

Accreditation in Adult Transthoracic Echocardiography (TTE) Information Pack

This pack is for the use of all candidates undergoing the accreditation process and becomes effective as of

1st June 2024

This document supersedes all previous versions.

This document is a guide to completing BSE TTE accreditation

Submission, assessment criteria and portal user guide are included



Contents

Welcome message from the Chair of Accreditation	3
Introduction & aims	4
Summary of process requirements	4
Exam fees	5
Extensions	5
Appeals	5
Mentor	5
Written Theory Examination	6
Multiple-choice section	6
Image reporting section	6
Practical Assessment	7
Logbook submission	7
Other information regarding the logbook:	8
Scanning Assessment	9
Viva case submission	9
Practical assessment - outcomes and process for re-attempts (resubmissions)	10
Appendix 1: Training Syllabus	11
Appendix 2: Curriculum-based Competency Tool	31
Appendix 3: Reading List	35
Appendix 4: Written Examination Registration Guidance	36
Appendix 5: Written Exam multiple-choice questions examples	38
Appendix 6: Written Exam image reporting questions examples	39
Appendix 7: BSE logbook portal user guidance	40
Appendix 8: Logbook guidance and marking criteria	48
Appendix 9: Guidance for the removal of patient identifiable data	53
Appendix 10: Practical scanning mark scheme	54
Appendix 11: Patient case studies viva marking criteria	55
Accreditation Process Overview	60
Useful Links & Contacts	61



Welcome message from the Chair of Accreditation

Dear Candidate,

Welcome to the British Society of Echocardiography (BSE). The process underlying accreditation is set up to assist the echocardiographer in training with the ultimate aim of achieving and maintaining a high standard of clinical echocardiography for the benefit of our patients.

The accreditation process is regulated to ensure high proficiency and professional standards. We aim to enable as many members as possible to achieve accreditation. A list of accredited members is maintained on the BSE website.

Please remember that we are here to support you throughout this process. If you need any assistance or have constructive feedback to offer the accreditation committee, please don't hesitate to let us know. We are committed to your success.

Good luck with your accreditation process.

Best wishes,

Rennett

Sadie Bennett

Chair, BSE Accreditation Committee



Introduction & aims

- Accreditation is a service for BSE members and is not a compulsory or regulatory certificate of competence or excellence.
- Accredited members are expected to be able to perform and report echocardiographic studies unsupervised.
- The Accreditation process consists of a written theory examination and a practical assessment. This pack provides further instructions for both.
- Accreditation is a minimum requirement and cannot be regarded as a guarantee of competence.
- Echocardiography skills can only be maintained by continued education and practical involvement in echocardiography. This is underlined by limiting accreditation to five years, after which reaccreditation must be sought. Further details surrounding re-accreditation are available on the BSE website.
- Accredited members are expected to uphold the BSE code of conduct standards. Where concerns about an accredited member's echocardiography practice arise, this should be dealt with locally in the first instance and should only be escalated to the Accreditation Chair if improvement in echocardiography practice has not been demonstrated.
- > Return to practice routes for re-accreditation are available for previously accredited members.

Summary of process requirements

- 1. The candidate must be a member of the BSE.
- 2. Candidates must have a designated mentor to assist them through the accreditation process.
- **3.** The accreditation process has two compulsory elements: a written theory examination and a practical assessment. **You must pass both elements to become an accredited member.**
- 4. The written theory exam comprises A multiple-choice question (MCQ) theory section and a "best answer" image reporting section.
- 5. The practical assessment consists of a logbook, a practical scanning assessment, and a viva assessment of five patient case studies.
- 6. The candidate must pass the written assessment before registering to attend the practical assessment.
- 7. The logbook should be collected within 24 months of the written examination.

Any queries regarding the accreditation process should be addressed to the BSE Accreditation Department; contact details and registrations are available at www.bsecho.org.

Tel: 0208 065 5794 (lines open from 09:00-17:00 Mon-Fri), mail: accreditation@bsecho.org.



Exam fees

A fee of ± 375 is charged for the complete accreditation process. This fee is payable upon registration for the written section of the examination and covers the practical assessment. There is a non-refundable booking fee of ± 50 upon registering for a secured placement at the practical assessment.

Candidates who are unsuccessful in the written section of the examination will be charged a reduced fee of £187.50 to re-sit this section. This reduced fee only applies to the second attempt if taken within 12 months of an unsuccessful first attempt.

Candidates are entitled to one re-attempt at the practical assessment. A re-attempt at the practical assessment is subject to an additional fee of £187.50.

Fee increases may occur annually.

Extensions

Extensions to the 24-month deadline may be granted. Extension request forms must be submitted **before the submission deadline**. Requests received after the case deadline may not be granted.

Less-than-full-time extensions are available for up to 24 months for candidates working less than fulltime as stipulated by their contracted hours. Further information can be found on the BSE website's <u>extension request</u> page.

Appeals

Candidates can appeal the decision on a practical assessment result. There is no appeals process for the written section of the examination. Further information on applying for an appeal can be found on the <u>practical assessment</u> page of the BSE website.

Mentor

A mentor is an experienced echocardiographer who can successfully guide a candidate through the BSE accreditation process. If the echocardiographer is BSE accredited, this is an advantage but not essential.

The mentor should understand the accreditation process, including the training syllabus (see Appendix 1) and all relevant assessment criteria.

The mentor must assess the candidate's ability to perform an echocardiogram proficiently. Once a proficient level of ability is achieved, the mentor must complete the curriculum-based competency tool and the mentor statements. These can be accessed and completed via the online logbook portal. The curriculum-based competency tool can also be found in Appendix 2.

Candidates who cannot find a mentor should <u>contact us</u>; we will do our best to help them find a suitable mentor.



Written Theory Examination

Appendix 1 contains the whole training syllabus for this accreditation process, and Appendix 3 includes a recommended reading list.

The written theory examination is held twice a year, usually in the Spring and Autumn. It is held at various Pearson VUE centres across the UK, the Republic of Ireland, and some overseas locations. Registration dates are announced on the written assessment section of the BSE website. See Appendix 4 for registration guidance.

The written examination has two parts: an MCQ theory section and an Image reporting section. To pass the written examination overall, it is necessary to pass both parts at the same exam sitting.

If the first attempt is unsuccessful, candidates may be eligible to retake the exam at a reduced rate.

Reduced rate: This only applies to a second attempt if it is taken within 12 months of the first attempt. If the second attempt is unsuccessful, the next attempt will be charged at the full fee.

There is no bar to re-sitting the written examination any number of times.

The pass mark for the MCQ is 70%, and the pass mark for the image reporting section is 60%. Following moderation, the Accreditation Chair may decide to vary these slightly.

Accreditation is awarded once a candidate has successfully completed the practical assessment. Satisfactory performance at the written assessment alone does not allow 'partial accreditation.'

Multiple-choice section

- Consists of 25 questions that must be answered within 60 minutes.
- Questions are designed to test the knowledge of echocardiographic findings, basic cardiology and the physics of ultrasound.
- Each question comprises a brief statement followed by five questions. Candidates are required to answer 'true' or 'false' to each question. Example questions are provided in Appendix 5.
- This part of the examination will be marked +1 for correct answers and 0 for incorrect or unanswered questions (no negative marking).
- There are no 'trick' questions.
- There are no fixed number of correct answers, i.e. for each question, every answer can be false or, every answer to be true or any combination of true or false.
- The maximum possible mark is 125.

Image reporting section

- Consists of 50 questions centred around 10 patient case studies that must be answered within 90 minutes.
- The candidate will be presented with 10 patient case studies. Each case study will consist of
 relevant patient details and various echocardiographic images.



- For each case study, the candidate must answer five questions. Each question will have four possible answers; the candidate must select the best single answer. An example case study and questions are provided in Appendix 6.
- The maximum possible mark is 50.

Practical Assessment

The practical assessment is held up to five times per year. Dates, locations and online registration instructions are announced on the practical assessment section of the BSE website.

The practical assessment has three parts: a 250-case logbook, a practical scanning assessment, and a viva assessment of five patient case studies.

All candidates must attend an assessment within 26 months of starting the accreditation process (i.e., within two months of their case collection deadline). A two-month grace period gives the candidate time to review, prepare, and submit the logbook and five viva cases.

- Registration for the practical assessment should ONLY be sought after collecting the logbook and patient case studies.
- It is the candidate's responsibility to complete online registration forms and update personal information correctly.
- If you have any concerns about the information provided, you should contact the <u>accreditation team</u> for guidance and support.

Logbook submission

The logbook should demonstrate the candidate's ability to meet the competencies, as shown in Appendix 2. The specific case mix of the logbook is shown below.

It should consist of **250 reports** personally **performed and reported** by the candidate during the specified 24-month period.

If the candidates hold BSE or EACVI TOE Accreditation, the logbook is reduced to 150 reports. The logbook numbers are not reduced for candidates holding EAVCI TTE accreditation.

The logbook format is copies of the actual clinical report. The reports must be uploaded and submitted via the BSE online logbook portal. Please see the portal user instructions in Appendix 7.

Non-portal logbooks will not be accepted.

Please see Appendix 8 for full details of what is expected in reports and how the logbook is marked.

Duplicate reports are not acceptable.

If you have problems finding enough specific cases, discuss this with your mentor, who may consider arranging for you to attend a nearby centre.

The logbook should reflect the candidate's best clinical practice. As such, targeted scans should not be included unless they show significant and rare pathology.

Competencies and mentor statements are to be completed via the BSE online logbook portal.

Fully subscribed BSE members can request access to the portal before taking the written examination by emailing accreditation@bsecho.org. See Appendix 7 for further guidance.



The logbook should reflect the normal case-load of a general adult department with the following constraints:

- > At least 25 cases should be for left ventricular abnormality assessment*
- At least 50 cases should be for valve disease assessment**
- > At least 10 cases should be for replacement/repaired valves
- At least 10 cases should be for right ventricular abnormality assessment***
- > At least 5 cases should be for pericardial disease/effusion assessment
- At least 5 cases should be for abnormalities of the aorta
- > At least 2 cases should be for confirmed endocarditis, mass or thrombus
- At least 5 cases should be for left ventricular hypertrophy assessment, at least 2 should be for hypertrophic (-/+ obstructive) cardiomyopathy
- > At least 3 cases should be for simple congenital disease (e.g. ASD, VSD, PDA, BAV)
- A maximum of 15 cases should be for specialised studies (i.e. bubble echo and contrast studies). This section is not compulsory.
- A maximum of 30 cases should be for no significant abnormality.

* This section should demonstrate a candidate's ability to assess for left ventricular abnormalities (normal / dilated cavity size, systolic impairment with global or regional wall motion abnormalities or diastolic impairment). At least half of the reports in this section should include left ventricular ejection fraction measurements (by manual, semi-automated or automated tools).

** This section should demonstrate a candidate's ability to assess for all severities of valve pathology and not primarily mild disease. The majority of these studies should consist of moderate to severe pathology.

*** This section should demonstrate a candidate's ability to assess for right ventricular abnormalities (normal / dilated cavity size, systolic impairment with global or regional wall motion abnormalities).

Other information regarding the logbook:

- All patient-identifiable data needs to be removed. This may require the manual removal of identifiable data. See <u>Appendix 9.</u>
- At least the final 150 cases should be reported primarily by the candidate, although another operator may check them.
- Logbook reports should reflect the latest <u>BSE guidance</u>. Where local policy deviates from this, a supporting letter (and current standing operating procedure) from the department echo lead stating local policy should be included. This should be submitted under the "optional supporting information" section on the BSE logbook portal.
- The candidate's name must appear on the report as the performing and reporting echocardiographer/sonographer. Where local policy deviates from this, a supporting letter and current standard operating procedure from the department's Echo lead stating local policy should be included. This should be submitted under the "optional supporting information" section on the <u>BSE logbook portal</u>.



• The department's echo lead completes the logbook's final sign-off / validation. The portal user guide is in Appendix 7.

Scanning Assessment

This part of the assessment is designed to assess a candidate's practical scanning ability and ability to perform basic image optimization.

A candidate must acquire up to 10 different echocardiographic imaging views within 20 minutes. A real-life model or simulator may be used.

- All imaging views used in this assessment are taken from the minimum BSE transthoracic echocardiography dataset.
- A pass mark/trigger score of 66% is used. Once obtained, the candidate will be deemed successful at this part of the assessment process.
- The candidate is not expected to be familiar with the equipment. The Assessor will alter the equipment setting as directed by the candidate.
- For full details of the practical scanning marking criteria, please see Appendix 10.

Viva case submission

Consists of a viva assessment of five separate patient case studies. See below for the required case mix.

The candidate will be expected to discuss their patient cases with the Assessor. All five cases may be reviewed.

For full details of the viva case marking criteria, please see Appendix 11.

The case studies should be assessed using the most up-to-date BSE guidance. Candidates are expected to know this guidance, and local deviations in practice **will not** be accepted.

The cases must represent a complete, high-quality study. They should be accompanied by a printed report that is complete and comprehensive and reflects the patient case study presented.

The candidate must ensure that at least one complete cardiac cycle is recorded. The cases must play automatically / continuously within a PowerPoint presentation (or equivalent). Cases that do not play appropriately may be classified as an unsuccessful attempt.

Candidates must bring and present their patient case studies on their own laptop. It is the candidate's responsibility to ensure these are anonymised and can be viewed in a manner that allows an assessment of the cases being presented.

The viva case studies should include one of each of the following:

- 1. A study showing no significant abnormality.
- 2. Moderate or severe aortic stenosis.
- 3. Moderate or severe mitral or aortic regurgitation.
- 4. Regional wall motion abnormality.
- 5. The fifth case should show an example of one of the following (which has not previously been shown in the cases above):



- a) Valve repair/replacement.
- b) Mass or thrombus.
- c) Simple congenital heart disease.
- d) Significant left ventricular hypertrophy.
- e) Significant pericardial effusion, mitral stenosis or right heart disease.

Tips on getting suitable video cases are available online- check the <u>TTE accreditation</u> page.

Patient case studies may be used in subsequent BSE written exams, educational and training sessions

Practical assessment - outcomes and process for re-attempts (resubmissions)

A candidate will have two attempts at passing the practical assessment part of the accreditation process. A second attempt (referred to as resubmission) at the practical assessment is subject to a fee of £187.50.

- If a candidate is successful in all three parts of the practical assessment, the candidate will be awarded BSE accreditation and will join the <u>accredited member list</u>.
- If a candidate is unsuccessful in any of the three parts of the practical assessment, the candidate will be deemed unsuccessful at this first attempt. The candidate will be given constructive feedback to facilitate a re-attempt. The candidate may be requested to resubmit logbook reports/patient case studies. These must be new reports / patient case studies. A candidate is not permitted to resubmit previously assessed work under any circumstance.
- If a candidate fails the second attempt (resubmission), the accreditation process must start over, with the candidate undertaking the written examination again.

In the event of an unsuccessful attempt, the candidate is required to:

Attend another practical assessment and re-attempt **ONLY** the parts of the practical assessment that the candidate was unsuccessful at in the first attempt. The pass marks from the remaining practical assessment elements will be upheld.

The timescale allowed for re-attempts (resubmissions) will depend on which elements were unsuccessful and the candidates' current and future work commitments. This will be discussed with the candidate during the first attempt. Typical timeframes may include 3-9 months and can be up to 12 months following the first attempt.

Our feedback consistently demonstrates that non-face-to-face feedback does not adequately equip a candidate to pass at the next sitting. Therefore, all re-attempts at the practical assessment require the candidate's attendance in person to facilitate adequate and helpful face-to-face feedback*

*We may authorise virtual or remote submissions, subject to committee approval.



Appendix 1: Training Syllabus

The following sections form the minimum suggested training syllabus for this accreditation process.

Candidates should use this as a guide to prepare for this accreditation process's written and practical assessments.

1 General Concepts

1.0 The place of echocardiography

- Information that echocardiography can, and cannot provide
- 'Ruling out' pathology (sensitivity, specificity & Baye's theorem)
- Likelihood of findings influencing patient management
- Undesirable outcomes: inaction while waiting for results, clinical 'red herrings'
- Indications for echocardiography
- Competing and complementary technology
- Cardiac catheterisation (ventriculography and coronary angiography)
- C-T imaging
- Magnetic resonance imaging
- Nuclear Cardiology

Service Provision

- Considerations of Physiologist-led versus physician-led service
- Costs: fixed and variable
- Provision and indication for specialised techniques, e.g. TOE. Stress echo, Contrast echo
- Availability and access
- Controlling workload
- Training & motivation of staff
- Audit, Quality Control, Clinical Governance
- Infection control

Relationship with patients

- Explaining the procedure in terms relevant to the particular patient
- Respect for patients' dignity and cultural backgrounds
- Relationships with colleagues
- Handling requests for information about the study findings



Reporting and Documentation

- Standard methods & terminology
- Distinction between Technical and Clinical reports
- Responsibility for reporting Medico-legal considerations (Data Protection Act)

2 Clinical role of echocardiography

2.0 Imaging Physics & Instrumentation

- Concepts and terminology
- Concepts of compression waves
- Definitions: frequency, wavelength, propagation velocity, amplitude?
- Units of measurement: Hz and MHz
- Decibel Comparison of Ultrasound with audible sound.

2.1 Propagation of ultrasound through tissues

- Speed of sound in different body tissues.
- Frequency range used for diagnostic imaging
- Distinction between specular reflection and backscatter
- Principles of attenuation and scattering

2.2 Ultrasound Transducers

- Piezo-electric effect
- General concepts of 2D and 3D transducer construction
- Characteristics of the ultrasound beam: Far (Fraunhofer) & Near (Fresnel) zones, side lobes
- Beam steering methods: mechanical & electronic
- Focusing methods, including dynamic receive focusing
- Focus position and use of dual focus
- The role of intracardiac echocardiography

2.3 Imaging physics

- Factors affecting choice of imaging frequency: typical practical values for adults & children
- Broad-band imaging



- Harmonic imaging
- and M Mode methods.
- Curved Anatomical M Mode
- Scanning speed limitations, relationships between pulse repetition frequency, frame rate, scan lines per frame, field of view, and depth to be imaged.
- Concept of Parallel Processing and its influence on frame rate and image quality
- Effect on evaluation of rapid motion
- Temporal resolution.
- Greyscale and dynamic range
- Measurement and optimisation of Resolution: axial, azimuthal and elevation
- Lateral resolution and side-lobe/grating artefacts
- Reverberation artefacts
- Limiting factors for detecting small targets

2.4 Echo Instrumentation

- Function of machine controls: Transmit power; overall gain; time gain compensation; reject, logarithmic compression, signal processing, dynamic range, pre-processing; post-processing
- Optimisation of imaging parameters, including transducer frequency, scan angle, gamma correction, spatial and temporal smoothing
- Optimisation of 3D volume acquisitions including frame/volume rate, cropping and manipulation of viewing plane
- The advantages of 3D echocardiography over 2D echocardiography, e.g. appreciation of mitral valve pathology, elimination of geometric assumptions in cardiac chamber volume estimations

2.5 Optimising Images

- Use of gel (infection risk from transducer, operator)
- Positioning of the subject
- Standard views: Parasternal, apical (4, 5 and 2-chamber, long axis), subcostal, suprasternal, right parasternal, long and short axis.
- Use of non-standard views
- Adapting for subjects with difficult echo windows, ventilated patients, ward-based studies, emergency room studies

2.6 Storage and Display of Images

- Basic concept of digital acquisition and storage systems. Scan converters and digital memories.
- Display devices and controls, recording techniques



Basic understanding of digital image processing and recording methods: pixel density, volume
of data, the DICOM standard, concept of data compression (JPEG, AVI, etc.) archiving of
echocardiographic studies on magneto-optical discs, CD/DVD, portable solid-state memories,
ECG-gated acquisitions vs. continuous recording, facility to review acquired loop prior to
storage, facility to choose the number and type of cardiac cycles to be recorded, facility for
offline image properties adjustment and further quantitative analysis.

3 Doppler Physics & Fluid Dynamics

3.0 Basic Fluid Dynamics

- Fluid flow: significance of peak & mean velocities
- Determination of volumetric flow, Continuity equation
- Laminar & turbulent flow: Reynolds' equation (qualitative)
- Transition from Laminar to turbulent flow: inlet jet Bernoulli equation
- Bernoulli principle for fluid dynamics relationship of fluid speed and statics pressure/potential energy
- Coanda effect

3.1 Principles of Doppler

- Interaction of ultrasound waves with moving blood: The Doppler effect
- The Doppler equation: factors influencing magnitude of Doppler shift
- Spectral analysis: fast Fourier transform (qualitative)
- The spectral Doppler display: determination of mean, modal and peak velocities
- Limitation of CW Doppler caused by lack of depth discrimination
- Audible range of Doppler shift frequencies
- The effect of beam angle errors on Doppler velocities
- Aliasing: how it is caused and how it manifests in practice: The Nyquist limit
- Influence on aliasing of: transducer frequency; sample depth (range x velocity product); and beam angle
- High pulse repetition frequency (extended range) PW Doppler and the phenomenon of range ambiguity
- Relative advantages and disadvantages of CW, PW and HPRF modes
- Concept of colour flow imaging as multi-sampled PW
- Velocity estimation by moving target indication and autocorrelation (qualitative)
- Limitations of mean velocity: use of velocity variance to show high velocities/ turbulence
- Aliasing in colour Doppler and the effect of scan frequency on the Nyquist limit
- The principles of pulse wave tissue Doppler
- Packet size, colour mode and sector size and their effect on frame rate and aliasing



4 Deformation Analysis

4.0 Principles of Myocardial Deformation

- The definition of displacement, velocity, strain and strain rate
- The cardiac ultrasound coordinate system for describing motion and deformation: longitudinal, radial, circumferential and rotational axes
- Quantifying myocardial deformation as opposed to velocity or displacement
- Concept of shear deformation; rotation of the base and apex of the left ventricle, and the resultant twisting deformation or torsion

4.1 Quantifying myocardial strain and strain rate by tissue Doppler

- The concept of the myocardial velocity gradient
- The concept of strain and strain rate to define deformation
- Tissue Doppler imaging for deriving strain and strain rate: practical parameters in measuring strain and strain rate (e.g. sample size and shape, offset distance, drift compensation, spatial and temporal averaging, tracking of sample volume)
- Reproducibility issues

4.2 Speckle Tracking Echocardiography/2D strain

- Familiarity with the concept of speckles and speckle tracking in greyscale 2D loops
- Speckle tracking for angle-independent derivation of velocities, displacement, strain and strain rate
- The impact of frame rates on the quality of speckle tracking
- Speckle tracking vs. tissue Doppler techniques for assessing myocardial motion and deformation
- Speckle tracking for measuring left ventricular rotation and torsion
- Kindred technologies

5 Doppler instrumentation

5.0 Spectral Doppler Instrumentation

- Duplex Doppler using imaging transducers
- The 'Stand-alone' Doppler probe
- Features of the spectral display: positive & negative velocities; scale & baseline controls.
- Effect of high-and low-pass filter and intensity threshold ('reject') settings
- Pulsed Doppler sample volume: influence of gate length and distance (beam width)
- Representation of signal strength by image intensity
- How aliasing manifests on the spectral display.



5.1 Colour Flow Instrumentation

- The colour display: BART convention
- Colour maps to show velocity scales
- Image domination and additive colour modes
- Difference between velocity and power (signal amplitude) displays
- Basic principles of Tissue Doppler Imaging, including optimisation of filters for detecting tissue versus blood velocities, sample volume and size, impact of interrogation angle on measured velocities, minimising aliasing, and maximising frame rates to detect short duration myocardial motion
- Differences between colour Doppler tissue Doppler Imaging and pulsed wave tissue Doppler imaging
- Minimisation of myocardial translational movements during acquisition.
- The concept of tracking on colour Doppler tissue Doppler imaging to ensure that sample volume remains in the region of interest
- Parametric (curved M-mode) display of tissue Doppler images
- The relevance of importing cardiac cycle time points, such as aortic valve closure, into tissue Doppler traces

6 TOE Instrumentation

6.0 General concepts

- Transducer types: single plane, biplane, multiplane
- Optimising machine settings for TOE Patient monitoring for TOE and general safety considerations
- Control of infection
- General indications and recognition of the limitations of TTE.

7 Safety of ultrasound

- Potential hazardous biological effects: heating, resonance and cavitation effects
- Measurement of beam intensity (SPTA)
- Practical precautions: power levels, use of colour and CW Doppler

8 Cardiac Anatomy and Physiology

8.0 Anatomy of the thorax

- Thorax contained by rib cage & diaphragm
- Lungs & pleura; heart & pericardium; mediastinum
- Blood vessels within the thorax



8.1 Gross anatomy of the heart

- Basic cardiac embryology
- Nomenclature of chambers and valves
- Major relationships of chambers, valves and blood vessels
- Distinguishing features of valves and chambers as related to echocardiography
- The pericardial sac

8.2 Cardiac anatomy and physiology as demonstrated by echocardiography

- Detailed structural anatomy of the heart, great vessels and pericardium
- Visualisation of normal cardiac anatomy and normal variants in standard echocardiographic planes
- 2D/3D, M-mode and Doppler features of normal valve anatomy (aortic, mitral, tricuspid and pulmonary), function and normal variants
- The phases of atrial function: reservoir, conduit and contractile phases
- The LV remodelling process in response to disease: eccentric (chronically elevated preload) vs. concentric hypertrophy (chronically elevated afterload)

8.3 The Cardiac Cycle

- Temporal relationships of the ECG, chamber pressures and valve movements
- Typical values for intracardiac pressures
- Relationship of valve movements to heart sounds
- Identification of valve opening and closure signals on Doppler recordings
- The timing of aortic valve closure as a marker of end-ejection, as derived from M- mode, blood flow Doppler or tissue Doppler

9 Cardiac functional parameters

9.0 Measurements and calculations

- On-screen measurement of length, slope, area, volume and time interval, and their significance for 2-D, 3D images, M-mode and spectral Doppler displays
- Knowledge, appreciation and limitations of M-mode measurements (including MAPSE and TAPSE, Ao / LA dimensions)
- Derivation of stroke volume, ejection fraction, relative wall thickness, LV Mass and indexed LV
 mass
- 2D / 3D methods of measuring LV volume
- Limitations of single plane estimations of LV ejection fraction e.g. Teicholtz formula method



- Limitations of single plane measurements of LA size
- Geometric assumptions used in estimation of cardiac chamber volumes with M-mode and 2D imaging
- The advantages of deriving volumes and ejection fraction by 3D echocardiography
- Limitations of measurement and/or calculation validity in the presence of poor quality and/or off-axis images

9.1 Doppler determination of cardiac output, ejection time, valve function and velocity acceleration

- Methods for assessing normal valve function (aortic, mitral, tricuspid and pulmonary) to include: peak and mean velocities, peak and mean gradients, pressure half time and flow rate assessments (where appropriate)
- Methods of measuring diastolic function: E/A ratio, deceleration time, pulmonary venous flow patterns, the ratio of the peak early diastolic transmitral velocity and the peak early diastolic tissue velocity of the mitral valve annulus (the E/E' or E/Ea) ratio methods for estimating LV filling pressures, mitral valve propagation velocity
- Peak and mean pressure gradient measurements by Doppler and their relationship to catheterisation data
- Measurement of pulmonary pressures from tricuspid and pulmonary regurgitant flow velocities and assessment of inferior vena cava contraction during inspiration

10 Contrast and bubble contrast studies

10.0 Bubble contrast studies

- Main indications for a bubble contrast study: Diagnosis of intracardiac shunts, diagnosis of intra-pulmonary shunts, improved assessment of tricuspid regurgitation velocities.
- Technique for performing a hand-agitated contrast bubble study
- Acoustic views required and optimisation of machine control settings for bubble contrast studies
- Patient manoeuvres to provoke right -to-left passage of bubbles during assessment for PFO
- Knowledge of findings consistent with a positive and negative intra-cardiac and intrapulmonary shunt.
- Relevance of injecting bubble contrast through upper arm vein vs. femoral vein
- Technique for performing a hand-agitated contrast study
- Contra-indications and clinical precautions



10.1 Awareness of encapsulated contrast agents and techniques

- Knowledge of available contrast agents
- Knowledge of contrast agent's characteristics including interaction of ultrasound
- Generation of harmonic energy by bubble distortion and fracture
- Optimisation of machine control settings for contrast agents
- Indications for administration of contrast agents to include: Enhancing endocardial definition for assessment of regional contractility and accurate cardiac volume estimations, detection of intracardiac masses, distinguishing thrombus from a vascular tumour, diagnosis of cardiomyopathies (e.g. non-compaction), arrhythmogenic right ventricular dysplasia, myocardial perfusion assessment.
- Use of contrast in stress echocardiography for improving detection of wall motion abnormalities and for assessment of myocardial perfusion
- Appreciation of contrast administration bolus vs infusion
- Contra-indications and clinical precautions

Valve pathology

11 Mitral valve disease

11.0 Mitral Stenosis

- Aetiologies and typical 2D/3D echocardiographic features: rheumatic, calcific, myxoma / tumours, cor-triatriatum, congenital
- Qualitative description of valve and sub-valve calcification and fibrosis
- Assessment of mitral stenosis severity to include: mean gradient, planimetry (2D and 3D), pressure half time method, continuity equation, PISA method: techniques and limitations
- Factors favouring successful balloon valvuloplasty: Wilkins score
- Role of exercise stress echocardiography to evaluate for changes in mean trans-mitral gradient, PA systolic pressures, exercise tolerance and symptomatic status with exercise to aid in the timing of surgery/balloon valvuloplasty
- Role of echocardiography in assessment and follow-up

11.1 Mitral regurgitation

 Aetiologies and typical 2D / 3D echocardiographic features of primary mitral regurgitation: mitral valve prolapse (fibro-elastic deficiency, myxomatous mitral valve, flail leaflet, Barlow's disease), ruptured chordae, infective endocarditis, mitral annular calcification, rheumatic, congenital causes, drug and / or radiotherapy-induced



- Aetiologies and typical 2D / 3D echocardiographic features of secondary mitral regurgitation: Ischemia and impairment/dysfunction of sub-valvular apparatus (i.e. papillary muscle dysfunction or rupture), LV dilatation, leaflet tethering, LA dilatation, annular dilatation
- Chronis vs acute mitral regurgitation and differential diagnosis
- Awareness of Carpentier classification
- Consequences of mitral regurgitation on left and right heart chamber sizes and right heart pressure
- Assessment of mitral regurgitation severity to include: vena contracta, PISA (2D and 3D), effective regurgitant orifice area, regurgitation volume, regurgitation fraction, MV VTI/AV VTI, increased antegrade E wave velocity and shape and density of contour of Doppler signal: techniques and limitations
- Pulmonary vein flow patterns seen in all mitral regurgitation severity ranges: techniques and limitations
- Role of echocardiography in determining timing of surgery for primary mitral valve disease: ejection fraction, end-systolic LV diameter, EROA, resting PA pressure.
- Role of TOE in assessing mitral valve pathology and in determining likelihood of repair as opposed to replacement
- Role of exercise stress echo to evaluate for MR severity changes, LV function assessment, PA systolic pressures, symptomatic status and exercise tolerance during exercise to aid in the timing of surgical intervention.
- Role of echocardiography in assessment and follow-up

12 Aortic valve disease

12.0 Aortic stenosis

- Aetiologies and typical 2D / 3D echocardiographic features of aortic stenosis: rheumatic, bicuspid (and classification of type), senile degenerative, sub-and supra-valve obstruction
- Assessment of aortic stenosis severity to include: maximum velocity, mean gradient, aortic valve area by continuity equation (including indexed values), valve planimetry, dimensionless index: techniques and limitations.
- Consequences of aortic stenosis on cardiac chamber size and function
- Appreciation of the causes of discordant parameters when assessing aortic stenosis and potential remedies.
- Use of apical, right parasternal and suprasternal positions to obtain optimal AV Doppler parameters.
- Definition of low flow low gradient severe aortic stenosis
- Concept of flow-rate and effect on transvalvular velocities
- Use of stress echocardiography for distinguishing pseudo-severe stenosis vs truly severe stenosis in low flow aortic stenosis



- Use of stress echocardiography in patients with low flow low gradient severe AS and assessing for LV contractile reserve
- Difference between transaortic pressure gradients derived from echocardiography and from cardiac catheterisation
- Role of echocardiography in assessment and follow-up

12.1 Aortic Regurgitation

- Aetiologies and typical 2D / 3D echocardiographic features of aortic regurgitation: rheumatic, bicuspid, aortic root disease, infective endocarditis (including root abscesses).
- Assessment of aortic regurgitation severity to include: Colour Doppler size of jet relative to left ventricular outflow tract diameter, vena contracta, effective regurgitant orifice area, regurgitant volume, diastolic flow reversal in descending aorta, diastolic flow reversal in the abdominal aorta (including BP pulse pressure width), indirect effects on LV size and function: techniques and limitations.
- Consequences of aortic regurgitation on cardiac chamber size and function
- Role of echo in determining timing of surgery
- Role of TOE in assessing aetiology and severity
- Role of echocardiography in assessment and follow-up

13 Tricuspid valve disease

13.0 Tricuspid stenosis

- Aetiologies and typical 2D / 3D echocardiographic features of tricuspid stenosis: rheumatic, prolapse, congenital, infective endocarditis, carcinoid, functional.
- Assessment of tricuspid stenosis severity to include: Mean pressure gradient, inflow velocitytime integral, pressure half time and valve area by continuity equation: techniques and limitations.
- Consequences of tricuspid stenosis on cardiac chamber size and function
- Role of echocardiography in assessment and follow-up

13.1 Tricuspid regurgitation

- Aetiologies and typical 2D / 3D echocardiographic features of tricuspid regurgitation: rheumatic, prolapse, congenital, infective endocarditis, carcinoid, functional /secondary (RA, RV, annular dilatation), trauma and device leads.
- Assessment of tricuspid regurgitation severity to include: Colour Doppler shape and density of continuous Doppler signal, effective orifice area, regurgitation volume (by PISA), colour flow area, PISA, vena contracta, tricuspid inflow, colour Doppler signal, hepatic vein flow



pattern, indirect effects on RA, RV, IVC and intraventricular septal motion: techniques and limitations

- Consequences of tricuspid regurgitation on cardiac chamber size and function
- Role of echocardiography in assessment and follow-up

14 Pulmonary valve disease

14.0 Pulmonary stenosis.

- Aetiologies and typical 2D / 3D echocardiographic features of pulmonary stenosis: rheumatic, congenital, infective endocarditis, carcinoid, sub-valvular and supra-valvular obstruction, infundibular obstruction
- Assessment of pulmonary stenosis severity to include: peak velocity, peak gradient: techniques and limitations
- Consequences of pulmonary stenosis on cardiac chamber size and function
- Role of echocardiography in assessment and follow-up

14.1 Pulmonary regurgitation

- Aetiologies and typical 2D / 3D echocardiographic features of pulmonary regurgitation: congenital, endocarditis, carcinoid, rheumatic, secondary (post valvuloplasty, post TOF repair)
- Assessment of pulmonary regurgitation severity to include: jet width percentage of RVOT, vena contracta percentage of pulmonary valve annulus, deceleration time, pressure half time, 3D vena contracta, Doppler pulmonary regurgitation index, origin of PR jet in relation to pulmonary artery bifurcation: techniques and limitations
- Consequences of pulmonary regurgitation on cardiac chamber size and function
- Role of echocardiography in assessment and follow-up

15 Infective endocarditis

- Typical bacteraemia / fungal causes of infective and non-infective endocarditis
- Use of Duke criteria for infective endocarditis
- Typical 2D / 3D echocardiographic features of vegetations for bacteraemia / fungal causes of infective and non-infective endocarditis
- Typical and atypical locations of vegetations
- Complications of endocarditis to include: abscess, fistula, perforation, valve destruction and regurgitation, prosthetic valve dehiscence, new paravalvular regurgitation, healed/ chronic vegetations
- Infective and non-infective endocarditis associated with congenital heart disease and hypertrophic cardiomyopathy



- Role of TOE in suspected endocarditis
- Role of echocardiography in assessment and follow-up

16 Prosthetic heart valves

- Typical 2D/3D, M-mode and Doppler features of the main types of replacement / repaired valves to include: Mechanical (tilting disc, bilealfet and ball and cage), bio-prosthese (stented and stentless), leaflet repair ± annuloplasty rings, percutaneous valve intervention (mitral clip and TAVI).
- Assessment of age-related deterioration of bioprostheses
- Assessment of artefacts, pannus, thrombus and vegetations (and associated complications) on prosthetic valves
- Role of TOE in examining normal and malfunctioning prosthetic valves
- Assessment of prosthetic valve stenosis to include: 2D, M-mode and Doppler assessment, use of continuity equation and indexed values, the phenomenon of pressure recovery
- The assessment of normal and abnormal aortic prosthetic valve function and differentiation between high flow states, patient-prosthesis mismatch and insignificant / significant stenosis. To include the use of maximum velocity, acceleration time: techniques and limitations.
- The assessment of normal and abnormal mitral prosthetic valve function and differentiation between normal, possible and significant prosthetic stenosis. To include: peak velocity, mean gradient, VTi, effective orifice area and pressure half time: techniques and limitations.
- The assessment of normal and abnormal tricuspid prosthetic valve function. To include: mean gradient, pressure half time, tricuspid valve E velocity, VTi: techniques and limitations
- The assessment of normal and abnormal pulmonary prosthetic valve function. To include: mean gradient, peak velocity: techniques and limitations: techniques and limitations
- Assessment of consequences of prosthetic valve dysfunction, to include: chamber dilatation, progression to pulmonary hypertension
- Assessment of prosthetic valve regurgitation to include: trans-versus para-valvar regurgitation, normal versus abnormal prosthetic valve regurgitation, assessment by CW, PW and Colour Doppler: techniques and limitations.
- Role of echocardiography in assessment and follow-up

17. Cardiomyopathies

17.0 Dilated cardiomyopathy

- Aetiologies and typical 2D/3D echocardiographic features of dilated cardiomyopathies
- Detection and assessment of associated lesions to include: functional valve regurgitation, thrombus in cardiac chambers, pericardial effusions, pulmonary hypertension
- Role of echocardiography in assessment and follow-up



17.1 Hypertrophic cardiomyopathies

- Aetiologies and typical 2D/3D echocardiographic features of hypertrophic cardiomyopathies
- Techniques for measurement of left ventricular wall thickness, detection of left ventricular outflow tract obstruction and intracavity gradient (including the use of breath hold and provocative manoeuvres to assist in the detection of inducible gradients)
- Assessment of right ventricular involvement
- Associated abnormalities to include: systolic anterior motion mitral valve and associated mitral regurgitation, apical aneurysms and associated thrombus, abnormal papillary muscle location
- Differentiation from other causes of hypertrophy, e.g. hypertension, athletic heart', amyloidosis, Fabry's disease, Friedreich's ataxia cardiomyopathy
- Role of echocardiography in assessment and follow-up

17.2 Restrictive cardiomyopathy

- Aetiologies and typical 2D/3D echocardiographic features of restrictive cardiomyopathies: storage / infiltrative disorders (e.g. Fabry's, Danon disease and Friedrich ataxia), amyloidosis, sarcoidosis, idiopathic, endomyocardial fibrosis, carcinoid heart disease, radiation or drug induced
- Assessment of restrictive cardiomyopathies to include: 2D findings, Doppler & TDI features (small to normal LV cavity size, normal wall thickness, normal or near normal LVEF, reduced GLS, associated GLS patterns, dilated atria, increased E/A ratio, reduced deceleration time, reduced early diastolic velocities, increased E/E' ratio, reduced S velocities)

18 LV non-compaction

- Aetiology and typical 2D/3D echocardiographic features of LV non-compaction
- Assessment of LV non-compaction to include: Visual assessment of prominent LV trabeculation and deep recesses. Non-compacted: compacted wall ratio of >2:1, colour Doppler flow within deep recesses, global LV systolic function assessment, thrombus assessment, abnormal papillary muscle structure
- Role of contrast agents

19 Intra-cardiac masses

- Aetiology and typical 2D/3D echocardiographic features and locations of masses to include: thrombus, cardiac tumours (primary and secondary) and myxoma's.
- Differentiation of myxoma from other cardiac tumours



- Features suggestive of malignancy
- Role of TOE in the assessment of intracardiac masses
- Role of contrast in the assessment of intracardiac masses

20 Pericardium and pericardial pathology

20.0 Pericardium

- Anatomy of the normal pericardium
- Relationships of serous pericardium to heart and great vessels
- Transverse and oblique sinuses of the pericardium

20.1 Echocardiographic features of pericardial fluid

- Location of fluid in relation to patient position and fluid volume
- Differentiation from pleural effusion
- Assessment of volume of pericardial fluid
- Role of echocardiography in pericardiocentesis

20.2 Features of tamponade

- Collapse of RA and/or RV walls
- Effect on IVC and hepatic vein flow pattern
- Effect on A-V valve flow velocities during respiratory cycle

20.3 Features of pericardial constriction

- Pericardial thickening/ appearance
- Effect on A-V valve flow velocities
- Effect of respiration

21 Coronary artery disease and LV systolic function

- Assessment of global LV systolic function to include: 2D/3D, M-mode and Doppler indices, Simpson's biplane ejection fraction, stroke distance, stroke volume, stroke volume index and cardiac output.
- Use of tissue Doppler and speckle tracking echocardiography for assessment of regional
- myocardial velocities and deformation in ischaemic heart disease, at rest and with stress



- Longitudinal function of the left ventricle, as assessed by M-mode (MAPSE), tissue Doppler of the mitral valve annulus and global longitudinal strain analysis
- The concept of post-systolic contraction
- The concept of isovolumic acceleration by tissue Doppler
- Left ventricular torsion and its implications for systolic function of the LV
- Appreciation of assessing for cardiotoxic effects of cancer therapy and the impact of LV systolic function

22 Diastolic function of the left ventricle

• The 4 stages of diastolic function as assessed by transmitral flow Doppler, deceleration time, E': techniques and limitations

Doppler for assessing diastolic dysfunction:

- Effect of LA size
- Pressures and pseudo-normalisation, effect of mitral regurgitation
- The use of Valsalva manoeuvre in reducing LA pressures to differentiate normal from pseudo normal transmitral Doppler patterns
- The use of left atrial size, IVRT, tissue Doppler (diastolic longitudinal velocities of the mitral valve annulus, the E/E' ratio), pulmonary vein flow pattern and mitral propagation velocity
- for assessing diastolic function
- The importance of untwisting in left ventricular filling
- Assessment and knowledge of LA strain (reservoir)

23 Left ventricular dyssynchrony and assessment by echocardiography

- Techniques for measuring interventricular and intraventricular dyssynchrony for predicting response to cardiac resynchronisation treatment
- Tissue Doppler quantitation of intraventricular dyssynchrony and their limitations
- Techniques for optimising settings of the cardiac resynchronisation device after implantation

24 Stress Echocardiography

- Indications and basic knowledge of techniques for exercise, dobutamine or vasodilator stress echocardiography
- Exercise or pharmacological stress echocardiography for diagnosis of ischaemic heart disease and myocardial viability



- The concept of viable and hibernating myocardium, and the relevance of the various responses of the myocardium to stress
- The concept of contractile reserve and flow reserve (for AS)
- The American Society of Echocardiography regional wall motion scoring system
- Dobutamine stress echo in 'low flow' aortic stenosis
- Exercise stress echo in valvular heart disease and pulmonary hypertension

25 Myocardial Infarction and its sequelae

- Ability to assess for regional wall motion abnormalities (hypokinesia, akinesia and dyskinesia) and global LV systolic function/impairment
- Knowledge of 2D, M-mode and Doppler features of MI complications to include: postinfarction VSD, mitral papillary muscle rupture, cardiac tamponade, mural thrombus, myocardial scarring, Dressler's syndrome, left ventricular aneurysm (true aneurysm vs. pseudoaneurysm)
- Assessment of the main features of stress-induced (takotsubo) cardiomyopathy as a deferential diagnosis to acute myocardial infraction

26 Pulmonary Hypertension and functional assessment of right ventricle

- Aetiologies of pulmonary hypertension to include: primary, post pulmonary embolism, lung disease, connective tissue disease and secondary to left-sided lesions
- Assessment of pulmonary hypertension by 2-D, M-mode and Doppler features of pulmonary hypertension
- Assessment of regional and global RV systolic function to include: TAPSE, RV S velocity, fractional area change of the RV, RV free wall 2D global longitudinal strain.
- Assessment of right ventricular dysfunction in acute pulmonary embolism (McConnell's sign and 60/60 sign) and chronic pulmonary embolism

27 Diseases of the Aorta

- Assessment of the aortic root (sinuses of valsalva and ST junction), proximal ascending, aortic arch, descending thoracic aorta and abdominal aorta by 2-D, M-mode and Doppler.
- Assessment of Marfan syndrome, sinus of Valsalva aneurysm, thoracic aortic aneurysm, aortic dissection (to include aortic cusp prolapse, aortic regurgitation, pericardial fluid) by 2D, M-mode and Doppler.

28 Adult Congenital Heart Disease



- Anatomy, pathophysiology and natural history of common congenital lesions present in adults to include; 2-D, M-mode and Doppler features of the following (pre-operatively and post-operatively, as seen in the older child or adult):
- Atrial septal defects (primum, secundum and sinus venous)
- Perimembranous and muscular ventricular septal defects
- Partial and complete atrio-ventricular septal defects
- Persistent ductus arteriosus
- Bicuspid aortic valve and associated aortopathy
- Sub-and supra-valve aortic stenosis
- Aortic coarctation
- Pulmonary stenosis
- Ebstein's anomaly
- Fallot's tetralogy
- Transposition and corrected transposition of the great arteries
- Role of contrast echocardiography in evaluating shunts in adults
- Calculation of shunts
- Role of TOE in adult congenital disease

29 Echocardiographic assessment of common clinical presentation of patients

- Heart failure or breathlessness
- Arrhythmia
- Ejection systolic murmur
- Hypertension
- Collagen abnormalities (including systemic sclerosis)
- Renal failure
- Stroke

30 Emergency and ICU Echo

30.0 General

• Constrained environment (multiple arterial/venous lines, ventilator, lighting issues etc)

31 The hypotensive/shocked patient and post-cardiac arrest

- Role of focused peri-arrest study and appreciation of limited echo windows
- Evaluation of LV (systolic and diastolic) and RV systolic function.
- Exclusion of severe valve disease (e.g. severe AS, endocarditis) and acute aortic dissection
- Assessment for pericardial effusion and cardiac tamponade, hypovolaemia and under-filling, and high output cardiac failure



- Septic shock assessment of for LV systolic and diastolic function
- Value of repeated echo studies to assess any deterioration/improvement in underlying state

32 Suspected acute pulmonary embolus

• Assessment of acute pulmonary embolus to include 2D, M-mode and Doppler assessment of RV size and function, tricuspid regurgitation and pulmonary artery systolic pressure assessment, IVC size and respiratory variation, thrombus presence in IVC/RA

33 Blunt and penetrating cardiac trauma

• Typical echocardiographic features including pericardial effusion, right and left ventricular contusion, acute valve lesions, aortic dilation and dissection/transection, VSD, pleural effusion

34 Echo in the ventilated patient

- Echocardiographic assessment for common clinical presentations to include 2D, M-mode and Doppler of acute arrhythmias such as fast AF, cardiac source of embolus CVA/peripheral embolic event in ventilated patients
- Value of TOE in ventilated patients (if poor transthoracic echo window)

35 Post surgery patient

- Appreciation of effects of general anaesthesia and cardio-pulmonary bypass on LV function
- Assessment of post-surgery haemodynamic compromise/ acute deterioration to include: common findings post cardiac surgery (tamponade, wall motion abnormalities and valvular dysfunction) and general surgery (air/fat embolism, venous thromboembolism, acute MI, volume overload)

36 Assessment of filling status

- Awareness of the role of echocardiography in assessing filling using LV and RV systolic and diastolic function, IVC, SVC and hepatic vein size and reactivity, atrial septal motion, chamber sizes and variation in Doppler velocities.
- Role of repeated echo studies in assessing effects of fluid challenge and inotropes



37 Additional topics

The level of knowledge expected is that of a competent echocardiographer performing transthoracic studies and sustaining knowledge through the <u>BSE and other educational resources</u>, including issues relevant to clinical scanning and practice raised in the <u>BSE Newsletter</u> (E-news).



Appendix 2: Curriculum-based Competency Tool

The following competency assessment tool should be used to ensure all knowledge and practical experience is covered during the candidates' training period.

The competency tool must now be completed by the candidate's mentor via the BSE <u>online logbook</u> <u>portal</u>.

Competency	Date achieved
1. BASIC ECHOCARDIOGRAPHY	
Knowledge	
Basic principles of ultrasound	
Basic principles of spectral Doppler	
Basic principles of colour flow Doppler	
Basic instrumentation	
Ethics and sensitivities of patient care	
Basic anatomy of the heart	
Basic echocardiographic scan planes	
Parasternal long axis standard, RV inflow, RV outflow	
Parasternal short axis including aortic valve, mitral valve and papillary muscles	
Apical views, 4- and 5-chamber, 2-chamber and long-axis.	
Subcostal and suprasternal views	
Indications for transthoracic and transoesophageal echocardiography	
Normal variants and artefacts	
Practical competencies	
Interacts appropriately with patients	
Understands basic instrumentation	
Cares for machine appropriately	L
Can obtain standard views	
Can optimise gain setting, sector width, depth, harmonics, focus, sweep speed,	
Doppler baseline and scale, colour gain	
Can obtain appropriate images and undertake accurate measurements	
Can recognise normal variants – Eustachian valve, chiari work, LV tendon	
Can use colour examination in at least two planes for all valves optimising gain	
and box-size	
Can obtain pulsed wave Doppler at:	
a. left ventricular inflow (mitral valve)	
b. left ventricular outflow tract (LVOT)	
c. right ventricular inflow (tricuspid valve)	
d. right ventricular outflow tract, pulmonary valve & main pulmonary artery	



2.LEFT VENTRICLE Knowledge

Coronary anatomy and correlation with 2D views of left ventricle Segmentation of the left ventricle for regional wall motion assessment Measurements of global systolic function. (LVOT VTI, stroke volume, fractional shortening, ejection fraction using Simpson's rule, S velocities) Doppler mitral valve filling patterns & normal range Appearance of complications after myocardial infarction:

- a. Aneurysm, pseudoaneurysm
- b. Ventricular septal rupture and papillary muscle rupture
- c. Ischaemic mitral regurgitation

Features of dilated, and hypertrophic cardiomyopathy Common differential diagnosis Athletic heart, hypertensive disease

Practical competencies

Can differentiate normal from abnormal LV systolic function Can recognise large wall motion abnormalities Can describe wall motion abnormalities and myocardial segments Can obtain basic measures of systolic function VTI, FS, LVEF, S velocities Understands & can differentiate diastolic filling patterns Can detect and recognise complications after myocardial infarction Can recognise features associated with dilated cardiomyopathy Can recognise features associated with hypertrophic cardiomyopathy Can recognise hypertensive heart disease Can recognise athletic heart

3. MITRAL VALVE DISEASE

Knowledge

Normal anatomy of the mitral valve and sub-valvular apparatus and their relationship with LV function Causes of mitral stenosis and regurgitation Ischaemic, functional, prolapse, rheumatic, endocarditis

Practical competencies

Can recognise rheumatic disease Can recognise mitral prolapse Can recognise functional mitral regurgitation Can assess mitral stenosis 2D planimetry, pressure half-time, gradient Can assess severity of regurgitation, chamber size, signal density, proximal flow acceleration & vena contracta

4. AORTIC VALVE DISEASE and AORTA Knowledge

Causes of aortic valve disease Causes of aortic disease Methods of assessment of aortic stenosis and regurgitation Basic criteria for surgery to understand reasons for making measurements

Practical competencies

Can recognise bicuspid, rheumatic, and degenerative disease Can recognise a significantly stenotic aortic valve Can derive peak & mean gradients using continuous wave Doppler Can measure valve area using the continuity equation



	1
Can recognise severe aortic regurgitation	
Can recognise dilatation of the ascending aorta	
Knows the echocardiographic signs of dissection	
5. RIGHT HEART	
Knowledge	
Causes of tricuspid and pulmonary valve disease	
Causes of right ventricular dysfunction	
Causes of pulmonary hypertension	
The imaging features of pulmonary hypertension	
The estimation of pulmonary pressures / probability of pulmonary	
hypertension	
Practical competencies	
Recognises right ventricular dilatation	
Can estimate PA systolic pressure / probability of pulmonary hypertension	
Can estimate right atrial pressure from the appearance of the IVC	
6. REPLACEMENT / REPAIRED HEART VALVES	
Knowledge	
Types of valve replacement / repair	
Criteria of normality	
Signs of failure	
Practical competencies	
Can recognise broad types of replacement valve	
Can recognise repaired valves	
Can recognise para-prosthetic regurgitation	
Can recognise prosthetic / repaired obstruction	
7. INFECTIVE ENDOCARDITIS	
Knowledge Duke criteria for diagnosing endocarditis	
Echocardiographic features of endocarditis	
Criteria for TOE	
Practical competencies	
Can recognise typical vegetations	
Can recognise an abscess	
Can recognise complications just on valve regurgitation	
8. INTRACARDIAC MASSES	
Types of mass found in the heart	
Features of a mxyoma	
Differentiation of atrial mass	
Normal variants and artifacts	
Practical competencies	
Can recognise a LA myxoma	
Can differentiate LV thrombus and trabeculation	



9. PERICARDIAL DISEASE	
Knowledge	
Features of tamponade	
RV collapse, effect on IVC, A-V valve flow velocities and respiratory variation.	
Features of pericardial constriction	
Differentiation of pericardial constriction from restrictive myopathy	
Differentiation of pericardial construction non restrictive myopathy	
Practical competencies	
Can differentiate a pleural and pericardial effusion	
Can recognise the features of tamponade	
Can judge the route for pericardiocentesis	
Can recognise restrictive physiology	
Differentiation of pericardial constriction from restrictive myopathy	
10. ADULT CONGENITAL HEART DISEASE	
Knowledge	
Anatomy and echo features of basic congenital disease:	
ASD, VSD, partial & complete atrio-ventricular defects	1 A A A A A A A A A A A A A A A A A A A
Patent ductus arteriosus	
Sub and supravalvar aortic stenosis	
Sub valvar, valvar and supra-valvar pulmonary stenosis	
Ebstein's anomaly	
Fallot's tetralogy	
Role of contrast	
Estimation of pulmonary artery pressure	
Practical competencies	
Can recognise a secundum ASD	
Can recognise a patent ductus arteriosus	
Can recognise sub and supravalvar aortic stenosis	
Can recognise sub valvar, valvar and supra-valvar pulmonary stenosis	
Can recognise Ebstein's anomaly	
Can recognise Fallot's tetralogy	
Can estimate pulmonary artery pressure	



Appendix 3: Reading List

The Accreditation Committee of the British Society of Echocardiography provides the reading list and represents only a handful of available textbooks for candidates to learn from.

- Textbook of Clinical Echocardiography (5th edition, June 2013), (6th edition, May 2018) -Catherine Otto
- Echocardiography Review Guide: Companion to the Textbook of Clinical Echocardiography (3rd edition) – Catherine Otto et al. (2015)
- Feigenbaum's Echocardiography (7th edition) William Armstrong and Thomas Ryan (2010)
- Echocardiography: A Practical Guide for Reporting and Interpretation (3rd edition) –
- Helen Rimington and John Chambers (Nov 2015)
- Echocardiography (Oxford Specialist Handbooks in Cardiology (2nd edition) Paul Leeson et al. (2012)
- Making Sense of Echocardiography: A Hands-on Guide (2nd edition) Andrew Houghton (2013)

Protocols and the most up-to-date BSE guidelines are available under the <u>Education</u> tab of <u>www.bsecho.org</u>.

Please note that only fully subscribed BSE members are granted full access to all education and exam content.



Appendix 4: Written Examination Registration Guidance

<u>BSE written exams</u> are delivered in partnership with Pearson VUE testing services. Candidates can sit the exam at local centres throughout the UK, the Republic of Ireland, and some overseas areas.

Pre-registration (through the BSE website)

- 1. Candidates must have an active BSE membership (fully paid and up to date).
- 2. Candidates must register their interest in taking the written exam by completing an **online pre-registration form** via the accreditation section of <u>www.bsecho.org</u>. The pre-registration window is open for up to four weeks.
- 3. Candidates' registered names should appear like their photo identification. Pearson VUE follows a strict admission policy.
- 4. BSE will transfer your data and requirements to Pearson VUE, who will contact all preregistered candidates with further information on confirming placements for the exam.

Delivery methods: Candidates can take the exam in two ways: in a **Test Centre (recommended)** or online proctored exam (OnVUE), which allows them to sit the exam from home (subject to system requirement).

Please note: Candidates who take the exam from home agree to take full responsibility for any technical issues, such as device updates, popup blocking, connection errors, and internet bandwidth. Even if the system checks before the exam are successful, faults may occur during the exam. It's important to understand the potential risks of using this method.

Special accommodations

Pearson VUE can provide <u>special accommodations</u> to candidates with official requirements, such as extra time, a reader, or medication during the examination.

All requests must be in writing and supported by documents from a healthcare professional/provider detailing the requirements and reason for the request. The BSE will approve requests at its discretion and must be submitted within the pre-registration window. To submit such requests, forward them to accreditation@bsecho.org.

Registration (through Pearson VUE)

Pearson VUE will manage all registration and payments after the pre-registration stage.

Candidates in need of special accommodations should notify the BSE during pre-registration.

Cancellations made in less than seven days do not qualify for a refund. All cancellations must be processed through Pearson Vue.

> On the day of the exam

Instructions will be given on the day of the exam via a video tutorial at the test centre. The instructions can also be accessed through Pearson VUE's online resources before the exam. Candidates will complete the exam on a computer at the test centre.



The online exam already includes a basic calculator and a whiteboard application. The examining test centre will give candidates an erasable sheet.

If the candidate chooses to take the exam from home using online proctoring (OnVUE), a calculator and whiteboard are built into the exam as an online app for the candidate to use at their convenience. Therefore, no form of stationery is permitted when taking the exam.

Candidates are required to bring a government photo ID and another form of identification. Please ensure that the registration details match your photo ID exactly; otherwise, you will be refused entry. If denied entry, candidates should contact BSE immediately.

The test centre will not facilitate any last-minute requests for special accommodations.

Results

Results are released 5-6 weeks after sitting the exam. Scores will be uploaded to BSE personal profiles. Both sections must be passed to achieve a complete pass grade.

Pass: Candidates can request login details to the portal to begin uploading logbook reports. The submission deadline will appear under 'Practical submission deadline' after the Written exam scores within the 'Participation' tab of the BSE member profile. This information is also emailed to the candidate (subject to account status).

Fail: candidates can register interest to sit in the next sitting of the exam.

- The reduced fee only applies to candidates who physically sat the exam (for the first time) and were unsuccessful; the second attempt must be taken at the next sitting (within 12 months).
- Results cannot be appealed or 'remarked' as the tests are computer-based.

Please watch the demo available via Pearson VUE: http://www.pearsonvue.com/demo/

> Additional Information

Candidates are advised to check the security procedures in the "What to expect section" of the Pearson VUE/BSE guide page: <u>https://home.pearsonvue.com/Test-takers/Resources.aspx</u>.

Pearson VUE has a strict admissions policy. Candidates' registered names should be exactly as they appear on their government photographic ID.



Appendix 5: Written Exam multiple-choice questions examples

Answer 'True' (T) or 'False' (F) to each of the following.

There is no negative marking—one mark is added for a correct answer, and no mark is deducted for an incorrect answer.

Q1	In an ultrasound imaging system:	~		
a)	Sector width, sector depth and frame rate can all be controlled independently.	F		
b)	Frame rate falls as sector width increases			
c)	Using a lower-frequency transducer improves the frame rate	F		
d)	Frame rate increases as sector depth increases	F		
e)	Using Colour Flow Doppler reduces the frame rate	Т		

Q2	On a spectral Doppler display:			
a)	The velocity at which aliasing occurs increases at higher ultrasound frequencies	F		
b)	The velocity at which aliasing occurs increases at greater depths			
c)	The velocity at which aliasing occurs increases at greater sector angles	F		
d)	At 2MHz the aliasing velocity at 10cm is approximately 1.5m/s	Т		
e)	The aliasing velocity can be increased by increasing the pulse rate (high PRF)	Т		

Q3	An Atrial Septal Defect (ASD) may be associated with:	
a)	Paradoxical interventricular septal motion	Т
b)	No obvious defect of the atrial septum on imaging	Т
c)	Right ventricular dilatation	Т
d)	Left ventricular dilatation	F
e)	Flow of blood from left atrium to right atrium	Т

Q4	Regarding assessing aortic stenosis:				
a)	Aortic valve maximum velocity of 5.2m/s is consistent with severe AS				
b)	A mean gradient of 30mmHg and a valve ratio of 0.20 is consistent with severe AS	F			
c)	In severe AS there is rapid acceleration and early peaking of the Doppler waveform				
d)	A rate of change of >0.9m/s/year is associated with poor patient prognosis	F			
e)	An aortic valve velocity ratio of 0.34 and a maximum velocity of 3.8m/s is consistent with moderate AS	Т			



Appendix 6: Written Exam image reporting questions examples

Several moving clips and stills will be included in each question. Although these can be viewed and replayed as many times as the candidate wishes, the candidate should be mindful of the time spent on each question.

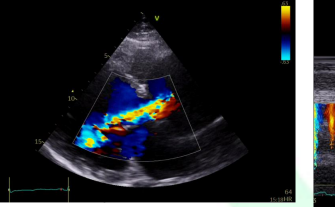
The **SINGLE BEST ANSWER** should be selected.

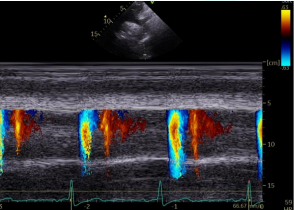
There is no negative marking—one mark is added for a correct answer, and no mark is deducted for an incorrect answer.

Case 1

Request: male, 42-year-old, admitted with chest pain radiating into back, SOBOE.

Data: LVIDd: 7.4cm, SoV dimension: 7.0cm, STJ: 6.9cm, proximal ascending aorta: 7.4cm, TAPSE: 1.4cm. proximal RVOT dimension: 4.2cm. Descending aorta end diastolic velocity: 0.30m/s. TR Vmax: 3.2m/s, right atrial area: 26cmsq, pulmonary valve acceleration time: 100ms, AR Pressure half time: 149msec.





1.1	Regarding the severity of the aortic regurgitation	Answer
а	There is moderate central aortic regurgitation	
b	There is moderate eccentric aortic regurgitation	
С	There is severe central aortic regurgitation	Т
d	There is severe eccentric aortic regurgitation	

1.2	Regarding the echo probability of pulmonary hypertension	
а	There is no echo probability of pulmonary hypertension	
b	There is low echo probability of pulmonary hypertension	
С	There is intermediate echo probability of pulmonary hypertension	
d	There is high echo probability of pulmonary hypertension	Т



Appendix 7: BSE logbook portal user guidance

1. User Login Details:

- Request login details by emailing the accreditation team-<u>accreditation@bsecho.org</u>.
- Provide your BSE ID number, the type of *accreditation you are pursuing.
- Also, inform us of your mentor's name and email address- we will assign them to your logbook.
- An automated message from the portal will be emailed to you with your login details.
- o Link to the portal: <u>https://logbook-v2.bsecho.org/login</u>

British Society of Echocardiography	,
Username or Email	
accreditation@bsecho.org	••••
Password	
•••••	•••
Remember me	
Forgot your password?	gin

a. If you have forgotten your password, please click the link titled Forgot your password?

Forgot your password? No problem. Just let us know yo email address and we will email you a password reset link that will allow you to choose a new one.
Email



2. Update your profile

• Click on your name, then **'Profile'** to update your name, email and password.

Candidate Dashboard Mentor Dashboard A	Assessor Dashboard Admin Dashboard	Jo Vashishta 🗸	
		Manage Account	
Profile	C	Profile	
		Logout	
Profile Information Update your account's profile information and email address.	Membership Number		
	BSE Staff		
	Username		
	thanjjo		
	First Name		
	Jo	۵.	
	Surname		
	Vashishta		
	Email		
	jo@bsecho.org		
		_	
		Save	

Enter new password and click 'save.'

Update Password Ensure your account is using a long, random password to stay secure.	Current Password New Password	
	Confirm Password	
Browser Sessions Manage and logout your active sessions on other browsers and devices.	If necessary, you may logout of all of your other browser sessions across all of your devices. Some of your recent sessions are listed below; however, this list may not be exhaustive. If you feel your account has been compromised, you should also update your password.	



3. User dashboard (e.g. Candidate, Mentor or Assessor)

Click on the visible heading to access your dashboard

Candidate Dashboard	Mantor Dashboard Assessor Dashboard	Admin Dashboard			Jo Vashishta 🗸
Candidate Dashboard					
ACCREDITATION	WRITTEN EXAM DATE	LOGBOOK	COMPETENCIES	MENTOR STATEMENTS	STATUS
TTE Test version for upgrade	× No date set	0 of 1 0%	0 of 3 0%	0 of 7 0%	In Progress

a. Enter Written Exam Date

Click on \times No date set to bring up the calendar and select the date you sat the written exam.

Level 1 candidates should add the date they started the accreditation.

Candidate Dashboard Mento	TTE Test version for upg Written Exam Date	Irade	Close Save	MENTOR
TTE Test version for upgrade	× No date set	0 of 1 0%	0 of 3 0%	0 of 7 0%

b. Click the box under the Logbook title to begin uploading PDF reports. The portal will take only PDF uploads.

\bigcirc	Candidate Dashboard	Mentor Dashboard	Assessor Dashboard	Admin Dashboard		
Candic	late Dashboard					
ACCRE	DITATION	WRITTEN EXA	M DATE	LOGBOOK		
TTE Tes	st version for upgrade	10/11/2021		0 of 1 0%		



To add a new case, click on 'Add a new Case', give it a Title, enter the date of the case and Choose File.

Candidate Dashboard Mentor Dashboard Assessor Dashboard Admin Dashboard	d	Jo Vashishta 🗸
TTE Test version for upgrade \rightarrow Case 1	😡 Lagbook Comments 🛛 🔍 Case Comments	ê ବ
Case 1 O You haven't added any cases yet Add a new Case	View Annotate E A A A F T C & No Presets 5 C 2	Q 7 8
Candidate Dashboard Mentor Dashboard Assessor Dashboard TTE Test version for upgrade > Case 1 Case 1 OT	Add a new Case	

- Explore the features and tools by hovering over the icons to find what they can do.
- To save your work, click 📀, to delete click 🧰

Candidate Dashboard	d Mentor Dashboard Assessor Dashboard Admin Dashboard J	lo Vashi	shta ~
TE Test version for upg	rade > Case 1 > Test 🖄 🔍 Legbook Comments	۵	ዋ
ase 1	Image:	2 🗖	ı Ø
est O			
Add a new Case	British Society of Echocardiography		
	Appendix 8: Report format		
	THIS IS A SUGGESTED FORMAT FOR A REPORT WITHIN THE WORKPLACE. PLEASE NOTE – ALL REPORTS SUBMITTED MUST BE ANONYMIZED AS PER <u>APPENDIX 15</u>		
	The report should comprise the following sections:		
	Demographic and other Identifying Information		
	Chlipptony information		



The 'Rectangle' tool allows masking over unwanted data. Click the Save button to keep the anonymised changes.

ase 1 1/1	▲ 202% → ④ ④ ▲ U □ ↓ View Annotate	Q 🛱
est	EI A A F T Image: A for the second seco	
Add a new Case	This important section should contain final commer by the TTE request. This may comprise simple repet the main part of the report (e.g. "severe LV dysfund report's technical aspects, particularly for abnormal previous echocardiographic studies or reports shou similarities) highlighted. Technical limitations of the included.	

You can add a logbook or case comments to share with your mentor.

	0	Candidate Dashboard	M E	lentor Dashboc	Logbook Cor	nments				
	TTE Test	t version for upg	rade	> Case 1	ਿ	You can add comments when discussing work with your mentor		Comments	😔 Case Comments	
>		1/1			4	VASHISHTA, JO 29/11/2021	V	- D - 5	c 2	
					Post a comn	nent	Send	10		
	Ad	id a new Case			Ammondia	British Soci of Echocard	lety diograp	hy		

4. Competencies

Your mentor will access your portal via their login and sign off on each of the competencies.

Candidates can view their progress on the dashboard.

\bigcirc	Candidate Dashboard	Mentor Dashboard	Assessor Dashboard	Admin	Dashboard					Jo Vashishta 💊	-
Candia	late Dashboard										
ACCRE	DITATION	WRITTEN EX	AM DATE		LOGBOOK		COMPETENCIES	MENTOR STATEMENTS	ST	ATUS	
TTE Te	st version for upgrade	10/11/2021			1 of 1 100%	\langle	0 of 3 0%	0 of 7 0%		n Progress	



a. Mentor view:

The mentor clicks the sections below the' DATE SIGNED OFF' header to sign off competencies by clicking on 'Sign off.'

TTE Test version for upgrade - Vashishta, Jo

COMPETENCY	SIGNED OFF BY	DATE SIGNED OFF
Received correctly		
1a. Basic Echocardiography - Knowledge		
a. Basic principles of ultrasound		Sign Off 🗸
b. Basic principles of spectral Doppler	😿 Vashishta, Jo	29/11/2021 ×
c. Basic principles of colour flow Doppler	😿 Vashishta, Jo	29/11/2021 ×

When the mentor has completed the competency sign-off, they must do the same for the 'Mentor statement.'

2. I certify I have observed the candidate scanning and I am satisfied that he/she is competent at completing a full transthoracic echo x Vi x tudy. 3. I certify that the candidate has reached a standard of training to be able to independently perform and report a transthoracic echocardiographic study. He/she has reached all of the mandated competencies. I have signed off the candidate's competency sheet.		
study. 3. I certify that the candidate has reached a standard of training to be able to independently perform and report a transthoracic echocardiographic study. He/she has reached all of the mandated competencies. I have signed off the candidate's competency sheet. 4. I certify that the candidate above has performed and reported the cases included in the accompanying Log Book within a 24-month period (or the timeframe as agreed by the BSE). 5. I certify that this is a demonstration of the logbook portal for testing purposes only and does not constitute BSE TTE accreditation and that	🛚 Vashishta, Jo	29/11/2021 ×
echocardiographic study. He/she has reached all of the mandated competencies. I have signed off the candidate's competency sheet.	v Vashishta, Jo	29/11/2021 ×
period (or the timeframe as agreed by the BSE).	🛚 Vashishta, Jo	29/11/2021 ×
	🛚 Vashishta, Jo	29/11/2021 ×
	<	Sign Off 🗸

5. Candidate logbook submission

Candidates can check the progress of their logbooks in the dashboard by clicking the arrow after 'In Progress'.

Candidate Dashboard

ACCREDITATION	WRITTEN EXAM DATE	LOGBOOK	COMPETENCIES	MENTOR STATEMENTS	STATUS
TTE Test version for upgrade	10/11/2021	1 of 1 100%	3 of 3 100%	7 of 7 100%	In Progress



a. Verify and submit

Check you have completed the requirement before clicking 'Continue.'

Candidate Dashboard Mentor De	Verify & Submit TTE Test version for upgrade		
Candidate Dashboard	 Have the correct number of cases been added to your logbook? Did your written exam fall within the correct timeframe of your earliest uploaded case? 		
ACCREDITATION	 Are your accreditation competencies complete? Are your mentor statements complete? 		
TTE Test version for upgrade	Close	7 of 7 100%	In Progress

b. Enter the Head of Department Email Address and click submit:

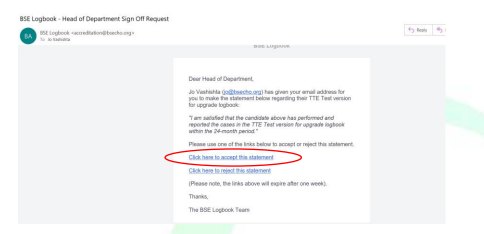
Candidate Dashboard Mentor D.	Verify & Submit TTE Test version for upgrade		Jo Vashishta 🗸
Candidate Dashboard	Your logbook has been verified and is ready for submission. Please provide your Head of Department email address to continue.		
	Head of Department Email Address		
ACCREDITATION		MENTOR STATEMENTS	STATUS
TTE Test version for upgrade	Close Submit	7 of 7 100%	In Progress >

- d. Contact <u>accreditation@bsecho.org</u> to inform you that you have entered your HOD's email address and clicked submit.
- e. We will send the email to your HOD so that they can validate your logbook. Please ask your HOD to check their junk mail if the email is not visible.

andidate Dashboard					
ACCREDITATION	WRITTEN EXAM DATE	LOGBOOK	COMPETENCIES	MENTOR STATEMENTS	STATUS
TTE Test version for upgrade	10/11/2021	1 of 1 100%	3 of 3 100%	7 of 7 100%	Validating Head of Department



6. Validate logbook: Your Head of Department must click the link to accept the statement.



a. Head of Department validated: After clicking the statement, the Head of Department receives the message below.

Some NHS emails may block messages from the logbook portal- <u>accreditation@bsecho.org</u>. In this case, candidates should consider providing an alternative email address, e.g. non-NHS email addresses.

7. Logbook submitted: Once the logbook has been validated, it is ready for an assessor to mark.

ACCREDITATION	WRITTEN EXAM DATE	LOGBOOK	COMPETENCIES	MENTOR STATEMENTS	STATUS
TTE Test version for upgrade	10/11/2021	1 of 1 100%	3 of 3 100%	7 of 7 100%	Submitted

- No further action is required from this point.
- Candidates will be notified when marking is complete.

Candidate Dashboard



Appendix 8: Logbook guidance and marking criteria

To meet all competencies of this accreditation process, the logbook should represent good/excellent examples of a candidate's daily workload. Ideally, it should reflect the most up-to-date BSE guidance (see page 7 if your department has different locally agreed-upon working practices).

Whilst we encourage the use of good/excellent work to be included in the logbook, it is acknowledged that not every report will meet this standard. Therefore, when considering whether to include a report, please refer to the following as an absolute minimum.

If a report does not meet the below, it should not be included as a logbook report

Clinical question: Must be stated.

<u>Patient Ht, Wt and BSA</u>: should be given unless it is not possible to obtain these measurements (reasons for this should be stated, e.g. patient in bed)

<u>BP:</u> Should be included particularly where clinically relevant (i.e. aortic stenosis, LVH).

Left ventricle:

Descriptive section:

- Comment on left ventricular cavity size, absence/presence (and degree) of hypertrophy.
- Comment on global left systolic function, including regional wall motion abnormalities if present.
- Comment on left ventricular diastolic function.

Measurements/analysis section:

- LV diastolic and systolic dimensions, LV wall thicknesses.
- Visually estimated ejection fraction.
- E wave velocity, A wave velocity, E wave deceleration time, e' velocities, E/e' (can be reported under mitral valve section), mitral annular S'.

Mitral valve:

Descriptive section:

- Comment on mitral valve structure, leaflet thickness and mobility.
- Comment on absence/presence of mitral stenosis.
- Comment on absence/presence of mitral regurgitation.



Measurements/analysis section: E wave velocity, A wave velocity, E wave deceleration time (can be reported under LV section).

If stenosis is present: A range of measurements taken from BSE guidelines.

If more than mild regurgitation is present: A range of measurements taken from BSE guidelines.

<u>Left atrium:</u>

Descriptive section: Visually comment on left atrial size.

Measurements/analysis section: If seen, monoplane volume (A4C or A2C).

Aortic valve:

Descriptive section:

Comment on aortic valve structure, leaflet thickness and mobility.

Comment on absence/presence of aortic stenosis.

Comment on the absence/presence of aortic regurgitation.

Measurements/analysis section:

Aortic Vmax, maximum and mean gradient, aortic VTI.

Left ventricular outflow tract Vmax, left ventricular outflow tract max and gradient, left ventricular outflow tract VTI.

If stenosis is present: Measurements as above plus: aortic valve area or dimensionless index

If more than mild regurgitation is present: A range of measurements taken from BSE guidelines.

<u>Aorta:</u>

Descriptive section:

Comment on aortic root, proximal ascending aorta, and aortic arch size or "not well seen to assess" if it is more appropriate.

Measurements/analysis section: If seen; sinuses of valsalva dimension, sino-tubular junction dimension, proximal ascending aorta dimension.



Right ventricle:

Descriptive section:

- Comment on right ventricular size
- Comment on global right ventricular systolic function.

Measurements/analysis section:

- RVD1
- TAPSE

<u>Right atrium:</u>

Descriptive section: Visually comment on right atrial size.

Measurements/analysis section: If seen, RA area.

Tricuspid valve:

Descriptive section:

- Comment on tricuspid valve structure, leaflet thickness and mobility, or "not well seen" if appropriate.
- Comment on absence/presence of tricuspid stenosis.
- Comment on absence/presence of tricuspid regurgitation.

Measurements/analysis section: Tricuspid regurgitation Vmax (if tricuspid regurgitation present) or "no measurable tricuspid regurgitation Vmax" if more appropriate.

If stenosis is present, a range of measurements will be taken from BSE guidelines.

If more than mild regurgitation is present - A range of measurements per BSE guidelines.

Pulmonary valve:

Descriptive section: Comment on pulmonary valve leaflet thickness and mobility or, "not well seen" if more appropriate.

Measurements / analysis section: If seen, pulmonary valve Vmax.

If stenosis is present - A range of measurements taken from BSE guidelines.



If more than mild regurgitation is present - A range of measurements taken from BSE guidelines.

Pulmonary hypertension:

Descriptive section:

Comment on echocardiography probability of pulmonary hypertension or "unable to comment" if more appropriate.

Measurements / analysis section:

Tricuspid regurgitation Vmax (if present), at least one further echocardiography parameter to help quantify the descriptive statement.

Inferior Vena Cava:

Descriptive section:

Comment size and collapsibility or, "not well seen" if more appropriate.

Measurements / analysis section: If seen, IVC max dimension.

Pericardium:

Descriptive section:

Comment on absence / presence of pericardial fluid. If present: A comment on location, size and hemodynamic effects.

Measurements / analysis section:

If no pericardial fluid – N/A

If pericardial effusion is present: Effusion dimensions and assessment of haemodynamic effects, including MV, TV, LVOT inflow variation with respiration, presence or absence of cardiac chamber collapse, IVC size and collapsibility assessment.

If a valve replacement is present:

Descriptive section:

 Replacement valve location, type and size (if known, or stipulate not known if more appropriate) and stability of replacement valve.



- Comment on absence / presence of prosthetic stenosis.
- Comment on absence / presence of prosthetic regurgitation. Including the likely origin of the regurgitant jet.
- Comparison to previous study where possible.

Measurements / analysis section:

- A range of hemodynamic parameters assessing forward flow that is relevant to the replacement valve position.
- A range of parameters to assess regurgitation severity as per valve native disease.

If a valve repair is present:

Descriptive section:

- Location and type of valve repair (if known, or stipulate not known if more appropriate) and stability of repaired valve.
- Comment on absence / presence of repaired valve stenosis.
- Comment on absence / presence of repaired regurgitation. Including likely origin of the regurgitant jet.

Measurements / analysis section:

- A range of hemodynamic parameters assessing forward flow that is relevant to the position of the repaired valve.
- A range of parameters to assess regurgitation severity as per valve native disease.

Conclusion:

- Must summarise main findings.
- A comparison to previous studies should made where possible.



Appendix 9: Guidance for the removal of patient identifiable data

The duty of confidentiality arises from the common law of confidentiality, professional obligations and staff employment contracts. Breach of confidence may lead to disciplinary measures, question professional reputation and possibly result in legal proceedings.

Guidance is provided to Healthcare Professionals in the 'NHS Code of Practice on Confidentiality' (November 2003):

http://www.dh.gov.uk/prod_consum_dh/groups/dh_digitalassets/@dh/@en/documents/digitalasset/ dh_4069254.pdf

Patient information that can identify individual patients is confidential and must not be used or disclosed in any part of the submission required for this accreditation process. In contrast, anonymised information is not confidential and may be used.

Key identifiable information includes:

- a. Patient's name
- b. Address
- c. Full post code
- d. Date of birth
- e. NHS number and local identifiable codes

Key identifiable information may also include information that can be used to identify a patient directly or indirectly. For example, rare diseases, drug treatment, or statistical analyses involving very small numbers within a small population may allow individuals to be identified.

Guidance to candidates submitting Logbooks and Cases for Accreditation

The NHS Code of Practice on confidentiality means that evidence submitted for this accreditation process must have removed **ALL** patient identifiable information beyond gender and age/year of birth.

Reports – Please use the BSE <u>online portal</u> and electronically delete all patient information except age and gender.

We advocate against using other electronic anonymisation methods as sometimes data is still present. If in doubt, manually remove patient identification information before use.

Video cases—We appreciate that removing patient IDs may be difficult. Therefore, it is advised that the video cases are specifically collected and the data inputs made relevant to your cases (E.g., the Patient Name could be 'BSE Case 1', and the Patient Number could be your membership number followed by the case number, '1111-1').

The final decision remains at the discretion of the Chair of the Accreditation Committee.



Appendix 10: Practical scanning mark scheme

The marking criteria used for the practical scanning assessment can be seen below.

2 minutes:	 Familiarisation of echo machine / equipment. Assessor will be on hand if assistance is required.
20 minutes:	 Candidate to have 2 minutes to obtain and acquire each image. The Assessor will instruct the candidate on the images to acquire. The Assessor can alter echo machine / equipment setting to optimise images at the direction of the candidate.

The pass mark is set at 66%. Once this mark is achieved, the candidate will be deemed successful at this station.

Each image the candidate acquires is scored as per the marking scheme below.

F = Fail = 0 points:	unable to demonstrate appropriate skill set
BF = Borderline Fail = 1 point:	unable to demonstrate appropriate skill set, is able to describe reasons how improvement could be achieved
BP = Borderline Pass = 2 points: quality	able to acquire/demonstrate skill set although fails to optimize image acquisition
P = Pass = 3 points: optimization of images	able to fully demonstrate high quality image acquisition with appropriate

All images used in the practical scanning assessment are taken from the BSE minimum dataset. An example of the imaging list used in this assessment can be seen below.

Image List One

2 minutes per acquisition Encourage candidates to move on if necessary	Image (Score Weighting)
1	2D Parasternal Long Axis (5)
2	2D Parasternal Short Axis Left Ventricle (5)
3	2D modified Short Axis demonstrating Main Pulmonary Artery (3)
4	PW Doppler RVOT (1)
5	2D Apical 4 Chamber (5)
6	PW Doppler Mitral Valve (1)
7	2D Apical 2 Chamber (5)
8	2D A4C modified to show RV, with Colour Doppler, demonstrating TR if present (3)
9	2D Subcostal 4 Chamber (3)
10	Blind CW Doppler Descending Aorta (3)
	Modification of Patient Position to Optimise Image Quality (5)
	Image Optimisation (3)



Appendix 11: Patient case studies viva marking criteria

The next few pages show the individual marking criteria for each of the patient video case studies. All criteria must be met to a satisfactory standard for the patient case study to be passed.

A minimum of two patient case studies will be assessed. The British Society of Echocardiography reserves the right to assess all five patient viva cases.

Case 1 – No significant pathology. Practice must be s Evidence of satisfactory practice	Tick	Evidence of unsatisfactory practice	Tick
ECG		ECG	
Largely present throughout without 2D image		Unstable or frequently absent, making timings	
interference		inaccurate	
Optimization Infrequent, non-repetitive optimization errors which do not detract from the case conclusion		Optimization Frequent, repetitive optimization errors which detract from the case conclusion	
Complete study Images are complete enough to allow full assessment of the selected pathology, including Doppler study and measurements		Incomplete study Images are missing which are relevant to the accurate assessment of the selected pathology, including inadequate Doppler study or relevant measurements quoted in report but not demonstrated.	
2D measurements Accurate throughout with minor errors that do not change the categorisation of the chosen pathology		2D measurements Frequent inaccuracies or isolated inaccuracies that change the categorisation of the chosen pathology	
Colour Doppler Accurate box size, gain, scale and baseline settings demonstrating anatomy clearly		Colour Doppler Frequent inaccuracies of box size, gain, scale and baseline settings which prevent clear demonstration of the anatomy	
Spectral Doppler Accurate use with good cursor alignment and optimised waveforms		Spectral Doppler Inaccurate use with poor cursor alignment or waveform optimization altering pathology assessment	
Pathology assessment No images missing which are key to pathology assessment No measurements significantly inaccurate that are key to pathology assessment		Pathology assessment Poor quality or missing images missing which are key to pathology assessment Measurements key to pathology assessment are significantly inaccurate and change the categorisation of the pathology	
Report is complete and accurate Comprehensive/accurate description of all parts of the heart Correct categorisation of chosen pathology (NB none significant abnormalities may be included in this case) Correct interpretation of findings in the clinical context		Report is incomplete or inaccurate Partial/inaccurate description of parts of the heart Incorrect categorisation of chosen pathology Incorrect interpretation of findings in the clinical context	



Evidence of satisfactory practice	Tick	Evidence of unsatisfactory practice	Tick
ECG Largely present throughout without 2D image		ECG Unstable or frequently absent making	
interference		timings inaccurate	
Optimization Infrequent, non-repetitive optimisation errors which do not detract from the case conclusion		Optimization Frequent, repetitive optimisation errors which detract from the case conclusion	
Complete study Images are complete enough to allow full assessment of the selected pathology, including Doppler study and measurements		Incomplete study Images are missing which are relevant to the accurate assessment of the selected pathology, including inadequate Doppler study or relevant measurements quoted in report but not demonstrated	
2D measurements Accurate throughout with minor errors that do not change the categorisation of the chosen pathology		2D measurements Frequent inaccuracies or isolated inaccuracies that change the categorisation of the chosen pathology	
Colour Doppler Accurate box size, gain, scale and baseline settings demonstrating anatomy clearly		Colour Doppler Frequent inaccuracies of box size, gain, scale and baseline settings which prevent clear demonstration of the anatomy	
Spectral Doppler Accurate use with good cursor alignment and optimised waveforms		Spectral Doppler Inaccurate use with poor cursor alignment or waveform optimization altering pathology assessment	
Pathology assessment Good quality CW from the A5C and stand-alone CW from at least one other window.		Pathology assessment Missing, poor quality or significantly lower stand-alone CW signal.	
No images missing which are key to pathology assessment		Images missing which are key to pathology assessment	
No measurements significantly inaccurate that are key to pathology assessment (LVOT diameter, LVOT VTi and aortic VTi)		Measurements key to pathology assessment significantly inaccurate and change the categorisation of the pathology (LVOT diameter, LVOT VTi and aortic VTi)	
Report is complete and accurate Comprehensive and accurate description of all parts of the heart		Report is incomplete or inaccurate Partial and inaccurate description of parts of the heart	
Correct categorisation of chosen pathology (NB trivial abnormalities may be included in this case)		Incorrect categorisation of chosen pathology	
Correct interpretation of findings in the clinical context		Incorrect interpretation of findings in the clinical context	



Adult Transthoracic Accreditation. Case 3 – Regurgitati	on. Pra	ctice must be satisfactory in all areas to pass	Tic
Evidence of satisfactory practice		Evidence of unsatisfactory practice	
ECG		ECG	
Largely present throughout without 2D image		Unstable or frequently absent making timings	
interference		inaccurate	
Optimization		Optimization	
Infrequent, non-repetitive optimisation errors which		Frequent, repetitive optimisation errors which	
do not detract from the case conclusion		detract from the case conclusion	
		Incomplete study	
Complete study		Images are missing which are relevant to the	
Images are complete enough to allow full		accurate assessment of the selected pathology,	
assessment of the selected pathology, including		including inadequate Doppler study or relevant	
Doppler study and measurements		measurements quoted in report but not	
		demonstrated	
2D maggiramente		2D measurements	
2D measurements		Frequent inaccuracies or isolated inaccuracies that	
Accurate throughout with minor errors that do not		change the	
change the categorisation of the chosen pathology		categorisation of the chosen pathology	
Calaura Danadara		Colour Doppler	-
Colour Doppler		Frequent inaccuracies of box size, gain, scale and	
Accurate box size, gain, scale and baseline settings		baseline settings which	
demonstrating anatomy clearly		prevent clear demonstration of the anatomy	
Spectral Depaler		Spectral Doppler	
Spectral Doppler		Inaccurate use with poor cursor	-
Accurate use with good cursor alignment and		alignment or waveform optimisation altering	
optimised waveforms		pathology assessment	
Pathology assessment		Pathology assessment	
Good assessment of regurgitation. Understanding of		Poor or inadequate assessment of severity. Failure	
the methods available to assess severity and		to return Doppler baseline to normal after PISA	
accurate demonstration if appropriate (eg PISA/Vena		assessment	
contracta/RV/ERO/PV flow)		Images missing which are key to pathology	
No images missing which are key to pathology		assessment	
assessment		Measurements key to pathology	
No measurements significantly inaccurate that are		assessment significantly inaccurate and	
key to pathology assessment		change the categorisation of the pathology	
Report is complete and accurate Comprehensive and		Report is incomplete or inaccurate Partial and	
accurate description of all parts of the heart		inaccurate description of parts of the heart	
Correct categorisation of chosen pathology		Incorrect categorisation of chosen pathology	
Correct interpretation of findings in the clinical		Incorrect interpretation of findings in the clinical	
context		context	



Evidence of satisfactory practice	Tick	Evidence of unsatisfactory practice	Tick
ECG Largely present throughout without 2D image interference		ECG Unstable or frequently absent making timings inaccurate	
Optimization Infrequent, non-repetitive optimisation errors which do not detract from the case conclusion		Optimization Frequent, repetitive optimisation errors which detract from the case conclusion	
Complete study Images are complete enough to allow full assessment of the selected pathology, including Doppler study and measurements		Incomplete study Images are missing which are relevant to the accurate assessment of the selected pathology, including inadequate Doppler study or relevant measurements quoted in report but not demonstrated	
2D measurements Accurate throughout with minor errors that do not change the categorisation of the chosen pathology		2D measurements Frequent inaccuracies or isolated inaccuracies that change the categorisation of the chosen pathology	-
Colour Doppler Accurate box size, gain, scale and baseline settings demonstrating anatomy clearly		Colour Doppler Frequent inaccuracies of box size, gain, scale and baseline settings which prevent clear demonstration of the anatomy	
Spectral Doppler Accurate use with good cursor alignment and optimised waveforms		Spectral Doppler Inaccurate use with poor cursor alignment or waveform optimisation altering pathology assessment	
Pathology assessment Appropriate measurement of Simpson's biplane showing systolic and diastolic measurements in both A4C and A2C which correlates with visual impression		Pathology assessment Incomplete assessment of Simpson's or measured inaccurately which leads to a change the categorisation of the reported LVEF	
No images missing which are key to pathology assessment		Images missing which are key to pathology assessment	
No measurements significantly inaccurate that are key to pathology assessment		Measurements key to pathology assessment significantly inaccurate that change the categorisation of the pathology	
Report is complete and accurate Comprehensive and accurate description of all parts of the heart including RWMAs Correct categorisation of chosen pathology Correct interpretation of findings in the clinical context		Report is incomplete or inaccurate Partial and inaccurate description of parts of the heart including RWMAs Incorrect categorisation of chosen pathology Incorrect interpretation of findings in the clinical context	

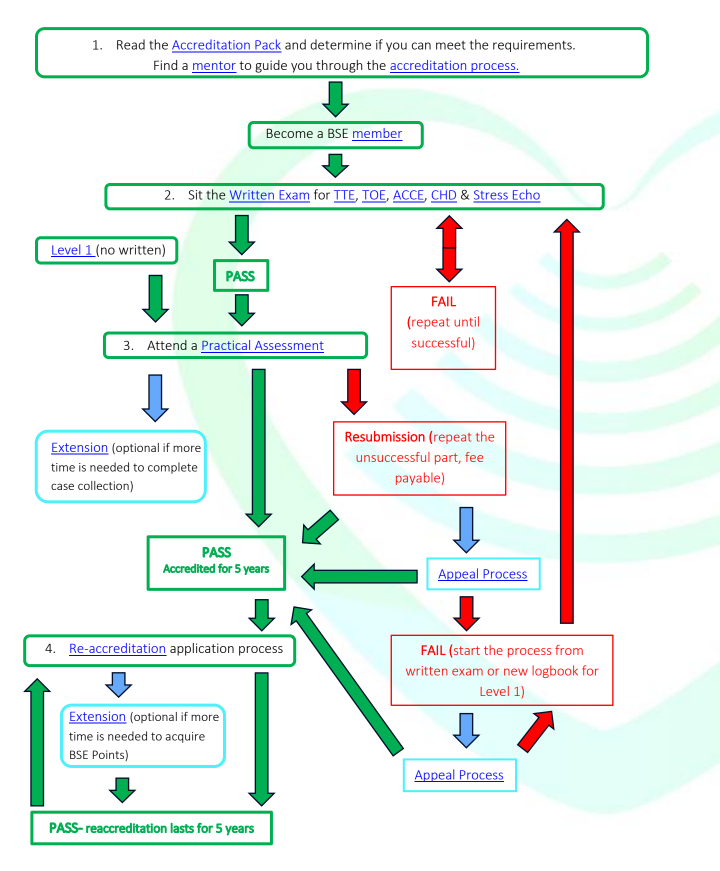


Evidence of satisfactory practice	Tick	Evidence of unsatisfactory practice	Tick
ECG Largely present throughout without 2D image interference Optimization Infrequent, non-repetitive optimisation errors which do not detract from the case conclusion		ECG Unstable or frequently absent making timings inaccurate Optimization Frequent, repetitive optimisation errors which detract from the case conclusion	
Complete study Images are complete enough to allow full assessment of the selected pathology, including Doppler study and measurements		Incomplete study Images are missing which are relevant to the accurate assessment of the selected pathology, including inadequate Doppler study or relevant measurements quoted in report but not demonstrated.	-
2D measurements Accurate throughout with minor errors that do not change the categorisation of the chosen pathology		2D measurements Frequent inaccuracies or isolated inaccuracies that change the categorisation of the chosen pathology	
Colour Doppler Accurate box size, gain, scale and baseline settings demonstrating anatomy clearly		Colour Doppler Frequent inaccuracies of box size, gain, scale and baseline settings which prevent clear demonstration of the anatomy	
Spectral Doppler Accurate use with good cursor alignment and optimised waveforms		Spectral Doppler Inaccurate use with poor cursor alignment or waveform optimisation altering pathology assessment	
Pathology assessment No images missing which are key to pathology assessment		Pathology assessment Images missing which are key to pathology assessment	
No measurements significantly inaccurate that are key to pathology assessment		Measurements key to pathology assessment significantly inaccurate and change the categorisation of the pathology	
Report is complete and accurate Comprehensive and accurate description of all parts of the heart Correct categorisation of chosen pathology Correct interpretation of findings in the clinical context		Report is incomplete or inaccurate Partial and inaccurate description of parts of the heart Incorrect categorisation of chosen pathology Incorrect interpretation of findings in the clinical context	

Check the tips for getting the video cases online via the <u>Transthoracic accreditation</u> page.



Accreditation Process Overview





Useful Links & Contacts

Some pages are restricted to BSE paid members only and require login before accessing.

- Accreditation process- <u>https://www.bsecho.org/Public/Public/Accreditation/Personal-accred/Process.aspx</u>
- Education resources (protocols & guidelines) https://www.bsecho.org/Public/Public/Education/Protocols-and-guidelines.aspx
- Extension requests- <u>https://www.bsecho.org/Public/Public/Accreditation/Personal-</u> accred/Extension-requests.aspx
- Logbook portal- <u>https://logbook.bsecho.org/</u>
- Pearson VUE Testing- <u>https://home.pearsOnVUE.com/bse</u>
- **Practical assessments-** <u>https://www.bsecho.org/Public/Public/Accreditation/Personal-</u> <u>accred/Practical-assessment.aspx</u>
- Re-accreditation- <u>https://www.bsecho.org/Public/Public/Accreditation/Personal-accred/Re-</u> accreditation.aspx
- **Regional representatives map** <u>https://www.bsecho.org/Public/About-</u> <u>Us/Governance/Council-committees/Regional-representatives.aspx</u>
- Written examination dates- <u>https://www.bsecho.org/Public/Public/Accreditation/Personal-accred/Written-examination.aspx</u>

Join the Accreditation Clinics on the first Thursday of the month at 1 pm to ask your questions about accreditation. The Clinics are hosted by the Accreditation team with the support of a committee member involved in the assessment process.

Sign up for a clinic- https://www.bsecho.org/Public/Public/Events/Events_List.aspx

Contacts

- All accreditation queries (including exam registrations) and requests to access the portal should be made to <u>accreditation@bsecho.org</u>
- Membership questions should be sent to <u>membership@bsecho.org</u>
- Events, education and e-learning questions should be sent to events@bsecho.org
- Concerns or complaints should be directed to admin@bsecho.org
- Phone number for all areas: 0208 065 5794 (Mon-Fri 9 am-5 pm)