clarify the details of the review process. The CHI review manager will be available for support from that point on.
**APPENDIX E**

**EXAMPLE TIMETABLE**

Review weeks may vary in the way in which activities are structured. This table is an example of how a typical review week is organised.

<table>
<thead>
<tr>
<th>Time</th>
<th>MONDAY</th>
<th>TUESDAY</th>
<th>WEDNESDAY</th>
<th>THURSDAY</th>
<th>FRIDAY</th>
</tr>
</thead>
<tbody>
<tr>
<td>am</td>
<td>Arrive at trust</td>
<td>Clinical teams</td>
<td>Clinical teams</td>
<td>Corporate interviews continued</td>
<td>Final debriefing session with CHI review team</td>
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<tr>
<td></td>
<td>Meet with trust coordinator</td>
<td></td>
<td></td>
<td>Additional interviews</td>
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<td></td>
<td>Orientation to trust</td>
<td></td>
<td></td>
<td>Revisits</td>
<td></td>
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<tr>
<td></td>
<td>Meet executive team/members of trust board</td>
<td></td>
<td></td>
<td>Completion of site visit recording forms</td>
<td></td>
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<tr>
<td></td>
<td>Clinical team member interviews</td>
<td></td>
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<td></td>
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<tr>
<td></td>
<td>Lunch meeting for team</td>
<td>Lunch meeting for team</td>
<td>Lunch meeting for team</td>
<td>Lunch meeting for team</td>
<td>Lunch meeting for team</td>
</tr>
<tr>
<td>pm</td>
<td>Clinical team member interviews</td>
<td>Clinical teams</td>
<td></td>
<td>CHI review team prepare for feedback session</td>
<td>Final meeting with members of the trust board and</td>
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<td></td>
<td></td>
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<td></td>
<td></td>
<td>staff who attended the start-up meeting -</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>feedback given</td>
</tr>
<tr>
<td>Evening</td>
<td>Possible night visit Debriefing session</td>
<td>Debriefing session</td>
<td>Possible night visit Debriefing session</td>
<td>Debriefing session</td>
<td></td>
</tr>
</tbody>
</table>
