The British Society of Echocardiography ("BSE") is offering any Echocardiography Department in the UK ("Department") the opportunity to apply for accreditation in accordance with these guidelines ("Applicant"). The BSE grants such accreditation on a purely voluntary basis ("Accreditation"). There may be other Departments which have not sought Accreditation but which offer an excellent echocardiography service. The decision of the BSE to grant accreditation to a Department is only an indication of the BSE’s opinion of the standard of that Department. The BSE does not intend the outcome of the Departmental Accreditation process to be relied upon by any person under any circumstance. This shall include, but not be limited to, any Department, any individual medical practitioner, including GPs, any patient, referring health authority and any member of the public. The Accreditation shall be based on sufficient information being provided to the BSE by Applicants using application forms and an inspection of a Department.

1 Introduction

1.1 The BSE cannot guarantee, or be liable in respect of, the quality, efficacy or performance of an Applicant and/or Department either prior to Accreditation or otherwise.

1.2 Accreditation of an Applicant or Department shall be determined by all relevant information provided by such Applicant or Department and it is the responsibility of that Applicant or Department to ensure that such information is accurate and up-to-date. The BSE shall not be liable in respect of any decisions made based on such information received.

1.3 A process for accreditation in adult echocardiography has been in place since 1994 and for transoesophageal echocardiography since 2003.

1.4 However, accreditation of individual echocardiographers alone cannot guarantee a high quality Department. It is also necessary to have adequate machines, management and organisation.

1.5 The accreditation of Departments could provide the following advantages:

1.5.1 To devolve echocardiographic control giving more local autonomy
1.5.2 To empower an increasing number of Departments to mark log-books and videos for adult accreditation thus accelerating the process and also bringing it more into line with European practice
1.5.3 To facilitate the negotiation of contracts with commissioners and bodies such as Primary Care Trusts
1.5.4 To certify non-cardiac practitioners (e.g. GPwSI, PwSI, nurses) and maintain quality control and continued learning
1.5.5 To encourage improvements in standard by comparison against a nationally-agreed yard-stick

1.6 Departmental Accreditation may be applied for in respect of the following four modules:

1.6.1 Transthoracic echocardiography
1.6.2 Transoesophageal echocardiography
1.6.3 Stress echocardiography
1.6.4 Training to BSE adult proficiency standard
If the Department is carrying out any Transoesophageal Echocardiogram ("TOE") or Dobutamine Stress Echocardiogram ("DSE") then the relevant module form must be completed irrespective of whether specific accreditation in these areas is being sought.

1.7 Recommendations in respect of the standards required for each of these modules are given in sections 3-6. It should be noted that the recommendations contained in section 3-6 below are guidance only and that the standards required for each of the modules may be subject to change at the discretion of the BSE.

1.8 The standards to which the Departments will be compared against are set out below in the separate section ‘Criteria for Grading’. There are two levels of grading: standard and advanced. To achieve advanced Accreditation, all aspects of a Department’s work must be accredited at standard level and the requirements for advanced transthoracic accreditation met. Standard accreditation fulfills the basic requirements for an adequate echo service, while advanced accreditation offers a higher level of service including more rigorous quality control. Please note that, as stated above, Departmental Accreditation is a voluntary process. Whilst the BSE is keen to work with Departments to help them achieve these standards and does not expect all departments to reach all standards in all areas, it has the ultimate discretion as to whether to accredit a Department.

1.9 Throughout this document, the term ‘sonographer’ is used to mean a non-medical echocardiographer and subsumes the terms clinical physiologist, cardiac or echocardiography technician and radiographer.

1.10 It should be noted that Echocardiography is changing rapidly and it is expected that this document will be reviewed in response to developments including screening echocardiography, portable systems and the training of GPwSI and other non-cardiologists in echocardiography. This document should therefore only be viewed as a general guide and the guidance and criteria for Accreditation contained in this document may be subject to change at the ultimate discretion of the BSE.

1.11 The registration fee for Departmental Accreditation is £500. These fees will cover the cost of administration and also of organising accreditation visits where applicable.

2 Process

2.1 The Applicant completes an application form and pays a non-refundable registration fee.

2.2 An inspection will be undertaken at a mutually agreeable time in order to clarify or confirm information supplied on the application form. It should be noted that BSE has the ultimate discretion to decide whether a department deserves Accreditation. The ‘Criteria for Grading’ set out below should be viewed as guidance only.

2.3 The Applicant is informed whether or not Accreditation is awarded and at which level for each of the specialty areas for which it has been requested.

2.4 A Department can claim BSE Accreditation for those areas in which BSE is of the opinion that it has reached the required standard e.g. ‘BSE-Accredited department for the provision of transthoracic echocardiography in adults and for training to BSE adult proficiency level’.

2.5 An unsuccessful Applicant will be notified of the areas requiring improvement and invited to re-apply when these have been addressed. The BSE will use its best endeavours to offer advice and support to help achieve this. An unsuccessful applicant may Appeal the decision of the Accreditation Committee on the grounds, and following the process set out in section 8.

2.6 Occasionally a department may be advised of remedial measures that could be implemented within a set time period to achieve Accreditation. These will usually be within a focussed area and achievable within a few weeks. The Accreditation Committee will require evidence that the changes have been implemented, which may involve submission of paperwork, file uploads or a second inspection visit. The remedial process is entirely at the discretion of the Accreditation Committee and must be completed within a year of the original application.
2.7 Accreditation lasts for 5 years, and can then be re-applied for.

2.8 If there is a change of Technical or Clinical Head of Department during an Accreditation period, the BSE should be informed within 6 months.

2.9 The BSE has the right to suspend a Department’s accredited status under the following circumstances:

2.9.1 If it has reason to believe that that the safety of the public, or the public interest is seriously endangered. The BSE may suspend a Department while it decides whether or not to withdraw approval, where it has decided to withdraw approval but before the decision takes effect, or while any appeal is pending;

2.9.2 If it believes that its reputation is being damaged or likely to be damaged in any way whatsoever as a result of Accreditation of a Department;

2.9.3 If a Department or any member of such Department is subject to any investigation, formal or otherwise; or

2.9.4 If it does not reasonably consider a Department to be worthy of Accreditation, for any reason whatsoever.

2.10 The BSE may remove a Department’s accredited status in the following circumstances:

2.10.1 If it has reason to believe that that the safety of the public, or the public interest is seriously endangered;

2.10.2 If it believes that its reputation is being damaged or likely to be damaged in any way whatsoever as a result of Accreditation of a Department;

2.10.3 If a Department or any member of such Department is subject to any investigation, formal or otherwise; or

2.10.4 If it does not reasonably consider a department to be worthy of Accreditation, for any reason whatsoever.

2.11 A department whose Accreditation has been suspended or removed may appeal the decision, following the procedure laid down in section 8.

2.12 The BSE may inspect an Accredited Department at any time during the period of Accreditation following reasonable notice.

2.13 The BSE shall comply with the provisions of the Data Protection Act 1998 (“DPA”) and, to the extent that it is a Data Processor (which has the meaning given to it under the DPA), it shall put in place:

2.14 Appropriate technical and organisational measures against the processing of Personal Data (which has the meaning given to it under the DPA) and against unauthorised, accidental or unlawful access to the Personal Data (having regard to the state of technological development and the costs of implementing any such measures) as well as reasonable security programmes and procedures for the purpose of ensuring that only authorised personnel have access to the Personal Data processing equipment and that any persons whom it authorises to have access to the Personal Data shall respect and maintain due confidentiality;

2.15 A level of security programmes and procedures which reflect:

a) the level of damage which may be suffered by a Data Subject (which has the meaning given to it under the DPA) to whom the Personal Data relates as a result of unauthorised or unlawful possession of the Personal Data; and

b) the state of technological development and the costs of implementing such procedures and programmes; and
2.16 As required by the DPA, such security programmes and procedures which specifically address the nature of any Sensitive Personal Data (which has the meaning given to it under the DPA).

2.17 The Applicant acknowledges that it shall comply with the relevant data protection statement as set out in Appendix A.

3 Standard 1 - Transthoracic Echocardiography

3.1 Recommendations for staffing and training.

3.1.1 All centres must have both a specialist Technical and Clinical Head of Department.

3.1.2 The Clinical Head should be trained in clinical cardiology and specialist echocardiography and ideally hold individual Accreditation. His/her job description should include setting clinical guidelines and policy, performing studies, training doctors and sonographers, audit, clinical meetings and quality control. He/she should set up a system for reviewing requests and reports, and urgent clinical review in response to findings at echocardiography. In a District General Hospital performing 3000 studies per year, at least one session per week should usually be allocated directly to echocardiography.

3.1.3 The Technical Head should be responsible for performing studies, audit, service improvement, training doctors and sonographers and liaising with Occupational Health and the Works Department. The Technical Head must hold BSE accreditation and be graded at least Band 7.

3.1.4 Sonographers performing and reporting studies unsupervised should be BSE accredited and at least band 6 and typically band 7.

3.1.5 Continuing education should be provided (and funded) to fulfil BSE re-accreditation requirements or to a similar level. There should be a small library of relevant reference textbooks within the Department.

3.1.6 The job profile of a sonographer includes training, self-education, audit, and quality control in addition to performing echocardiograms.

3.2 Recommendations for organisation and equipment.

3.2.1 Echo rooms used for inpatients on beds should be at least 20 m² in area.

3.2.2 Ventilation, heating, lighting and ancillary facilities must be appropriate (see Appendix).

3.2.3 Echo machines must have the capacity for imaging including second harmonic imaging, colour mapping, pulsed Doppler and both steerable and stand-alone continuous wave Doppler. Ideally tissue Doppler should also be available.

3.2.4 A single echo machine can handle up to a maximum of 2500 studies each year but this figure will be lower if there is a significant ward-based or complex workload.

3.2.5 The machine should be serviced regularly, and be replaced or have a major upgrade at least every 5 years.

3.2.6 There must be consideration of patient comfort, privacy, dignity and provision of adequate information.

3.2.7 There must be awareness of health and safety issues especially relating to back and eye problems and adequate liaison with occupational health and risk management departments (see Appendix).

3.2.8 A report database should exist, with facilities for storing and retrieving specific echo studies.

3.2.9 A separate viewing room is recommended for reviewing studies and off-line reporting.
3.2.10 There should be appropriate storage space

3.2.11 A patient information leaflet should be available

### 3.3 Recommendations for performing studies.

3.3.1 A standard transthoracic study (one TTE equivalent) takes 30 minutes. Training may prolong this to 45 minutes (1.5 TTE equivalents). A complex study (e.g. including contrast injection or detailed valve haemodynamic assessment) may take up to 1 hour (two TTE equivalents). As an ideal, allowing for all aspects of the job profile, a sonographer will perform no more than 2000 studies per year.

3.3.2 A list of indications for echocardiograms should be agreed.

3.3.3 Prioritising and filtering, of inappropriate requests should be performed by sonographers supported by the Clinical Head.

3.3.4 Minimum standards for studies should be established. Study protocols appropriate to specific clinical conditions should be established.

3.3.5 A format for reports should be established, including who should issue conclusions and who is qualified to sign reports.

3.3.6 The requirements of the Data Protection Act 1998 must be complied with regarding data storage.

3.3.7 Reports from routine studies should usually be issued on the day of the examination. For urgent or inpatient studies, at least a preliminary report should usually be issued immediately.

3.3.8 A mechanism must be in place for reporting cases that require urgent clinical attention.

3.3.9 Regular meetings, ideally weekly, should be held to review unusual, challenging or otherwise difficult cases.

3.3.10 A formal quality assurance system should be in place with regular blind over reading of selected studies to ensure consistency of performance and interpretation. Meetings should take place at least 4 times per year with all echocardiographers attending at least 50%.

### 4 Standard 2 - Transoesophageal Echocardiography (TOE)

All the standards for transthoracic echocardiography in addition to the following:

#### 4.1 Recommendations for staffing and training.

4.1.1 All centres must have a designated Head of TOE. This will usually be the Clinical Head of Echocardiography.

4.1.2 Outpatient TOE studies require an operator with appropriate training, a cardiac-trained nurse with experience in managing airways and, ideally, a sonographer.

4.1.3 Continuing education must be provided for the operators.

4.1.4 Each operator should perform or directly supervise at least 50 studies per annum.

4.1.5 Ideally operators should have BSE/ACTA/EAE TOE accreditation.

4.1.6 A list of indications for TOE should be agreed.

#### 4.2 Process
4.2.1 Minimum standards for studies should be established and the head of TOE must be responsible for ensuring that all operators adhere to them

4.1.1 A preoperative checklist should be used

4.1.2 Whenever sedation is used, it should be in accordance with the recommendations given in ‘Implementing and Ensuring Safe Sedation Practice for Healthcare Procedures in Adults’ published by the Academy of the Royal Colleges in 2001 and regulations stated in the NPSA Risk of Overdose with Midazolam Injection in Adults 2008

4.1.3 The TOE probe must be checked electrically at a frequency dependent on usage. A log of these checks must be kept

4.1.4 The TOE probe should be cleaned regularly

4.2 Recommendations for organisation and equipment

4.2.1 There should be appropriate provision of:
- Room (ideally > 25 m² in area)
- Couch with facility for head-down tilt
- Facilities for cleaning and sterilising the probe.
- Storage cupboard for the probe
- Resuscitation apparatus and drugs
- Lockable drug cupboard
- Suction
- Oxygen
- Pulse oxymeter
- Sphygmomanometer
- Facilities for recovery of the patient
- Protocols for patient care

5 Standard 3 - Stress Echocardiography

All the standards for transthoracic echocardiography in addition to the following:

5.1 Recommendations for staffing and training.

5.1.1 All centres should have a designated Head of Stress Echocardiography, normally the Clinical Head of Department

5.1.2 Stress echocardiography studies require an experienced operator and a sonographer or trained nurse

5.1.3 The study reporter must be specially trained in stress echocardiography

5.1.4 There must be a mechanism in place for feedback to assess clinical correlation

5.1.5 Each operator/reporter should perform or directly supervise at least 100 studies per annum

5.1.6 Continuing education must be provided for the interpreter

5.1.7 At least one member of staff performing the study should possess at least Immediate Life Support (ILS) Training or be a specialist cardiologist

5.1.8 A list of indications for stress echocardiograms should be agreed

5.1.9 Appropriate protocols for studies should be established and the Head of Stress Echocardiography must be responsible for ensuring that all operators adhere to them

5.2 Recommendations for organisation and equipment
Ideally there should be appropriate provision of

- Designated room (size >25 m$^2$)
- Stress echocardiography software
- Contrast agents and contrast specific software
- Infusion syringe for pharmacological stress or equipment for exercise stress e.g. bicycle
- ECG monitor and recorder
- Sphygmomanometer
- Resuscitation apparatus and drugs readily available

6 Standard 4 – Training to BSE Proficiency Standard

The centre must be accredited in transthoracic echocardiography at standard level.

6.1 Recommendations for staffing and training

6.1.1 Each centre must have a BSE Accredited individual responsible for training. This person may be from a Clinical or Technical background

6.1.2 Staffing levels and workload appropriate to the number of trainees to ensure adequate clinical capacity. A guide would be two BSE accredited staff and 2000 echoes p.a. for a department to accommodate one trainee

6.1.3 At least one, and ideally two protected tutorial half-day sessions per week for both trainer and trainee

6.1.4 Access for both trainer and trainees to local, national and international meetings

6.1.5 Regular weekly Departmental case review sessions

6.2 Recommendations for equipment

6.2.1 Core library e.g. 3 up to date echo textbooks and 1 general cardiology textbook in the Department and access to cardiology journals electronically or within the hospital

6.2.2 Training material-tapes/CDs/digital cases, etc.

6.2.3 Internet access should be available to all staff

6.3 Performance

6.3.1 History of success in training students to BSE proficiency/similar level

7 Departmental Inspections (“Inspection”)

7.1 Purpose

The purpose of an Inspection is solely to determine whether or not information submitted on Accreditation Application Forms is currently accurate. It should be noted that Inspections will be undertaken at the ultimate discretion of BSE.

7.1.1 Process

7.1.2 Inspections will form part of the Accreditation process but may be undertaken if the Department appears to have changed significantly since the original submission.

7.1.3 At least 28 working days’ notice will be given of an Inspection visit. Proposed visits will be re-scheduled only in special circumstances (e.g. Technical Head on leave)

7.1.4 Inspections will normally be undertaken by two people nominated by the BSE Accreditation Committee, one of whom will normally be a doctor who is him/herself a
Clinical Head of Echocardiography in a department of similar size, and the other a sonographer who is him/herself a Technical Head of a similar department.

7.1.5 The costs of an Inspection initiated by the BSE will be borne by the society.

7.1.6 Inspectors will bring with them copies of Accreditation Forms submitted, together with any amendments. They will:

- Check that equipment and facilities are as claimed
- Observe selected patient studies
- Check a sample of clinical and technical reports
- Interview Clinical and Technical heads of Department, plus approximately one-third of other staff and trainees
- Inspect daybooks, appointment diaries, etc.
- At the end of the Inspection, hold a meeting with the Clinical and Technical Heads, at which any apparent discrepancies between data on the Accreditation Application and observed practice will be highlighted

This list is not exclusive and that Inspections will differ from time to time.

7.2 Assessment

7.2.1 The inspectors will submit a written report of their findings to the BSE Accreditation Committee. After discussion within the Committee, the Chairman will write to the Clinical and Technical Heads of the department concerned. This letter will be sent within 60 days of the Inspection and will offer one of the following conclusions for each category of Accreditation applied for or held:

7.2.1.1 Accreditation awarded at either standard or advanced level
7.2.1.2 Accreditation not awarded
7.2.1.3 Remedial measures required before Accreditation may be awarded

8 Appeals (“Appeals”)

8.1 Where a Department considers that the BSE has unfairly refused, regraded, suspended or removed Accreditation, written appeal to the BSE Council. Appeals may be made only on the following grounds:

8.1.1 That the decision was affected by bias or breach of the Society’s guidelines for Centre Accreditation
8.1.2 At an Inspection, the inspectors did not carry out the inspection in accordance with the procedure set out in section 7
8.1.3 That conditions prevailing at the time of the inspection were unusual and temporary (e.g. several key staff off sick)
8.1.4 That information provided to the inspectors was incomplete or inaccurate

8.2 An Appeal will be considered by the BSE Council at its discretion (or a panel nominated by the Council for this purpose and containing at least 3 elected Council members). No member of the Appeal panel shall have taken part in the Inspection, or have any current or past association with the Centre concerned. The findings of the Appeal panel will be sent to the Department within 90 days of the Appeal being lodged and shall be final.
## CRITERIA FOR GRADING

### TRANSTHORACIC

<table>
<thead>
<tr>
<th>Standard</th>
<th>Advanced</th>
</tr>
</thead>
<tbody>
<tr>
<td>Both clinical and technical heads of echocardiography</td>
<td>Both technical and clinical heads have BSE accreditation</td>
</tr>
<tr>
<td>Clinical head Has at least one PA dedicated to echo</td>
<td>Formal and systematic quality control in place</td>
</tr>
<tr>
<td>Technical head spends 5 or more sessions in echocardiographic activities (including management or quality control)</td>
<td>Technical Head spends 8 or more sessions in echocardiography</td>
</tr>
<tr>
<td>Agreed minimum standards for studies</td>
<td>Clinical Head spends 2 or more sessions in activities directly related to echocardiography</td>
</tr>
<tr>
<td>Patient information leaflet available including arrangements for chaperones</td>
<td>Digital archiving</td>
</tr>
<tr>
<td>List of indications for echo published internally</td>
<td>No machine in regular use older than 5 years</td>
</tr>
<tr>
<td>Triaging of requests</td>
<td>All trained sonographers BSE accredited</td>
</tr>
<tr>
<td>Report database</td>
<td>One or more rooms at least 20 m² in area</td>
</tr>
<tr>
<td>System of review for uncertain echocardiograms</td>
<td>One WTE sonographer for no more than 2000 standard and complex studies per annum</td>
</tr>
<tr>
<td>System of alerts for important pathology found at echocardiography</td>
<td>System of liaison with other departments to advise about timing of or results of studies</td>
</tr>
<tr>
<td>Provision for continuing education</td>
<td>A patient information leaflet available</td>
</tr>
<tr>
<td>Studies archived. Reports written that day</td>
<td>Adequate storage space</td>
</tr>
<tr>
<td>Most machines have 2nd harmonic imaging</td>
<td></td>
</tr>
<tr>
<td>All machines have colour and stand-alone Doppler</td>
<td></td>
</tr>
<tr>
<td>No machine in regular use upgraded more than 10 years ago</td>
<td></td>
</tr>
<tr>
<td>30-40 minutes allowed per standard study and up to 1 hour for a complex study</td>
<td></td>
</tr>
<tr>
<td>All sonographers reporting studies Band 6 or higher</td>
<td></td>
</tr>
<tr>
<td>Data protection act applied</td>
<td></td>
</tr>
<tr>
<td>Manual handling policy implemented</td>
<td></td>
</tr>
<tr>
<td>Rooms uncluttered and of adequate size</td>
<td></td>
</tr>
<tr>
<td>Appropriate provision of patient facilities</td>
<td></td>
</tr>
</tbody>
</table>
## TRANSOESOPHAGEAL

<table>
<thead>
<tr>
<th>Standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>Designated Head of TOE</td>
</tr>
<tr>
<td>Head of TOE has BSE/EAE accreditation</td>
</tr>
<tr>
<td>Each operator performs &gt; 50 studies each year</td>
</tr>
<tr>
<td>Designated person, usually a nurse, to manage airway and recover the patient</td>
</tr>
<tr>
<td>Recovery area</td>
</tr>
<tr>
<td>Minimum standards for studies established</td>
</tr>
<tr>
<td>List of indications for TOE agreed internally</td>
</tr>
<tr>
<td>Patient information leaflet sent out</td>
</tr>
<tr>
<td>The following equipment:</td>
</tr>
<tr>
<td>Resuscitation equipment</td>
</tr>
<tr>
<td>Omniplane probe</td>
</tr>
<tr>
<td>Routine use of:</td>
</tr>
<tr>
<td>Oxygen</td>
</tr>
<tr>
<td>Pulse oximeter</td>
</tr>
<tr>
<td>BP monitor</td>
</tr>
<tr>
<td>Patient preparation including letter and pre-procedure check-list</td>
</tr>
<tr>
<td>Appropriate arrangements for cleaning/sterilisation of probes</td>
</tr>
<tr>
<td>Room at least 25 m² in area</td>
</tr>
<tr>
<td>Provision for continuing education</td>
</tr>
<tr>
<td>Provision for quality control</td>
</tr>
<tr>
<td>Sedation used according to published guidelines</td>
</tr>
</tbody>
</table>
## STRESS ECHOCARDIOGRAPHY

<table>
<thead>
<tr>
<th>Standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>Designated Head of Stress Echocardiography</td>
</tr>
<tr>
<td>Head maintains CME for stress echo</td>
</tr>
<tr>
<td>Audit of results against angiography or other independent standard</td>
</tr>
<tr>
<td>More than 100 studies each year per reporting operator</td>
</tr>
<tr>
<td>Patient information leaflet sent out</td>
</tr>
<tr>
<td>Machine capable of changing MI and with digital stress echo package</td>
</tr>
<tr>
<td>Studies performed by at least two people At least one must have ILS or equivalent</td>
</tr>
<tr>
<td>If a doctor is not present they must be immediately available</td>
</tr>
<tr>
<td>Blood pool contrast available</td>
</tr>
<tr>
<td>Machine with 2nd harmonic imaging</td>
</tr>
<tr>
<td>Resuscitation facilities readily available</td>
</tr>
</tbody>
</table>
## BSE TRAINING

<table>
<thead>
<tr>
<th>Standard</th>
<th>Advanced</th>
</tr>
</thead>
<tbody>
<tr>
<td>All standard sections of transthoracic echocardiography</td>
<td>All advanced sections of transthoracic echocardiography</td>
</tr>
<tr>
<td>Designated Head of Training who is BSE accredited</td>
<td>Accreditation in TOE and DSE</td>
</tr>
<tr>
<td>One protected training sessions each week for each trainee</td>
<td>Full range of adult pathology on site</td>
</tr>
<tr>
<td>In house assessment programme</td>
<td>Extensive library including CD, or other case material</td>
</tr>
<tr>
<td>Departmental training or business meetings at least once each week</td>
<td>History of success in passing candidates for BSE adult accreditation, ideally more than 3 candidates within the last 3 years.</td>
</tr>
<tr>
<td>Internet access</td>
<td>Formal written training programme available</td>
</tr>
<tr>
<td>Textbooks of basic and advanced echocardiography and general cardiology</td>
<td></td>
</tr>
<tr>
<td>Full range of adult pathology either on site or supplemented by visits to a near-by centre</td>
<td></td>
</tr>
<tr>
<td>Attendance at one conference each year for trainer and trainee</td>
<td></td>
</tr>
</tbody>
</table>
Appendix

Room Specifications

Summary of Workplace Health, Safety and Welfare Regulations, 1992
(Published by Unison, Unison Centre, Holborn Tower, 137 High Holborn, London WC1V 6PL)

The Workplace Health, Safety and Welfare Regulations 1992
These regulations apply to existing echo scanning rooms and all new rooms. They arise out of the European Workplace Directive and are mainly concerned with minimum standards for the work place. Regulations with relevance to an echo scanning room or Department concern:

1 Ventilation (Regulation 6). In enclosed workplaces you must be provided with effective and suitable ventilation, which does not cause uncomfortable draughts e.g. ceiling-mounted.

2 Temperature (Regulation 7). During working hours the temperature in the workplace must be reasonable. The approved codes of practice (ACOP) do not give a maximum temperature but still recommend air cooling plants and shading windows. The minimum working temperature is at least 16 degrees C. A thermometer should be provided in the scanning room if the temperature is uncomfortable.

3 Lighting (Regulation 8). Every workplace should have suitable and sufficient lighting and where possible natural light.

4 Room dimension and space (Regulation 10). Every workroom should have sufficient floor area height and unoccupied space allowing for a minimum of 11 cubic metres per person (assuming the room is 3 metres high). Area taken up by equipment or furniture is additional to this space allowance.

5 Workstations and seating (Regulation 11). Every workstation must be suitable for the worker using it and the work being carried out. The echo machine and the patient on the couch are classed as the workstation. Suitable seating (with height adjustment and back support) should be provided when this work is done sitting down with a foot rest if necessary.

6 Toilets (Regulation 20). Staff toilets should be provided in readily accessible places. One toilet with washbasin should be provided for 1 to 5 members of staff. This may be more applicable to a suite of scanning rooms and may already be provided when the scanning rooms are situated in a cardiac Department.

7 Washing facilities (Regulation 21). Suitable and sufficient washing facilities should be provided at readily accessible places. Ideally a washbasin with hot and cold water, soap and a means of drying can be situated within the scanning room itself.

Health and Safety

Extracts from the Display Screen Equipment Regulations 1992
Published by Unison, Unison Centre, Holborn Tower, 137 High Holborn, London WC1V 6PL

Under these regulations the echo machine is classed as the workstation (and any PC used in the room e.g. for reporting echo scans) and the user is the employee who habitually uses the display screen equipment as part of his/her job. Within the definition of the workstation other equipment and factors are taken into account such as: telephones, printers and the environment around the display screen such as the noise, lighting, temperature etc. The particular regulations that are relevant to an echo scanning room or Department are:

1 Assessment of workstations (Regulation 2) The employer must carry out suitable and sufficient assessment of the workstation to identify risks to the user. This is usually done on a regular basis or when there has been a significant change in the workstation such as a new user of piece of equipment. Following the assessment the employer must act to reduce the risks identified. Particular emphasis is given to eyesight, stress and upper limb disorders.

2 Eye and eyesight tests (Regulation 5) Employers must provide on request and offer at regular intervals a free professional full eyesight test (not a simple ‘Keystone’ vision test) but the employer is not required to provide them automatically. Users must, therefore, request the tests if they experience sore eyes or headache during scanning or reporting on a PC. If the eyesight test reveals a problem the employer must provide special and normal corrective appliances such as spectacles. If the eyesight test does not reveal a problem then the workstation standards must be checked.

3 Health and safety training (Regulation 6) Adequate health and safety training must be provided for users of echo scanners and PCs. The training should cover recognition of hazards such as screen flicker and screen glare, the importance of good posture and regular breaks and how to

---

1 The NHS Estates ultrasound room data sheets specify a minimum temperature of 20°C
2 This is also covered in the EEC Display Screen Equipment Regulations, 1992
request an eye test and report problems with the equipment etc. The user should be encouraged to contribute to the assessment of the workstation as well.

**Provision and Use of Work Equipment Regulations, 1992**
Published by Unison, Unison Centre, Holborn Tower, 137 High Holborn, London WC1V 6PL

This document refers to general work environments. Work equipment must comply with requirements for suitability, maintenance, information and training. This document extracts aspects which are relevant to echocardiography

**Extracts from the Provision and use of Work Equipment Regulations, 1992**

Work equipment includes the echo machine and any PC used in the room and also the couch or bed and any wheelchair or aids used to get the patient in a position to be scanned (see also Appendix f).

The following extracts are relevant to echocardiography:

1. **Suitable work equipment** (Regulation 5) Employers must ensure that all work equipment is suitable for the work it is provided to do. This is confirmed by means of a risk assessment which should cover:
   - 1.1 The design and condition of the equipment e.g. could it cause strain injury or could it be modified or replaced with a better designed piece of equipment to prevent injury to staff or patients
   - 1.2 The working conditions where the equipment is used e.g. is the scanner electrically safe or is the floor where the scanner is used uneven making the heavy scanner prone to toppling over. This also applies to transferring a patient from a wheelchair to a scanning couch or transferring the patient from a bed following a transoesophageal scan
   - 1.3 The purpose of the equipment e.g. older fixed frame wheelchairs should not be used with an immobile patient; a lightweight deconstructable wheelchair would be needed in this situation.

2. **Maintenance** (Regulation 6) The work equipment must be maintained in an efficient working order and a maintenance log must be kept up to date. This may also include the power sockets to ensure a stable electrical supply and secure earth as the patient may have ECG cables attached during a scan. A calibration log of syringe drivers used in stress studies should also be kept.

3. **Information and instructions** (Regulation 8) All staff who use the work equipment must have available to them comprehensive and adequate health and safety information. The information should cover:
   - 3.1 Conditions and method of use e.g. most echo scanners cannot be used in the presence of inflammable anesthetics
   - 3.2 Foreseeable and abnormal situations e.g. emergency echo scan in cardiac arrest situations
   - 3.3 What to do in if there is an accident, breakdown or emergency e.g. patient falling because wheelchair brakes were not effective.

4. **Training** (Regulation 9) All staff who use the equipment must have adequate health and safety training. This training should cover the health and safety risks and precautions to be taken as well as supervisors receiving risk assessment training on the work equipment.

5. **Controls for starting equipment** (Regulation 14) The equipment used must have one or more controls for the starting and controlling of the equipment e.g. a syringe driver used in stress echo scans.

6. **Control for stopping equipment** (Regulation 15) Where appropriate the equipment must have a one or more controls to stop the equipment e.g. as above.

7. **Controls** (Regulation 17) All controls for work equipment must be clearly visible and identifiable e.g. resuscitation equipment, syringe drivers, drip pumps etc.

8. **Stability of equipment** (Regulation 20) The equipment should be stabilised to prevent it falling or overturning but this will also apply to additional equipment added to the scanner such as video recorders and printers. This can also apply to portable oxygen saturation monitors for transoesophageal echocardiography as well as resuscitation equipment and syringe drivers or drip pumps used in stress echocardiography.

9. **Lighting** (Regulation 21) Suitable and sufficient lighting must be provided. Problems may be encountered outside the normal scanning room such as the Intensive Care Unit. In these situations the control of ambient light may be difficult if there are inadequate blinds. Additional blinds may need to be installed for subsequent scans or the use of a temporary anti glare screen may help.

10. **Markings** (Regulation 24) The work equipment must carry clearly visible health and safety markings. These marking may indicate machine weight, whether the machine should not to be used with inflammable anaesthetics, prudent use of transducer power levels or maximum wheelchair carry weight.

11. **Warnings** (Regulation 25) Work equipment must incorporate warning or warning devices as appropriate. Audible warnings are usually present on syringe drivers, drip pumps and defibrillators and staff should be made aware of them. The addition of an emergency call button is advised in an echo scanning room especially if transoesophageal and stress studies are performed there.
Manual Handling Policy

Manual handling includes lifting, lowering, pushing, pulling, carrying and supporting loads and also patient handling. Between 1992 and 1995 nearly 14,000 manual handling accidents were reported to the Health and Safety Executive of which over 60% involved patient handling. There should be a manual handling policy in operation in all hospitals. The hospital occupational health Department should be to perform an assessment if the cardiology or echocardiography Department does not have a manual handling assessor or trainer. The health and safety officer and or manual handling assessor keep up to date with new manual handling devices and aids as well as improved techniques and regulations. They will be able to advise on changes to the Department based on the scientific study of ergonomics as well as staff training.

1. Risk assessment

The purpose of the manual handling risk assessment is to identify and extenuate possible problems e.g. install a more suitable scanning couch or have two people push the echo scanner to the ward. A record will be made of the findings and a follow up review will be made with revisions if necessary. The four main factors considered in the assessment are:

1.1 The task
Staff should be assessed as they perform echo scans in the Department and on the ward. Pushing and maneuvering the echo scanner as well as transferring the patient must also be assessed. Factors that should be taken into consideration are:

1.1.1 Holding the load (transducer, TOE probe or patient) at a distance from the trunk,
1.1.2 Unsatisfactory body movements (e.g. twisting the trunk),
1.1.3 Poor posture (e.g. stooping or over stretching),
1.1.4 Excessive movement of the load over distance (e.g. pushing the echo scanner or transferring the patient),
1.1.5 The risk of sudden movement of the load (e.g. patient falling)
1.1.6 Prolonged physical effort including a fixed posture and insufficient rest or recovery periods.

An aching back or limb at the end of the day should not be accepted as ‘part of the job’.

1.2 The load
The load that staff are likely to encounter should be assessed taking into consideration its weight, whether it is bulky or unwieldy, or difficult to grasp and whether it is or is likely to become unstable. When the object is a patient they may cooperate or hinder in the transfer or may find themselves suddenly unable to continue. Staff may react by trying to prevent the patient from falling which may cause injury to both. If staff are properly trained and positioned they may be able to allow a controlled fall by letting the patient slide down their body and onto the floor but with good risk assessment this situation should not arise.

1.3 The working environment
The environment in which the scans are performed should be assessed taking into account:

1.3.1 Prevention of good posture because of space constraints and inadequate work equipment e.g. un-adjustable couch
1.3.2 Slippery surfaces and uneven floors
1.3.3 Extremes of temperature noise etc.
1.3.4 Lighting condition
1.3.5 Poor storage facilities.

1.4 The individual staff capabilities
The principles of the manual handling operations regulations is that the job should be adapted to suit the employee who should not be subject to unrealistic physical demands. Examples of questions that should be addressed are:

1.4.1 Can the employee push a 200 Kg (400 lb.) echo machine to a ward across uneven surfaces
1.4.2 Can the employee maneuver the machine into a suitable position by the ward bed
1.4.3 Can the employee reach the echo videos on the shelf - should a footstool be provided in the echo scanning room / Department.
1.4.4 Is any member of staff at particular risk e.g. pregnancy or history of previous back problems.

2 Reducing risks
Following the assessment, action must be taken to reduce the risks identified. If at all possible the manual handling operation should be eliminated or minimized by reorganizing the task. Understanding the risks can help in finding the solution. Some factors contributing to risk are:

a) Number of years spent in echocardiography
b) Frequency of lifting
c) Scanning with the machine on the left, the transducer in the right hand and the patient on the right.
d) Poor job satisfaction

A solution should be sought in consultation with the manual handling assessor and the staff member. No member of staff in the echo Department should be expected to handle patients or loads where there is likelihood that they or the patient may be injured. The use of manual handling equipment may reduce the risk of injury significantly. Many suppliers loan such equipment for trial periods and provide initial training in its use. Examples of these aids are:

a) Electrically height adjustable couches
b) Roll boards which help transfer patients from bed to trolley
c) Two person sling for moving patient up beds / couches
d) Floor turntables which allow a standing patient to be swiveled from a wheelchair to a seated position on a couch
e) Adjustable wheelchair with removable sides
f) Slide board allowing patients to slide from a seated position in a wheelchair for example to a seated position on the couch
g) Grab handles on wall to allow patient to pull themselves onto their side for scanning (patients generally prefer to be independent and move themselves and this should be encouraged)
h) Electric hoists may be useful with very heavy immobile patients.

3 Training
Training of all staff (including agency staff) in manual handling techniques and use of equipment/aids must be undertaken following a baseline analysis of the needs. Information should be provided to staff enabling them to report faulty equipment and how to maintain the equipment. The training programme should include:

a) Ergonomics looking at the tasks and the environment and encouraging staff to alter their own environment to make the work safer
b) Spinal mechanics
c) Mechanical handling techniques
d) Demonstration of any mechanical equipment used in the Department.

4 Monitoring.
Following risk assessment there must be follow up monitoring to assess the effectiveness of the arrangements for reducing the risk. This must be recorded. Such monitoring should identify problems before something goes wrong and can be referred to in the event of an accident. Information about any accident is useful for determining whether the manual handling policy failed or was not applied correctly.