SCoR/BMUS Guidelines for Professional Ultrasound Practice. Revision 4, December 2019

SOCIETY AND COLLEGE OF RADIOGRAPHERS AND BRITISH MEDICAL ULTRASOUND SOCIETY

Guidelines for Professional Ultrasound Practice

DECEMBER 2015
Revision 4, December 2019.

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Due recognition is given to the editors and contributors to previous editions of United Kingdom Association of Sonographers (UKAS) Guidelines whose earlier work forms the foundation of this new document. UKAS merged with the Society and College of Radiographers in January 2009.
The previous 2008 UKAS ‘Guidelines for Professional Working Standards: Ultrasound Practice’ has been archived but is available on-line at http://www.sor.org/learning/document-library/guidelines-professional-working-standards-ultrasound-practice

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Subsequent revisions, new contributors named at first contribution.

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- AXREM (Association of Healthcare Technology Providers for Imaging, Radiotherapy and Care)

Note relating to December 2019 Revision
A new style of hyperlink is introduced that is integral with the text and the referencing within the document standardised. Section to section hyperlinks have been included and the document re-ordered. Thanks are due to Gill Harrison, Associate Professor, City, University of London for this work.
Foreword to the December 2015 edition

It is my pleasure to introduce the updated ‘Guidelines for Professional Ultrasound Practice’, in the past affectionately known to sonographers as the ‘UKAS Guidelines’. The United Kingdom Association of Sonographers (UKAS) was set up to support sonographers, provide advice and practice guidance and ultimately get sonography recognised as a profession in its own right. To this day the latter still remains a challenge! However, since the last edition of the Guidelines was produced in 2008, UKAS has merged with SCOR, so, while UKAS no longer exists, its legacy lives on in this revised document. It is a testament to the quality of the original Guidelines that some sections are relatively unchanged. The advice is as equally sound and relevant today as it was then. Guidelines, however, need to keep in step with evolving technology, changes in practice and professional progression. For this reason, it was decided to produce the revised version as a web-based document that can be regularly updated, amended and expanded as and when required.

As with all previous editions, these Guidelines are not designed to be prescriptive but to inform good practice. May they continue to be used in departments across the United Kingdom for years to come.

Wendy Williams
Member, Ultrasound Advisory Group, Society and College of Radiographers.
Former UKAS committee member
December 2015

Rationale and Terminology used within the document

These current Guidelines are a collaboration between the Society and College of Radiographers (SCoR) and the British Medical Ultrasound Society (BMUS). The format of a web-based document provides easier access to relevant sections and hyperlinks, whilst allowing for annual updates. Some links within the Guidelines are to members only SCoR or BMUS content and may require additional log-on.

The document has been written to complement the 2014 joint document by the Royal College of Radiologists (RCR) and the Society and College of Radiographers entitled “Standards for the Provision of an Ultrasound Service”1. It provides guidance on topics that were not included in the joint RCR/SCoR Standards document and provides further detailed advice on some areas of practice that were.

There can be overlap between the terms ‘Standards’, ‘Guidelines’ and ‘Protocols’ and this can cause confusion. For the purposes of this document, the definitions used are the same as those in the 2014 Royal College of Radiologists (RCR) and Society and College of Radiographers (SCoR) document1.

**Standard:**
‘A required or agreed level of quality or attainment. A standard is a way of ensuring optimum levels of care or service delivery. Standards promote the likelihood of an ultrasound examination being delivered safely and effectively, are clear about what needs to be done to comply, are informed by an evidence base and are effectively measurable’2.

**Guideline:**
‘A general rule, principle or piece of advice. Guidelines provide recommendations on how ultrasound examinations should be performed and are based on best available evidence.’
They help ultrasound practitioners in their work but they do not replace their knowledge and skills.

Protocol:
An agreement, preferably based on research, between practitioners to ensure the delivery of high quality standardised ultrasound examinations.

These Guidelines, which are not prescriptive, are made available to be used as recommendations for good practice. Since the first publication of the UKAS ‘Guidelines for Professional Working Practice’ in 1993, service provision, technology and patient expectations in medical ultrasound have been transformed. The examination-specific section, including guidelines and common clinical scenarios (ref: section 4) has been compiled by the British Medical Ultrasound Society Professional Standards team and is presented as examples of best practice. They have been included so that departments can use them as a basis to generate their own departmental examination protocols when there are no nationally agreed ones available. There are also sections giving general guidance and advice, including reporting and audit. Hyperlinks have been extensively used within the text to give access to the many relevant documents already published on a wide range of topics by organisations other than the SCoR and BMUS.

IMPORTANT: These guidelines do not and cannot cover all elements of an ultrasound examination and, in addition, ultrasound practitioners are advised to access standard texts, documents and research in order to fully inform local departmental protocols and procedures.

There are no guidelines included for obstetric ultrasound within these Guidelines. Practitioners are referred to publications from other sources such as the national fetal anomaly screening programmes, the Royal College of Obstetricians and Gynaecologists (RCOG) (especially their Greentop Guidelines), National Institute for Health and Care Excellence (NICE), the Fetal Medicine Foundation (FMF), Association of Early Pregnancy Units (AEPU), British Society of Gynaecological Imaging (BSGI), and the International Society of Ultrasound in Obstetrics and Gynaecology (ISUOG).

The term patient has been used throughout the document in preference to other terms such as client or service user.

Several professional titles are used by those who practice ultrasound and this can lead to considerable confusion. The term ultrasound practitioner is used throughout this document when appropriate to do so. This is consistent with use of this term within the 2014 RCR/SCoR document ‘Standards for the provision of an ultrasound service’, which uses the definition:

Ultrasound Practitioner:
A healthcare professional who holds recognised qualifications in medical ultrasound and is able to competently perform ultrasound examinations falling within their personal scope of practice. The professional background of ultrasound practitioners can be very varied and will include radiologists, radiographers, sonographers, midwives, physiotherapists, obstetricians and clinical scientists.

A definition of ‘sonographer’ that is used in connection with the Public Voluntary Register of Sonographers (PVRS) which is administered by the SCoR can be found in Section 1. This definition
makes a distinction between those ultrasound practitioners who are registered with the General Medical Council (GMC) and those who are not. These Guidelines will be of relevance to all, hence the use of the term 'ultrasound practitioner' whenever possible.

Occasionally the term ‘operator’ is used. This term is defined within the Glossary of the 2014 RCR/SCoR Standards for the Provision of an Ultrasound Service document\(^1\) as:

‘A generic term used for someone who uses ultrasound equipment. It does not imply that they hold recognised ultrasound qualifications as would an ultrasound practitioner’\(^1\).

It is the nature of any document whether published in a traditional format or on-line that it can very quickly become out of date. It is the intention of BMUS and the SCoR that this document will be regularly updated but it is the responsibility of the ultrasound practitioner to ensure that they research and apply the most up to date evidence in association with the contents of this document. At the time of publication (Revision 4, December 2019), all hyperlinks have been checked and are complete. Please report any broken links to the following contact addresses: https://www.sor.org/contact-us or https://www.bmus.org/contact-us/

Comments and feedback are also very welcome and will guide us in the further development of these Guidelines.

The Society and College of Radiographers and the British Medical Ultrasound Society would like to thank all who have contributed to this new on-line edition of what was previously the UKAS Guidelines. Please see acknowledgements section.

We would also like to again take this opportunity thank all the contributors and editors of previous editions of the Guidelines who have provided us with such a firm foundation on which to build.

References:
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Document History:

Prior to the SCoR and BMUS collaboration, to produce the original version of this document, the United Kingdom Association of Sonographers (UKAS) published the following documents:

i) Guidelines for Professional Working Practice, published in December 1993
SECTION 1: Introduction

1.1 Explanation of the Professional Title ‘Sonographer’

Although it is the intention within this document to use wherever possible the term ‘ultrasound practitioner’, a full explanation of the term ‘sonographer’ will be helpful for context and important in terms of professional recognition and recommended qualifications.

Sonographers are qualified healthcare professionals who undertake, analyse, interpret, report and take responsibility for the conduct of diagnostic, screening and interventional ultrasound examinations. Their individual scope of practice can be wide and varied. Sonographers also perform advanced diagnostic and therapeutic ultrasound procedures such as biopsies and joint injections. Sonographers are either not medically qualified or they hold medical qualifications but are not registered as a doctor with a licence to practice with the General Medical Council (GMC).

The following definition of ‘sonographer’ is used in connection with the Public Voluntary Register of Sonographers (policy and processes PDF, at the bottom of the page). The definition was amended in 2018 to take into account proposed changes and additions to ultrasound education pathways.

‘A healthcare professional who undertakes and reports diagnostic, screening or interventional ultrasound examinations. They will hold qualifications equivalent to a Postgraduate Certificate or Diploma in Medical Ultrasound, BSc (Hons) clinical ultrasound or an honours degree apprenticeship that has been accredited by the Consortium for the Accreditation of Sonographic Education (CASE). They are either not medically qualified or hold medical qualifications but are not statutorily registered with the General Medical Council.’ PVRS: Policies and Processes.

In addition to university based postgraduate and undergraduate medical ultrasound programmes an integrated BSc (Hons) degree apprenticeship pathway, leading to qualification as a sonographer has now been developed by the Trailblazer group and approved for delivery by the Institute for Apprenticeships and Technical Education. [2019]

Individuals without a recognised qualification, including student sonographers should always be supervised by qualified staff.

At the time of this document’s publication Health Education England are working with a wide range of stakeholders (including the SCoR and BMUS) to develop a new career framework for sonography and to establish new educational routes, including pathways at undergraduate level. These plans are likely to come to fruition over the next few years.

It should be noted that statutory registration as, for example, a radiographer, nurse, midwife or doctor does not in itself mean that appropriate ultrasound qualifications are held. [2018]

Membership of professional bodies such as the SCoR or scientific organisations such as BMUS is separate to registration. There is general and widespread confusion between the terms ‘membership’ and ‘registration’. A sonographer can be a member of the SCoR and/or BMUS. They may be statutorily registered with the HCPC, NMC or other statutory regulator. They may be voluntarily registered with the Public Voluntary Register of Sonographers which is administered by the SCoR. [2018]
The title of ‘Dr’ is not a protected one and may refer to medical or academic awards obtained in the UK or overseas. Care does need to be taken so as not to unintentionally mislead a patient with respect to GMC registration being held when it is not. [2018]

For those sonographers coming to work in the UK from overseas whose first language is not English the HCPC requirements for English proficiency can be found on the HCPC website. The Public Voluntary Register of Sonographers, which is administered by the SCoR, follows the HCPC requirements with respect to English language proficiency.

The CASE website has a list of accredited medical ultrasound programmes. CASE accredit programmes of education, but do not accredit individual sonographers. [2018]

The British Society of Echocardiography (BSE) and Society for Vascular Technology of Great Britain and Ireland (SVT) accredit individual ultrasound practitioners working within their respective specialties. [2018]

The Society and College of Radiographers (SCoR) can provide accreditation of advanced and consultant practitioners and practice educators for sonographer members (SCoR member log-in required).

A sonographer should:

i) recognise and work within their personal scope of practice, seeking advice as necessary;
ii) ensure that a locally agreed and written scheme of work is in place;
iii) work with reference to national and local practice and guideline recommendations;
iv) ensure they hold appropriate professional indemnity insurance or obtain this by virtue of their employment (ref: section 2.2).

See also section 2.1 on codes of professional conduct for sonographers.

The general standards of education and training for ultrasound practitioners are set out on page 12 of the 2014 Royal College of Radiologists/Society and College of Radiographers document ‘Standards for the Provision of an Ultrasound Service’.

### 1.2 Registration for Sonographers

This section uses the professional term ‘sonographer’ instead of the generic ‘ultrasound practitioner’ and refers to the long running campaign to have ‘sonography’ recognised as a profession and for the professional title of ‘sonographer’ to be legally protected.

The registration situation for sonographers is complex (Thomson and Patterson, 2014)³.

The majority of sonographers are statutorily registered, but this will depend on their professional background and is not achievable for all. Statutory registration will most likely be as a radiographer or clinical scientist with the Health and Care Professions Council (HCPC) or as a midwife or nurse with the Nursing and Midwifery Council (NMC) and not as a sonographer, which is not a protected title. Whether statutorily registered or not, sonographers are encouraged to apply to register with the Public Voluntary Register of Sonographers (PVRS) which is administered by the Society and College of Radiographers.
For some sonographers, this will be the only register available to them. For those sonographers who are already statutorily registered, applying to register with the PVRS will help to protect the public and support the case for statutory regulation. This was recommended by the then Health Professions Council (HPC) to the Secretary of State for Health in 2009 but has not progressed. **Government policy since 2011** has been not to bring further aspirant groups into statutory registration unless there is a clear evidence of clinical risk that requires this.4

The HCPC provide guidance on their site about “Regulating further professions” and NHS employers have advice on sonographer registration, on the site “Medical radiography and ultrasound workforce”.

The Society and College of Radiographers have produced a document giving advice on “Ultrasound training, employment, registration and professional indemnity insurance (2019)” and the British Medical Ultrasound Society has information on “How to become a sonographer” which includes working in the UK and qualifications.

For some sonographers working in areas of practice coming within the remit of the Academy for Healthcare Science (AHCS), statutory registration may be available either by following approved education and training routes as a clinical scientist or by being able to demonstrate ‘equivalence’. Statutory registration, if it is obtainable, will be with the HCPC as a clinical scientist. The AHCS also administers a voluntary register which is accredited by the Professional Standards Authority (PSA).

The Registration Council for Clinical Physiologists runs a PSA accredited voluntary register that is relevant for professionals specialising in echocardiography.

References:


1.2.1 Professional Standards Authority report on sonographer regulation [2019]

As part of a major project including the SCoR, BMUS and a wide range of stakeholders to develop a new career structure for sonography, Health Education England (HEE) commissioned the Professional Standards Authority (PSA) to analyse the evidence available under the criteria outlined in the document “Right-touch assurance: a methodology for assessing and assuring occupational risk of harm” (2016) and provide advice on the most appropriate form of assurance for the role. Evidence was presented to the PSA by the SCoR, BMUS, RCR and CASE along with other interested parties in December 2018 “Summary of the rationale for the statutory registration of sonographers”.

The PSA report “Right-touch assurance assessment for sonographers” was made available to the public in July 2019. The SCoR, BMUS and other stakeholders were disappointed and concerned by the conclusions which overturned the recommendations of the PSA’s own internal public advisory panel that had been constituted for the review and had recommended statutory registration.
Following its publication the SCoR and BMUS in conjunction with other stakeholders has been in contact with senior NHS and government officials to try and achieve statutory registration for all sonographers and recognition of title. This is to both protect the public and to allow the proposed HEE career structure for sonographers to be fully realised.

1.3 Profession vs Tool

There are many healthcare professionals working within the UK who use ultrasound as a ‘tool’ to assist with their overall treatment or evaluation of patients. There is published advice on education and training available to those who use ultrasound in this way but whose main work and role is not that of an ultrasound practitioner. For those who use the professional title of ‘sonographer’, ultrasound is their daily work and their primary profession. When used as a ‘tool’, ultrasound aids and assists a healthcare practitioner with their wider examination and treatment, but in overall terms, ultrasound is only a small part of their work. It is important for safe and effective service delivery that all ultrasound examinations are undertaken by appropriately trained and competent personnel and that there is associated audit and continuing professional development (CPD) in the use of ultrasound.

The following links provide further information about ultrasound education and standards

i) CASE accredited focused courses can be found in the “Directory of CASE accredited courses” along with details of the “Standards for sonographic education”.

ii) Royal College of Radiologists (RCR) have published “Ultrasound training recommendations for medical and surgical specialities” and “Focused ultrasound training standards”

1.4 Screening examinations using ultrasound

The United Kingdom National Screening Committee advises ministers in all four countries and resides within Public Health England, an executive agency of the Department of Health and Social Care.

Before any pathology or condition is accepted for national screening there is a full evaluation against the NSC published criteria.

The NSC website has details of membership and links to additional sites for further information. Details of the evidence review process can also be found on the NSC guidance website. It should be noted that there may be variations in the screening programmes that operate across the four countries of the UK and ultrasound practitioners should contact the relevant organisations for current advice.

- Scotland “National Screening Programmes”
- Northern Ireland “Screening”
- Wales “Screening”

In England, all public screening information is available on the NHS Choices website. Information for professionals is hosted on the gov.uk website, and learning resources are on the e-learning for healthcare website, developed by Health Education England.
The four national screening programmes that are of particular relevance to ultrasound practitioners are:

i) **Antenatal screening**

In England the Fetal Anomaly Screening Programme (FASP) is responsible for the two ultrasound scans that are offered to every pregnant woman in England. There are equivalent organisations to FASP in the devolved countries although the 11\(^{th}\) week to 14\(^{th}\) week scan is not offered as a screening scan in Northern Ireland.

The two ultrasound scans for which FASP (England) is responsible are the 11\(^{th}\) to 14\(^{th}\) week scan that includes the combined test for Trisomy 21, 13 and 18 and the 18 to 20\(^{th}\) week fetal anomaly scan.

FASP has published comprehensive information for professionals. Two key documents are the [NHS Fetal Anomaly Screening Programme Handbook](#) (2018) and [Fetal Anomaly Screening Programme: Handbook for Ultrasound Practitioners](#) (2015).

Equivalent organisations to FASP in Scotland and Wales:

- **Scotland**: National Services Division: [Pregnancy and Newborn Screening](#)
- **Wales**: [Antenatal Screening Wales](#)

Independent providers offering screening ultrasound scans to NHS patients during pregnancy must work within the published screening programme standards for the country in question.

ii) **NHS Abdominal Aortic Aneurysm (AAA) Screening Programme**

The **Abdominal Aortic Aneurysm screening programme** has now successfully completed its roll out across the UK.

Information on AAA screening in the devolved countries is available for:

- **Scotland**
- **Wales**
- **Northern Ireland**

iii) **NHS Breast Screening Programme**

The [NHS breast screening (BSP) programme](#) website has resources relating to the programme and education. Although ultrasound is not part of the initial screening examination, specialists in breast ultrasound will use ultrasound techniques for further evaluation and biopsy.

iv) **NHS Neonatal and Infant Physical Examination (NIPE) screening programme**

This national screening programme is responsible for issuing guidance and standards regarding the screening pathways for physical examination of the newborn in England. There is currently no equivalent screening programme in the devolved countries.
Guidance on when ultrasound examinations of the neonatal hip should be performed can be found on the website [Newborn and infant physical examination: programme handbook](#) (section 6.5). Standards for the NIPE programme can be found in the [Newborn and infant physical examination screening: standards](#) documentation.

It is important to note that the neonatal hip ultrasound examination itself is a post-screening examination and is outside the direct remit of the NIPE Screening Programme.

**Public Heath England advice on private screening for different conditions and diseases.** Information outlining the advantages and disadvantages of screening outside the national programmes, along with information for health care professionals and patient information leaflets can be found on the website [‘Private screening for health conditions: NHS recommendations’](#).

**Duty of candour guidance in the screening programmes**
The NHS published a document [‘NHS screening programmes: duty of candour’](#) in October 2016 (See also section 1.17)

**Public Health England e-learning platform**
The Public Health England screening programmes provide links to educational resources and updated information in their [blog](#).
SECTION 2: Governance and Safety

2.1 Code of Practice for Sonographers

This code of practice has been included to support the use of the professional title of ‘sonographer’ and the future development and regulation of the sonography profession. Hence the use of this term rather than the generic ‘ultrasound practitioner’ (ref: section 1)

A Code of Practice can be defined as a set of written rules which explains how people working in a particular profession should behave. It is designed to cover all circumstances, is written in broad terms and expresses ethical principles.

The statements below are from the Code of Practice for Sonographers which were previously published by the United Kingdom Association of Sonographers (UKAS) in the Guidelines for Professional Working Standards – Ultrasound Practice (2008) and are as equally relevant today. These statements that reflect best practice are a guide and offer advice to sonographers, educationalists, students of medical ultrasound and other health care practitioners. They are statements of professional conduct that reflect the individual's rights, local and national changing patterns of ultrasound service delivery and the requirement of sonographers to demonstrate continuing competency through personal and professional development.

There are other codes that sonographers and ultrasound practitioners may need to follow. If statutorily registered with the GMC, HCPC or NMC, these regulators have themselves published codes of conduct and ethics. Professional bodies such as the Society and College of Radiographers and the Royal College of Midwives, for example, also have published codes of conduct. The Public Voluntary Register of Sonographers has associated with it published “Standards of Conduct, Performance and Ethics and Standards of Proficiency” (scroll down for the relevant PDF documents)

A code of practice for sonographers

1. Sonographers have a duty of care to their patients with respect to the minimisation of ultrasound exposure consistent with diagnostic needs.

2. Sonographers are ethically and legally obliged to hold in confidence any information acquired as a result of their professional and clinical duties, except where there is a legal obligation for disclosure.

3. Sonographers must be committed to the provision of a quality ultrasound service having due regard for the legislation and established codes of practice related to health care provision in order to minimise risk to patients, patients’ carers and other professionals.

4. Sonographers are legally and professionally accountable for their own practice and must not be influenced by any form of discrimination.

5. Sonographers must identify limitations in their practice and request training and support to meet their perceived needs.
6. Sonographers will take all reasonable opportunity to maintain and improve their knowledge and professional competency and that of their peers and students.

7. Sonographers must pay due regard to the way in which they are remunerated for their work.

8. Sonographers have a duty of care to work collaboratively and in co-operation with the multi-disciplinary health care team in the interests of their patients and patients’ carers.

9. Sonographers must act at all times in such a manner as to justify public trust and confidence, to uphold and enhance the reputation of sonography and to serve the public interest.

10. Sonographers must ensure that unethical conduct and any circumstances where patients and others are at risk are reported to the appropriate authority.

11. Sonographers who are held accountable in another area of health care must relate this Code to others that govern their practice.

12. Student sonographers pursuing a qualification in medical ultrasound must adhere to their University or Higher Education Institution’s Codes of Conduct that relate to all elements of their ultrasound education and training.

2.2 Professional Indemnity

The UK government introduced legislation in 2014 which requires ultrasound practitioners who are statutorily registered with the Health and Care Professions Council (HCPC) (e.g. as a radiographer, physiotherapist or clinical scientist), Nursing and Midwifery Council (NMC) (e.g. as a nurse or midwife), or other statutory regulator, to have a professional indemnity arrangement as a condition of their statutory registration. The majority of statutorily registered ultrasound practitioners will already meet this requirement and will not need to take any further action. They will either work in an employed environment where their employer will indemnify them, and / or if they undertake self-employed work, they will have already made their own professional indemnity arrangements. However, some statutorily registered ultrasound practitioners may need to take steps to make sure that they have appropriate professional indemnity arrangements in place.

Registrants and applicants for statutory registration will be asked to confirm that they do, or will, meet this requirement by completing a professional declaration when renewing or registering for the first time. The HCPC have published guidance “Professional indemnity and your registration” on the requirements along with an accompanying flow diagram.

The NMC have published guidance “Professional indemnity arrangements” as have the GMC “Insurance indemnity and medico-legal support”.

In addition to working in an employed environment, professional indemnity insurance can be obtained through membership of trade unions and professional bodies or by purchasing from medical defence unions or commercial insurers. Ultrasound practitioners should carefully review and follow the terms of any indemnity insurance they have.
Ultrasound practitioners who are self-employed or who work in a part employed/ part self-employed environment are particularly advised to read the guidance published by their statutory regulator.

There is no professional indemnity insurance associated with voluntary registration on the Public Voluntary Register of Sonographers. If an ultrasound practitioner is not statutorily registered, it is clearly good practice to ensure that they have appropriate professional indemnity arrangements in place both to protect the public and themselves.

2.3 Safety of Medical Ultrasound

The BMUS “Statement on the safe use, and potential hazards of diagnostic ultrasound” states that ‘Ultrasound is now accepted as being of considerable diagnostic value. There is no evidence that diagnostic ultrasound has produced any harm to patients in the time it has been in regular use in medical practice. However, the acoustic output of modern equipment is generally much greater than that of the early equipment and, in view of the continuing progress in equipment design and applications, outputs may be expected to continue to be subject to change. Also, investigations into the possibility of subtle or transient effects are still at an early stage. Consequently, diagnostic ultrasound can only be considered safe if used prudently’.

Ultrasound exposure depends on many factors, including the examination type, patient body habitus and equipment settings. Some modes, such as B-mode have a lower potential for tissue damage than Doppler, with pulsed Doppler techniques having the potential for the highest exposure levels. Recommendations related to ultrasound safety assume that the equipment being used is designed to international or national safety requirements and that it is operated by competent and trained personnel.

It is the responsibility of the operator or ultrasound practitioner to be aware of, and apply, the current safety standards and regulations and to undertake a risk/benefit assessment for each examination.
Key principles for the safe use of ultrasound are published in the document “Guidelines for the safe use of diagnostic ultrasound equipment”:

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<tbody>
<tr>
<td>i)</td>
<td>Medical ultrasound imaging should only be used for medical diagnosis and /or as an aid to medical/surgical interventions.</td>
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<tr>
<td>ii)</td>
<td>Ultrasound equipment should only be used by people who are fully trained in its safe and proper operation. This requires:</td>
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<td></td>
<td>• an appreciation of the potential thermal and mechanical bio-effects of ultrasound;</td>
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<td></td>
<td>• a full awareness of equipment settings;</td>
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<td>• an understanding of the effects of machine settings on power levels.</td>
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It is however recognised that ultrasound is a useful tool used for improving patient safety during procedures such as line and needle placement. Whilst users may not have a full understanding of the physical properties of ultrasound imaging, they must be aware of the need to limit examination times and only use equipment for the proposed medical purpose.

|iii) | Examination times should be kept as short as is necessary to produce a useful diagnostic result. |
|iv) | Output levels should be kept as low as is reasonably achievable while producing a useful diagnostic result. |
|v) | The operator should aim to stay within the BMUS recommended scan times (especially for obstetric examinations). |
|vi) | Scans in pregnancy should not be carried out for the sole purpose of producing souvenir videos or photographs. |

The British Medical Ultrasound Society has UK leading advice on ultrasound safety that all ultrasound practitioners should be familiar with ‘Physics and safety’.

BMUS have also published guidance for the use of volunteers and patients for demonstrations “Guidelines for the management of safety when using volunteers & patients for practical training and live demonstration in ultrasound scanning and consent” This includes scanning during pregnancy for demonstration and teaching purposes. [2019]

Reference:
2.4 Medico-legal Issues

The place of work should have a written set of protocols that accurately describes the range of ultrasound examinations undertaken. Their content should address the ultrasound examinations, their reporting and the appropriate referral pathways for patients with normal and abnormal ultrasound findings. The details in the protocols should be such that a new staff member, having read them, could carry out and report these examinations and appropriately refer the patient, after the examination, to the expected standard. Protocols should be updated regularly and their review date should be included in their content. Superseded protocols should be kept on file permanently.

Records are currently required by law to be kept for a number of years as specified by Department of Health and Social Care advice (ref: section 2.13).

The following guidance should be considered:

- ultrasound practitioners should be aware that they are legally accountable for their professional actions, including the reporting of ultrasound examinations, in all circumstances;
- the report is a public document and part of the patient’s medical record, together with any images, and/or video recordings which may accompany it;
- when a patient consents to an ultrasound examination, they have the right to expect it to be performed and reported by a competent ultrasound practitioner;
- a competent ultrasound practitioner is one who works to the standards defined by the guidelines of their place of work, the code of conduct of their professional body, the guidelines of that and other relevant bodies and of the regulatory body where appropriate;
- the standard of care provided by a competent ultrasound practitioner is that which the majority of similar individuals would provide and/or which a significant body of similar individuals would provide in similar and contemporaneous circumstances;
- images that accompany an ultrasound examination carried out by a competent ultrasound practitioner evidence the assumption that the necessary standard of care has been delivered (ref: section 4.15);
- all images must be capable of being attributed to the correct examination and should include the patient identifier(s), examination date and time;
- nationally published requirements for the storage of images must be followed. Examples would be the image storage requirements of the abdominal aortic aneurysm and fetal anomaly screening programmes and those published by the Department of Health and Social Care (ref: section 4.2).

See also Duty of Candour (ref: section 2.12.4)
Reference is also made to medico-legal considerations relating to consent in section 2.9.
2.5 Transducer and Equipment Cleaning and Disinfection

In addition to the following guidelines ultrasound practitioners should refer to published local infection control protocols and procedures. Many organisations also have infection control leads who should be consulted as necessary. Compared to just a few years ago there is now a wide range of information available; departments and sonographers will need to make their own assessments as to what are the most appropriate cleaning and disinfection methods for transducers and equipment given their own circumstances.

There have been two previous Medicines and Healthcare Products Regulatory Agency (MHRA) alerts relating to transducer cleaning and disinfection:

- Reusable transoesophageal echocardiography, transvaginal and transrectal ultrasound probes (transducers) – failure to appropriately decontaminate (2014)
- Ultrasound transducer probes with an internal lumen used for taking transrectal prostate biopsies (2009)

The following documents and statements are available:


Definitions of terms relating to sterilisation and disinfection including ‘critical’ and ‘semi-critical’ are given in the following NHS Scotland advice: “Guidance for the decontamination of semi-critical ultrasound probes, semi-invasive and non-invasive ultrasound probes” (2017). There is also important guidance within the document on the use of covering sheaths.


The British Medical Ultrasound Society also has the following advice (2017):

‘All ultrasound transducer probes should be cleaned immediately after a scan to remove all organic residues and body fluids. This involves removal of the used probe cover (if used), wiping off the gel followed by thorough cleaning with probe compatible cleaning agents as per probe manufacturer’s instructions.

Ultrasound probes should then undergo appropriate disinfection or sterilisation. All critical probes (probes contacting sterile tissues or blood) should be preferably sterilised, but if sterilisation is not possible, they should be minimally high level disinfected and used with a sterile sheath. All semi-critical probes (both semi-invasive probes contacting mucous membranes and non-invasive probes contacting non-intact/broken skin) should be high level disinfected either manually or with automated systems. High level disinfection is still required when using a sheath as sheaths can have micro-perforations or can break. All non-critical probes contacting only intact skin may be low level disinfected.

Only probe manufacturer recommended, and probe compatible disinfection products should be used to avoid any damage to the probe. After reprocessing the probes should be stored so as to prevent recontamination’


2.6 Ergonomic practice including managing the high BMI patient

Prevention and management of work related musculoskeletal disorders

Work related musculoskeletal disorders (WRMSD) are known to be associated with ultrasound practice. There are several causative factors including high workloads, increasing patient body mass index, poor equipment, room design and organisation of list, stress, poor posture and ergonomics when scanning. It is important that ultrasound practitioners take care of themselves and their working environment whilst scanning.

Employers have a legal duty of care to their employees and should be guided in ways to avoid potential work related injuries i.e. by supplying equipment fit for purpose and being realistic about time management. Departmental guidelines should include strategies to minimise the risk of WRMSD, including appropriate management of workload (ref: section 2.8).

A range of advice and guidance documents have been published, to which ultrasound practitioners are referred:

Health and Safety Executive

- Risk management of musculoskeletal disorders in sonography work (2012)

Society and College of Radiographers

- The causes of musculoskeletal injury amongst sonographers in the UK (2002)
Factors to consider when scanning patients with a high BMI:

The following points are all particularly relevant when scanning high BMI/bariatric patients and are in addition to general good practice methods of reducing the incidence of WRMSDs. All trusts and health boards should have policies relating to care and manual handling associated with high BMI/bariatric patients, and these should also be available and consulted.

**Equipment**

- Use a ‘high BMI’ preset on the machine as a starting point for manipulating the image. Manufacturers can set these up to your requirements at the time of installation and will optimise features such as transducer frequency and harmonics.
- Use good-quality equipment with good harmonics. Use the equipment settings to optimise the image wherever possible.
- Do not exceed the couch weight limit, which should be clearly posted.
- Use available moving and handling aids when necessary; scan in-patients in their beds rather than transferring them to an examination couch.

**Working practices**

- Wherever possible, the sonographer workforce should be rotated to ensure that it is not the same sonographer group exposed to risk. This will, of course, depend on the skill mix of the local sonographer workforce.
- Try standing to scan, placing one leg in front of the other in a static lunge if necessary, to ensure that your arms and shoulders are not under strain and your back remains straight (image 1). If standing, ensure the weight is evenly balanced between both feet.
- Remember to use micro breaks to relax muscles and tendons during the scan. When measurements are taken, remove the probe from the patient and rest the scanning hand for a few seconds.
- Do not extend the examination time beyond what is normally allowed if there is unlikely to be any gain.
  - It may be that a second appointment is necessary in some cases.
  - FASP provides guidance with respect to repeat examinations on those women attending for the 18–20 weeks’ fetal anomaly scan and where the image quality is compromised by an increased BMI. “The woman should be informed that the screening is incomplete and this should be recorded.” Sensitivity and honesty are needed when discussing this with the woman.
There is also ‘twice on the couch’ advice for the 11\(^{2}\) to 14\(^{1}\) week scan, which forms part of the combined screening test “\textbf{timing is everything in FASP screening}” (2018). Women should be referred for second trimester screening if the nuchal translucency cannot be accurately measured at the second attempt.

- Avoid pressing unnecessarily hard and for too long. This may increase the risk of WRMSDs and it can be uncomfortable for the patient. Increased transducer grip or pressure can increase the chance of a WRMSD. A power grip is recommended to reduce the risk of injury.
  - Firm pressure may be contra-indicated for some types of pathology or clinical situations.

- Try alternative techniques such as:
  - Use a helper to support tissue/fatty aprons (panniculus) and generally assist with the examination.
  - Lift the panniculus or scan above or to the side. Often scanning above and angling the probe inferiorly can be useful, rather than trying to scan underneath.
  - Decubitus scanning can assist by moving the panniculus to the side away from the region of interest.
  - The \textbf{Sims position} can also help. This involves having the patient almost prone on the couch and scanning around the side, thus reducing the tissue mass to be scanned (\textbf{Harrison and Harris, 2015}).

\textbf{Image 1: A static lunge to reduce arm abduction}
2.7 Intimate examinations and chaperones

The definition of an intimate examination may differ between individual patients for ethnic, religious or cultural reasons. In addition, some patients may have a clear preference for a health carer of specific gender due to their ethnic, religious or cultural background, because of previous experiences or in view of their age. Where possible, such individual needs and preferences should be taken into consideration.

When conducting an intimate examination, the ultrasound practitioner should:

- act with propriety and in a courteous and professional manner;
- communicate sensitively and politely using professional terminology;
- follow published guidance for use of a chaperone (links below);
- respect the patient’s rights to dignity and privacy;
- comply with departmental schemes of work and protocols.

Patients should not be asked to remove clothing unnecessarily. When required, private, warm, comfortable and secure facilities for dressing and undressing should be provided. Care should be taken to ensure privacy in waiting areas used by patients not fully dressed in their own clothes. During the ultrasound examination, only those body parts under examination should be exposed.

Care must be taken to maintain confidentiality when non-health care personnel are nearby.

Patients should be given the opportunity to have a chaperone, irrespective of the ultrasound practitioner’s gender and the examination being undertaken. The ultrasound practitioner should give equal consideration to their own need for a chaperone, again, irrespective of the examination being undertaken or the gender of the patient.

A record should be made in patients’ records when chaperones are offered and used, and when they are declined. The record should include the name and designation of the chaperone. Chaperones should normally be members of the clinical team who are sufficiently familiar with the ultrasound examination being carried out to be able to reliably judge whether the ultrasound practitioner’s actions are professionally appropriate and justifiable. Chaperones should be trained in the role.

Patients’ privacy and dignity should be maintained throughout the examination which should be conducted without interruption. Only personnel essential for carrying out the examination should be in the room.

It is good practice (as for any examination) to ensure that, when possible, hand washing and equipment cleaning are carried out in full view of the patient at the beginning and end of the examination to reassure them that effective infection control procedures are being applied.

Advice on students/trainees and intimate examinations is given in the 2016 SCoR document and in the 2015 RCR guidance (see below).

There are several organisations that have produced advice on the conduct of intimate examinations and also on the use and role of chaperones.

- General Medical Council (2013) [Intimate examinations and chaperones](#)
- Medical Defence Union (2018) [Protecting yourself from a sexual assault allegation](#)
- Royal College of Radiologists (2015) [Intimate examinations and the use of chaperones](#)
- Society and College of Radiographers (2016) [Intimate Examinations and Chaperone Policy](#)
2.8 Examination times

The time allowed for an ultrasound examination should take into account the fact that the actual transducer time is only a component of the overall examination.

Time needs to be allowed for room preparation, assessing the ultrasound request, introductions and explanations, obtaining valid consent and assisting the patient when necessary on to and off the examination couch. Post-procedure time is required to discuss the findings with the patient, write the report, archive the images and attend to the after-care of the patient, including making arrangements for further appointments and/or further investigations. Equipment will also need cleaning and disinfecting as required post examination.

An ultrasound practitioner has a professional responsibility to ensure that the time allocated for an examination is sufficient to enable it to be carried out competently. It is critical to patient management that no ultrasound examination is compromised by departmental and/or government targets.

The allocated appointment time will vary depending on the type and complexity of the ultrasound examination. It may also be influenced by the expertise of the ultrasound practitioner and training commitments within the department. In addition, the duration of the examination will be influenced by the scan findings and/or the physical condition of the patient. The quality of equipment and general support available to the sonographer are also relevant.

Examination times should be determined with reference to national standards such as those published by the Fetal Anomaly Screening Programme (FASP), by organisations such as NICE and by proper evaluation of the local working arrangements and resources that will be different for each service provider. Examination times will need to take into account whether there are trainees present and their stage of training, if teaching is to be effective.

Although obstetrics is not included in the scope of this document the following recommended times are included in this section for completeness.

Times for the two obstetric ultrasound screening scans are now within NHS England FASP Service Specifications Nos 16 and 17.

- **Service specification 16**, page 10 (the ultrasound component of the combined screening test) states a minimum of 20 minutes. Follow link for full wording.
- **Service specification 17**, page 10 (18+0 to 20+6 weeks ultrasound scan) states a minimum of 30 minutes for a singleton pregnancy and 45 minutes for a multiple pregnancy. Follow link for full wording.

**NICE Guidelines on Multiple Pregnancy (CG 129)** published in September 2011 state that:

- 45 minutes should be allowed for anomaly scans in twin and triplet pregnancies (section 1.3.3.3, page 19), concurring with FASP guidelines
- 30 minutes should be allowed for growth scans on twins and triplets (section 1.3.3.4, page 19)

**The Abdominal Aortic Aneurysm Screening Programme** (2017) has guidance on clinic booking times and overall session numbers in its Standard Operating Procedures. Appointments are at 5- to 10-minute intervals, with short breaks within each session, to include 15 to 18 scans in a 3-hour session.
The Society and College of Radiographers has published guidance “Ultrasound examination times and appointments”. In the absence of any local evidence-based determination of examination times or national standards, the SCoR advises that 20 minutes should be the minimum for a general abdominal ultrasound examination. The complete document should be read for full context.

Individual departments can determine examination times taking into account local circumstances. A tool to help evaluate these is the former “NHS Improvement - Examination Times Assessment Tool”. Full details of how to access are in Appendix 3 of the SCoR examination times document.

Many request forms are very non-specific in terms of the patient’s symptoms and due allowance may need to be made for this in schedule planning if it is decided to proceed (Ref: section 4.4 and section 4.5). For example, it may be necessary to perform both transabdominal and transvaginal scans to fully evaluate the female abdomen and pelvis with ultrasound.

2.9 The 6 C’s, patient identification, communication and consent

Compassion in Practice (the 6 C’s).

The 6 C’s are a set of values that underpin ‘Compassion in Practice’, a vision and strategy for nursing, midwifery and all health and care staff. This is part of the “Leading change, adding value” framework.

They are Care, Competence, Compassion, Communication, Courage and Commitment.

While undertaking any ultrasound examination and working in accordance with locally agreed practice, ultrasound practitioners should:

• Greet and correctly identify the patient using their name, address, date of birth;
• introduce self and others using ‘hello my name is…….’ and explaining the title and role of those present;
• obtain sufficient verbal and/or written information from the referring clinician to undertake correctly the examination requested (Ref: section 4.4 and section 4.5);
• ensure the proper care and maintenance of equipment and not use damaged equipment or equipment that is not fit for purpose;
• be mindful of the need to use appropriately trained professional interpreters as and when necessary to communicate adequately with the patient;
• be able to discuss the relative risks and benefits of the examination with the patient;
• explain the scanning procedure appropriately to the patient;
• obtain valid, informed consent from the patient or their representative being mindful of their capacity to understand;
• be aware of the individual patient’s special needs including chaperoning and privacy during the examination (ref: section 2.7);
• be professional and understanding throughout the examination; manage the interaction between the patient and others in the room, in a way that enables the examination to be carried out to a competent standard;
• explain and discuss the findings with the patient within local guidelines;
• interpret and communicate the findings appropriately and in a timely fashion to the referring clinician;
• ensure appropriate arrangements have been made for further care before the conclusion of the examination as necessary.

Valid, informed consent must be obtained before commencing any ultrasound examination or procedure. Ultrasound practitioners who do not respect the right of a patient to determine what happens to their own body in this way may be liable to legal or disciplinary action.

The consent process is a continuum beginning with the referring health care professional who requests the ultrasound examination and ending with the ultrasound practitioner who carries it out. It is the responsibility of the referring professional to provide sufficient information to the patient to enable the latter to consent to the ultrasound examination being requested. It is the responsibility of the ultrasound practitioner to ensure that the patient understands the scope of the ultrasound examination prior to giving their consent. See the 2019 case Mordel v Royal Berkshire NHS Foundation Trust [2019] EWHC 2591 (QB). The Montgomery ruling (2015) highlights the need to ensure that patients are fully involved in decision making processes, including providing evidence based information that a ‘reasonable person’ in their position might ‘attach significance to the risk’.

Verbal valid, informed consent must be obtained for all examinations and should be recorded in the ultrasound report. Additional valid, informed verbal consent should be obtained where a student ultrasound practitioner undertakes part or all of the ultrasound examination under supervision. Some categories of ultrasound examination (interventional ultrasound, guided procedures e.g. biopsy) will require written consent.

Literature which explains the scope of the examination clearly and accurately should be made available to patients prior to the ultrasound examination. NHS Choices carries information on a wide range of topics, for example ‘Ultrasound scan’.

The national screening programmes have explanatory literature available for patients obtainable via the NHS Choices website and have published consent standards which are usually now included with the various NHS England service specifications. An examples is NHS Choices information on abdominal aortic aneurysm screening.

There is much information that has been published on valid, informed consent. The following are all relevant:

British Medical Ultrasound Society (BMUS)
• Statement on patient information and informed consent

General Medical Council
• Consent: Patients and Doctors making decisions together

Health and Care Professions Council
• Guidance on confidentiality
• GDPR added to Confidentiality guidance for registrants

Royal College of Radiologists
• Standards for patient consent particular to radiology (2012)
2.10 Clinical Governance

Clinical governance is defined in the 1998 consultation document “A First Class Service in the New NHS” and also in 1998 by Scally and Donaldson in the British Medical Journal as:

“A framework through which NHS organisations are accountable for continuously improving the quality of their services and safeguarding high standards of care by creating an environment in which excellence in clinical care will flourish.”

As clinical governance is based on professional values and concern for others, the ultrasound practitioner is actively involved in this process of accountability as part of their daily activities. By safeguarding high standards of care and seeking to continuously improve its quality, it ensures that health care provision is patient-centred which is central to the concept.

The main components of a clinical governance framework can be summarised as follows: ³

i) Risk management
ii) Clinical audit
iii) Education, training and Continuous Professional Development
iv) Patient and carer experience and involvement
v) Staffing and staff management

An example of published Trust information on clinical governance can be found at University Hospitals Birmingham site “The main components of clinical governance”.

Public Health England have produced guidance “Clinical governance” (2019).

For the ultrasound practitioner, clinical governance involves:

i) clinical effectiveness: taking part in personal, departmental and wider audit programmes to evaluate clinical practice and service to patients. This will include audit of ultrasound examinations and reports, participation in multi-disciplinary team meetings and radiology discrepancy meetings;

ii) patient identification, communication and consent: (ref: section 2.9);

iii) patient safety: including avoiding physical injury and following published ultrasound safety guidelines (ref: section 2.3 and section 2.6);

iv) ensure the proper care and maintenance of equipment and not use damaged equipment or equipment that is not fit for purpose;

v) risk management: ultrasound practitioners have a duty to participate in education and training offered by employers on subjects such as back care, health and safety and infection control;

vi) education, training and Continuing Professional Development (CPD) (ref: section 3.3);
vii) team working. See RCR/SCoR document “Team Working in Clinical Imaging” (2012) (Under Review);

viii) patient, public and carer involvement;

ix) being accountable for one’s own actions;

x) the implementation of national clinical guidance which reflects the best standards of care. Examples would include implementing NICE Guidelines and national screening programme guidance and requirements;

xi) incident reporting and raising concerns. This is of particular importance following the publication of the Francis Report in 2013. (ref: section 2.12)

In 2008 the National Ultrasound Steering Group published a document entitled “Ultrasound Clinical Governance”. The National Ultrasound Steering Group was a short-term sub-group of the National Imaging Board.

In June 2018 BMUS published a “position statement on NHS ultrasound services” that refers to the National Imaging Board document. [2018]

The following ‘four layer’ advice from the General Medical Council6 for those who work in an employed environment as part of wider clinical teams is also relevant to the topic of clinical governance and is included for consideration:

‘The first layer (of patient protection) is the individual practitioner and their commitment to a common set of ethics, values and principles which puts patients first. Next is team-based regulation which reflects the importance of acting if a colleague’s conduct or performance is putting patients at risk. After that comes workplace regulation which reflects the responsibilities of NHS and other healthcare providers and finally, the regulator, through work on standards, education and fitness to practise6.

Relevant websites include:
- Royal College of Radiologists (2014) Standards for the provision of an ultrasound service
- NHS Wales. Ultrasound Clinical Governance in Wales
- Royal College of Radiologists (2014) Standards for learning from discrepancies meetings. (Under Review)

Reference:

### 2.11 Ultrasound equipment and quality assurance testing

An ultrasound practitioner is expected to:

i) Have detailed knowledge of ultrasound equipment in order to ensure that it is appropriate for purpose;
ii) Manipulate the equipment correctly so that patient diagnosis and management are not compromised;

iii) Take care when using the equipment in order to avoid damage;

iv) Ensure that regular planned preventative maintenance is carried out by qualified personnel;

v) Ensure that an agreed quality assurance programme is in place that incorporates the regular inspection of ultrasound machines and ancillary equipment.

The stated aims of quality assurance procedures applied to ultrasound equipment are to ensure consistent and acceptable levels of performance of the imaging system and image recording facilities and to ensure the safety of the patient. The foundation of a good quality assurance programme is regular visual inspection of the equipment and the reverberation pattern of each probe by the users, since the majority of faults may be detected in this way. Common faults are damage to probes, which may present an electrical or infection hazard and/or affect the efficiency of all or part of the probe. Formal quality assurance protocols focus on the consistency of specific features of image quality over time. The acceptability of image quality may not be apparent from measurable changes in the parameters tested. The issue of what constitutes unacceptable equipment performance is still very difficult to assess objectively, but there is evidence that probe faults, such as drop out affecting more than 1 element, compromise diagnostic quality.

There is a section relating to image quality requirements, quality assurance and equipment replacement in the RCR/SCoR document “Standards for the provision of an ultrasound service” (2014) (section 2).

The British Medical Ultrasound Society has recommended Quality Assurance testing and monitoring advice available in the article by Dudley et al (2014) “The BMUS guidelines for regular quality assurance testing of ultrasound scanners”.

2.12 Professional responsibilities and safeguarding

2.12.1 Raising concerns

The executive summary of the February 2013 Report of the Mid Staffordshire NHS Foundation Trust Public Enquiry (‘The Francis Report’) is relevant to this sub-section.

NHS Trusts and Health Boards will have their own published policies on ‘raising concerns’ following the publication of the Francis report. All healthcare professionals have a professional duty to report concerns they may have about the safety of patients and of service delivery.

The following advice will also be of help if needing to raise concerns:

Care Quality Commission:
• Report a concern if you are a member of staff

Health and Care Professions Council:
• Fitness to practise - Raising concerns

NHS England:
• Freedom to speak up: raising concerns (whistleblowing) policy for the NHS (2016)
NHS Resolution

- ‘Delivering fair resolution and learning from harm Our strategy to 2022’ (2017) provides information on their strategy to help support NHS staff when things go wrong and help identify ways to learn from errors and prevent future errors.
- ‘Five years of cerebral palsy claims’ (2017) reviewed cerebral palsy claims as part of the NHS target to reduce still birth, neonatal and maternal deaths, to share learning from such cases.

Society and College of Radiographers:

- Raising concerns in the workplace: guidance for SoR members (2015) [SCoR member log-in required]

A “Duty of Care of healthcare professionals” (2013) document provides information for health care professionals about protecting patients and staff.

2.12.2 Safeguarding

Ultrasound practitioners have a duty to report concerns relating to children and vulnerable adults. Employers will have available advice and policies as to the pathways that ultrasound practitioners are required to follow. Training and updating in local safeguarding procedures and policies is a mandatory requirement of the employer.

Further information can be found at:
- Public Health Wales (2017) National Safeguarding Team (NHS Wales)
- Heath and Social Care Board (no date) Northern Ireland adult safeguarding partnership
- NICE (2009, updated 2017) Child maltreatment: when to suspect maltreatment in under 18s
- Dearnley, R (2019) Preventing modern slavery: The role of the NHS

2.12.3 Female Genital Mutilation (FGM) - statutory requirements of practitioners

From 31st October 2015 (England and Wales) there is a legal requirement for all statutory registered ultrasound practitioners to report female genital mutilation in those less than 18 years of age to the police. This duty is on the individual ultrasound practitioner and not the employer; it cannot be transferred. Complying with the duty does not breach any confidentiality requirement or other restriction of disclosure that might apply. The same principle also applies to ultrasound practitioners who are not statutorily registered. Ultrasound practitioners in Scotland and Northern Ireland should use their established safeguarding protocols as required.

Useful information:
- NHS England “FGM: mandatory reporting in healthcare” – has flow diagrams and posters
• NMC (2019) “Additional information on female genital mutilation cases”
• Education Authority Northern Ireland “Multi-agency practice guidelines: female genital mutilation”

2.12.4 Duty of Candour

All healthcare professionals have a responsibility to act in an open and transparent way, particularly when things go wrong. Ultrasound practitioners should also be aware of the requirements of their Professional Indemnity Insurer if asked to make any statements regarding patient care, complaints and claims.

Further information can be found at:
• NHS Resolution (2017) “Saying Sorry”
• General Medical Council and Nursing and Midwifery Council (2015) “Openness and honesty when things go wrong: The professional duty of candour”.

2.12.5 Conscientious objection

The following is the conscientious objection clause from the Standards of Conduct, Performance and Ethics that is associated with the Public Voluntary Register of Sonographers:

‘You must report in writing to your employing authority, at the earliest date in your employment, any conscientious objection that may be relevant to your professional practice. You should explore with them ways in which you can avoid placing an unreasonable burden on colleagues because of this. Your right to conscientious objection does not exempt you from providing service users with full, unbiased information; for example, prior to prenatal screening or testing. You do not have the right to refuse to take part in any emergency treatment’.

The NMC have advice ‘Conscientious objection by nurses, midwives and nursing associates’. This gives further details of the legal circumstances when such a clause might apply.

2.13 Acquisition, archiving and use of ultrasound data

Ultrasound data refers to ultrasound images, ultrasound reports and request forms. They include images captured and stored in digital, video, film and thermal paper formats and written reports/requests generated either in electronic form or on paper. The RCR published the second edition of ‘Guidance on maintaining patient confidentiality when using radiology department information systems’ in November 2019.

Image recording

The compilation of an appropriate number of annotated images that represent the entire ultrasound examination is good practice as it provides the following:

• support for the written report (NHS resolution, 2018);
• a second opinion to be given on those parts of the examination that have been imaged;
• a contribution to clinical governance through audit and quality assurance procedures (RCR and SCoR, 2014; BMUS audit tool (BMUS member log-in required);
• a teaching tool;
• evidence that the examination was carried out to a competent standard;
• evidence that local and national guidelines and protocols were followed.

Unless the entire examination is recorded it must be recognised that the ultrasound images cannot be fully representative of that examination. The stored images will have been chosen by the ultrasound practitioner as a reasonable selection to support the examination report only. If an abnormality or other pathology has been missed during the real time examination, it is unlikely to have been included on an image.

All images should have the following demographic and machine information correctly recorded on them:
• Patient identification
• Date of Examination
• Hospital/Trust/department/provider identification.

Ultrasound practitioners should be aware that the on-screen information is not always reproduced on the recorded images. For example, the safety indices displayed during the real time examination may not be replicated. This will depend on the machine in use.

Managers of ultrasound services should ensure that the local protocols and guidelines address the issue of:
• ultrasound data acquisition
• storage and archiving of specific ultrasound data in accordance with national guidance and current legislation including the data protection and freedom of information acts.

It is the ultrasound practitioner’s responsibility to ensure that they are familiar with local protocols for image recording and storage.

Selection of images

Recording of images should be done in accordance with agreed local protocols and / or as required by national protocols where these exist (e.g. within screening programmes). Ultrasound practitioners should ensure that stored images are correctly annotated and can be benchmarked against the national standard.

Professional judgement should be exercised in the selection and recording of any images in addition to those required by protocol to support the examination report or demonstrate that a measurement has been made. As technology develops, entire examinations may be able to be routinely saved, thus removing the need to select specific images for recording.

Screening programme requirements

The Fetal Anomaly Screening Programme and the Abdominal Aortic Aneurysm Screening Programme have published requirements for the images that must be recorded.
The equivalent screening programmes in the devolved countries will have their own recommendations (ref: section 1.4).

**Independent work (ref: section 2.16)**

Ultrasound practitioners working independently should be clear as to who owns the images, what their responsibilities are for image storage and have clear governance procedures relating to image recording and storage. If images are to be given to the patient at the end of the examination it is recommended that, at a minimum, an identical image set is stored by the provider for future reference.

Ultrasound practitioners working in England under the Care Quality Commission (CQC) ‘practising privileges’ exemption should be aware of the wording on page 8 of the CQC ‘Scope of Registration’. This states that the hosting provider will own all records and will have responsibility for ensuring that essential levels of quality and safety are met.

Only the CQC can give formal advice on registration requirements, the onus is on the independent provider to register unless advised otherwise by the CQC.

General advice on independent practice can be found at section 2.16 of these Guidelines and within the RCR and SCoR’s (2014) “Standards for the provision of an ultrasound service” document and the “NHS Litigation Authority’s (now NHS Resolution) report” (see page 24 onwards).

**Storage of images and data**

There are varying requirements for the storage of images and associated ultrasound data including the report and request form. For obstetric scans this can be up to 25 years or more. Scans on children and young people have similarly long retention times. Full details can be found at NHS Digital (2016) ‘Records management code of practice for health and social care’.

Trusts, Health Boards, departments and independent providers should include details of image storage requirements and responsibilities within their governance procedures.

**Reference:**

**2.14 Recording of images by patients during examinations**

The Society and College of Radiographers updated its guidance on the “recording of images and clinical discussions by patients during diagnostic imaging, interventional procedures and radiotherapy treatment” in January 2019.

There is also a document “NHS obstetric ultrasound examinations. Guidance on sale of images, fetal sexing, commercial considerations and requests to record” (2019).
2.15 ‘Have you paused and checked’ posters and prompt cards

‘Have you paused and checked?’ posters and a prompt card have been published to support ultrasound practitioners in clinical imaging services. They are designed to act as a ready reminder of the checks that need to be made when any ultrasound examination is undertaken. The posters are designed in PDF format to allow for easy downloading and printing. They can then be displayed in the department as required. There are A4 and A3 sizes available and also an A6 version that is small enough to be placed close to a machine or work station. The posters have been developed by a joint working party from the Society and College of Radiographers and the British Medical Ultrasound Society.

2.16 Independent Practice

Many ultrasound practitioners work independently where they are not directly employed by an organisation such as a Health Board, NHS Trust or independent company; some will combine independent work with an employed post. Some ultrasound practitioners set up private or other forms of companies, work as franchisors or franchisees or as a sole trader. There are a wide range of ways in which independent ultrasound practitioners can work. Ultrasound practitioners have also been successful in obtaining contracts via local NHS clinical commissioning groups and the NHS ‘Any Qualified Provider’ scheme, either individually or in partnership with other ultrasound practitioners. These Guidelines cannot provide specific advice although the following information may be of help.

The Care Quality Commission (CQC) regulations which are a legal requirement apply to England only. Ultrasound practitioners practicing independently in the devolved countries are advised to contact the equivalent organisations in their own countries for advice about any legal requirements that may apply.

i) Unless exempted, the legal body that provides a regulated activity (service provider) must by law register with the Care Quality Commission. Regulated activities include ultrasound. Service providers can be an individual, company, charity, partnership, NHS Trust or other organisation. The onus is on the service provider to register. The CQC produced a publication “The scope of registration” (2015)

The regulations relating to hosting arrangements, subcontracted services and ‘practising privileges’ are complex and can be found on pages 8 and 9 of “the scope of registration” document. Only the CQC can formally advise as to whether CQC registration is necessary.

CQC guidance “Independent healthcare services: information for providers” may help with additional advice.

Scotland has published legal requirements on the registration of independent clinics, “the regulation of independent healthcare in Scotland”

ii) Many contracts of employment with Trusts and Health Boards and independent companies require disclosure of independent work and activities that may have a bearing on the work of the employer and, even if not stated, may be implied or judged to be so once operational. There may be conflicts of interests that arise. Independent practitioners are advised to seek advice, for example, from their employer or seek independent legal advice.
iii) Ultrasound practitioners should be entirely clear on when they are working independently (self-employed) and when they are working as an employee. This should be made clear in any written contracts, but some are very poorly written and constructed and this is not always apparent. The Advisory, Conciliation and Arbitration Service (ACAS) have advice on employment contracts or independent legal advice can be sought. Independent practitioners should be aware of the differences between a ‘contract of service’ and a ‘contract for service’. An employee-employer relationship is a ‘contract of service’ and a contractor-client relationship is a ‘contract for services’. Further information can be found on the website “IR35 Explained: ‘Contract of service’ and ‘contract for services’”.

iv) It is a requirement of statutory registration with regulatory bodies such as the Health and Care Professions Council and the Nursing and Midwifery Council that a declaration of having professional indemnity insurance in place is made at the time of initial registration or at renewal or revalidation. (ref: section 2.2) Ultrasound practitioners working independently must be aware of, and follow the requirements and conditions of, their professional indemnity insurer. It is important to be fully aware of any terms and conditions that may apply and that it is applicable to all areas of the ultrasound practitioner’s work. Many professional indemnity policies will apply only to the individual practitioner and not to companies. Some policies such as the SCoR’s Professional Indemnity Insurance are secondary to a contract of employment which must be in place with the employer accepting primary vicarious liability.

Additional insurances for public liability, employer’s liability and the ultrasound equipment itself may also be required depending on circumstances. From April 1st 2013 independent sector providers of NHS services have been able to join the Clinical Negligence Scheme for Trusts (CNST).

v) The safety of patients is paramount and ultrasound practitioners must have documented evidence of their competencies, continuing professional development and reflective practice and show evidence of audit of all aspects of the service(s) they provide. Governance arrangements should include protocols and procedures for image acquisition, storage and retention (ref: section 2.13). See also the 2014 RCR and SCoR document “Standards for the provision of an ultrasound service”.

vi) Independent providers of NHS services are advised that there may be a requirement to hold an NHS Provider (Monitor) licence. Providers are exempt if their annual applicable turnover from the provision of NHS services is less than £10 million but there can be other factors affecting this such as if services are specifically designated by commissioners to require an NHS Provider licence to be held. Providers are advised to make their own enquiries with clinical commissioning groups with which they hold contracts as the rules are complex. These regulations apply in England.

The RCR and SCoR (2014) document “Standards for the provision of an Ultrasound Service” (RCR/SCoR, 2014) applies to independent practice as it does for Trusts, Health Boards and independent companies. There is guidance within on the qualificatory requirements of staff providing independent ultrasound services.

viii) See section 2.13 for guidance on image acquisition, retention and storage which includes reference to independent practice.
ix) Independent providers of national screening programme examinations (e.g. Fetal Anomaly Screening Programme) must follow their published requirements with regards to service delivery and quality assurance procedures.

x) Independent practitioners are encouraged to seek Quality Standard for Imaging (QSI) accreditation of their services (ref: section 3.2) [2018]

xi) General Data Protection Regulation [2018]

The General Data Protection Regulation (GDPR) came into effect on May 25th 2018. Independent sonographers are likely to have responsibilities for data under the Act as data controllers. There will probably be a requirement to register with the Information Commissioner. A registration self-assessment tool is available. The assessment tool takes approximately 5 minutes to complete. There is a small fee (for the majority likely to be £40) if required to register with the Information Commissioner.

xii) Marketing and advertising. [2018]

The SCoR within its “Code of Professional Conduct” allows for the practice of independent practitioners advertising their services, provided that any advertisement conforms to the British Codes of Advertising Practice and Sales Promotion. To achieve the standard, the following criteria apply:

- Advertisements should not be false, fraudulent, misleading, deceptive, self-laudatory, unfair or sensational
- Advertisements should be dignified and professionally restrained.

The health care practitioner/patient relationship is important, therefore due regard should be paid to the maintenance of the highest ethical standards in any advertising. Direct appeals to patients, either face to face or over the telephone, should be avoided. It is undesirable to use too many abbreviations which can be confusing to patients and clients. Comparative claims with other practitioners should not be made in respect of superiority of skills, equipment and/or facilities. The term ‘specialist’ should be restricted to those who have a defined specialist skill. While it may be correct and proper for independent practitioners to be able to publicise their service and practice, they should act in a restrained and professional manner at all times. Claims with regards to registration status should not mislead.

Further information can be obtained from the Advertising Standards Authority (ASA)

Independent practitioners who are registered with a regulatory body such as the HCPC, GMC or NMC must also comply with their requirements.

It is important that claims made about the performance of specific imaging modalities or methods for screening for various types of pathology are accurate. See the documentation ‘Private screening for health conditions: NHS recommendations’
SECTION 3: Education and Accreditation

3.1 E-learning for Healthcare

Image Interpretation is an e-learning programme developed by the Society and College of Radiographers, in collaboration with Health Education England’s (HEE) award-winning e-Learning for Healthcare (e-LfH) programme.

The e-learning provides continuing professional development (CPD) to support sonographers with interpreting ultrasound imaging and offering reports. The online training sessions complement existing teaching methods and provide a valuable reference point. They are designed to be engaging and interactive, using quality images, video and animation. Content is presented using various templates such as ‘real-life’ scenarios, case studies and self-assessment questions.

There are seven modules on ultrasound; each contains a number of sessions as indicated below, all ultrasound sessions were reviewed and updated in 2017:

- Module 15 - Gynaecological Ultrasound - 9 sessions
- Module 16 - Abdominal Ultrasound - 19 sessions
- Module 17 - Men’s Health Ultrasound - 3 sessions
- Module 18 - Vascular Ultrasound - 6 sessions
- Module 19 - Musculoskeletal Ultrasound - 11 sessions
- Module 20 - Head and Neck Ultrasound - 5 sessions
- Module 21 - Obstetric Ultrasound - 15 sessions and 15 multiple choice quizzes

The Image Interpretation sessions have been written by expert ultrasound practitioners to match the format used by e-LfH. This is a very valuable learning resource and can contribute to an ultrasound practitioner’s continuing professional development (Ref: section 1.18)

To access the free sessions, please register at http://portal.e-lfh.org.uk/. All e-LfH programmes are now also available to anyone with an NHS.ac.uk, .ac.uk or .gov.uk account.

For information about the programme please visit http://www.e-lfh.org.uk/programmes/image-interpretation/

In addition to ‘Image Interpretation’ there are a range of other e-LfH modules covering a very wide range of healthcare provision, leadership, communication and core skills. The National screening programme e-learning (FASP, AAA) (ref: section 4.2) is also now on the e-LfH site and available after registration and log in.

3.2 Quality Standard for Imaging (QSI) [updated 2019, previously ISAS standard]

All providers of ultrasound services are encouraged to align their services to the QSI, (jointly owned and developed by the Royal College of Radiologists and the College of Radiographers). Support to meet the standard is available through the 5 staged approach found on the RCR/SCoR websites (www.rcr.ac.uk/qsi or www.sor.org/qsi) or by contacting the Quality Improvement Partner at QSI@sor.org. Services can be independently assessed by the United Kingdom Accreditation Service (UKAS) against that standard. Accreditation provides strong and independent confirmation that high quality services are being delivered, please access the webpages for further information.
QSI is now recognised by the Care Quality Commission and has been approved for use within CQC hospital inspection methodology. Details are on the United Kingdom Accreditation Service site.

3.3 Continuing Professional Development

Continuing professional development (CPD) has been defined as follows:

"...the process by which health professionals keep updated to meet the needs of patients, the health service, and their own professional development. It includes the continuous acquisition of new knowledge, skills, and attitudes to enable competent practice..."\(^8\)

It is a process that seamlessly continues from starting as a student, through successful completion of a first qualification and lasts throughout a healthcare professional’s life. CPD is embedded in the NHS clinical governance strategy (ref: section 2.10). All ultrasound practitioners must be engaged with continuing professional development.

Many ultrasound practitioners are registered with the Health and Care Professions Council (HCPC) as a radiographer or the Nursing and Midwifery Council (NMC) as a nurse or midwife. Medically qualified staff will be registered with the General Medical Council (GMC). These regulators all have their own requirements for CPD. Some regulators use an ‘outputs’ based model based on learning and reflection (e.g. HCPC). Other regulators use an ‘inputs’ based model which will place emphasis on certificates and attending study days.

Statutory regulators provide information on CPD for their registrants, for example the HCPC document “Continuing professional development and your registration”, which applies to many ultrasound practitioners.

If an ultrasound practitioner is not statutorily registered, the same good practice principles of CPD will still apply. For those ultrasound practitioners voluntarily registered with the Public Voluntary Register of Sonographers (PVRS), they will have signed at the point of application to state that they undertake CPD activities. Audit of CPD, based on HCPC principles and systems, was introduced from 1 March 2017 renewal of voluntary registration. Further information can be found in the document “CPD audit requirements for voluntary registrants who do not hold statutory registration”.

Evidence of CPD is also a requirement to maintain accreditation with the Society of Vascular Technology of Great Britain and Ireland (SVT).

Evaluation and reflection on the learning gained from the following activities can all be used as evidence to meet HCPC and PVRS CPD requirements. Other regulators may have a different
emphasize or procedures; ultrasound practitioners are advised to consult the website of their own regulator where information will be available.

i) Successfully completing a programme of study
ii) Attendance at, and participation in, appropriate professional workshops and conferences
iii) Defining and implementing a departmental audit programme
iv) Implementing a change process in practice
v) Mentoring an ultrasound student in practice
vi) Participation in an ultrasound or professional research project
vii) Attendance at and participation in case reviews
viii) Attendance at radiology discrepancy meetings and multi-disciplinary team meetings
ix) Submission of a paper to a journal
x) Critical evaluation of a peer reviewed research paper
xi) Peer review of an article for a journal
xii) Teaching peers and students both formally and informally
xiii) Completion of E-Learning for Health modules and other on-line learning activities
xiv) Active engagement with a professional ultrasound group, professional or regulatory body or scientific society
xv) Promoting the practice of ultrasound to other health care professionals within a wider context
xvi) Communicating with, and imaging, patients

The list of possible activities above is for example only. There are many and varied other activities that can also contribute to an individual’s CPD.

It is important that suitable records are maintained, and evidence is compiled on a regular basis. Evidence should not only include attendance certificates at events but also written records of personal learning, reflection and details of how the learning will improve your practice, the service and experience of service users.

Society and College of Radiographers members have access to “CPD Now”. This is a user-friendly on-line tool for identifying, planning, evaluating and recording CPD that will also help the individual if asked to present evidence of CPD by the HCPC. A ‘CPD Now’ app is available for Android and IOS (Apple) devices.

A document published by the Interprofessional CPD and Lifelong Learning UK Working Group “Principles for continuing professional development PD and lifelong learning in health and social care” was published in January 2019. This provides additional guidance on CPD.

Reference:
3.4 Audit and learning from discrepancy

Introduction

As outlined in the Royal College of Radiologists/Society and College of Radiographers 2014 document “Standards for the provision of an ultrasound service”, objective and robust audit and review of non-obstetric ultrasound imaging is difficult owing to the very nature of the imaging specialty. Ultrasound is an operator dependent imaging modality where image assessment and diagnosis occur in real-time. The recorded still images are a record of the examination that was performed but they do not necessarily reflect the quality of the examination undertaken. That said, assessment of hard copy images can be an indication of whether any imaging parameters have been altered and technique modified in response to the conditions found while undertaking the scan. There are multiple limiting factors affecting the quality and outcome of any ultrasound examination. In addition to patient factors, age and quality of the machine being used, experience of the operator and not least, an understanding of the clinical question being asked all have an impact in the final outcome of any examination. Undertaking quality assurance of ultrasound studies under these conditions is challenging.

The subjectivity and operator dependence of ultrasound imaging needs to be recognised within any audit programme being implemented. Prior to implementation, it is essential that there is an understanding of what the audit programme is trying to achieve. It is also essential that staff members within the team in which the audit programme is being implemented understand the rationale for this programme, understand the process and engage with the process itself.

Audit programmes should be viewed as a process rather than simply a means to an end. It should highlight areas where improvements can be made while recognising that resources and support will be necessary to ensure improvements can be implemented. Learning outcomes and actions points are an essential part of this audit programme so that improvement to clinical practice can be monitored, focused and reviewed.

Actionable reports are required for safe patient management and audit of the reporting outcomes is strongly advised by the RCR (“Actionable reporting”). This method could be used in conjunction with the peer review audit tool outlined below.

For any ultrasound practitioner working in isolation or within a team it is important to gain an understanding of the expected and achievable standard of practice. Knowing and understanding this standard will provide evidence for commissioners of any contracts but, more importantly, it will provide a benchmark against which practitioners can be measured and can measure themselves. While issues of failing competency are rare, they are incredibly difficult to deal with if there is no known standard of practice within a service (Parker and Byass)\(^9\).

Reviewing images and reports is the essential first step in an ultrasound audit programme. A range of practices are already in place which may be used to assess quality or inform services of their performance. These often include Continuing Professional Development (CPD) activities such as follow-up of individual cases, image/discrepancy review sessions and attendance at MDT meetings. While such CPD activities are vital in developing the education of ultrasound practitioner they should be used in conjunction with, rather than instead of a more formal quality assurance programme.\(^10\)

When undertaking an audit programme within an ultrasound service, a holistic view of the quality of an ultrasound examination is required. While image quality and overall report accuracy are important, it is essential that other factors such as clarity, content, readability and relevance of the report should be assessed. A limiting factor to the clarity of the report may be due, in part, to the
quality of the referral. It is recommended that any audit programme assesses the referral for relevance and clarity of clinical question in addition to assessing the ultrasound examination itself.

Audit programme

There are various methods of audit process proposed in the literature and currently being undertaken in practice, however the British Medical Ultrasound Society (BMUS) have devised a universal 'peer review audit tool' that can be used to evaluate the referral, the image quality and report. While this tool may not encompass all audit programmes, it is recommended as a starting point from which in-house audit tools can be developed to meet local needs.

As yet, there are no national standards for expected quality of images and reports for non-obstetric ultrasound. The benchmark standard against which images and reports will be assessed will be limited by the individuals or department standard. An optimum programme would be to recruit external auditors to review practice, be this neighbouring Trusts or colleagues but it is recognised that this may encounter financial and time constraints.

In line with the Royal College of Radiologists publications 'lifelong learning and building teams using peer feedback' (2017) and 'standards of learning from discrepancy meetings' (2014) it is strongly recommended that this peer review audit is undertaken in conjunction with a discrepancy meeting. It is recommended that each service agrees a tolerance level of acceptable quality and any cases falling below this tolerance level should be discussed openly within a discrepancy meeting and learning points and further action agreed within the team of peers.

Recommendations for use of the BMUS audit tool

It is acknowledged that a peer review of images and reports takes time. A reasonable estimation of time required is to allocate an average of 5 minutes per case reviewed. It is recommended that services should aim for a review of 5% of all examinations and reports (RCR, 2014; RCR, 2017).

A timely retrospective audit of cases is required. Services may wish to allocate time on a daily, weekly or monthly basis. Access to image and report storage facilities are required and often assistance from IT departments or PACS managers is required to retrieve retrospective data of examinations performed.

A randomised sample of examinations will reduce bias between reviewers and users of this tool are advised to determine a reliable method to both retrieve data and ensure it is randomised. Some users may prefer that the cases are anonymous, but this can be detrimental to the service being able to identify learning needs of individual practitioners.

Individual services will need to agree who within the team is to perform reviews bearing in mind that an individual’s inclusion into audit programme often improves compliance with them and enhances a sense of ownership and responsibility. It is strongly suggested that all practitioners within a team participate in the process.

A database for identifying cases that have been reviewed and storing the audit data will need to be established. This can be an electronic database, a system linked to hospital patient records or a paper filing system. Services need to be mindful of data protection and information governance guidance and legislation with regards to storing patient data.
Review process

Once cases for review have been identified, the reviewer will need to access the referral information, the stored images and the issued report.

- All three aspects of the examination are reviewed.
- Initially the clinical question should be reviewed. Is it clear and appropriate?
- The images are then reviewed and categorised into good, acceptable or poor. These judgements will be based on the reviewer's own standard of practice.
- The report will then be reviewed and again categorised into good, acceptable or poor. BMUS recommended reporting standards and best practice is documented in the following article (‘What makes a good ultrasound report’ 2014) available from the Ultrasound journal.
- Finally, the reviewer should determine if the clinical question has been answered and whether appropriate advice or a conclusion has been given where appropriate. This can include a statement of normality or 'no cause of symptoms demonstrated' and may be dictated by departmental practice. It is recognised that in some specialised cases, or cases including intervention, a conclusion may not be desirable or helpful.

All scores should be recorded on an appropriate database. It is recommended that cases falling below the departments predetermined minimum standards level are discussed with the individual practitioner before being discussed openly at a discrepancy meeting. It is strongly recommended that any significant errors, such as unreported pathology or significant typographical errors are rectified immediately by issuing either a supplementary report or recalling the patient following discussion with the practitioner and / or clinician referring the examination.

Learning from discrepancies

An ultrasound disagreement is identified by a person second reviewing images and / or a report and their opinion is different to the original in retrospect. This review may occur as part of peer review, MDT or X-ray discussion, in-house audit or as part of routine work.

A reporting discrepancy occurs when a retrospective review, or subsequent information about patient outcome, leads to an opinion different from that expressed in the original report. Not all reporting discrepancies are errors.

It is recommended that local Terms of Reference are agreed when establishing the meetings. The terms of reference should include as a minimum:

- the membership of the meeting;
- the chair of the meeting;
- confidentiality of the cases and discussions;
- agreement as to whether the cases are anonymous or not;
- the review process for cases being discussed;
- how the cases are documented;
- any grading and / or scoring system being implemented;
- any voting process being used;
- who is eligible to vote;
- who is the arbitrator of any discussion;
- how learning outcomes are recorded;
• how action points are recorded;
• how data is recorded for review;
• duty of candour actions;
• how serious errors or discrepancies are escalated.

Suggestions for holding ultrasound discrepancy meetings

It is recommended that discrepancy meetings are held every month. Those present with appropriate qualifications vote on the case using the following system.

Grade of disagreement

The aim of the outcome of discussions at the discrepancy meeting is to grade the severity of the disagreement. The following grades are suggested:

<table>
<thead>
<tr>
<th>Grade</th>
<th>Radiological significance not clinical significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>No discrepancy- simply a disagreement!</td>
</tr>
<tr>
<td>1</td>
<td>Understandable miss Disagreement with report – no action required</td>
</tr>
<tr>
<td>2</td>
<td>Disagreement with report – report amended (should be called most of the time)</td>
</tr>
<tr>
<td>3</td>
<td>Significant disagreement with report – action required (should be called almost every time)</td>
</tr>
</tbody>
</table>

The majority view is taken.

Grade 0 is not a discrepancy but simply a disagreement of opinion between healthcare professionals.

Grade 1 is regarded as an understandable miss. Two examples are detailed below:

• Images of the examination demonstrate that the para-aortic region was not imaged or commented on. Operators need to be aware of reason for scanning para-aortic region. This is done to assess for enlarged lymph nodes, but it is understandable why this was not imaged given the clinical details and presence of gas in the abdomen.

• A liver that appears to be hyperechoic compared to the kidney on review of the images may have appeared normal at the time of the examination. The appearances may be due to the machine quality and setting. Only with additional clinical information such as increased LFT’s would the possibility and significance of fatty infiltration be considered.

These are contentious points, and this is why we take a majority view from a group of qualified peers.

Grade 2 and 3 are reporting discrepancies. A reporting discrepancy occurs when a retrospective review, or subsequent information about patient outcome, leads to an opinion different from that expressed in the original report. Not all reporting disagreements are errors.

Grade 2 may be a discrepancy in which the wording of the report is ambiguous or additional statements are required to improve diagnosis. An example of a grade 2 discrepancy is detailed below:
On review of the images, there is apparent right renal pelvis fullness but no other evidence of hydronephrosis or renal obstruction. No evaluation of resistance indices made which may have improved diagnostic confidence. The report states no cause for obstruction seen implying the kidney is obstructed. On discussion with the peers, the majority would have worded the report differently which may have affected future patient management but not significantly affected outcome.

Grade 3 may be a discrepancy in which an abnormality is demonstrated on the images and not documented on the report or where the abnormality imaged is interpreted differently from the majority view. In the vast majority of these cases the patient is recalled for a subsequent follow up ultrasound examination or alternative imaging to evaluate whether an abnormality has been missed or misinterpreted. The findings of any such review examinations are fed back to the individual operator via a one to one discussion and the team at the next disagreement meeting.

Type of disagreement

If a grade 3 discrepancy has occurred, it is recommended that the discrepancy is benchmarked to promote personal and corporate reflection.

<table>
<thead>
<tr>
<th>Type of Discrepancy</th>
<th>Observation</th>
<th>Interpretation</th>
<th>Poor imaging technique</th>
<th>Poor wording</th>
</tr>
</thead>
</table>

Following discussion and agreement of the grade and type of discrepancy, the meeting should agree learning outcomes and action points for the individual and team. This may include a change to guidelines, additional training, a revision of pathologies and conditions occurring in certain patient groups or simply a review of a single case. Any such learning outcomes and action points should be recorded, with a copy being sent to the individual practitioner and a copy stored securely within the department for future reference.

Duty of Candour

A process needs to be established as to how errors and / or disagreements are communicated to the referrer and subsequently the patient. It is recommended that ultrasound practitioners and departments seek advice regarding local practice and guidelines within their local service. (ref: section 2.12.4).

The NMC and GMC have advice on the Duty of Candour in their document “Openness and honesty when things go wrong: the professional duty of candour” (2015)

Ultrasound Practitioners should also be aware of the requirements of their Professional Indemnity Insurer if asked to make any statements regarding patient care, complaints and claims.
References:

Further information can be found at:
SECTION 4: The Ultrasound Examination

This section includes examination-specific guidelines and common clinical scenarios. They have been compiled by the British Medical Ultrasound Society Professional Standards team and are presented as examples of best practice which it is hoped will be of value to departments. Guidelines or links to information on the vetting and justifying of ultrasound requests, reporting and audit are also included. The Guidelines presented here are not prescriptive. These Guidelines do not and cannot cover all elements of an ultrasound examination and ultrasound practitioners are advised to access additional published information and research in order to fully inform their own local departmental protocols and procedures when there are no nationally agreed ones available.

4.1 Overview of ultrasound examination procedures

Relating to all ultrasound examinations, the ultrasound practitioner should be aware of locally agreed standards of practice and current guidelines of other professional bodies and organisations. The following points should be considered for all ultrasound examinations:

- the patient is correctly identified following required departmental procedures;
- the clinical details provided are sufficient to carry out the examination requested and the correct examination has been requested;
- relevant information is available from the case notes, previous investigations and other sources;
- the role of the ultrasound examination is understood in the clinical context for the patient;
- the potential risks involved in the procedure are explained to the patient;
- valid, informed consent is obtained before proceeding with the examination;
- requirements for chaperones are followed where applicable;
- due consideration is given to the need for an interpreter;
- a systematic scanning approach is adopted that can be modified as required;
- requirements and recommendations should the examination be incomplete;
- the need to extend the ultrasound examination, and/or proceed to additional imaging techniques where necessary in accordance with locally agreed protocol;
- the aftercare of the patient;
- appropriate national and local health and safety regulations including infection control are applied.

4.2 Obstetric, Vascular, Echocardiography and Breast examinations

In the 2015 edition and the subsequent revisions there are no practice guidelines included for obstetric and vascular ultrasound examinations, echocardiography or for ultrasound of the breast.

Early pregnancy and obstetrics

Ultrasound practitioners are referred in particular to publications from the national fetal anomaly screening programmes, the Royal College of Obstetricians and Gynaecologists (especially their Greentop Guidelines), the Fetal Medicine Foundation, the Association of Early Pregnancy Units, British Society of Gynaecological Imaging, the International Society of Ultrasound in Obstetrics and
Gynaecology, the National Institute of Health and Care Excellence (NICE) and RCR (2017) ‘Ultrasound training recommendations for medical and surgical specialities’.

Echocardiography

British Society of Echocardiography (BSE) have published protocols for the various procedures that fall within their remit.

Vascular

For information on this aspect of ultrasound practice, please see the website of the Society for Vascular Technology of Great Britain and Ireland (SVT)

The SVT have published protocols for the various procedures that fall within their scope of practice: http://www.svtgbi.org.uk/professional-issues/ (SVT member log in required)

National recommendations for carotid ultrasound examinations can be found in the article ‘Joint recommendations for reporting carotid ultrasound investigations in the United Kingdom’.

For abdominal aorta see section 4.8.6

4.3 NICE and other guidelines

The National Institute for Health and Care Excellence (NICE) publish a wide range of Guidelines, many of which have implications for ultrasound practice.

Other organisations publishing Guidelines include the following (the list is for example only):

- Royal College of Obstetricians and Gynaecologists (particularly the ‘Greentop Guidelines’)
- British Society of Gynaecological Imaging
- Royal College of Radiologists
- British Medical Ultrasound Society
- Society and College of Radiographers
- British Society of Echocardiography
- Society for Vascular Technology of Great Britain and Ireland
- International Society of Ultrasound in Obstetrics and Gynaecology
- National Screening Committee and the individual national screening programmes (ref: section 1.4)
- Fetal Medicine Foundation
- Clinical Imaging Board
- Association of Early Pregnancy Units

Anyone undertaking an ultrasound examination should meet knowledge, understanding and performance criteria defined by the Skills for Health National Occupational Standards

- CI.C.2019 Perform, interpret and report on ultrasound examinations
- CI.I Perform image guided procedures and/or interventions
4.4 Vetting of ultrasound requests

Introduction

An ultrasound department or provider may receive requests from many different sources including wards, outpatient departments and primary care. Some departments and providers will also accept self-referrals for certain types of examination.

A fully completed ultrasound request in either paper or electronic form will normally be required for every examination undertaken. Departments and providers should make clear within their local requesting protocols who may request an ultrasound examination. This may, for example, be restricted to a medically qualified person or a qualified and registered healthcare practitioner. It is advised that non-medical requestors of NHS ultrasound scans have an agreed scheme of work approved by the imaging department or provider and by their relevant clinical lead prior to referrals being made. If self-referrals are accepted by the department or provider, the circumstances when this may occur should be recorded within the local requesting protocols.

The ultrasound scans themselves may be performed by a variety of staff, in a variety of locations, both in and out of normal working hours. It is essential that ultrasound departments are proactive in managing workload to ensure that the right scan is performed in the right place, by the right person and at the right time. Protocoling of ultrasound requests by an ultrasound practitioner is therefore important.

Aim of Vetting

- To ensure that ultrasound scans are justified and that the correct scan has been arranged with the correct patient preparation.
- To ensure that clinically urgent requests are undertaken in a timely manner.
- To ensure that the scans are undertaken by the right person, in the right place and at the right time. The request should be checked to ensure that it is filled out correctly and complies with individual department policies. The vetting practitioner should be confident that the ultrasound request provides sufficient clinical information and is appropriate to answer the clinical problem posed. There should be an agreed departmental mechanism for dealing with inappropriate requests and requests for which the vetting practitioner is uncertain.

It is recommended that there is a procedure for flagging clinically urgent requests together with a mechanism for dealing with such requests.

4.5 Justification of ultrasound requests

Introduction

Advice has been produced by BMUS in the document “Justification of Ultrasound Requests” (2017) to give best practice guidance to ultrasound providers. It is intended to support primary care physicians and ultrasound providers in the appropriate selection of patients for whom ultrasound (US) would be beneficial in terms of diagnosis and or disease management. (BMUS log-in required)

This document has been written to aid ultrasound providers in justifying that an ultrasound examination is the best test to answer the clinical question posed by the referrer. While it is primarily aimed at primary care, the guidance is also relevant for other referrer groups. This document has been compiled by a panel of ultrasound experts to support good practice in vetting.
and justifying referrals for US examinations. It has been written with a pragmatic approach to managing referrals based on the panel’s expert opinion. This document can be used to assist and underpin any local guidelines that are produced. Reference is made to the evidence-based iRefer publication (Royal College of Radiologists) and should be used in conjunction with this. The NICE guidance NG12. “Suspected Cancer: Recognition and Referral” published in June 2015 (updated 2017) has also been considered in the production of this updated publication.

In many instances, NICE advises urgent direct access CT but if this is unavailable, it advises that patients are referred for an urgent ultrasound examination. Local practice will dictate appropriate pathways following consideration of capacity and demand issues in each Trust.

Principles

This document is based on several non-controversial principles:

- Imaging requests should include a **specific clinical question(s)** to answer, and
- contain **sufficient information** from the clinical history, physical examination and relevant laboratory investigations to support the suspected diagnosis(es).
- The majority of US examinations are now performed by ultrasound practitioners such as sonographers who are not medically qualified. Suspected diagnoses must be clearly stated, not implied by vague, non-specific terms such as “Pain query cause” or “pathology” etc.
- Although US is an excellent imaging modality for a wide range of abdominal diseases, there are many for which US is not an appropriate first line test (e.g. suspected occult malignancy).
- Given sufficient clinical information, most NHS providers will re-direct US requests to CT or MR where appropriate with the agreement of local commissioners.

This general guidance is based on clinical experience supported by peer reviewed publications and established clinical guidelines and pathways. Individual cases may not always be easily categorised and local arrangements for prompt access to specialist advice are essential.

Local guidelines should include identification of who justifies the referral, timescales for vetting and appropriate training for individuals undertaking this process.

Changes to guidelines and pathways should be approved by local trust governance processes. It is recommended that any referrals returned to the referrer have an accompanying letter explaining the rationale behind this. All actions should be documented and recorded on the local radiology information system.

Implementation

For practitioners working in England, most primary care ultrasound services are commissioned via the Clinical Commissioning Group (CCG). Services that have adopted the BMUS Good Practice Guidelines have initially negotiated with the CCGs regarding the service that is to be commissioned. This can be informed by service and good practice guidelines should be incorporated into any contracts.
The suggested steps are given as guidance to service leads to aid implementation of the Good Practice Referral Guidelines:

1. Be clear as to why guidelines are to be introduced in addition to these being best practice guidelines. They may also represent best use of limited resources or may be required to limit demand.
2. Discuss BMUS document with local service colleagues, amending as necessary to suit local practice.
3. Discuss edited BMUS document with local secondary care providers and consultants to ensure that this sits within local referral pathways as some examinations may be better suited to primary or secondary care depending upon local practice.
4. Once agreed, discuss with service provider contracts lead (this may be a team in a large trust or a local manager if an independent provider) and enter into discussions with local CCG.
5. Engage local CCG and GP representatives, attend local learning events, meetings or committees to present case and need for change. Be prepared to edit document further to reflect local requirements.
6. Once agreed, plan a start date for implementation. A lead-in period may be considered where referrals are accepted but where the referrers are informed that they do not fit proposed guidelines and why.
7. Plan vetting time into schedules as declining referrals and communicating with GPs is time consuming.
8. Ensure a clear and robust communication pathway is in place to ensure the referrer is informed if the referral is declined. It is vital that the reasons for declining are clear so that referrers may learn the new guidelines.
9. Ensure there is a system in place to record all referrals, including those declined. This may be the development of a local code on the patient information system that the service uses or a password protected database.
10. Feedback to CCG the impact that the implementation of guidelines is having on waiting lists, referral quality and of any non-compliance.

4.6 Recommendations for the production of an ultrasound report

When considering ultrasound report writing, the following documents are important:

- RCR/SCoR (2014) “Standards for the provision of an ultrasound service” (section 5)

- The Royal College of Radiologists (2015) “position statement on recording the identity of healthcare professionals who report ultrasound examinations”

Safety statement

It is recognised, in the book “the safe use of ultrasound in medical diagnosis” (2012) that all personnel undertaking ultrasound examinations should be aware of the potential biological effects, situations in which examinations may be contra-indicated and be aware of the nationally recommended mechanical and thermal indices and the ALARA principle to reduce ultrasound patient dose.

General comments

- The ultrasound report should be written and issued by the operator undertaking the ultrasound examination and viewed as an integral part of the whole examination.
- The report should be written as soon as possible after the examination has been completed.
- The name and status of the operator issuing the report should be recorded on the report. Where applicable it is good practice to include the statutory regulatory body and registration number of report author.
- The report author should take responsibility for the accuracy of the report and ensure that the report is communicated to the appropriate personnel.
- The report author should be aware of his/her limitations and consequently seek clinical advice when necessary.
- The report author should be aware at all times of the implications for the patient of the contents of the report and act in accordance with local guidelines, policies and procedures.

RCR (2018) “Standards for interpretation and reporting of imaging investigations” highlight nine essential steps for production of an ultrasound report

1. Understanding clinical information

   Sufficient clinical information should be provided by the referring clinician or be available to allow relevant and appropriate interpretation of the images. The person interpreting the images and then producing the report must understand the referring clinician’s information and request. They should ensure that they fully understand the aim of the ultrasound examination in order to affect a clinically useful and relevant report. The diagnostic importance of the report in the clinical management of the patient should be understood.

2. Technical knowledge

   Ultrasound is an operator-dependent technique and the diagnostic quality of the images is very dependent upon the skills of the operator. The person interpreting and reporting the images must be able to reflect critically upon the image quality and appraise the impact on diagnostic accuracy. Where the images are technically sub-optimal (e.g. high patient Body Mass Index), the reporting author must decide whether the patient requires recall and re-scan. It is recommended that any technical limitations of a scan are clearly recorded in the report.
3. Observation

To minimise the possibility of error in producing a report for the wrong patient, it is essential that the report author cross-checks the unique patient identification with the date and type of ultrasound examination performed. (ref: section 1.12)

Observations may be sub-classified as:
- normal findings;
- abnormal findings: expected or unexpected;
- equivocal findings: may be normal or abnormal;
- normal variants.

TOP TIP: It is recommended that a systematic approach to image scrutiny is used for both live and post-scan analysis to ensure that the Shape, Size, Outline, Texture and Measurements of any findings are appropriately considered (Acronym SSOTM).

4. Analysis

Detailed critical analysis of the images should be undertaken, taking into account the observations and clinical reasoning to formulate a clinical opinion and to consider its diagnostic implication.

Examples:
- are the findings abnormal and do they directly relate to the clinical question?
- are the appearances simply normal age-related changes?

If the appearances represent active pathology, then further critical analysis is required to identify the most likely diagnosis and/or to provide a list of differential diagnoses.

5. Medical interpretation

The interpretation of the findings and subsequent report must be considered in the light of the wider clinical picture. In order to produce a relevant diagnostic report, review and understanding of any previous imaging or relevant investigations may be required. The Royal College of Radiologists (2018) states that “a clinically relevant opinion encompasses all the known factors about the patient, as well as the imaging findings”. In this context, it is essential that the report author has extensive medical knowledge to reach a diagnosis or a series of ranked differential diagnoses on which clinical decisions can be made.

Whether the report is produced by a medically qualified or non-medically qualified ultrasound practitioner, it is essential that there are governance procedures in place to ensure that the individual does possess the knowledge, skills and competence to affect this duty without detriment to the patient outcome. In addition, there must be a fail-safe mechanism to ensure that, when required, a radiological or equivalent expert opinion can be obtained promptly to allow issue of the report without undue delay.
6. **Advice**

It is essential that the report author is aware of the diagnostic accuracy of the examination related to the individual patient. The level of certainty or doubt in the diagnosis should be made evident. If a definitive diagnosis cannot be made then advice on further appropriate imaging investigations should be provided, when required. Local policy should exist clearly stating mechanisms for advising on relevant further investigations that takes account of the professional background of the report author, which may be a non-medical one.

7. **Communication with the referrer**

There needs to be an effective mechanism in place locally to ensure timely dissemination of the ultrasound report. RCR (2016) “Standards for the communication of radiological reports and fail-safe alert mechanisms”

The report should be issued in line with Royal College of Radiologists’ recommendations (RCR 2016; RCR 2018) and must provide information which relates to the original clinical question, including expected and unexpected findings and their clinical relevance.

There should be an awareness of the referrer’s understanding of such reports and they should be worded to ensure that the referrer is able to understand the clinical information expressed, including their access to complementary investigations (examples CT/MRI).

A mechanism to allow the referring clinician to discuss the report findings with the report author to ensure better understanding of the clinical impact of the report on management is appropriate. For example, a dedicated department email address may be set up to allow communication between referrers and reporters for such queries.

8. **Taking appropriate action**

The report author also has a duty of care to the patient to ensure that when immediate or urgent action is required that this information is delivered to the referring clinician promptly. This should be agreed at local level through appropriate ‘alert mechanisms’. Such urgent communication(s) should be recorded in the report. (RCR 2016)

9. **Communication with the patient**

It may/may not be possible for the report author to pass on information regarding the outcome of the examination to the patient, particularly if there has not been time to review the wider clinical aspects of the case.

Caution is recommended when discussing the clinical findings with the patient, particularly in respect of possible management strategies where the report author may not have adequate knowledge. Ultrasound practitioners should also adhere to the guidance provided on communication by their professional body.

Appropriate training should be undertaken prior to giving any difficult or unexpected news to ensure that such news is given sensitively and effectively and without ambiguity.
Report authors must be aware that patients may have access to the report and therefore caution in the wording is advocated.

**Report content**

It is recommended that an ultrasound report be divided into the following sections:

- **Type of examination performed**
  The type of ultrasound examination performed should be stated at the top of the report to ensure there is a true record.

  Examples:  
  * Ultrasound examination of the upper abdomen  
  * Transvaginal ultrasound examination of the pelvis

  This is particularly important in the latter example where it is essential to document the intimate nature of the procedure. Care is also required where a decision may have been made for appropriate reasons not to examine the pelvis when an abdomen and pelvis was requested. Standard generated headers e.g. “Ultrasound abdomen and pelvis” must be amended appropriately and the reasons identified in the report.

- **Summary of clinical details**
  It is often helpful to record an overview of key clinical history and findings at the start of the report. If the referring clinician has only provided scant information and further relevant information has been gleaned from the patient by the operator then this should be recorded accordingly. This will assist in setting the report in its true clinical context.

  In cases where the clinical question within the referral is unclear, it may be useful to state the inferred clinical question which the report then sets out to answer.

- **Descriptive element**
  It can be helpful to include a description of the observations and findings to include analysis of the Size, Shape, Outline, Texture and any Measurements of the structures examined.

  For example:
  
  “A well defined mass with mixed echoes is present in the left rectus sheath. The lesion is exquisitely tender. The mass measures 5.2 x 4.6 x 3.6 cm. Appearances and findings are in keeping with a rectus sheath haematoma”.

  Any technical difficulties encountered must be noted, together with their impact on diagnostic accuracy.

  For example:
  
  “Only limited intercostal views of the liver obtained owing to the presence of bowel gas obscuring access. However, where seen, the liver is normal in size and appearance”.

- **Conclusion**
  The standard use of a conclusion is good practice and should include an interpretation of the observed examination findings set into the clinical context. The main principal diagnosis should be given and where possible the RCR recommend that you come to a single diagnosis.
Where this is not possible then the most likely diagnosis should be highlighted, with other options listed in rank order of likelihood (RCR 2018).

- **Differential diagnoses**
  Where these are necessary, they should be limited in number and brief and should include a statement as to why these are less likely than the primary diagnosis.

- **Recommendations for further investigations / management**
  To include any appropriate recommendations for further investigation (e.g. CT/MRI/drainage/biopsy) dependent upon local department guidelines and practice.

- **Identification of the author and contributors**
  The name, position and profession of the person performing and interpreting the ultrasound examination should be clearly stated. (RCR 2018; RCR 2016a; RCR 2016b) This should also apply to anyone involved in second party reporting. It is good practice to state the report author’s GMC, HCPC or any other statutory regulatory body number. The specific action of any secondary involvement should also be stated e.g. “Dr --------, Consultant Radiologist reviewed the images and agrees with the above interpretation”.

- **Identification of Chaperones**
  Details of any chaperones or third parties in the room should be documented giving name and professional capacity. Patient’s actions regarding whether a chaperone was accepted or offered and declined should be documented. (ref: section 2.7).

- **Consent**
  Details of any consent gained prior to the procedure should be documented. For instance, verbal consent gained for intimate (transvaginal) examinations or written consent for biopsy procedures should be documented. (ref: section 2.9).

- **Additional Relevant Information**
  Any further information gained during the examination which may affect future investigations or may have resulted in a non-standard procedure being undertaken should be included. For instance, “Due to latex allergy a latex free probe cover was used”. (N.B. Local guidelines will need to be considered if such statements are to be used).

**Report style** (Edwards et al1 ‘What makes a good ultrasound report’) 

- Reports should be succinct, clear, unambiguous and relevant wherever possible.
- A standard reporting template is helpful for both the ultrasound operator and referring clinicians to establish a ‘house style’ which is clearly understood by all parties involved. However, standard reports which are understood and accepted by staff within a hospital may need to be modified for outside referrals. For example, reports to GPs may require additional advice regarding patient management than reports being written for hospital consultant referrals.
- Abbreviations are not recommended in order to avoid ambiguity and potential confusion.
- Reports should be written in plain English language and should be free from the use of any ultrasound terminology (e.g. transonic, echogenic etc.) as such phrases are
generally meaningless to non-ultrasound users and, as such, may be subject to misinterpretation. Short paragraphs and appropriate layout should be used.

- The report should address the clinical question and generally pertain to the reason for referral.

For example:

“The gallbladder is very tender and cholecystitis is the likely cause of the right upper quadrant pain”.

- Where the examination is extended to examine other areas other than those in the primary request then the structures examined and the subsequent findings must be clearly documented. The examination and the technique(s) used should be documented.

For example:

“A large left sided varicocele is present and, in view of this, the kidneys were examined. Both kidneys appear normal and in particular the left kidney is normal on ultrasound examination.”

- Any limitations (technical or clinical) of the examination should be stated and, if a relevant organ has not been fully examined/assessed, the reason(s) should be indicated.

For example:

“The pancreas is obscured by bowel gas and not seen adequately enough to assess. The gallbladder is contracted as the patient is not fasted therefore the presence of small gallstones cannot be confirmed or excluded with confidence”.

- Diagrams may be useful in describing the findings to the referring clinician (examples: vascular studies and gynaecology examinations) and a standard template can be used as an adjunct to the written report.

- The report should be conclusive where possible, indicating when the appearances are consistent with a specific diagnosis. Where no conclusion is possible, alternative explanations for the ultrasound appearances may be offered.

- Any relevant actions undertaken should be reported.

For example:

“I have informed the patient that she has an ovarian cyst which requires follow up. I have arranged a follow up scan in six weeks’ time. The patient is aware of this appointment. In view of the findings I have personally discussed these results with the referring clinician, Dr XXX by telephone”.

- If second opinions have been sought and given, or if other personnel have scanned the patient, their status, actions and opinions should be stated.

For example:

“Case discussed with Dr XXX, Consultant Radiologist, who agreed with findings and report.”
Patient also scanned by Mrs XXX, Lead Sonographer, who agrees with the findings and report”.

In conclusion, a good report is a report that answers the clinical question. The clinical question is ideally given in an appropriate request. Good reporting avoids confusion, clearly identifies the appropriate findings and gives a correct interpretation in a clear and unambiguous format. Ultimately, good reporting equates to good communication skills and, in the clinical context, will avoid error and potential harm to the patient.

Reporting Audit [2018]

Actionable reports are required for safe patient management and audit of the reporting outcomes is strongly advised by the RCR (2018). This method could be used in conjunction with the peer review audit tool outlined in section 3.4 of this publication.

Reference:

4.7 Gynaecological ultrasound examinations

Gynaecological ultrasound is a useful imaging technique for female patients of all ages and has a role in identifying a wide range of pathology. It is particularly useful for imaging women of child-bearing age and younger patients since there is no associated radiation burden. For maximum diagnostic accuracy the ultrasound practitioner must be aware of age-related appearances, the menopausal status of the patient and expected physiological changes of the menstrual cycle.

4.7.1 General scanning principles

Clinical history

Prior to the examination, the ultrasound practitioner should consider the clinical referral and understand the clinical question posed. Additional essential information may be obtained from the patient. Before commencing the examination the practitioner should be aware of:

- Reason for referral, age, menstrual history, obstetric history, symptoms, relevant medication, previous relevant imaging, previous gynaecological surgery/treatment.

Technique

It is considered best practice for all initial gynaecological ultrasound examinations to be performed via the transabdominal (TA) approach using the distended bladder as an acoustic window. It should be noted that scanning transabdominally with an empty bladder can also be useful. This gives a good overview and can be better in many cases than a scan with an over-filled bladder. Transabdominal assessment allows the whole of the pelvic contents to be imaged. Large or deep pelvic abnormalities may be identified but often with limited resolution.

The patient should be invited to empty their bladder to undergo transvaginal (TV) ultrasound assessment for an alternative and often more detailed evaluation of the pelvic structures. Valid, Informed consent for TV scan must be obtained. It is accepted that some patients may decline this, or it may be inappropriate. Details of the type of examination and patient consent or decision to decline should be documented in the report.
The transvaginal examination allows only a limited area to be imaged but with higher resolution when compared with transabdominal imaging. Large or deep pelvic abnormalities may not always be demonstrated adequately.

Colour Doppler and/or power Doppler may be relevant in appropriate clinical presentations e.g. the assessment of myometrial vascularity, ovarian tumour angiogenesis, endometrial perfusion. Setting a low pulse repetition frequency to maximise Doppler sensitivity is essential.

3D/4D ultrasound offers the ability to assess the uterus and ovaries in multisectional and volumetric reconstruction and may be useful in the assessment of pathology and congenital malformations.

4.7.2 Structures to examine and evaluate

The practitioner should examine structures in at least two planes. The size, outline, echotexture and vascularity, where relevant, of normal anatomy, anatomical variants and abnormal findings should be considered, taking into account menstrual status and age-related appearances.

The following structures should be examined:

- **Bladder** - wall and contents

- **Cervix** - internal os, external os, cervical canal, continuity with uterus, assessment of size, outline, echotexture

- **Vagina** – The vagina may be seen but ultrasound is only of limited value in the assessment of pathology. Limited assessment of outline and echotexture is possible for some pathologies and congenital malformations

- **Uterus** - size, shape, echotexture and position (e.g. anteversion, retroversion) should be noted. The uterine length is dependent upon age and parity. The outline should be smooth. Assess echotexture and uniformity of the myometrium. Consider focal areas/masses. Consider particularly in the transverse plane whether congenital anomalies are present.

- **Endometrium** - thickness, echotexture and junctional zone should be noted.
  - **Premenopausal endometrium**: Consider menstrual-related appearances, thickness, uniformity and junctional zone. The endometrium measures up to approximately 16 mm in normal premenopausal women\(^\text{12}\).
  - **Postmenopausal endometrium**: Consider thickness uniformity and junctional zone. If asymptomatic, up to and including 11 mm\(^\text{13,14}\) is acceptable although features such as increased vascularity or inhomogeneous endometrium should be noted and referral to a gynaecologist may be considered. Protocols should be agreed with local clinicians.

- **Ovaries** - size (in relation to reproductive status), shape and appearance should be noted. A volume calculation obtained from measurements in three different planes should be offered when ovarian volume is considered abnormal. The number and size of follicles may
be relevant depending on hormone status, fertility treatment or use of specific contraceptives.

• **Adnexae** - structures including the broad ligament, pelvic muscles, main blood vessels, and fallopian tubes may be identified.

• **Rectouterine pouch (pouch of Douglas)** - if fluid is present, consider whether is it more likely physiological or pathological.

### 4.7.3 Common Pathologies

#### 4.7.3.1 Uterus

**Fibroids (leiomyoma)** occur commonly, especially in women of black ethnicities, and have an estimated prevalence of up to 70%. They are clinically apparent in 25% of women of reproductive age and are benign tumours of fibrous and smooth muscular tissue. Some contain areas of calcification and/or necrosis. Location is variable and may be intramural, subserosal, submucosal, pedunculated and cervical. If fibroids occur near uterine cornua, they may impinge on the fallopian tube and impair patency.

The common presentation is menorrhagia and/or dysmenorrhoea. Fibroids appear on ultrasound as well-defined areas of altered echogenicity. The fibrous content attenuates the ultrasound beam at a higher rate than the surrounding tissue often resulting in multiple lines of shadowing being visible behind the fibroid.

When fibroids are identified the practitioner should evaluate the number, location(s) and size. Where previous imaging is available, it is important to compare interval changes in size and appearance. Rapid growth is associated with malignancy. Vascular patterns vary considerably and are unreliable for diagnosis, therefore colour Doppler has limited value.

**Uterine leiomyosarcoma (LMS)** is a rare carcinoma of uterine muscle that occurs in one to five of every 1000 women with fibroids. There are no reliable methods to diagnose LMS before surgery and ultrasound appearances may be non-specific but can include a very heterogeneous myometrium with cystic spaces. LMS should be suspected if fibroids change rapidly in size, although it is worth emphasising that even in cases where there has been rapid growth, most fibroids will still be benign.

**Adenomyosis** is diffuse or nodular deposits of endometrium within the myometrium. Presentation is often menorrhagia and/or dysmenorrhoea in multiparous women in their 40s. Ultrasound appearances may comprise an enlarged bulbous uterine fundus and body, and a coarse, heterogeneous myometrial echotexture, often containing small cystic spaces. Frequently, the posterior myometrium is thicker than the anterior myometrium. Acoustic linear striations and an indistinct endometrial/myometrial interface may be visible. Concurrent discrete fibroids may also be seen. However, careful evaluation is required. Previously adenomyosis has been misdiagnosed as ‘early fibroid change’. If the appearance of the uterus is diffuse, with asymmetry of the myometrial walls and linear, rather than peripheral vascularity the appearances is more likely to be adenomyosis.
Endometriosis is the presence of ectopic endometrial tissue at sites in the pelvis other than the myometrium. Fragments of endometrium have the same cyclical menstrual changes as those that are normally sited. Patients may experience cyclical menstrual pain.

The gold standard for detection is laparoscopy. Although ultrasound is now being used more frequently in the diagnosis of endometriosis, however it is acknowledged that a requirement of specialist knowledge and operator dependence often limits the more subtle ultrasound findings.

On transvaginal ultrasound, endometriosis may be seen as discrete nodules in the rectouterine pouch and other peritoneal folds but more commonly, ultrasound practitioners rely on the presence of endometriomas to form a diagnosis.

Endometriomas on ultrasound appear as smooth, thick walled ‘cysts’ with a homogeneous mid-grey ‘ground glass’ echotexture. There are no loculations or solid elements.

Adherent “kissing” ovaries, retroflexion /anteflexion of the uterus or absent ‘sliding side’ on palpation with the TV probe may be secondary features of endometriosis.

TOP TIP: Gently push with the TV probe, after warning the patient, to assess for movement of the pelvic organs.

2.7.3.2 Endometrium

Benign endometrial hyperplasia (defined as > 15 mm to 16 mm\textsuperscript{12} in the secretory phase or more than 11 mm in asymptomatic postmenopausal women)\textsuperscript{13–15}.

This is a common condition, often in response to oestrogen stimulus, which usually regresses spontaneously or responds well to treatment with hormones or minor surgery. Drugs such as HRT and Tamoxifen may induce endometrial hyperplasia. A very small percentage of women with endometrial hyperplasia go on to develop endometrial cancer.

Carcinoma

Almost all (95%) cases of endometrial cancer present as abnormal uterine bleeding or unscheduled postmenopausal bleeding (PMB). In postmenopausal women with PMB, a well-defined endometrium measuring 5 mm or less is unlikely to be cancerous\textsuperscript{14,15}.

Note: reference values vary, and some centres use a threshold of 4 mm\textsuperscript{15,16}. Agree PMB protocols locally with clinicians.

Polyps

These benign growths are common in pre- and postmenopausal women. Most are asymptomatic but may present with intermenstrual bleeding (IMB) or PMB. In cases of thickened endometrium and a suspected polyp, it is often helpful to rescan premenopausal women in the first week of their menstrual cycle.
Ultrasound appearance:
- focal endometrial thickening or a hyperechoic mass within the endometrial cavity
- low flow colour Doppler may help identify a 'feeder vessel' supplying the polyp.

**Asherman's syndrome**
This condition involving scar tissue/adhesions in the uterine cavity may be seen in women with a history of severe pelvic infection or repeated dilatation and curettage procedures. Presentation includes pain, dysmenorrhoea and subfertility.

On ultrasound there may be foci of increased echogenicity in the border between the endometrium and the myometrium (junctional zone). Fibrous connections, or synechiae, may be seen to criss-cross the uterine cavity.

2.7.3.3 **Ovaries**

**Simple cyst** - simple ovarian cysts are common and in premenopausal women most are physiological. The ultrasound appearance is of a well-defined, anechoic, unilocular cyst with thin, smooth walls and good through transmission of sound. Most are asymptomatic, are managed conservatively and resolve spontaneously.

**Haemorrhagic cyst** - ultrasound appearances vary depending on the time of the event causing haemorrhage, but these cysts are commonly homogeneous with thin fibrous internal strands initially but become heterogeneous as clots form and then retract. Presentation is usually pain.

**Ovarian torsion** - the ultrasound appearances are complex and may develop over time. They include:
- enlarged, congested, oedematous ovary
- appearance of a complex adnexal mass
- reduced vascular perfusion
- transient vascular occlusion in cases of intermittent torsion
- ischaemia, infarction
- free fluid in the pelvis

**Polycystic ovarian syndrome (PCOS)**
PCOS is a syndrome of ovarian dysfunction with multiple clinical signs including menstrual irregularities, menstrual infrequency, signs of androgen excess (hirsutism, acne), and obesity. No single diagnostic criterion, such as hyperandrogenism or ovaries with polycystic appearances on ultrasound, is sufficient for clinical diagnosis. It is acknowledged that there are challenges in the diagnosis of PCOS and there should be an aim to prevent over-diagnosis on the strength of ultrasound features especially in the adolescent age group, therefore ultrasound not indicated in adolescents, due to overlap with normal reproductive physiology. According to the “International evidence based guideline for the assessment and management of polycystic ovary syndrome” (2018) exclusion of thyroid disease (thyroid stimulating hormone), hyperprolactinemia (prolactin), and non-classic congenital adrenal hyperplasia (17-hydroxy progesterone) is recommended with further evaluation recommended in those with amenorrhea and more severe clinical features including consideration of hypogonadotropic hypogonadism, Cushing’s disease, or androgen producing tumours. In patients with irregular menstrual cycles and hyperandrogenism, an ovarian ultrasound is not necessary for PCOS diagnosis; however, ultrasound will identify the complete PCOS phenotype.
Two of the following three criteria are required for a diagnosis of PCOS:

- Oligo- or anovulation
- Clinical and/or biochemical signs of hyperandrogenism
- Features of polycystic ovarian morphology (PCOM) on ultrasound

Recommendations from the "International evidence based guideline for the assessment and management of polycystic ovary syndrome" (2018) include:

1. “Ultrasound should not be used for the diagnosis of PCOS in those with a gynaecological age of < 8 years (< 8 years after menarche), due to the high incidence of multi-follicular ovaries in this life stage.
2. The threshold for PCOM should be revised regularly with advancing ultrasound technology, and age-specific cut off values for PCOM should be defined”.
3. Transvaginal ultrasound is the preferred method for the diagnosis of PCO, subject to appropriate consent.
4. Transvaginal ultrasound transducers “with a frequency bandwidth that includes 8MHz” PCOM can be diagnosed when:
   - ≥ 20 follicles on either or both ovaries
   and/or
   - ovarian volume ≥ 10ml if no corpora lutea, cysts or dominant follicles are present.
5. “If using older technology, the threshold for PCOM could be an ovarian volume ≥ 10ml on either ovary.
6. In patients with irregular menstrual cycles and hyperandrogenism, an ovarian ultrasound is not necessary for PCOS diagnosis; however, ultrasound will identify the complete PCOS phenotype.
7. In transabdominal ultrasound reporting is best focussed on ovarian volume with a threshold of ≥ 10ml, given the difficulty of reliably assessing follicle number with this approach.
8. Clear protocols are recommended for reporting follicle number per ovary and ovarian volume on ultrasound. Recommended minimum reporting standards include:
   - last menstrual period
   - transducer bandwidth frequency
   - approach/route assessed
   - total follicle number per ovary measuring 2-9mm
   - three dimensions and volume of each ovary
   - Reporting of endometrial thickness and appearance is preferred – 3-layer endometrial assessment may be useful to screen for endometrial pathology
   - other ovarian and uterine pathology, as well as ovarian cysts, corpus luteum, dominant follicles ≥ equal 10mm
9. There is a need for training in careful and meticulous follicle counting per ovary, to improve reporting.”
Ovarian masses

Ultrasound has an important role to play in managing a suspected ovarian mass. According to NICE (2011) “Ovarian cancer: recognition and initial management”, if the woman has a serum CA125 value of 35 IU/ml or greater, an ultrasound scan of the abdomen and pelvis should be requested.

- A combination of the transvaginal and transabdominal routes may be appropriate for the assessment of larger masses and extra-ovarian disease
- Use of colour Doppler has generally not been shown to significantly improve diagnostic accuracy
- Combined use of the transvaginal route in combination with colour Doppler and 3D imaging may improve sensitivity, particularly in complex cases

**TABLE 1: Ovarian masses in premenopausal women**

<table>
<thead>
<tr>
<th>Description</th>
<th>Suggested management (From RCOG GG62)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Simple. Less than 30 mm diameter. No septa or a single thin septum of less than 3 mm thick. No or minimal calcification in the wall.</td>
<td>Almost certainly physiological. No follow-up required.</td>
</tr>
<tr>
<td>Simple. Measuring between 30 mm and 49 mm.</td>
<td>Almost certainly benign. No follow-up required.</td>
</tr>
<tr>
<td>Simple. Measuring 50 mm to 70 mm.</td>
<td>Almost certainly benign. Yearly ultrasound follow-up. If cyst is reduced in size at the next scan, no further follow-up. If unchanged or larger, gynaecological referral suggested.</td>
</tr>
<tr>
<td>Simple. Measuring more than 70 mm.</td>
<td>Consider MRI and gynaecological referral/surgical intervention.</td>
</tr>
<tr>
<td>Features suggesting dermoid cyst (mature cystic teratoma): solid foci, fluid layering, shadowing and/or 'mesh' appearance, measuring &gt;50 mm</td>
<td>Consider gynaecological referral/surgical intervention.</td>
</tr>
<tr>
<td>Complex mass which is indeterminate or solid, multilocular, thick vascular septa, papillary nodules (+/- vascularity). Ascites may be present.</td>
<td>Tumour markers and gynaecological referral</td>
</tr>
</tbody>
</table>
**TABLE 2: Ovarian masses in postmenopausal women**

Agree all protocols with local clinicians but the following information may be helpful.

<table>
<thead>
<tr>
<th>Description</th>
<th>Suggested management (From RCOG GG34)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asymptomatic, simple, unilateral, unilocular, &lt;50 mm</td>
<td>Low risk of malignancy.</td>
</tr>
<tr>
<td></td>
<td>If the CA125 serum value is normal, 4 - 6 monthly ultrasound surveillance for one year.</td>
</tr>
<tr>
<td></td>
<td>If cyst is unchanged or smaller at the end of this time, and CA125 remains normal, the woman may be discharged.</td>
</tr>
<tr>
<td>Persistent complex adnexal mass - solid, multilocular, focal thickening</td>
<td>Gynaecological referral</td>
</tr>
</tbody>
</table>

**The International Ovarian Tissue Analysis (IOTA) group** has developed some simple ultrasound ‘rules’. These can be used to classify masses as more likely benign (B-rules) or more likely malignant (M-rules). Refer to the IOTA website or to the RCOG (2011) “Ovarian Masses in Premenopausal Women, Management of Suspected (Green-top Guideline No. 62)”.

**B-rules**
- unilocular cysts
- presence of solid components where the largest solid component <7 mm
- presence of acoustic shadowing
- smooth multilocular tumour with a largest diameter <100 mm
- no blood flow

**M-rules**
- irregular solid tumour
- ascites
- at least four papillary structures
- irregular multilocular solid tumour with largest diameter ≥100 mm
- very strong blood flow

Further information on managing ovarian cancer can be found in NICE (2011) guidelines “Ovarian cancer: recognition and initial management”.

**Adnexal masses** include pelvic abscess and/or pyo- and hydrosalpinx. Ultrasound appearances include incomplete septa from dilated serpiginous fallopian tubes, cog-wheel appearance in cross section, debris and low level echoes within.
### 2.7.4 Extending the examination

The ultrasound practitioner may need to consider proceeding to an abdominal ultrasound examination when indicated. For example, examining the kidneys in the presence of a large fibroid to exclude hydronephrosis or to confirm/exclude abdominal ascites where a complex ovarian mass has been seen.

### 2.7.5 Pelvic ultrasound reporting

The report should contain the following information:

- Name and status of the practitioner as well as details of any chaperone present
- Summary of clinical details
- Type of examination performed i.e. transvaginal and/or transabdominal and valid, informed consent
- Any limitations to the examination
- Interpretation of findings, including description and analysis
- Conclusion with recommendation, where appropriate

It may be useful to have a standardised reporting format for normal gynaecological scans, which includes the organs routinely examined and which is acceptable to the imaging department and referring clinicians.

### 2.7.6 Reporting examples

#### Referral for post menopausal bleeding

**Clinical details:** Age 57yrs. Approx. 6 years post menopause – intermittent bleeding for one month. Transabdominal and transvaginal scans of the pelvis performed with verbal consent.

*The uterus is normal in size but there is a 6 mm x 4 mm polyp within the endometrium. The endometrial thickness is 3 mm and is distended by 4 mm of fluid. No other abnormality detected. Both ovaries are of normal appearance.*

**Conclusion:** Endometrial polyp, otherwise normal uterus and ovaries.

*Chaperone Mrs XX, Imaging Assistant, was present during examination with patient’s consent*

__________________________________________________________

#### Referral for pelvic pain with palpable mass

**Clinical details:** 25 days post LMP. Irregular cycle 4-6 weeks. Intermittent pain and O/E pelvic mass palpated.

*Transabdominal and transvaginal scans of the pelvis performed with verbal consent.*
Anteverted uterus containing several submucosal fibroids on the anterior wall, the largest of which is \(X\) mm in diameter. The endometrium is not clearly visualised. Ultrasound appearances of both ovaries are normal with a corpus luteum in the left ovary.

**Conclusion:** Fibroid uterus

Chaperone Mrs XX, Imaging Assistant, was present during examination with patient’s consent

---

**Referral for deep dyspareunia**

**Clinical details:** LMP - unsure six weeks ago. Irregular cycle. Complains of deep dyspareunia of two months duration.

Transabdominal ultrasound of the pelvis performed with verbal consent. Transvaginal scan declined.

Normal anteverted uterus and endometrium, with endometrial thickness \(X\) mm. Ultrasound appearances of the left ovary and adnexa are normal. The right ovary demonstrates normal ultrasound appearances. Adjacent to the right ovary is a complex tubular structure measuring \(Y\timesY\timesY\) mm containing low level echoes. Small volume of fluid noted in the rectouterine pouch.

**Conclusion:** These ultrasound appearances are consistent with pyosalpinx or tubo-ovarian abscess.

Chaperone Mrs XX, Imaging Assistant, was present during examination with patient’s consent

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**References:**


17. The Rotterdam ESHRE/ASRM-sponsored PCOS consensus workshop group (2004). Revised 2003 consensus on diagnostic criteria and long-term health risks related to polycystic ovary


Further information can be found at:

Please refer to advice published by organisations such as:

- National Institute for Health and Care Excellence;
- International Ovarian Tumor Analysis Group;
- British Society of Gynaecological Imaging;
- Royal College of Obstetricians and Gynaecologists;
- International Society of Ultrasound in Obstetrics and Gynaecology.

The YouTube channel ‘Gynaecology Ultrasound’ contains many short gynaecology ultrasound videos, containing still images and ultrasound video clips, on a wide range of subjects including ovarian masses, postmenopausal bleeding and case of the week. If you subscribe to the YouTube channel, you will receive an email alert every time a new video is uploaded.

Dr Susannah Johnson, FRCOG offers a useful website ‘Gynaecology Ultrasound’
4.8 Abdominal ultrasound examinations

4.8.1 General principles

During an abdominal ultrasound examination, the anatomical structures which the ultrasound practitioner should normally examine must be in accordance with the clinical information given and are shown in the following table.

TABLE 1: Structures for Abdominal Ultrasound Examination

<table>
<thead>
<tr>
<th>STRUCTURES</th>
<th>EVALUATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liver</td>
<td>Size, shape, contour and ultrasound characteristics of all segments, appearance of intrahepatic vessels and ducts, porta hepatis and adjacent areas. Portal venous, hepatic venous and arterial systems</td>
</tr>
<tr>
<td>Diaphragm</td>
<td>Contour, movement, presence of adjacent fluid, masses, lobulations</td>
</tr>
<tr>
<td>Ligaments</td>
<td>Appearance of falciform ligament, ligamentum teres and venosum</td>
</tr>
<tr>
<td>Gallbladder</td>
<td>Size, shape, contour and surrounding area. Ultrasound characteristics of the wall and the nature of any contents</td>
</tr>
<tr>
<td>Common duct</td>
<td>Maximum diameter and contents; optimally it should be visualised to the head of pancreas</td>
</tr>
<tr>
<td>Pancreas</td>
<td>Size, shape, contour and ultrasound characteristics of head, body, tail and uncinate process; diameter of main duct</td>
</tr>
<tr>
<td>Spleen</td>
<td>Size, shape, contour and ultrasound characteristics including the hilum. Assessment of splenic vein blood flow and presence/absence of collateral vessels</td>
</tr>
<tr>
<td>Aorta</td>
<td>Diameter, course and branches including the bifurcation, appearance of its walls, lumen and para-aortic regions. See also section 2.8.6</td>
</tr>
<tr>
<td>IVC</td>
<td>Patency, diameter, appearance of its lumen and para-caval regions</td>
</tr>
<tr>
<td>Adrenals</td>
<td>Not routinely viewed but any apparent abnormality of size and ultrasound characteristics should be noted</td>
</tr>
<tr>
<td>Kidneys</td>
<td>Size, shape, position and orientation, outline and ultrasound characteristics of cortex, medulla, collecting system, main and intra-renal arteries and veins</td>
</tr>
<tr>
<td>Ureters</td>
<td>Assessment of the presence/absence of dilatation/reflux/ureteric jets</td>
</tr>
<tr>
<td>Urinary bladder</td>
<td>Appearance of wall and contents. Assessment of volume pre- and post-micturition</td>
</tr>
<tr>
<td>Prostate</td>
<td>Size and shape</td>
</tr>
<tr>
<td>Gastro-intestinal tract</td>
<td>Wall thickness, contents, diameter of lumen, motility, presence/absence of masses</td>
</tr>
<tr>
<td>Other structures</td>
<td>Where relevant include: omentum, muscles, abdominal wall, possible hernias, lymph nodes sites for potential fluid collection (including upper/ lower abdomen and the thorax)</td>
</tr>
</tbody>
</table>

Proceed to examination of the pelvis where necessary (ref: section 2.7)
General Guidance.

Where possible, relevant previous imaging and reports should be reviewed, prior to the ultrasound scan. Document in the ultrasound report if they are not e.g. during electronic downtime or when the original scan was performed elsewhere.

The ultrasound practitioner should tailor the examination according to the clinical presentation. The emphasis of examination of the abdominal structures may be altered according to the clinical scenario and patient history. If targeted ultrasound has been performed, this should be documented within the report, making explicit which structures have and have not been examined.

All abdominal organs should be examined in two planes i.e. longitudinal section (LS) and transverse section (TS), with additional views as required. The purpose of the scan is to survey the entire organ, if possible, with representative images of normality and any pathology being taken. The images should document all abnormal features mentioned in the report.

The abdominal ultrasound examination is inevitably a clinical examination and any tenderness found during a scan should be stated in the report, indicating where possible whether it is organ-specific or diffuse. The absence of tenderness should also be documented where relevant e.g. sonographic Murphy’s sign is negative.

During the examination the ultrasound practitioner should demonstrate:

- normal anatomy/variants of abdominal organs and structures including age-related appearances of each organ in at least two planes. This should include assessment of size, outline and ultrasound characteristics;
- relative echogenicities of abdominal organs;
- pathological findings including focal and diffuse processes and associated haemodynamic findings (pre- and post-operative assessments);
- the presence of any intra-abdominal fluid, focal fluid collections or masses;
- where clinically relevant: vascular anatomy including position, course and lumen of relevant vessels (haemodynamic observations including the presence/absence of flow, its direction, velocity and Doppler waveform).

Further information can be found at:

- British Medical Ultrasound Society (2017) BMUS Recommended good practice guidelines: Justification of ultrasound requests
4.8.1.1 Reporting examples

General principles of reporting apply and reference is made to the reporting section of this document. (ref: section 2.6).

Sample abdominal ultrasound reports

Outlined below are sample reports for various common clinical scenarios. These are provided as guidance with an aim of standardising and improving reporting skills in this important field of practice.

---

**Referral for upper abdominal pain**

**Clinical details:** RUQ pain with occasional vomiting and fatty intolerance, ?gallstones.

Upper Abdominal ultrasound performed with verbal consent:

Normal liver.

The gallbladder is tender, has a thickened, oedematous wall and contains several stones. The CBD is dilated measuring 9 mm, but the lower end of the duct is not demonstrated due to overlying duodenal gas. No intrahepatic duct dilatation.

The pancreas is poorly visualised, despite a water load.

Normal spleen, both kidneys and abdominal aorta.

**Conclusion:** Acute cholecystitis with gallstones and a dilated common bile duct. The lower end of the duct is not seen and MRCP is advised as the next step.

---

**Referral for painless jaundice**

**Clinical details:** Painless jaundice. Bilirubin 400 µmol/L

Upper Abdominal ultrasound performed with verbal consent:

There is intrahepatic duct dilatation around the porta hepatis and into the left lobe of liver. The liver otherwise appears normal. Although no liver lesions are seen on this baseline scan, a non-contrast scan does not exclude the presence of metastases.

The gallbladder is non-tender and contains some small stones.

The CBD is dilated measuring 10 mm down to the head of pancreas, where there is a 20 mm mass. The pancreatic duct distal to the mass is also dilated at 3-4 mm.

No ascites or peri-pancreatic fluid demonstrated.

**Conclusion:** 20 mm mass in the head of the pancreas causing biliary obstruction. This is likely to be malignant. Urgent CT is advised for staging. Report faxed to referring clinician or Pancreatic MDT informed or....

---
**Referral for chronic liver disease**

**Clinical details:** Known chronic alcoholic with liver cirrhosis. For surveillance

Upper Abdominal ultrasound performed with verbal consent:

There is a 1.5 cm nodule in segment 6, which is a new finding since the previous scan of x/x/x.

The portal and splenic veins remain patent with hepatopetal flow and there are varices around the splenic hilum which have increased since the previous examination.

Splenomegaly - 15.5 cm.

No ascites

**Conclusion:** A new liver lesion suspicious for hepatocellular carcinoma is present. An urgent MRI is recommended.

Relevant clinical team informed

**Referral for liver lesion assessment**

**Clinical details:** focal liver lesion on CT scan. ? nature

Upper Abdominal ultrasound performed with verbal consent:

A recent CT KUB demonstrates a 4 cm liver lesion in segment 6. This is confirmed on ultrasound to be a hyperechoic solid lesion. No other liver lesions are present.

2mls of Sonovue contrast agent administered. No known contraindications.

The lesion demonstrates peripheral nodular arterial enhancement with rapid centripetal filling and good contrast take-up in the sinusoidal phase.

**Conclusion:** Benign incidental haemangioma of no clinical significance. The liver is otherwise normal.

**Referral for abnormal liver function tests (LFTs)**

**Clinical details:** Palpable liver edge with abnormal LFTs, H/O alcohol abuse. Smoker.

Upper Abdominal ultrasound performed with verbal consent:

(Additional information from patient records: AST xxx, ALT xxx, Bili xxx)

Fatty liver with several areas of fatty sparing in segments 4 and 8. There are no focal lesions but the liver texture is diffusely nodular and the liver capsule is irregular.

The non-tender gallbladder is contracted and contains several stones. No biliary duct dilatation.

Enlarged spleen - 16 cm.

Patent portal and splenic veins with hepatopetal flow.

Normal pancreas, both kidneys and abdominal aorta.

**Conclusion:** Probable cirrhosis with signs of portal hypertension. Referral to a hepatologist is recommended.
4.8.2 Ultrasound examination of the liver

The liver is a large organ and complete examination requires supine, subcostal and intercostal access, supplemented by repeating in the left lateral decubitus / left posterior oblique positions in order to cover the entire organ.

Imaging should document the liver size, reflectivity, parenchymal echo-texture and surface regularity.

Size

Use of formal liver measurements is controversial as alterations in liver volume may not be reflected in isolated measurements. Moreover, the anthropometry of the patient will affect liver size and shape. Early enlargement may be more sensitively identified by the blunting of the free inferior edge of the right liver. Variation in size and position of liver segments should be documented e.g. enlargement of the caudate (segment 1) in established cirrhosis.

Echotexture

The liver is minimally hyperechoic or isoechoic compared to the normal renal cortex. Where abnormality is suspected, or diffuse liver disease is the clinical indication for the scan, then a split screen comparison image of liver/kidney and spleen/ kidney should be taken.

Where the liver texture suggests steatosis (fatty change) then the following assessments should be specifically made:

- loss of signal in deep liver due to increased attenuation;
- loss of prominence of intrahepatic portal vein branch walls;
- altered liver surface (steatosis and fibrotic change often coexist);
- color and pulse wave Doppler analysis of portal and hepatic veins.

Liver surface

Subtle alterations in liver texture may be confirmed if the liver capsule can be demonstrated to be irregular rather than smooth. Images of the anterior aspect of the right lobe should be acquired intercostally with a high frequency linear probe. A left side down decubitus position and/or left posterior oblique position may be helpful. Careful evaluation of the hepatic veins may also show irregularity relating to hepatic nodularity.

Liver vessels and blood flow

Images should routinely include the hepatic veins draining into the IVC and the portal vein at the liver hilum.

If the liver texture is diffusely abnormal, or if portal hypertension /or chronic hepatitis is mentioned on the request, then Doppler studies of the portal vein and hepatic venous waveform should be obtained. Record the peak velocity and the direction of flow in the portal vein (PV) and the hepatic vein (HV) waveform pattern. Normal lower range of peak velocity in the main portal vein is 12 cm/s in the fasted patient. Higher velocities are not generally of significance except in liver transplants where it may indicate vessel stenosis and needs reporting if above 40cm/s (mean flow velocity).

Normal waveform of hepatic veins should be triphasic. Bi or monophasic flow indicates a loss of liver compliance (report as loss of elasticity to GPs) but this is a very non-specific sign which is also present in other conditions. The use of elastography is useful in assessing the liver stiffness. A section on elastography is available (section 2.13).
Highly pulsatile waveforms in both or either the portal vein or selected hepatic vein is indicative of congestive cardiac failure and this should be reported as may be suggestive as a cause of abnormal LFTs.

Further information can be found at:


2.8.3 Ultrasound examination of the gallbladder and biliary tree

The gallbladder is normally situated inferior to the right lobe of the liver. Its size and shape vary. The gallbladder neck usually sits in the gallbladder fossa and the fundus is frequently mobile, dependent upon patient position.

The gallbladder should be scanned following a period of fasting (drinking clear fluids only) to aid distension. It should be examined at least two patient positions, for example, supine, left side down decubitus, left posterior oblique and/or erect to establish movement of any contents and to unfold the organ. It is good practice to examine the gallbladder both along its long and transverse axes.

The shape and measurements of the gallbladder vary enormously. The gallbladder is normally pear-shaped when optimally dilated, with a narrow neck, widening towards the fundus. A very rounded shape may imply tense dilatation. Some gallbladders may be folded or have a Phrygian cap at the fundus. These shapes can conceal small stones if care is not taken to ‘unfold’ the organ and examine it comprehensively.

The wall of the neck of the gallbladder is slightly thicker than the wall of the body and fundus in a normal organ. Frequently the neck describes a ‘J’ or reversed J shape, and attention should be paid to this area to exclude pathology such as trapped stones. The bile contained in the gallbladder should be anechoic.

The wall should be thin (no more than 3 mm), smooth and well-defined. Measurements of wall thickness should ideally be taken with the wall perpendicular to the beam to reduce artefact due to beam thickness.

In patients attending with RUQ pain, the transducer may be used to gently explore the exact site of the pain. This information may be useful in confirming a diagnosis of cholecystitis. Gallstones are frequently asymptomatic, so it should not be assumed that the finding of gallstones establishes the cause of pain and a full scan should always be conducted.
Attention to equipment settings is important in demonstrating tiny stones, as posterior enhancement from the bile within the GB may obscure shadowing from small stones if incorrectly set.

TOP TIP: Use of a high frequency ultrasound probe is useful when the gall bladder is in superficial location, just below the skin surface.

Bile ducts

The common duct normally lies anterior to the main portal vein and is best imaged using the liver as an acoustic window at this point, with the beam perpendicular to the vein. The duct should then ideally be traced distally to the head of pancreas, (allowing for duodenal gas, which can sometimes be moved by altering patient position and/or gentle pressure from the transducer).

The measurement is taken from inner-lumen wall to inner-lumen wall, measured at the porta hepatis. Duct measurements should be taken in clinical context with reference to previous history, including previous cholecystectomy, and liver function tests.

The normal common duct has a variable diameter. The upper limit of normal calibre is generally accepted at 6mm in the adult, but this can increase after cholecystectomy and in older adults due to loss of elasticity of the duct wall. Individual ultrasound units may choose to use different parameters for normal bile duct size dependent upon the trade-off between sensitivity and specificity for biliary pathology. However, it is recommended that individual ultrasound units work in conjunction with local biliary surgeons where significant deviance from accepted normal ranges is proposed. It is worth noting that a duct less than 6 mm diameter does not necessarily imply normality as biliary obstruction can occur with a non-dilated duct, particularly if diffuse liver disease prevents dilatation or if the patient has cholestatic disease.

Bile duct wall thickening is an abnormal sign, even in the absence of dilatation, and is frequently associated with cholangitis or cholestatic disease.

It is difficult to demonstrate normal intrahepatic ducts past the first order of duct (right and left hepatic ducts) but they can be seen in young, thin patients with good equipment and should not be confused for intrahepatic duct dilatation. Comparison should be made with the accompanying veins. In health, the bile duct is narrower than the parallel portal vein.

In cases of intrahepatic duct dilatation, efforts should be made to establish the cause by tracing the system down to the point of obstruction. Isolated segments of intrahepatic duct dilatation, with a normal calibre extra-hepatic biliary tree, raises the possibility of serious disease such as primary sclerosing cholangitis (PSC) or cholangiocarcinoma. MRI is useful in further evaluation.

Further information can be found at:

4.8.4 Transabdominal ultrasound examination of the pancreas

CT and MRI are the principal imaging modalities for assessing the pancreas. Conventional ultrasound has a significant false negative rate for detecting serious pancreatic disease. However, when the pancreas is well seen, ultrasound may give better detail than CT and MRI.

While ultrasound scanning can image the pancreas completely in exquisite detail under optimal conditions, the result is often disappointing and the published performance data for ultrasound in the detection of treatable pancreatic cancer and chronic pancreatitis is too poor to recommend it as a first line technique.

However, it is expected that the pancreas will be scanned fully in all general upper abdominal examinations. The entire gland should be imaged in a combination of transverse, longitudinal and oblique views.

The pancreas is slightly hyperechoic compared with the liver and this generally increases with age-associated loss of glandular elements and increasing fibro-fatty proliferation. The size of the pancreas is variable and is not routinely measured in practice. As rule of thumb, if measurements are required, the maximum antero-posterior measurement of the head of pancreas is about 3.0 cm the neck and body 2.5 cm and the tail 2.0 cm. The main pancreatic duct can usually be identified and measured in the pancreatic head or body.

A pancreatic duct diameter of 2 mm or less is normal but some increase with age is associated with gland atrophy up to a maximum of 3 mm.

Vetting ultrasound requests for pancreatic imaging (ref: section 2.4 and section 2.5)

For the reasons given above, CT or MRI are usually the most appropriate first line modalities for suspected pancreatic cancer or chronic pancreatitis. Requests for US under these circumstances should be brought to the attention of a radiologist who will decide if CT or MRI is most appropriate.

High quality ultrasound imaging of the pancreas is appropriate for non-specific indications such as epigastric pain, but additional references to loss of appetite or weight loss should be interpreted as indicators of suspected cancer and managed accordingly.

Scanning technique

The adequacy of the scan will be dependent on good technique as much as body habitus. Improving access through these obstacles will depend upon patient preparation (4-6 hrs fasting to reduce bowel gas), use of graded compression, supplementing supine scanning with decubitus or erect positions, and the experience / determination of the operator.
Technique adjustments routinely used by experienced ultrasound practitioners include:

1. using the left lobe of liver as an acoustic window in suspended inspiration;
2. scanning in decubitus and erect positions;
3. intercostal scanning through the spleen to image the pancreatic tail at the splenic hilum;
4. giving water orally to create an acoustic window in the gastric antrum.

The texture, size and contour of the pancreas should be evaluated. The pancreatic duct, distal common bile duct, splenic/superior mesenteric/portal veins and the coeliac axis/superior mesenteric artery should be identified.

The head/uncinate process, neck, body and tail of the pancreas should be identified. The pancreatic tail may be best demonstrated coronally through the spleen, and part of the tail may be seen anteriorly through the gastric body/fundus. The diameter of the pancreatic duct should be assessed.

It is important to document any focal or diffuse change in echogenicity or duct calibre. The presence of parenchymal atrophy should be noted.

If ultrasound does demonstrate suspected inflammation or a mass, it is useful to use Doppler to verify the patency of the splenic and portal veins.

Reference:

4.8.5 Ultrasound examination of the spleen

The spleen is a homogenous organ with a smooth echotexture and border. It is hypoechoic compared to the liver but hyperechoic compared to the kidney.

Size

The spleen can vary greatly in size and the normal range is from 5 cm to 12 cm. It is important to note that spleen size will vary according to the anthropometry of the patient. As a rule of thumb, the spleen length should be the same or less than the left kidney, assuming the kidney is normal. Small spleens are rarely commented upon and are of doubtful significance.

Careful assessment of the splenic hilum should be made as this is a common area for splenunculi to develop. A splenunculus will be of the same smooth, homogenous echotexture as the spleen itself and is not pathological. It is valuable to report however as in patients who have a splenectomy and co-existing splenunculi are commonly reported to hypertrophy and replace the native spleen.
Common pathologies

Splenomegaly

The most common pathological condition of the spleen is splenomegaly. This can occur due to portal hypertension, haematological conditions or underlying infection. Correlation with other ultrasound findings, previous medical history and the patient’s clinical presentation is required. In the presence of ascites, hepatic perfusion with Doppler assessment of the portal and hepatic veins should be carried out. Assessment of the splenic vasculature is required to evaluate the presence of varices commonly associated with portal hypertension. In severe cases spleno-renal varices and shunts may develop.

Trauma

The spleen is the most frequently injured intra-abdominal organ. Ultrasound is a rapid, non-invasive imaging modality and focused assessment with sonography for trauma (FAST)\textsuperscript{21} is an accepted method for evaluating the unstable blunt trauma patients presenting in the emergency department. However, contrast-enhanced CT remains the gold standard for evaluating splenic injury, as well as injury elsewhere within the abdomen or chest.

Solitary Splenic Lesions

<table>
<thead>
<tr>
<th>Appearance</th>
<th>Likely diagnosis and comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cysts</td>
<td>Epidermoid cysts have calcified walls in 10% of cases. Hydatid cysts may be anechoic or of mixed echogenicity due to the presence of hydatid sand or infolded membranes. Multiple, small, internal or subjacent daughter cysts may arise.</td>
</tr>
<tr>
<td>Haemangioma</td>
<td>Usually solitary. Multiple lesions may be associated with Klippel–Trenaú–Weber, Beckwith–Wiedemann or Turner syndrome. Large lesions may result in portal hypertension or splenic rupture.</td>
</tr>
<tr>
<td>Lymphangiomas</td>
<td>Rare, slow-growing, vascular lesions that may be single or multiple.</td>
</tr>
<tr>
<td>Infarction</td>
<td>May be difficult to identify with ultrasound in the acute setting. Contrast enhanced ultrasound imaging can aid diagnosis.</td>
</tr>
</tbody>
</table>
### Multiple Lesions

<table>
<thead>
<tr>
<th></th>
<th>Appearance</th>
<th>Likely diagnosis and comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Infection</strong></td>
<td>Ill defined, hypoechoic lesions, which may contain echogenic debris and internal septations</td>
<td>Splenic abscesses occur most frequently in immunocompromised patients, and may be bacterial, fungal or granulomatous.</td>
</tr>
<tr>
<td><strong>“Spotty” Spleen</strong></td>
<td>Multiple, small (2–3 mm), highly echogenic foci with or without acoustic shadowing and may correspond to calcified lesions seen on plain film or CT</td>
<td>Multiple highly reflective focal lesions within the spleen gives rise to the so-called ‘spotty spleen’ appearance. It is usually secondary to previous granulomatous infection with histoplasmosis or tuberculosis.</td>
</tr>
<tr>
<td><strong>Lymphoma</strong></td>
<td>Multiple ill-defined and hypoechoic lesions, however, hyperechoic lesions of a complex nature and target lesions have also been described</td>
<td>Hodgkin’s and non-Hodgkin’s lymphoma account for the majority of splenic malignancies</td>
</tr>
<tr>
<td><strong>Metastases</strong></td>
<td>Variable appearance, ranging from hypoechoic poorly-defined lesions to hyperechoic lesions, with or without a hypoechoic rim or halo</td>
<td>Uncommon site for metastatic disease. The commonest primary tumours are malignant melanoma, breast and bronchogenic carcinoma. CT remains gold standard for imaging metastatic disease.</td>
</tr>
<tr>
<td><strong>Sarcoidosis</strong></td>
<td>Multiple hypoechoic nodules measuring up to several centimetres.</td>
<td>Hepatosplenomegaly and lymphadenopathy are common associated findings. Multisystem granulomatous disease of unknown origin and splenic involvement is relatively uncommon.</td>
</tr>
<tr>
<td><strong>Malignant Infiltration</strong></td>
<td>Splenomegaly with no discernible alterations in splenic echo texture</td>
<td>Diffuse leukaemic or lymphomatous involvement of the spleen. Clinical correlation is required. Doppler evaluation of the portal vein may be valuable</td>
</tr>
</tbody>
</table>

**Reference:**

**Further information can be found at:**
4.8.6 Ultrasound examination of the abdominal aorta [2018]

Please see standard text books and “NHS Abdominal Aortic Aneurysm screening programme” information for guidance on ultrasound examinations of the abdominal aorta.

In 2017 the SCoR was contacted by a coroner after the death of an 80 year old woman from rupture of a thoracic aortic aneurysm. She was on surveillance for a suspected supra-renal abdominal aortic aneurysm. The ultrasound appearances were reviewed following her death and considered to be those of a thoracic aortic aneurysm extending into the abdomen. The SCoR were asked by the coroner to advise members to consider the possibility of a thoracoabdominal aortic aneurysm via these guidelines. The following advice has been agreed by SoR’s UK Council:

‘When imaging the abdominal aorta sonographers should be alert to the possibility of a thoracic aortic aneurysm extending into the abdominal aorta and creating a thoracoabdominal aortic aneurysm. If a suprarenal abdominal aortic aneurysm and/or dilatation of the upper abdominal aorta is identified the possibility of a thoracic aortic aneurysm extending into the abdomen should also be considered. This should be flagged up in the scan report with wording such as ‘Suprarenal aneurysm/dilatation seen that may extend above diaphragm, further imaging to clarify and exclude a thoracic aortic aneurysm recommended’. Departmental protocols may, however, require the advice of a radiologist or consultant sonographer to be obtained with respect to the precise wording of the report and recommendations for further imaging. An intimal flap may be visible indicating dissection and advice on further imaging and referral on this life threatening pathology must be urgently sought’.

The Abdominal Aortic Screening Programme also has the following information on the extension of a thoracic abdominal aorta into the abdominal aorta (Page 12).

**TOP TIP:** If a suprarenal aortic aneurysm is seen, check

- Is the aneurysm extending from/into the thorax?

Report to suggest that a thoracic aortic aneurysm is a possibility. Recommend further imaging e.g. CT to assess

Reference:

Further information can be found at:

- The Abdominal Aortic Screening Programme (2016) also has information on the extension of a thoracic abdominal aorta into the abdominal aorta (page 12).
4.8.7 Ultrasound examination of the bowel

Bowel symptoms are a common cause for patients presenting to their doctors and being referred for imaging.

Suspected acute appendicitis (AA), diverticulitis and inflammatory bowel disease (IBD) are among the common conditions where ultrasound can make a major contribution in diagnosis and disease monitoring.

**Ultrasound technique**

After a six hour fast, the small bowel is quiet with reduced bowel gas; no other preparation is required.

A general abdominal curvilinear probe is useful to quickly assess the layout of large and small bowel. Particular attention should be given to deeper recesses (e.g. rectovesical pouch) which are inaccessible to the higher frequency probes used for detailed interrogation of the bowel wall.

The bowel diseases of interest are those which thicken and displace bowel contents, making them stand out against normal bowel segments.

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**TOP TIP:** Inflamed bowel is easier to identify than normal bowel.

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Focal bowel masses, segments of wall thickening, or dilated loops may be apparent even at lower frequencies, but high frequency probes are essential to characterize changes in the layers of the bowel wall.

Fortunately, the common bowel diseases most frequently involve bowel segments lying in the iliac fossae which are relatively close to the anterior abdominal wall (2-5 cm) and accessible to higher frequency probes.

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**TOP TIP:** Use Higher frequency linear probes

- the higher frequency the better
- 2-3 focal bands close together in the target zone
- harmonics ON
- edge enhancement
- minimal compounding
- graded compression

Pressure applied to the intestines is uncomfortable. Complaining of discomfort is not an indicator of disease, though patients with IBS may be particularly sensitive.
A gradual progressive increase in pressure (graded compression) brings the probe closer to the bowel, displacing bowel gas and overlying bowel loops and assessing the compressibility / rigidity of normal and abnormal bowel loops and mesenteric fat.

**TOP TIP:** Use left decubitus scanning

The caecum and appendix are on a variable length mesentery with some freedom of movement. Decubitus scanning is routinely recommended as overlying loops may fall away and the caecum position adjusts to reveal the appendix.

In this position the “retro caecal” appendix is often accessible.

Surveying the entire intestine within the abdominal cavity requires a systematic technique such as the use of overlapping vertical sweeps of a high frequency probe up and down the abdomen like a lawnmower.

**TOP TIP:** Use a “mowing the lawn” technique.

### Normal bowel - ultrasound features

The ultrasound appearance of the bowel wall largely reflects the anatomical layering and comprises five alternating bands of higher and lower echogenicity. The outer and inner bright bands are extremely fine and may not be seen.

1. superficial mucosa (fine bright line)
2. deep mucosa including the lamina propria (grey)
3. submucosa (bright)
4. muscularis propria (dark)
5. serosa (fine bright line)

### Abnormal bowel - ultrasound findings

Once the appendix or a suspect bowel segment has been identified, the following aspects should be assessed:

- wall thickening
- altered wall layers
- bowel lumen
- bowel plasticity/mobility/ peristalsis
- altered blood flow
- extramural mesenteric/ interloop changes
Wall thickening

Thickening of the bowel wall is the feature most commonly identified and measurements should be taken. Stomach wall is thickest (~<7 mm). Large and small bowel wall is usually < 3 mm. > 4 mm is usually considered abnormal. Thickening may be due to the presence of oedema, haemorrhage, inflammation, tumour growth or infiltration.

Altered bowel wall layers (gut signature)

Depending on the disease process, the gut signature may be preserved, exaggerated, distorted, diminished or obliterated.

Bowel lumen

Commonly when the bowel wall is thickened, the bowel lumen is compromised, becoming narrowed or strictured. However ultrasound may identify a dilated, fluid-filled, obstructed appendix. In the absence of clinical or sonographic evidence of acute inflammation, this may be a mucocele.

Bowel plasticity / mobility / peristalsis

Most diseases cause stiffening of the affected bowel segment with reduced or absent peristalsis.

Altered blood flow

As a rule, Doppler signals are not evident in healthy bowel wall. With acute inflammation, vascularity increases and dots, lines and branches of vessels can be recorded in the wall (mainly submucosa) and in the adjacent mesentery.

Extramural / mesenteric changes

Bowel wall disease may extend to involve peri-intestinal structures, adjacent loops or solid organs. Around inflammatory bowel lesions, collections and abscesses, mesenteric fat becomes oedematous (swollen and hyperechoic) displacing adjacent structures.

Ultrasound of the normal appendix

The normal appendix can be difficult to visualise. It varies greatly in size (average length 8 cm; range 1-24 cm) and position (pelvic/descending and retrocaecal being the most common). The appendix is identified as a thin, blind-ending tube, with a normal gut signature, in continuity with the caecal pole, arising approximately two centimetres from the ileocaecal valve.

The appendix wall has the same layered structure as the other parts of the intestine and the same thickness of <2 mm. The lumen is continuous with the caecum via a smaller orifice. Gas, fluid, semisolid residue and appendicoliths are often seen in the normal appendix lumen. However, obstruction of the orifice is a common cause of appendicitis.

Acute appendicitis (AA)

Sonographic signs:

- local tenderness
- wall thickening >2 mm (for single wall)
- wall layers may thicken at first, then be lost in transmural low echoes
- the lumen may distend
• the maximum outer diameter (MOD) includes the two walls plus the lumen. A MOD of >6mm does not mean appendicitis but a MOD of < 6 mm makes AA unlikely.

• Hypervascular on Doppler

• mesenteric oedema: the inflamed appendix may be surrounded by hyperechoic swollen fat.

• loculated periappendiceal fluid suggests imminent or actual perforation

TOP TIP: In focal appendicitis the maximum outer diameter may not exceed 6 mm and the diagnosis may be missed if the entire appendix is not visualized

Appendicoliths are frequently identified in asymptomatic patients with otherwise normal ultrasound appearances and are not a reliable indicator of inflammation.

The perforated appendix is even more difficult to find but may most reliably be identified by loss of the hyperechoic appendiceal wall layer (indicating transmural inflammation) and loculated periappendiceal or pelvic fluid collections.

The appendix may lie low in the pelvis or be obscured by bowel. If a normal appendix is not identified or a sonographic diagnosis of acute appendicitis or convincing alternative diagnosis made, further imaging (CT) is indicated in a timely fashion.

**Diverticulitis**

**US features of diverticula**

Diverticula appear as bright “ears” outside the bowel wall with acoustic shadowing due to the presence of gas or inspissated faeces. At higher probe frequencies, a thinned diverticular wall may be demonstrated with a reduced gut signature owing to the absence of muscularis propria. The neck of a diverticulum may be identified as an echogenic band traversing hypoechoic circular muscle which is often thickened.

An inflamed diverticulum is identified as an enlarged echo poor protrusion from the colon wall, with an ill-defined margin surrounded by hyperechoic, non-compressible fat. The gut signature is obliterated by the inflammation. Inspissated faeces may be seen as a central shadowing echogenicity.

Inflammation will commonly extend along the bowel producing asymmetrical or circumferential hypoechoic mural thickening that may be hyperaemic on Doppler scanning. An intramural or pericolic abscess may be identified as an anechoic collection that may contain pockets of air or debris.

**Reference:**


**Further information can be found at:**

4.9 Uro-genital system including testes and scrotum

4.9.1 Ultrasound examination of the kidneys [2018]

Kidneys should be examined in two planes and representative images recorded; images to include the adrenal areas for possible pathology. The renal length/size should be assessed and measured and reported, especially in paediatric cases and urological referrals. Refer to the normal measurements of height/age charts and report if out of the normal range in paediatric patients. Exophytic (protruding) cortical lesions and/or cysts should not be included in the length measurement.

Cortical thickness measurements

The texture, echogenicity and thickness of the renal cortex should be noted. Increased echogenicity and cortical thinning are indicative of renal parenchymal disease and chronic renal impairment.

Cortical thickness measurements should more accurately be called the parenchymal thickness. The parenchymal thickness includes the cortex and medulla. Parenchymal thickness is the measurement taken from the border of the sinus fat and parenchyma to the outer capsule of the kidney. The measurement is taken in longitudinal section at the mid pole (or where it is at its thinnest), being careful to avoid any column of Bertin. The lower limit of normal parenchymal thickness is 10 mm.²

The cortical thickness (yellow line) is from the renal capsule to pyramid. The parenchymal thickness (red line) is from the renal capsule to the sinus echo.
In cases of a suspected renal mass (other than simple cysts) it is useful to assess the contra-lateral normal kidney for cortical thickness. Prominent column of Bertin may be bilateral and if present in the contra-lateral kidney it may guide diagnosis. The use of contrast enhanced imaging should be considered as this is a useful tool in determining altered perfusion within suspicious lesions or the absence of follow in complex haemorrhagic cysts.

As a useful guide remember suspicious renal lesions may demonstrate any of the following:

- Predominantly solid but can be complex cystic masses (refer to Bosniak Classification of Cystic Renal Masses, Version 2019: An Update Proposal and Needs Assessment)
- Usually Exophytic – protruding outside of the renal capsule
- Different echogenicity from the surrounding cortex
- Well demarcated
- Vascularity – either altered or vessels deviated
- Indentation of the renal sinus (hypertrophied column of Bertin might mimic tumour)

Renal masses can range from benign cysts, angiomyolipoma through to malignant lesions. Service providers are strongly advised to develop management guidelines which include necessary follow up, onwards referral or alternative imaging in conjunction with their urological surgical colleagues and radiologists.

Common clinical scenarios

Haematuria

The ultrasound practitioner should establish whether haematuria is microscopic or macroscopic and be aware of the place of ultrasound in the diagnostic testing of a patient with haematuria. The investigation of haematuria may be best performed in the context of a dedicated haematuria clinic according to local practice.

Both kidneys, ureters (if visible) and urinary bladder should be assessed.

Look for renal lesions, hydronephrosis, stones, and lesions within the urinary bladder. The practitioner should be aware of the limitations of ultrasound in detection of transitional cell tumours of the renal pelvis. The patency of the renal veins should be assessed when a solid renal lesion is suspected.

Renal cysts should be documented and assessed for complexity. Ultrasound of simple/minimally complex cysts is sufficient. More complex renal cysts require formal Bosniak grading, either with contrast enhanced ultrasound or CT according to local guidelines.

The practitioner should be aware of normal anatomical variants that may mimic renal lesions such as hypertrophied renal columns (columns of Bertin), splenic humps etc. While the majority of normal anatomical variants should be correctly recognised with ultrasound, contrast enhanced ultrasound, CT or MRI should be considered where there remains doubt over a potential lesion.
The practitioner should be aware that ultrasound has limited sensitivity in assessment for transitional cell tumours in the renal collecting system, ureters or urinary bladder. Patients with unexplained macroscopic haematuria should be considered for further tests to better demonstrate these regions e.g. flexible cystoscopy and CT urogram.

**Acute/chronic kidney injury (AKI – formerly acute renal failure)**

The main aim of ultrasound is first to determine whether renal dysfunction is a medical or surgical (obstructive) problem.

Assess both kidneys for size, parenchymal thickness and cortical reflectivity. The practitioner should be aware of the changes in renal appearances with age. The urinary bladder should be assessed for distension, presence of tumours, trabeculation, wall thickness and diverticulum formation. The size of the prostate gland should be estimated in males and assessment of bladder emptying should be performed (where possible).

Renal cortical echogenicity should be assessed by comparison with adjacent liver/spleen and internally by comparison with medullary pyramids. Increased renal cortical echogenicity implies ‘medical’ renal disease but is a non-specific finding.

Where the kidneys are enlarged, the patency of the renal veins should be confirmed.

The renal collecting system should be examined for dilatation. In cases of pelvi-calyceal dilatation, the collecting systems, ureters, and urinary bladder should be examined to determine the level and cause of any obstruction.

The practitioner should be aware of common causes of bilateral renal obstruction such as tumours of the urinary bladder, pelvis and retroperitoneum; inflammatory conditions such as retroperitoneal fibrosis, endometriosis, and bladder outflow obstruction. The practitioner needs to be aware of physiological renal pelvis dilatation secondary to a full bladder and, in such cases, rescanning following micturition is useful.

In cases of acute kidney injury (AKI), pre-examination preparation with fluid loading should be avoided. In some centres, in-patients are requested to attend with an empty bladder or are catheterised.

**Loin/renal angle pain/obstruction**

Look for ultrasound features of renal obstruction and renal lesions.

The ultrasound practitioner should be aware of the limitations of ultrasound in assessing high grade urinary obstruction. The presence of pelvi-calyceal dilatation is not always due to urinary obstruction, neither does the absence of pelvi-calyceal dilatation rule out high grade obstruction. In the obstructed kidney, the degree of obstruction does not correlate well with the degree of dilatation. Chronic hydronephrosis may be associated with loss of parenchymal thickness.

The renal pelvis and calyces should be assessed for the presence of renal calculi. While larger calculi may be visible, ultrasound is of limited sensitivity in the detection of small calculi. The presence of acoustic shadowing is helpful and the practitioner should be aware of image processing technologies which may reduce the presence of shadowing, particularly spatial compounding. Colour Doppler may be useful to assess for the presence of ‘twinkle’ artefact.
Assess for ureteric distension and level/cause of obstruction. Note that the mid-ureter is frequently not visible due to overlying bowel gas but the ureter at the level of the pelvi-ureteric junction (PUJ) and vesico-ureteric junction (VUJ) is usually amenable to ultrasound assessment.

Secondary signs of obstruction may include presence/absence of ureteric jets within the bladder and increased vascular resistance to intra-renal arterial blood flow; the practitioner should be aware that changes in intra-renal blood flow patterns within an obstructed kidney follow a specific time schedule and the practitioner should be conversant with this.

The practitioner should be aware of the utility of other imaging tests, plain x-ray and CT KUB and the correlative nature of these tests.

**Urinary tract infection (UTI)**

In acute infection, the kidneys are frequently normal.

The kidneys should be assessed for size, position, morphology and echogenicity. The practitioner should be alert to the signs of acute renal infection such as small amounts of perinephric fluid (renal sweat), hydronephrosis/pyonephrosis, diffuse renal enlargement or focal cortical abnormality to suggest focal pyelonephritis.

The kidneys should be examined for anatomical variants which may predispose to urinary tract infection such as duplex kidneys, horseshoe kidney and renal ectopia.

The ureters should be assessed for dilatation. The urinary bladder should be examined for distension, wall thickening, and other signs of bladder outflow obstruction. Assess prostate gland for size. Bladder emptying should be assessed by estimating post-micturition residual bladder volume.

**Renal transplant (immediate post-operative period)**

The objective is to demonstrate any surgical complications which may require immediate intervention.

The size and morphology of the transplant kidney should be examined to establish a baseline for subsequent scans. The practitioner should establish the presence/absence of perinephric haematoma/collection and exclude pelvi-calyceal dilatation.

Colour Doppler of the entire kidney should be performed to establish that blood flow is present to all areas of the kidney. Pulsed wave Doppler of the inter-lobar arteries should also be performed to assess blood flow patterns which may give indirect evidence of arterial (inflow) or venous (outflow) problems of the graft. Doppler indices (resistance or pulsatility indices) should be recorded. The presence/absence of reversed arterial diastolic flow should be established and the practitioner should be aware of the possible causes for this. Finally, the renal vein should be examined with colour Doppler ultrasound to ensure patency throughout its length.

The practitioner should be aware of ultrasound appearances which require immediate surgical intervention such as arterial or venous occlusion of the transplant kidney and discuss with the surgical team when these are suspected.

**Reference:**

4.9.2 Ultrasound examination of the testes and scrotum [2018]

Sonographically examine the testes and epididymes in both a longitudinal and transverse plane, noting the echotexture and echogenicity of each. A full upper abdominal study should also be performed when testicular lesions and suspicious epididymal lesions are demonstrated.

A varicocele is reported if the AP diameter of the peri-testicular veins are more than 3mm\textsuperscript{23,24}. The venous flow is assessed using colour Doppler\textsuperscript{24}. Demonstration of the presence or absence of reflux is made using the valsalva technique and should be documented on the report. If a varicocele is present the upper renal tract and aorta should be examined to exclude the presence of masses compromising venous return\textsuperscript{24}.

**TOP TIPS:** In cases of complex epididymitis / epididymo-orchitis a follow-up scan in 6 weeks is advised, to avoid missing small underlying lesions.

Common clinical scenarios

**Scrotal mass**

The aim is to localise the scrotal lump (intra- or extra-testicular) and characterise if possible.

Both testes should be assessed for size, morphology and the presence/absence of a testicular lesion. The testes should be assessed for echogenicity by comparing both testes on a side-by-side view on the same image. The practitioner should follow local guidelines for referral/alerting clinical teams of suspected testicular cancer.

Assessment of the epididymal heads, bodies and tails should be made for thickening, presence of lesions, cysts etc. Be aware of the presence of normal mild epididymal thickening after vasectomy together with the common appearances of sperm granulomas.

The tunical space should be examined for the presence of hydrocoele and the scrotal wall should be examined for the presence of oedema or thickening.

Every effort should be made to visualise and characterise a scrotal mass. The patient should be asked to localise the mass with direct scanning over this area where the practitioner is unable to visualise the mass during normal scrotal scanning.

**Suspected testicular torsion**

In clinically suspected testicular torsion, ultrasound investigation should not delay surgical exploration to offer the best chance of preserving testicular viability. Ultrasound cannot confidently exclude torsion, but an examination may be performed in cases of acute testicular pain. Practitioners need to be aware of the ultrasound features of torsion and if in any doubt, urgent urological advice should be sought.
Assess the testes, epididymides and tunical space. Colour Doppler of both testes should be undertaken, assessing presence and symmetry of blood flow within the testes. See BMJ Best Practice ‘Testicular torsion’.

The absence of discernible blood flow within a testis (where flow is visible on the contra-lateral normal side) is highly predictive of testicular torsion. However, it should be noted that the presence of blood flow does not exclude torsion, particularly intermittent torsion. The spermatic cord should be examined to assess for the presence of twisting of vessels within the spermatic cord (the whirlpool sign).

The ultrasound practitioner should be aware that testicular torsion can also cause epididymal swelling and hydrocele formation, mimicking epididymo-orchitis, although the absence of intra-testicular flow may be diagnostic. However, intermittent testicular torsion can demonstrate increased intra-testicular blood flow on colour Doppler examination which may be indistinguishable from epididymo-orchitis.

The Healthcare Safety Investigation Branch report ‘Management of acute onset testicular pain’ should be read in conjunction with these guidelines.

Microlithiasis

Microlithiasis (defined as more than 5 echogenic foci per viewable image sector)24,26

Risk factors for microlithiasis include26:

- Previous germ cell tumour
- History of maldescent of the testis / cryptorchidism
- History of orchidopexy (surgery for cryptorchidism)
- Testicular atrophy (<12mls volume)
- Family history of germ cell tumour (1st degree relative)

Current follow-up management of microlithiasis in the asymptomatic and apparently healthy male, with no relevant risk factors26 is as follows:

- Regular testicular self-examination for compliant individuals, with advice to seek early medical attention if necessary;
- Patient information leaflet (appendix 1) ”Testicular microlithiasis imaging and follow-up: guidelines of the ESUR scrotal imaging subcommittee”.

TOP TIPS: For reporting

The report of the initial findings could advise:

“The patient should be educated regarding regular self-examination and to seek immediate medical attention if there are any palpable changes or masses detected. However, if this is a new finding the patient can be referred to urology for management advice”
Follow-up scrotal ultrasound and serum tumour marker testing are NOT recommended\textsuperscript{24,26,27}.

Service providers are strongly advised to develop management guidelines which include necessary follow up, onwards referral or alternative imaging in conjunction with their urological surgical colleagues and radiologists.

References:

4.10 Ultrasound examination of the adult head and neck

General principles
As in all areas of ultrasound, practitioners need to be aware of the various clinical scenarios that can propagate ultrasound requests. In the head and neck region the clinical question posed can be unfamiliar to the occasional operator/reporter. It is essential, in order to carry out a competent examination and issue a helpful report, that the ultrasound practitioner understands the question that is being asked. For example, in cases where a head and neck malignancy is suspected, a full examination of all the major lymph node territories is required in order to accurately stage and optimally manage the patient.

The three most common masses that present to head and neck lump clinics are: lymph node, thyroid and salivary. Examination and reporting needs to be tailored to the request received and the findings on ultrasound examination e.g. if a parotid mass is identified then examination of the contralateral parotid is mandatory to look for potential contralateral tumours (Warthin’s) and the findings should be recorded in the report. Examination may be tailored to a specific area of the neck (e.g. a palpable mass); depending on the findings, the examination may focus on the area in question alone, or may need to be expanded to assess the neck as a whole. For example: in a patient with a posterior triangle mass, if the finding is that of a small superficial lipoma then the examination can be contained to that area in question. The extent of the study should be recorded in the report e.g. “I have not examined the remainder of the neck”. However, if the ultrasound examination identified a
necrotic lymph node which is potentially metastatic then a full assessment of the neck in its entirety is mandatory.

Anatomical structures

1. Lymph nodes

Knowledge of the major lymph node territories and chains is essential and an understanding of the interchange between terminology used in the description of lymph node groups/chains (e.g. deep cervical /jugular chain) and the commonly used Level classification\textsuperscript{28} that is fundamental to head and neck cancer management. Confusion in this area can be easily created, both in requesting and reporting for the unwary. Knowledge of the typical criteria for the ultrasound differentiation between benign and malignant lymphadenopathy is essential. Descriptive reports that fail to classify the nodes into benign, equivocal or possibly/probably malignant are unhelpful.

Benign nodes are classically fusiform in shape, contain an echogenic hilus and possess a central hilar blood flow pattern. Whereas malignant lymph nodes tend to be rounded, the central hilus tends to be absent, contain areas of coagulation or cystic necrosis and exhibit deranged blood flow pattern with areas of vascular sparing and peripheral vessels. Nodal metastases from papillary carcinoma of the thyroid are typically more echogenic and contain punctate micro-calcification. Lymphoma classically presents as marked hypoechoic nodes (pseudo-cystic appearance), rounded with plethoric blood flow which commonly displays a benign central hilar pattern. As the architecture of lymphomatous nodes is preserved, this is mirrored in preservation of the “benign” echogenic hilus sign.

2. Salivary masses

Ultrasound is the optimal initial investigation for a patient with a salivary mass and therefore the practitioner needs to be aware of the common appearances of salivary tumours. 80% of salivary tumours will be benign pleomorphic adenomas and occur in the superficial parotid. They are typically lobular and hypoechoic, often with a pseudo-cystic appearance. Cystic change is suggestive of Warthin’s tumours as are multiplicity and contralateral tumours. An irregular, spiculated outline would suggest a carcinoma.

3. Thyroid

Practitioners should be aware of the typical features that enable a diagnosis of a benign thyroid nodule to be made and those features that indicate a potential malignancy. These have been outlined in the latest issue of the British Thyroid Association Guidelines\textsuperscript{29} and how they can be used to classify the thyroid mass into a benign, equivocal/indeterminate or (suspicious) malignant category (U1–U5) with FNA or core biopsy required for those in the indeterminate of malignant categories, if indicated (i.e. U3–U5).

Reports should therefore outline the features displayed and indicate in which category the findings sit – allowing appropriate management.

Benign nodules may show micro-cystic or cystic change with ring down signs of colloid, egg shell calcification and peripheral colour flow. They are typically hyper-echoic or iso-echoic in relation to the background echo texture whereas a solid hypo-echoic nodule which contains micro-calcification is highly suggestive of a thyroid carcinoma – typically a papillary carcinoma. The shape of the nodule (“taller rather than wide”) is also a sign of potential malignancy. When a carcinoma is suspected, a
search for potential lymph node metastases is required, together with a decision to proceed to FNA or core biopsy in line with guidelines.

4.10.1 Reporting examples

Outlined below are examples of sample reports for various common clinical scenarios:

**Referral for palpable mass (a)**

**Clinical details:** Mobile mass left posterior triangle. 2.5 cm mass. No known primary tumour, patient fit and well.

*Cervical ultrasound performed with verbal consent:*

The palpable mass in the left mid posterior triangle is identified as a fusiform shaped lymph node measured at 2.6 cm x 0.4 cm. It displays an echogenic hilus, the appearances are typical of a benign lymph node – no sinister features identified.

The remainder of the left neck looked unremarkable; I have not examined the right side of the neck.

**Conclusion:** benign left posterior triangle lymph node.

**Referral for palpable mass (b) (Primary care referral)**

**Clinical details:** Right upper/ mid cervical mass, increasing in size for past three months. Smoker. O/E hard mass in right upper cervical region? nodal. Nil else on examination.

*Cervical ultrasound performed with verbal consent:*

The palpable mass is identified as a 3.4 cm diameter lymph node mass in the right upper deep cervical chain. There are signs of coagulation necrosis and possible extra capsular spread – consistent with a metastatic lymph node from a potential squamous cell carcinoma primary.

Further rounded suspicious nodes are seen in the right mid deep cervical chain. Nodes identified in the right submandibular region and lower deep cervical chain and right posterior triangle - but these all appear benign.

The left side of the neck looks clear. The salivary glands and thyroid looked unremarkable. Some incidental benign nodules are seen within the left lobe of thyroid - but no signs of anything sinister.

I have performed a FNA (21g) on the right upper deep cervical lymph node, specimen obtained and sent for cytology. No complications identified.

I have asked the patient to contact your surgery in one weeks’ time to make an appointment with you to discuss the results. I suggest that he is referred for an urgent ENT review.

**Conclusion:** probable metastatic right upper and mid deep cervical chain lymphadenopathy, FNA performed. Urgent ENT referral recommended. Report to be faxed through to surgery.
Referral for right parotid mass. (ENT referral)

Clinical details: Patient noticed a soft lump in parotid region whilst shaving, unsure how long present. O/E 2 cm soft mass in right parotid region, VII exam normal. Nil else to find.
Diagnosis: ? Lymph node  ? salivary mass

Ultrasound cervical region performed with verbal consent:

A 4.6 cm (coronal) x 3.4 cm (AP) x 3.7 cm (sup/inf) mass is identified in the posterior aspect of the tail of the right parotid gland, it is predominantly solid with a small cystic element posteriorly. The mass is well encapsulated and hypo-echoic. No extension into the deep aspect of the right parotid seen. Apart from some benign looking intra parotid nodes, the remainder of the right parotid looks normal. No significant lymphadenopathy within the right cervical region.

A contralateral tumour is identified in the inferior aspect of the left superficial parotid. Again a small cystic element is present, the tumour measures 2.2 cm (AP) x 1.3 cm (coronal) x 2.1 cm (sup/inf) in diameter with no deep lobe involvement. Remainder of the left parotid looks unremarkable, no significant left cervical lymphadenopathy seen.
Both submandibular glands look normal. Nil else of note.

21g FNA performed of the right parotid tumour performed, mucoid material obtained and sent for cytology. No complications identified.

Conclusion: probable bilateral Warthin’s tumours, FNA performed of the right parotid tumour.

Referral for thyroid mass (a)


Ultrasound cervical region performed with verbal consent:

Multiple small iso-echoic nodules are identified in both lobes of the thyroid, several of the nodules exhibit cystic change with ring down signs indicative of colloid. Multi-cystic change also seen. No signs of anything sinister identified.
Thyroid is mildly increased in size, no significant associated lymphadenopathy and no retro-sternal extension present.

Conclusion: small benign multinodular goitre. No signs of anything sinister. Radiological classification U2 – benign.

Referral for thyroid mass (b) (ENT referral)

Clinical details: Left sided thyroid nodule, patient states has been present for two years. Thyroid function normal.
O/E firm nodule left lobe of thyroid, no retrosternal extension. No lymph nodes

Ultrasound thyroid performed with verbal consent:
Within the mid pole region of the left lobe of the thyroid there is a 2.6 cm solid hypo-echoic ovoid mass which contains some hyperechogenic foci – suggestive of micro-calcification. Several smaller nodules are seen in the right lobe but these display typical benign characteristics.

Within the left mid deep cervical chain there is a rounded hyperechoic lymph node just lateral to the IJV which also displays some hyperechoic foci. The remainder of the left neck looks clear as does the right neck.

The findings are highly suggestive of a small papillary carcinoma of the left lobe of the thyroid with a probable left mid deep cervical lymph node metastasis. I have therefore proceeded to a FNA (21g) of both the left thyroid nodule and the left mid cervical node.

Specimens obtained and sent for cytology. No complications identified.

**Conclusion:** probable papillary carcinoma of the left thyroid with left mid cervical lymph node metastasis, FNA performed.

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**Referral for staging ultrasound examination (Maxillofacial referral)**

**Clinical details:** Left lateral/posterior tongue primary SCC, T3. Smoker. Left Level II nodes on examination ?? right II nodes also, ? Bilateral metastases

**Ultrasound cervical region performed with verbal consent:**

The tongue primary tumour can be identified on ultrasound, within the posterior left tongue. It is measured at 1.6 cm (AP) by 1.1 cm (coronal) diameter but does not cross the midline. Superior/inferior diameter difficult to assess but measured at 1.9 cm.

There are multiple rounded lymph nodes with signs of coagulation necrosis in the left upper cervical region, largest measured at 2.1 cm in maximum diameter. There are also similar smaller nodes in the upper mid deep cervical chain. Features are those of metastatic lymphadenopathy. There are small benign looking nodes in the superior left submandibular region and within the lower left jugular (deep cervical) chain and posterior triangle.

Assessment of the right neck is unremarkable, benign nodes seen in the right upper deep cervical chain but no signs of contralateral lymph node metastases.

FNA of the largest node in the left upper deep cervical chain (Level II) performed, no complications identified. Haemorrhagic and necrotic material obtained and sent for cytology.

**Conclusion:** left tongue base/mid tongue tumour, with left upper and mid deep cervical chain (levels II &III) lymph node metastases. FNA performed on left upper deep cervical node. No contralateral lymph node metastases seen.

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**References:**


4.11 Paediatric ultrasound examinations

The Oxford Academic Health Science Network video provides an overview of ultrasound for children in the YouTube video ‘Having a children’s ultrasound appointment in hospital – patient guide’

Exam-specific guidelines and common clinical scenarios

4.11.1 Paediatric liver and biliary system

Liver: neonates and infants

Common clinical scenarios: jaundice, antenatal diagnosis of liver abnormality, RUQ mass, hepatomegaly.

Features to look for and include in report:
- liver shape, homogeneity and reflectivity;
- common duct (CD) normal or dilated (up to 1 mm in neonate), intrahepatic duct size. If CD enlarged are there stones/inspissated bile in it, is there a mass or enlarged lymph nodes compressing the duct?
- gallbladder (GB) shape, size and wall thickness and presence/absence of gallstones;
- portal and hepatic vein patency;
- spleen size, shape and position.

Abnormalities that may be seen include:
- an abnormally shaped liver, usually associated with biliary atresia if situs solitus, ambiguous, or inversus. Chronic liver disease features of left lobe hypertrophy and right lobe atrophy, will not have had time to evolve in a neonate;
- a fatty liver;
- focal abnormalities; benign - calcification, haemangioendothelioma, haemangioma, focal fatty change, cyst; malignant - hepatoblastoma;
- bile duct dilatation – owing to inspissated bile in the CD, choledochal cyst (not to be confused with a cyst at the porta that cannot be connected to the left and right ducts – found in some cases of biliary atresia);
- splenic size and position (? left or right) ?polysplenia. Left-sided polysplenia + situs solitus or ambiguous is diagnostic of biliary atresia.

If any abnormality is found, then a paediatric referral is necessary.

If a normal scan is found in an infant with conjugated hyperbilirubinaemia, then it is mandatory to refer the patient to a paediatric liver specialist as soon as possible so that biliary atresia may be confirmed or excluded and managed accordingly.
Liver: children

Examine: Liver, biliary tree, gallbladder and pancreas.

Common clinical scenarios: jaundice, painful RUQ, upper abdominal and/or back pain, abnormal LFTs, hepatomegaly.

Features to look for and include in report:

- liver shape, homogeneity and reflectivity;
- common duct (CD), normal or dilated (1 mm in neonate, up to 6 mm in teenager, relative to height), intrahepatic duct size. If CD enlarged are there stones in it, is there a mass or enlarged lymph nodes compressing the duct?
- gallbladder (GB) shape, size and wall thickness and presence/absence of gallstones;
- portal and hepatic vein patency;
- spleen size, shape and position;
- pancreas: relatively larger and of lower reflectivity than an adult pancreas, duct > 1mm.

Abnormalities that may be seen include:

- a liver of increased reflectivity (fatty liver) or decreased reflectivity (low fat content if child well, acute hepatitis if unwell);
- an abnormally shaped liver ?CLD;
- focal abnormalities - benign: cyst, calcification, haemangioma, focal fatty change, focal nodular hyperplasia, adenoma, abscess; malignant: hepatoblastoma, hepatocellular carcinoma (HCC) (usually in a cirrhotic liver), fibrolamellar carcinoma, sarcoma, metastases;
- bile duct dilatation;
- enlarged pancreas +/- duct dilatation.

GP patients should be referred to a paediatric specialist if any abnormality is found or if no abnormality is found and the patient remains jaundiced and/or has abnormal LFTs.

If the patient is pain-free and jaundice-free when the scan is performed, and the CD is found to be dilated then an MRCP is necessary to see if there is a choledochal cyst and common channel present.

Paediatric liver pathologies

Focal lesions: benign

- Abscess: an early abscess may be difficult to identify and the only clue may be posterior acoustic enhancement and clinical symptoms. The lesion then becomes echo-poor and more clearly defined and may possibly contain gas if the infection is caused by a gas-forming organism. Portal vein patency may be compromised.
- Adenoma: uncommon in children although they are associated with glycogen storage disorders. They may be of either increased or decreased reflectivity.
- Cysts: simple, choledochal (see biliary section).
• **Calcification:** this may be either incidental small foci that are a sequel of an intra-uterine event such as infection or it may be part of a larger solid lesion.

• **Focal nodular hyperplasia (FNH):** another uncommon lesion in children. They are associated with portosystemic shunts or portal atresia.

• **Haemangioma:** an incidental finding of a small lesion of increased reflectivity in either a subcapsular position or adjacent to a blood vessel. The differential is an area of focal fatty change.

• **Haemangioendothelioma:** found in neonates or infants. There may be multiple small focal lesions of reduced reflectivity throughout the liver or one large vascular lesion. The hepatic artery is large with high velocity intrahepatic flow and the diameter of the abdominal aorta decreases below the level of the coeliac axis. The infant may present in heart failure as most of the arterial blood is being shunted through the liver. These usually involute spontaneously.

• **Mesenchymal hamartoma:** a well-defined mainly cystic mass with multiple septa.

• **Trauma:** CT is usually the first investigation as early liver lacerations may not be visible on ultrasound. Ultrasound can be used in follow-up to measure fluid collections, but CT or contrast ultrasound is necessary to monitor for possible development of pseudo-aneurysms.

Focal lesions: malignant

• **Embryonal sarcoma:** this tumour has a variable appearance, sometimes solid and sometimes cystic.

• **Fibrolamellar carcinoma:** a rare variant of hepatocellular carcinoma (HCC) that usually occurs in older children.

• **Hepatoblastoma:** occurs in young children. The lesion may be solitary or multifocal, often poorly defined and may contain calcification. Adjacent vessels may become invaded.

• **Hepatocellular carcinoma (HCC):** often associated with a cirrhotic liver as found in biliary atresia or tyrosinaemia. The lesion may be solitary or multifocal, often poorly defined but does not usually contain calcification.

These tumours cannot be characterised on ultrasound and cross-sectional imaging +/- biopsy is necessary to confirm the diagnosis. CT is used under the age of one year as MRI contrast is not licensed for use in this age group.

• **Metastases:** these may be found in association with neuroblastoma, Wilms’ tumour, leukaemia, lymphoma.
One condition that can be diagnosed with ultrasound is neuroblastoma stage 4S. This usually occurs in infants under one year old and the adrenal primary together with the liver metastases of decreased reflectivity are characteristic of this condition.

Diffuse liver disease

- **Acute hepatitis:** Sometimes described as a ‘dark liver’ or a ‘starry sky’ appearance. The parenchyma is of reduced reflectivity causing the portal tracts to stand out more than normal.
  * the liver is enlarged with rounded inferior borders
  * the gallbladder wall may be oedematous
  * +/- ascites.

It is possible for a liver to have a low fat content and this appearance must not be confused with acute hepatitis; none of the secondary signs will be present and the child is usually well.

Another pitfall is acute hepatitis in a patient with a fatty liver. The parenchyma may appear of normal reflectivity, look for the other signs.

- **Fatty liver:** Fatty livers may be found in association with obesity, malnutrition, chemotherapy, steroids, storage disorders leading to a metabolic disturbance (glycogen storage disorders, tyrosinaemia (increased risk of HCC).

A ‘bright’ liver; the parenchyma is of increased reflectivity and there may be small areas of focal fatty sparing. These are not focal abnormalities, they are the only part of the liver with a normal appearance. The usual positions for focal fatty sparing are anterior to the right portal vein and superior to the gallbladder. If focal fatty sparing is thought to be in other positions, then it is worthwhile doing a contrast ultrasound scan as a focal abnormality may be present.

- **Chronic liver disease:** The liver has a finely or coarsely heterogeneous appearance with an irregular or nodular surface. A typical position for regenerative nodules is anterior to the right portal vein but further imaging must be performed to exclude malignancy.

There may be lobar atrophy/hypertrophy; often the left lobe is hypertrophied and the posterior right lobe atrophied and the para-umbilical vein may be patent. There may be an increase in periportal reflectivity due to fibrosis around the portal tracts.

Vascular-related pathology.

- **Budd-Chiari Syndrome:** The hepatic veins thrombose and the flow in the portal vein is reversed. There may be ascites and splenomegaly. In chronic Budd-Chiari, the parenchyma becomes heterogeneous and small serpiginous venous channels develop. The portal flow may revert to antegrade and the ascites may resolve.

If clinically indicated, a portosystemic shunt procedure may be performed that shunts the portal flow into the IVC.
**Patent ductus venosus (DV):** This normally closes soon after birth. It may remain patent because it is congenitally abnormal. If this is the case then it will be several millimetres wide, the flow in the left portal vein is reversed and there is no flow in the right portal vein as all the portal flow is shunted through the widely patent DV into the IVC. If left untreated then the right lobe atrophies and FNHs may develop.

The other secondary cause for a patent DV is liver cell failure and it is the liver’s way of trying to decompress the intrahepatic portal pressure. The lumen of the DV is much smaller, usually a millimetre or less, and if the liver recovers the shunt will close spontaneously. The flow in both left and right branches of the portal vein is antegrade. The para-umbilical vein may also be patent.

**Portal vein thrombosis:** This usually occurs if the child had an umbilical catheter inserted as a neonate. A typical ‘bag of worms’ appearance is seen at the porta anterior to the position that the portal vein should lie in. Occasionally there is cavernous transformation where there is one venous channel that may be in the position of the normal portal vein and this is not distinguishable on ultrasound.

**Vascular malformations:** There may be abnormal vascular connections between the portal vein and hepatic artery, with a large draining hepatic vein. Multiple vascular channels may be seen with both arterial and venous flow within them. CT +/- arterial embolisation or resection is normally performed.

Occasionally small shunts between portal and hepatic vein branches or between hepatic veins may be seen but these are usually haemodynamically insignificant and close spontaneously.

The portal velocity should be assessed and spleen length documented.

**Veno-occlusive disease:** This usually occurs after chemotherapy and affects the small vessels, not the main hepatic veins. In the acute phase there is no out-flow in the hepatic veins, they remain patent but the blood only oscillates during respiration. The portal flow reverses and there is ascites and splenomegaly. The liver becomes fatty. As the patient improves the hepatic vein flow re-establishes and the portal flow returns to antegrade flow.

### The biliary system

**Obstructive jaundice** in infants and children is usually caused by calculi, choledochal malformations, pancreatic masses, bile duct tumours (rhabdomyosarcoma), enlarged lymph nodes or sub-hepatic masses.

Neonates that present with conjugated hyperbilirubinaemia may have a dilated common duct due to inspissated bile within it (haemolysis, total parenteral nutrition [TPN], cystic fibrosis). The gallbladder may also contain inspissated bile/small calculi.

The common duct may be up to 1 mm in a neonate and 6 mm in an adolescent.

A choledochal cyst (cystic dilatation of the bile ducts):
Type I – dilatation of the extrahepatic common duct
Type II – diverticulum of the common duct
Type III – dilatation of the intra-duodenal common duct
Type IV – intra- and extrahepatic dilatation
Type V – intrahepatic dilatation only.

An MRCP scan should be performed if a choledochal cyst is found to see whether there is a common channel (the pancreatic duct opening into the lower common duct rather than the duodenum, leading to pancreatitis).

A neonate presenting with a cyst at the porta hepatis will have either a choledochal cyst or biliary atresia. A choledochal cyst can be seen to connect with the bile ducts, whereas the cyst found in some cases of biliary atresia cannot be seen to connect to the bile ducts. The gallbladder also has an abnormal appearance in 90% of cases of biliary atresia.

Cases of biliary atresia are associated with situs inversus, the liver may have an odd configuration, often lying centrally in the epigastrium with two equal-sized lobes. The portal anatomy may be unusual and often there is polysplenia.

Spontaneous perforation of the bile duct: the perforation occurs at the junction of the cystic duct/common hepatic duct and the infant presents with jaundice and ascites.

4.11.2 Paediatric urinary system

Neonates

Common clinical scenarios: Antenatal diagnosis of dilated renal pelvis and/or hydronephrosis, single kidney (pelvic kidneys are sometimes missed on antenatal scans), sepsis.

A baby should be scanned at least 48 hours after birth when following up an antenatal diagnosis of a dilated renal pelvis, so that the baby is not dehydrated at the time of the scan, thus masking any potential renal pelvic pathology.

Assessment:

- two kidneys that are normal in shape, size and position with no collecting system dilatation and normal cortical reflectivity. Neonatal kidneys may have increased cortical reflectivity up to 6 months of age and the pyramids appear relatively dark – not to be confused with cysts. Abnormal orientation (discussed in section on children)
- the renal pelvis can measure up to 6 mm
- a ‘cyst’ in the upper pole of a kidney – this may be an obstructed upper pole moiety in a duplex kidney, associated with a ureterocele
- no normal kidney and several cysts in the renal position- multicystic dysplastic kidney
- a normal-shaped bladder with a normal wall thickness and no evidence of lower ureteric dilatation or ureterocele.

If the renal pelvis is dilated then follow-up scans should be performed at 2 and 8 weeks of age. If the dilatation continues then further imaging will be required.
Children:

**Common clinical scenarios:** UTIs, loin pain, haematuria

**Assessment:**
- two kidneys that are normal in shape, size and position with no collecting system dilatation and normal cortical reflectivity.
- a normal-shaped bladder with a normal wall thickness and no evidence of lower ureteric dilatation or ureterocele. If the patient is able to micturate when asked then a post-micturition bladder scan should be performed to ensure complete emptying.

**Paediatric urinary tract pathologies:**
- only one kidney found: if it is a normal size then there should be an ectopic kidney, often pelvic. If the kidney is hypertrophied then the other kidney has involuted (multicystic dysplastic kidney found antenatally) or is absent.
- abnormal orientation: horseshoe kidneys have their lower poles more medial than the upper poles and there is a bridge of tissue connecting the two lower poles across the midline; cross-fused ectopia: both kidneys lie on the same side with the upper pole of the lower kidney fused with the lower pole of the upper kidney.
- a ‘cyst’ in the upper pole of a kidney, usually an obstructed upper pole moiety of a duplex kidney, often associated with a ureterocele.
- hydronephrosis: PUJ obstruction, VUJ obstruction.
- cysts: uncommon in children, associated with tuberose sclerosis.
- infantile polycystic kidneys: autosomal recessive disorder associated with congenital hepatic fibrosis.
- big, bright (enlarged and hyperechoic) kidneys with multiple cysts, (cysts may be tiny).
- adult polycystic kidney disease, autosomal dominant, may unusually present in childhood.
- the cortical reflectivity may be increased (normal finding up to 6/12 of age), a non-specific finding associated with medical renal disease as in adults.
- Wilms’ tumour: the most common paediatric abdominal tumour.

4.11.3 **Paediatric gastro-intestinal tract**

**Pyloric stenosis.** Projectile vomiting, non-bilious vomiting in an infant

**Assessment:**
- Scan the baby after a 4 hour fast.
- Turn the infant into the right lateral decubitus position so that any residual fluid in the stomach lies in the antrum over the pylorus and displaces any gas from this area. If there is a significant amount of fluid in the stomach, there is impaired gastric emptying. Watch to see if the canal opens.
- If the baby is gassy and an NG tube is present then inject 20 ml of sterile water into the stomach and watch the pylorus; aspirate after the scan if positive for pyloric stenosis.
- Upper limits for normal measurements are:
Single muscle wall thickness (not including mucosa or pyloric canal) = 3 mm
Canal length = 16 mm
TS diameter = 11 mm

**Intussusception**

Abdominal pain, redcurrant jelly stools, palpable abdominal mass.

An intussusception is usually ileo-colic. Most have no obvious pathology as the lead point, 10% are due to Meckel’s diverticula, polyps and duplications. It is also associated with lymphoma, haematoma and cystic fibrosis (CF).

Features to look for and include in report:

A segment of bowel prolapses into a more caudal segment and it is seen as bowel within bowel on cross section – appearances are like onion rings.

**Reference:**

**4.11.4 Neonatal hip**

Breech delivery and family history of developmental hip dysplasia are the most common referral criteria. However, local practice and the ‘Newborn and Infant Physical Examination screening programme (NIPE)’ guidelines should be taken into consideration when justifying referrals. Note that the conduct of the ultrasound examination of the neonatal hip is not itself part of the NIPE screening programme.

All babies with an abnormality detected on clinical examination should be scanned within two weeks of age. Babies with a known risk factor but no detectable abnormality are to be scanned by six weeks of age (NIPE). Whilst not mentioned in the NIPE standards, experience has shown that ‘abnormal’ features may be physiological before six weeks-of-age and therefore premature babies should be age-corrected before examination.

It is helpful to have a cradle to put the baby into as it keeps them in the lateral position comfortably.

Scan longitudinally over the greater trochanter parallel to the cradle to obtain a coronal image of the acetabulum at its maximum depth.

\( \alpha \) angle – this gives the depth of the acetabulum. The baseline goes along the straight lateral margin of the ilium. The second line goes from the inferior point of the iliac bone tangential to the bony acetabulum. An angle of greater than 60° is normal, a smaller angle indicates dysplasia.

A shallow acetabulum in a baby less than 3/12 old may be physiological immaturity but if found after 3/12 of age it signifies dysplasia.
β angle – this angle is useful in classifying the degree of dysplasia. The β angle is formed between the vertical cortex of the ilium and the triangular labral fibrocartilage. There is considerable variability in the measurement of this angle and it is, therefore, not always used.

Neither angle can be measured if the hip is dislocated.

Move the hip and watch it during manipulation to see if it subluxes. If so then follow up in 2 weeks. If the hip is still unstable then referral is necessary for a harness to be fitted.

Refer to textbooks/published articles for diagrams of the angles referred to above

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**TOP TIP:**
You can’t make a normal hip look abnormal, but you can make an abnormal hip look normal.

Femoral head ossification can be seen anywhere between 2 and 8 months.

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Further information can be found at:


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### 4.11.5 Neonatal intracranial ultrasound

The ultrasound practitioner should be aware of the potential bio-effects of diagnostic ultrasound, particularly heating at brain/skull interface, and ensure that the machine is set to the minimum power consistent with obtaining a high quality diagnostic ultrasound image. Colour Doppler should not be utilised except for clearly defined clinical reasons which provide additional diagnostic or prognostic information.

The ultrasound practitioner should be aware of common and less commonly used acoustic windows to the neonatal brain. While the anterior fontanelle is used as standard, the posterior fontanelle can be useful to examine the occipital horn of the lateral ventricles and the mastoid suture may be helpful in examining the posterior fossa, cerebellum, aqueduct of Sylvius and 4th ventricle.

A suggested approach for examining the neonatal brain is as follows:

**Presence of normal anatomy**
The ultrasound practitioner should be aware of normal brain anatomy in the neonate, to include changes with age, presence of midline structures, ventricular appearances, appearances of basal ganglia, periventricular white matter, cerebellum and extra-axial space.
Presence of intracranial haemorrhage
The ultrasound practitioner should be aware of common locations of intra-cranial haemorrhage, how this may vary according to gestational age, and how these may present on ultrasound. Grading of haemorrhage should be according to local agreement and protocol.

Ventricular size
The size of the lateral, 3rd and 4th ventricles should be assessed according to local protocol. Validated measurement techniques (e.g. ventricular indices) should be utilised in cases of ventriculomegaly. This can be useful to assess change in size over time and guide timing of intervention.

Periventricular white matter
The ultrasound practitioner should be aware of normal and abnormal appearances of the periventricular white matter and the limitations of ultrasound in examining this region. Follow-up should be considered in cases of periventricular flaring or suspected white matter damage.

Brain parenchyma
The ultrasound practitioner should be aware of normal sonographic appearances of the brain parenchyma and be aware of the potential appearances of acquired abnormalities.

4.12 Musculoskeletal ultrasound examinations

General principles.
Musculoskeletal ultrasound referrals, for the evaluation and treatment of joint and soft-tissue diseases, have increased rapidly over the last 10 years. The European Society of Musculoskeletal Radiology (ESSR) produced a consensus statement on ‘Clinical indications for musculoskeletal ultrasound’ (2017).

Advantages of ultrasound include detailed, real time, dynamic assessment of soft tissues and the ability to compare with the contralateral side. Doppler sonography has been proved to accurately assess the degree of activity in inflammatory disease.

One well documented drawback of sonography is operator-dependency. The quality and consistency of ultrasound examinations relies on the expertise of the sonographer both in ultrasound techniques and knowledge of anatomy and pathology of a wide range of musculoskeletal areas.

Contraindications for diagnostic shoulder scans are unlikely; however, some limitations exist and may include the following:

- patient obesity
- inability to see structures that lie deep to bone or intra-articular structures of the glenohumeral and acromioclavicular joints
- casts, dressings, open wounds/ulcers etc. can limit visualisation
- severe oedema/swelling
- patients who are unable to co-operate or provide a clinical history due to reduced cognitive functions e.g. Alzheimer’s or dementia and through involuntary movements.
As age-related changes are common in the musculoskeletal system\textsuperscript{31,32} and may not be the cause of the patient’s symptoms, ultrasound appearances must always be taken in clinical context and the referrer should be made aware of limitations in the report. For example:

- ‘ultrasound cannot exclude impingement’
- ‘ultrasound cannot exclude intra-articular pathology.’

Diagnostic ultrasound is often used as a precursor to therapeutic injections and care should be taken to assist in the direction of that injection. Some structural changes seen on ultrasound may not be currently relevant and may not be associated with pain. For example, effusion in the long head of biceps tendon sheath does not always reflect current tenosynovitis but may be an extension of a glenohumeral joint effusion. The addition of the site of current symptom on a report may help. Subacromial/subdeltoid bursal thickening may be present on the asymptomatic shoulder and does not always reflect current ‘bursitis’ so a comparison image and addition of current symptoms may help.

**Common musculoskeletal pathologies:**

**Bursitis:** thickening and inflammation of a bursa, often causing an effusion. Thickening of the bursa alone does not always constitute ‘bursitis’. Comparison with the asymptomatic side is important. Effusion +/- hyperaemia of the bursal wall using colour/power Doppler is often present in symptomatic bursitis.

**Calcific tendinopathy:** calcification within tendons. The presentation of calcific tendinopathy may vary from tiny calcific flecks to large conglomerates which may appear ‘soft’ and non-shadowing, or denser with a dense posterior acoustic shadow.

**Cuff arthropathy:** arthritis of the glenohumeral joint, particularly the humeral head, due to massive rotator cuff tears leaving a ‘bare’ humeral head.

**Effusion:** a collection of fluid in a tendon sheath, joint or bursa.

**Joint synovitis:** thickening and often hyperaemia of the synovium lining a joint.

**Tendinitis:** use of this term is discouraged as this suggests an inflammatory process that has not been proven.

**Tendinopathy:** A more general term for a diseased tendon although thought to be a ‘normal continuum’ as a result of aging. On ultrasound, loss of the normal fibrillar pattern, heterogeneity of the tendon, abnormally thickened or thinned, hypoechoic or hyperechoic in comparison with the asymptomatic side. May show neovascularity within the tendon on colour/power Doppler

**Tendinosis:** a degenerative process within a tendon, disordered biomechanics often leading to micro tears. Tendinopathy is a more familiar term now.

**Tenosynovitis:** inflammation of the tendon and sheath. Appearances should include tendon sheath thickening and hyperaemia on Doppler, pain on palpation. May also include effusion and tendinopathy but these may be present without current tenosynovitis. Detailed anatomical knowledge of the presence/absence of tendon sheaths is required.
Tendon or ligament tears: When reporting, it is important to describe the type of tear—partial, full or complete—and the dimensions and site of the tear. An opinion on the state of the remaining tendon or ligament tissue is also helpful. These details are important for subsequent treatment and surgical planning as tendons with severe tendinopathy are less likely to have a successful repair should surgery become appropriate.

**Reporting:**

The report is a recording and interpretation of observations made during the ultrasound examination. It should be written by the person undertaking the scan and viewed in clinical context.

The report is intended to answer a clinical question and to assist with the patient journey so local opinion from radiologists, rheumatologists, orthopaedic surgeons and physiotherapists should be taken into account in the construction of local report templates.

The standard report should include:

- documentation of the normal anatomy;
- details of any pathology including measurements and any increase in vascularity if appropriate;
- documentation of any limitation to range of movement and site and degree that pain or symptoms begin;
- any difficulties with interpretation of the ultrasound appearances.

Comparison with the contralateral side (assuming it is asymptomatic) will help when determining the clinical significance of age/activity related changes and should be imaged and documented in the report.

The pitfalls of ultrasound interpretation are widely documented and can be reduced by education and experience of the individual ultrasound practitioner.

**Examination specific guidelines and common clinical scenarios.**

4.12.1 Shoulder

**Purpose of scan**

To evaluate the following structures:

- main rotator cuff tendons – supraspinatus, infraspinatus and subscapularis tendons
- long head of biceps tendon
- posterior glenohumeral joint
- superficial acromioclavicular joint.
Common indications include:

- rotator cuff tear/tendinopathy
- restricted range of movement
- pain on abduction.

Scan protocol:

Numerous scan protocols for the shoulder are described in the literature including that from the European Society of Musculoskeletal Radiology’s (ESSR) protocol for the shoulder. Training protocols also exist and may be useful to standardise scanning and reporting such as that described by Smith et al.

The structures that should be identified as a minimum in a standard shoulder ultrasound scan are:

- long head of biceps tendon (LHBT) – for position, tendinopathy, tears, calcification, tenosynovitis, effusion;
- rotator interval (RI) – if LHBT is visible at the (RI) it is likely to be intact proximally;
- subscapularis tendon – for tendinopathy, tears, calcification;
- infraspinatus tendon – for tendinopathy, tears, calcification;
- supraspinatus tendon – for tendinopathy, tears, calcification;
- any visible bursae around the shoulder – subacromial/subdeltoid, subcoracoid;
- posterior glenohumeral joint – for effusion, cysts, joint synovitis;
- acromioclavicular joint – for effusion, cysts, joint synovitis.

Comparison with the contra-lateral side (assuming it is asymptomatic) will help when determining the clinical significance of age/activity-related changes and should be imaged and documented in the report.

Dynamic ultrasound assessment

Dynamic assessment around the shoulder may be controversial in some centres as some indications such as ‘impingement’ have multiple potential causes (some of which will not be seen using ultrasound) and may be considered to be a purely clinical diagnosis.

Dynamic assessment using ultrasound may be requested for the following:

- long head of biceps tendon: to assess its stability within the bicipital groove during external rotation;
- subscapularis tendon: to identify the myotendinous area which normally sits behind the coracoid process of the scapula;
- supraspinatus tendon: to assess for bunching of the tendon and/or overlying subacromial bursa against the acromion or coraco-acromial ligament during abduction which may be a cause of ‘impingement’;
- posterior joint recess: during internal/external rotation, to assess for a gleno-humeral joint effusion. This will be most evident during external rotation.
Imaging protocol

A standard shoulder series should include the following minimum images for a normal scan:

- long head of biceps tendon - longitudinal and transverse;
- rotator interval showing anterior portion of supraspinatus tendon, long head of biceps and lateral edge of subscapularis tendon;
- subscapularis tendon – longitudinal and transverse;
- supraspinatus tendon – longitudinal and transverse;
- infraspinatus tendon – longitudinal only;
- posterior glenohumeral joint recess;
- acromioclavicular joint.

Document the normal anatomy and any pathology found, including measurements and vascularity if indicated.

Common pathologies would include tendinopathy, bursitis and tendon tears.

Tendon tears

Rotator cuff tendon tears are often seen in asymptomatic individuals and are not always the cause of symptoms. It is sometimes difficult for the ultrasound practitioner to appreciate the significance of a tear on the patient’s current clinical symptoms and while it is important to evaluate and accurately describe tears, care should be taken if discussing scan findings with the patient.

When reporting, it is important to describe the type of tear: partial, full or complete and the dimensions and site of the tear. An opinion on the state of the remaining tendon tissue is also helpful. These details are important for subsequent treatment and surgical planning as tendons with severe tendinopathy are less likely to have a successful repair should surgery become appropriate.

Measurements

Type, size and location of tears is important, and measurements of full thickness tears should be made in two planes – anterior to posterior (transverse) and medial to lateral (longitudinal). The site of the tear measured from the rotator interval is also useful.

Site of supraspinatus tendon tears

The supraspinatus tendon can be divided into anterior, mid and posterior portions. The anterior free or leading edge, the mid-substance or footprint and the area that abuts the infraspinatus tendon - the distinction between the two tendons can be difficult to define.

Anterior leading edge tears involve the portion of the supraspinatus tendon that lies adjacent to the long head of biceps tendon at the rotator interval.

If there is still tendon tissue anterior to the tear, the tear is said to be mid-substance, crescent or footprint.
If the tear occurs at the supraspinatus/infraspinatus interface it is said to be posterior.
Types of tear

Partial thickness tear: A rotator cuff tendon tear that involves either the joint or bursal surface of a tendon and does not allow communication between the two compartments. Partial thickness tears may be articular surface, bursal surface or intrasubstance.

Under-surface or articular surface tear: A partial thickness rotator cuff tear involving the articular or joint surface of the tendon. In the supraspinatus, they can be called ‘rim rent’ or PASTA lesion – Partial Articular Supraspinatus Tendon Avulsion.

Bursal surface tear: A partial thickness rotator cuff tear involving the bursal surface of the tendon. In the supraspinatus tendon, this is sometimes called ‘scuffing’ as it is caused by friction from the under-surface of the acromion.

Intrasubstance tear: A partial thickness rotator cuff tear within the substance of the tendon – sometimes called concealed interstitial delamination (CID). A cleft is present within the tendon substance, but it does not reach either the bursal or articular surfaces of the tendon.

Full thickness tear: A rotator cuff tendon tear that extends across the whole depth of the tendon, involves both surfaces and resulting in communication between the joint and bursal compartments, regardless of the width of the tear. Some full thickness tears measure over 3 cm in diameter; some look like pinholes.

Complete full thickness tear: A full thickness tendon tear that extends across the whole width and depth of the tendon, usually resulting in retraction back from the insertion.

If the tear is complete, the entire width of the tendon will have torn, usually (but not always) at the insertion and there is retraction of the proximal stump. The degree of retraction will vary but, in the supraspinatus and subscapularis tendons, can result in the proximal stump sitting under the acromion/coracoid and so not visible with ultrasound. This is an important finding for surgeons because it is less likely that the tendon can be pulled back and repaired.

A supraspinatus tendon tear is often given an increased level of importance compared to the other rotator cuff tendons as it is more often symptomatic and amenable to surgery.

14.12.1.1 Reporting examples

The sample reports below are intended as a guide only as reporting style may be specific to individuals/departments.

Sample report for normal appearances

Clinical details: shoulder pain, ?rotator cuff tear

Ultrasound of the shoulder performed with verbal consent.

Normal appearances of the rotator cuff tendons. No evidence of tears or calcification or any bursal thickening or effusions. The long head of biceps tendon is intact and in situ.

No evidence of a gleno-humeral joint effusion.

Unremarkable ACJ noted.
Conclusion: Normal examination. No rotator cuff tear seen.

If appropriate, the report may include the following:

- the rotator cuff tendons have reasonable depth and texture;
- there is evidence of subacromial/subdeltoid bursal thickening but no more so than on the asymptomatic shoulder;
- there is good, pain free subacromial movement;
- non-tender ACJ OA noted.

Referral for tendinopathy

Ultrasound of the shoulder performed with verbal consent.

The supraspinatus tendon appears generally thickened/thinned and tendinopathic with loss of the normal fibrillar pattern. No tears seen.

Conclusion: Focal tendinopathic changes are noted at the anterior/middle/posterior portion of the supraspinatus tendon, no tears seen.

Referral for calcific tendinopathy

Ultrasound of the shoulder performed with verbal consent.

A x mm intratendinous calcium deposit is (calcific flecks are) noted within the ant/mid/post aspect of the supraspinatus tendon; no tears evident.

There is a 6mm densely shadowing calcific deposit within the supraspinatus tendon.

There is a 6mm non-shadowing deposit within the supraspinatus tendon likely to be soft calcific tendinopathy.

Conclusion: There is marked calcific tendinopathy of the rotator cuff tendons. The largest area of calcification in the supraspinatus tendon measures 15mm in diameter and the patient is tender on scanning.

Referral for bursitis

Ultrasound of the shoulder performed with verbal consent.

The subacromial/subdeltoid bursa contains an effusion and the wall is hyperaemic on power Doppler. The patient is tender to scan here and appearances are consistent with bursitis.

Referral for long head of biceps tendon

Ultrasound of the shoulder performed with verbal consent.

The long head of biceps tendon is intact but has dislocated from the bicipital groove medially and is sitting on the surface of the subscapularis tendon.
The long head of biceps tendon has subluxed onto the medial lip of the bicipital groove. The tendon is thickened and appears oedematous. The tendon sheath is hyperaemic and contains an effusion and the patient is tender to scan here. Ultrasound appearances suggest an intact subluxed long head of biceps tendon with evidence of tenosynovitis.

Referral for tendon tear

Partial thickness tear

Ultrasound of the shoulder performed with verbal consent.

There is a partial thickness tear of the bursal (or articular) surface (or intrasubstance) of the anterior aspect of the supraspinatus tendon which measures 2mm in longitudinal section and 3 mm in transverse section.

The supraspinatus tendon appears generally tendinopathic and there is a partial thickness (articular) (bursal) surface tear within the anterior/middle/posterior portion of this tendon. This tear measures XXmm x XXmm and affects over (or under) 50% of the tendon depth.

Conclusion: There is a partial thickness tear on the articular surface of the supraspinatus tendon lying Xmm posterior to the rotator interval. It measures XXmm in width and extends over (or under) 50% of the tendon depth. The remainder of the tendon has reasonable depth and texture.

Full thickness tear

Ultrasound of the shoulder performed with verbal consent.

There is a full thickness, insertional tear of the supraspinatus tendon lying XXmm posterior to the rotator interval. The tear measures XXmm in width and there is XXmm retraction from the insertion. The remainder of the tendon has reasonable depth and texture.

Complete full thickness tear

Ultrasound of the shoulder performed with verbal consent.

The supraspinatus tendon has completely ruptured and retracted; the retracted tendon end is not visualised. Cuff arthropathy noted.

Conclusion: There is a complete, full thickness tear of the supraspinatus tendon with proximal retraction of x mm from the insertion.

References:

4.12.2 Elbow

Purpose of scan

To evaluate any of the following structures where appropriate and symptomatic:

- common extensor and flexor tendons;
- medial ulnar collateral ligament;
- lateral radial collateral ligament;
- distal biceps tendon;
- triceps tendon;
- cubital tunnel and ulnar nerve;
- elbow joints/olecranon fossa.

Common indications

Common indications for the performance of an elbow ultrasound scan are:

- ‘tennis’ or ‘golfers’ elbow/epicondylitis;
- restricted range of movement? effusion/synovitis;
- distal biceps tendon tear;
- ulnar nerve compression;
- olecranon bursitis.

Scan protocol


Unlike the shoulder, diagnostic scanning of the elbow is usually focussed on a single structure, for example the common extensor tendon for ‘tennis elbow’, but other structures may be included if appropriate and symptomatic and within the clinical experience of the ultrasound practitioner.

According to Draghi et al (2007) “Ultrasound of the elbow: Examination techniques and US appearance of the normal and pathologic joint” these include:

- common extensor tendon - for tendinopathy, neovascularity, tears, calcification;
- common flexor tendon - for tendinopathy, neovascularity tears, calcification;
- distal biceps tendon - for tendinopathy, tears, calcification;
- olecranon bursa - for effusion/bursitis;
- anterior and posterior joint recesses - for effusion, joint synovitis;
- cubital tunnel - for ulnar nerve enlargement/flattening/subluxation.

Dynamic ultrasound assessment

Dynamic assessment using ultrasound may be requested for the following but depends on the skill and expertise of the ultrasound practitioner:

- ulnar nerve subluxation
- medial/lateral ligament tears
Imaging protocol

There is no standard elbow series as the areas scanned are dependent on symptoms and pathology found. The European Society of MusculoSkeletal Radiology guidelines ‘Musculoskeletal ultrasound technical guidelines II. Elbow’ provide further information on technique. The following protocols are a guide to each area:

- CEO (common extensor origin) / CFO (common flexor origin) longitudinal with colour box to show the absence of neovascularity;
- distal biceps tendon in longitudinal and transverse, distal tendon insertion in longitudinal and myotendinous area in transverse to exclude tear/ tendinopathy /calcification. Comparison with contralateral side;
- anterior and posterior joint recess within and without colour box to show the absence of effusion or synovitis;
- cubital tunnel with arm in flexion/extension to exclude subluxation of the ulnar nerve;
- medial/lateral ligaments in longitudinal in stressed and relaxed positions if appropriate.

Document the normal anatomy and any pathology found, including measurements and vascularity if indicated.

14.12.2.1 Reporting examples

The sample reports below are intended as a guide only as reporting style may be specific to individuals/departments.

Sample report for normal appearances

Ultrasound of the elbow with verbal consent

Normal appearances of the common extensor/flexor/distal biceps tendons of the elbow. No evidence of significant tendinopathy, tears or calcification.

No evidence of joint effusion or synovitis but ultrasound cannot exclude intra-articular pathology. The ulnar nerve appears normal in calibre around the elbow and is stable within the cubital tunnel on dynamic scanning’. No solid or cystic lesions seen in the cubital tunnel.

Conclusion: Normal examination. No cause for X [symptoms] seen.

Referral for tendinopathy

Ultrasound of the elbow with verbal consent

The tendon appears generally thickened/thinned with loss of the normal fibrillar pattern. There is evidence of neovascularity on Doppler and appearances are consistent with tendinopathy. No tears seen.

Conclusion: Focal tendinopathic changes are noted at the proximal/distal portion of the tendon, no tears seen.
Referral for calcific tendinopathy

Ultrasound of the elbow with verbal consent

A x mm intratendinous calcium deposit is (calcific flecks are) noted within the proximal/distal portion of the tendon, no tears evident.

There is a 6mm densely shadowing calcific deposit within the tendon.

Conclusion: There is a 6mm non-shadowing deposit within the tendon likely to be soft calcific tendinopathy.

Referral for olecranon bursitis

Ultrasound of the elbow with verbal consent

The olecranon bursa contains an effusion and the wall is hyperaemic on power Doppler. The patient is tender to scan here and appearances are consistent with bursitis.

Referral for tendon tears

When reporting, it is important to describe the type of tear – partial or complete – and the dimensions and site of the tear. An opinion on the state of the remaining tendon tissue is also helpful.

There is a well-defined, cystic area within the proximal portion of the tendon measuring XXmm in diameter and extending across approximately XX mm of the tendon depth. Appearances are consistent with cystic degeneration/partial thickness tear.

There is a complete insertional tear of the distal biceps tendon with retraction of XXmm from the insertion. There is loss of the normal fibrillar pattern in the detached tendon suggesting tendinopathy.

There is a large fluid collection around the distal pole of the biceps muscle likely to be a resolving haematoma. The distal biceps tendon is intact at the insertion but appears to have completely torn at the myotendinous junction.

4.12.3 Wrist and hand

Purpose of scan

To evaluate any of the following structures where appropriate and symptomatic:

- extensor and flexor tendons;
- joints of the hand/wrist and metacarpophalangeal (MCP), proximal interphalangeal (PIP), distal interphalangeal (DIP) and carpometacarpal (CMC) joints;
- ligaments of the thumb/finger joints;
- carpal tunnel;
- Guyon’s canal.

Common indications
Common indications for the performance of a wrist/hand ultrasound scan are:

- Swelling ?ganglion
- ? effusion/synovitis
- ? tendon tear

**Scan protocol**

Numerous scan protocols for the hand and wrist are described in the literature, including the European Society of MusculoSkeletal Radiology guidelines “Musculoskeletal ultrasound technical guidelines III. Wrist” and McNally (2008) “Ultrasound of the small joints of the hands and feet: current status”.

Unlike the shoulder, diagnostic scanning of the hand/wrist is usually focussed to a single area or pathology. For example, the dorsum of the wrist for ‘?ganglion’ or the joints of the hand and wrist for ‘? synovitis’. Other structures may be included if appropriate and symptomatic and within the clinical experience of the ultrasound practitioner. These include:

- extensor and flexor tendons for tendinopathy, tenosynovitis, tears, calcification;
- finger pulleys for triggering;
- joints of the hand/wrist for synovitis or effusion;
- bony cortex for erosions;
- ligaments of the thumb/fingers for sprains or tears;
- carpal or Guyon’s tunnel for nerve compression.

Comparison with the contralateral side (assuming it is asymptomatic) will help when determining the clinical significance of age/activity-related changes and should be imaged and documented in the report.

**Imaging protocol**

There is no standard wrist and hand series as the areas scanned are dependent on symptoms and pathology found. The European Society of MusculoSkeletal Radiology guidelines ‘Musculoskeletal ultrasound technical guidelines III. Wrist’ provide further information on technique. The following protocols are a guide to each area:

- tendons and sheaths - longitudinal with colour box to show the absence of neovascularity;
- tendons in longitudinal and transverse, distal tendon insertion in longitudinal to exclude tear/ tendinopathy /tenosynovitis/calcification. Comparison with contralateral side;
- dorsal and volar wrist joints with and without colour box to show the absence of effusion or synovitis;
- finger ligaments in longitudinal, in stressed and relaxed positions if appropriate.

Document the normal anatomy and any pathology found, including measurements and vascularity if indicated.
14.12.3.1 Reporting examples

The sample reports below are intended as a guide only as reporting style may be specific to individuals/departments.

Sample report for normal appearances

Ultrasound of the wrist/hand with verbal consent

Normal appearances of compartment one of the extensor tendons. The extensor pollicis brevis and abductor pollicis longus tendons appear normal. No evidence of De Quervain’s tenosynovitis.

No evidence of active synovitis or effusion seen arising from the wrists or within the MCP or PIPJs. No tenosynovitis. No erosions seen.

The ulnar nerve appears normal in calibre through Guyon’s tunnel and is symmetrical with the asymptomatic side. No evidence of compression or a solid or cystic lesion in the tunnel.

Referral for tendinopathy/tenosynovitis

Ultrasound of the wrist/hand with verbal consent

The tendon appears generally thickened/thinned with loss of the normal fibrillar pattern consistent with tendinopathy. No tears seen.

Conclusion: There is effusion and hyperaemia of the flexor tendon sheath consistent with tenosynovitis. No evidence of a tendon tear.

Referral for joint synovitis

These scans are usually limited to rheumatology patients and there are several grading systems in use. It is important that any grading system is discussed with rheumatology before being used.

Ultrasound of the wrist/hand with verbal consent

There is a moderate degree of active synovitis arising from the right wrist and within the right and left index and middle finger MCPJs. No evidence of active synovitis seen in the left wrist or the remainder of MCP or PIPJs. No tenosynovitis.

Referral for tendon tears

When reporting, it is important to describe the type of tear – partial or complete – and the dimensions and site of the tear. An opinion on the state of the remaining tendon tissue is also helpful.

Ultrasound of the wrist/hand with verbal consent

There is a complete full thickness tear of the middle finger flexor digitorum profundus tendon at the distal insertion. The tendon has retracted back to the level of the PIP joint and appears thin and tendinopathic. The flexor digitorum superficialis tendon remains intact and appears normal.
4.12.4 Hip

Purpose of scan

To evaluate any of the following structures where appropriate and symptomatic:

- anterior hip joint;
- proximal hamstrings;
- distal psoas tendon;
- adductor muscle/tendons.

Common indications

Common indications for the performance of a hip ultrasound scan are:

- ? effusion/synovitis
- ? hamstring tear
- ? enthesopathy

Scan protocol

Hip scanning protocols are described in the literature, for example The European Society of MusculoSkeletal Radiology guidelines ‘Musculoskeletal ultrasound technical guidelines IV. Hip’. Unlike the shoulder, diagnostic scanning of the hip is usually focussed to a single area or pathology. For example the anterior hip joint for ?effusion.

Other structures may be included if appropriate and symptomatic and within the clinical experience of the ultrasound practitioner. These include:

- anterior hip joint for synovitis or effusion;
- anterior hip/psoas tendon for tendinopathy, bursitis;
- greater trochanteric area for bursitis, gluteal enthesopathy, tendinopathy /tears/calcification;
- adductor tendons for enthesopathy, tears, tendinopathy, calcification;
- anterior thigh tendons for enthesopathy, tears, tendinopathy, calcification;
- posterior hamstring tendons enthesopathy, tears, tendinopathy, calcification.

Comparison with the contralateral side (assuming it is asymptomatic) will help when determining the clinical significance of age/activity related changes and should be imaged and documented in the report.

Imaging protocol

There is no standard hip imaging series as the areas scanned are dependent on symptoms and pathology found. The European Society of MusculoSkeletal Radiology guidelines ‘Musculoskeletal ultrasound technical guidelines IV. Hip’ provide further information on technique. The following protocols are a guide to each area:

- anterior hip joint longitudinal with and without colour box to show no effusion or synovitis (ensure Doppler settings capable of detecting deep flow);
• anterior hip/psoas tendon in longitudinal and transverse to show normal tendon with no bursal effusion;
• greater trochanteric area in longitudinal and transverse to show normal tendons with no bursal effusion;
• adductor tendons in longitudinal to show no tears;
• anterior thigh tendon insertions in longitudinal and transverse;
• posterior hamstring tendons in longitudinal and transverse.

Document the normal anatomy and any pathology found, including measurements and vascularity if indicated.

14.12.4.1 Reporting examples

The sample reports below are intended as a guide only as reporting style may be specific to individuals/departments.

---

Sample report for normal appearances

Ultrasound of the hip with verbal consent

*No evidence of a joint effusion or synovitis seen arising from the anterior hip joint although intra-articular pathology cannot be excluded with ultrasound.*

*Normal appearances of the distal iliopsoas tendon with no evidence of bursitis.*

*Normal appearances of the posterior hamstring origin at the ischial tuberosity.*

---

Referral for tendinopathy

Ultrasound of the hip with verbal consent

*The gluteus medius tendon appears generally thickened/thinned with loss of the normal fibrillar pattern consistent with tendinopathy. No tears seen. No evidence of a trochanteric bursal effusion.*

---

Referral for joint effusion

Ultrasound of the hip with verbal consent

*There is an effusion in the anterior hip joint capsule measuring XXX mm in depth compared to X mm on the asymptomatic side. The effusion does not appear to be complex but infection cannot be excluded.*

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Further information can be found at:

4.12.5 Knee

Purpose of scan

To evaluate any of the following structures where appropriate and symptomatic:

- quadriceps and patellar tendons;
- supra-patellar, pre-patellar, superficial and deep infra-patellar bursae;
- medial and lateral collateral ligaments;
- pes anserine tendons and bursa;
- iliotibial band;
- biceps femoris insertion;
- popliteal fossa;
- semimembranosus/medial gastrocnemius tendons/bursa.

Common indications

Common indications for the performance of a knee ultrasound scan are:

- effusion/synovitis;
- bursitis;
- popliteal cyst;
- quadriceps/patellar tendon tear.

Scan protocol

Knee scanning protocols are described in the literature, for example The European Society of MusculoSkeletal Radiology guidelines ‘Musculoskeletal ultrasound technical guidelines V. Knee’. Unlike the shoulder, diagnostic scanning of the knee is usually focused to a single area or pathology. For example, the popliteal fossa for Baker’s cyst.

Other structures may be included if appropriate and symptomatic and within the clinical experience of the ultrasound practitioner. These include:

- anterior knee for synovitis or effusion in the suprapatellar, pre patellar or infrapatellar bursae;
- popliteal fossa for Baker’s cyst or to exclude a popliteal artery aneurysm;
- extensor tendons or distal hamstrings for tendinopathy/tears/calcification/enthesopathy;
- medial or lateral collateral ligaments for strains, tears or calcification.

Comparison with the contralateral side (assuming it is asymptomatic) will help when determining the clinical significance of age/activity-related changes and should be imaged and documented in the report.

Imaging protocol

There is no standard knee imaging series as the areas scanned are dependent on symptoms and pathology found. The following protocols are a guide to each area:
• anterior knee longitudinal images with and without colour box to show no effusion or synovitis in the suprapatellar, prepatellar or infrapatellar bursae (ensure Doppler settings capable of detecting flow in deep structures);
• longitudinal and transverse images of quadriceps and patellar tendons to show normal tendons and their insertions;
• longitudinal images of medial and lateral collateral ligaments;
• longitudinal images of popliteal fossa to show posterior joint.

Document the normal anatomy and any pathology found, including measurements and vascularity if indicated.

4.12.5.1 Reporting examples

The sample reports below are intended as a guide only as reporting style may be specific to individuals/departments.

---

**Sample report for normal appearances**

No evidence of effusion or synovitis seen within the suprapatellar, prepatellar or infrapatellar bursae although intra-articular pathology cannot be excluded with ultrasound.

No evidence of a solid or cystic lesion seen in the popliteal fossa. No Baker’s cyst. The popliteal artery is of normal calibre.

Normal appearances of the quadriceps and patellar tendons. Normal fibrillar pattern. No evidence of tendon tears, calcification or enthesitis.

---

**Referral for quadriceps tendon tear**

Ultrasound of the knee with verbal consent

There is a complete tear of the quadriceps tendon at its insertion onto the superior pole of the patellar. With the knee joint extended, the tendon has retracted proximally by xx mm. The patellar tendon is intact.

---

**Referral for knee effusion**

Ultrasound of the knee with verbal consent

There is a small/medium/large effusion in the suprapatellar bursa with evidence of synovitis. The effusion does not appear to be complex.

---

**Referral for Bakers cyst/semimembranosus**

Ultrasound of the knee with verbal consent

There is a well-defined, simple cystic lesion in the medial popliteal fossa measuring XX mm. It lies between the medial head of gastrocnemius and the semimembranosus tendon, measured at 5.6 cm in maximum axial diameter. The site and appearances of this lesion are consistent with a Baker’s cyst.

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**Reference:**

4.12.6 Foot and Ankle

Purpose of scan

To evaluate any of the following structures where appropriate and symptomatic:

- anterior ankle joint;
- medial ankle – posterior tibial, flexor digitorum longus and flexor hallucis longus tendons and neurovascular bundle
- lateral ankle tendons – peroneus brevis and longus;
- anterior tendons – anterior tibial extensor hallucis longus and extensor digitorum longus tendons;
- Achilles tendon and insertion, retrocalcaneal and pre-Achilles bursa;
- plantar fascia;
- dorsal surface of tarsal joints;
- MTP joints;
- interdigital spaces;
- some ligaments – anterior talofibular, anterior tibiofibular, calcaneofibular, deltoid.

Common indications

Common indications for the performance of a foot/ankle ultrasound scan are:

- Morton’s neuroma;
- medial ankle pain. ?posterior tibial rupture/dysfunction;

Scan protocol

Numerous scan protocols for the foot and ankle are described in the literature for example The European Society of MusculoSkeletal Radiology guidelines ‘Musculoskeletal ultrasound technical guidelines VI. Ankle’. Unlike the shoulder, diagnostic scanning of the foot/ankle is usually focussed to a single area or pathology. For example, interdigital spaces for Morton’s neuroma. Other structures may be included if appropriate and symptomatic and within the clinical experience of the ultrasound practitioner. These include:

- extensor and flexor tendons for tendinopathy, tenosynovitis, tears, calcification;
- joints of the foot/ankle for synovitis or effusion;
- ligaments of the ankle for sprains or tears;
- tarsal tunnel for nerve compression;
- interdigital spaces 2/3 and 3/4 for Morton’s neuromata;
- plantar fascia for fasciopathy;
- Achilles tendon for tendinopathy, enthesis, enthesopathy, tears, calcification.

Comparison with the contralateral side (assuming it is asymptomatic) will help when determining the clinical significance of age/activity-related changes and should be imaged and documented in the report.
Imaging protocol

There is no standard foot and ankle series as the areas scanned are dependent on symptoms and pathology found. The following protocols are a guide to each area:

- Tendons and sheaths/paratendon - longitudinal with colour box to show the absence of neovascularity;
- Tendons in longitudinal and transverse, distal tendon insertion in longitudinal to exclude tear/tendinopathy/tenosynovitis/calcification. Comparison with contralateral side;
- Anterior ankle recess or MTP joints with and without colour box to show the absence of effusion or synovitis;
- Ankle ligaments in longitudinal in stressed and relaxed positions if appropriate.

Document the normal anatomy and any pathology found, including measurements and vascularity if indicated.

4.12.6.1 Reporting examples

The sample reports below are intended as a guide only as reporting style may be specific to individuals/departments.

Sample report for normal appearances

Normal appearances of the posterior tibial tendon. Good depth and normal fibrillary pattern with no evidence of tears significant tendinopathy or intratendinous calcification. No evidence of tenosynovitis.

No evidence of active synovitis or effusion seen arising from the anterior ankle joint. The anterior talofibular, calcaneofibular and tibiofibular ligaments appear intact but intra-articular pathology cannot be excluded.

Referral for tendinopathy/tenosynovitis

Ultrasound of the ankle with verbal consent

There is a moderate fusiform thickening (AP diameter measured at 7mm) of the Achilles tendon with loss of the normal fibrillar pattern and a moderate degree of neovascularity within the middle third. Normal appearances of the proximal and distal thirds of the tendon. No evidence of enthesitis or bursitis.

Conclusion: Appearances are consistent with a moderate middle third Achilles tendinopathy. No signs of tear identified.

Ultrasound of the foot with verbal consent

The posterior tibial tendon appears generally thickened/thinned with loss of the normal fibrillar pattern, but no tear seen. There is effusion and hyperaemia within the tendon sheath and appearances are consistent with posterior tibial tenosynovitis.

Conclusion: Posterior tibial tenosynovitis.
Note: there is no clear guidance on the best classification system for tendinopathy. Based on expert experience, fusiform thickening can be categorised into mild/moderate/severe. Generally mild 4 – 6 mm, moderate 7 - 10 mm and severe anything above 10 mm.

Referral for Morton’s neuroma/interdigital bursitis

Ultrasound of the foot with verbal consent

There is a well-defined hypoechoic, but solid, lesion in the left 2nd web space (i.e. between 2nd and 3rd metatarsal heads). No evidence of vascularity within the lesion on Doppler. It measures XX mm in transverse diameter (full measurements mm x mm x mm) and appearances are consistent with a Morton’s neuroma which is symptomatic on scanning. The other web spaces look normal.

Conclusion: Morton’s neuroma

There are large, ovoid, partially compressible lesions in the 2nd and 3rd web spaces of the right foot. They measure xx mm in transverse (full measurements mm x mm x mm), show increased peripheral vascularity and are symptomatic on scanning. Ultrasound appearances suggest second web space interdigital bursitis, but an adjacent Morton’s neuroma cannot be excluded.

Note: absolutely clarity is required in relation to location, e.g. avoid the term 2/3 web space: state ‘between 2nd and 3rd web spaces’.

Referral for tendon tears

When reporting, it is important to describe the type of tear – partial or complete – and the dimensions and site of the tear. An opinion on the state of the remaining tendon tissue is also helpful.

There is a complete full thickness tear of the posterior tibial tendon at the level of the medial malleolus. The tendon ends have retracted by 6 mm and the remaining visible tendon appears thin and tendinopathic.

There is a complete tear of the Achilles tendon lying 34 mm proximal to the upper border of the calcaneum. With the ankle in neutral, there is retraction of the proximal stump by approximately 24 mm. With the ankle in dorsiflexion, the tendon ends do not oppose – there is herniation of Kager’s fat pad in between the tendon ends. The gap in dorsiflexion is measured at 14 mm. There is relatively normal fibrillar pattern of the distal stump of tendon but the proximal stump appears severely tendinopathic.

Referral for plantar fasciitis/fasciopathy

The proximal plantar fascia is hypoechoic with loss of the normal fibrillar pattern. It is thickened to xx mm (over 4 - 4.3 mm) and is tender on scanning.

Conclusion:Appearances are consistent with plantar fasciitis.
4.12.7 Rheumatology ultrasound examinations

Examination specific guidelines and common clinical scenarios for inflammatory arthritis.

These guidelines are aimed at those scanning patients for ‘? Inflammatory arthritis’ who are not directly involved in a Rheumatology service – i.e. sonographers scanning within Radiology departments.

Rheumatologists often have local guidelines for their own scanning of these patients.

Purpose of an ultrasound scan

To evaluate the following structures:

Hands and wrists, feet and ankles for:
- synovial hypertrophy – synovial proliferation
- synovitis – synovial hypertrophy with vascularity on Doppler
- erosions
- tenosynovitis – inflammation of the tendon and sheath
- enthesitis

If these scans are carried out within a rheumatology department setting, they will be focussed to exclude or confirm evidence of inflammatory arthritis, not necessarily to determine the cause of hand or foot pain. This should be made clear on the request form and the report should include the clinical question to be answered.

Inflammatory arthritis

The diagnosis of any disease usually progresses along a well-defined path that has three parts: a history of the complaint, blood tests and, usually, imaging (x-rays or scans).

"Seropositive/seronegative" is a term that refers to the results of a blood test to help establish the diagnosis of rheumatoid arthritis (RA). This looks for the presence of two proteins in the blood37.
Rheumatoid factor (RF) is a very old but tried and tested investigation that was first introduced into rheumatology in the 1940s. Anti-cyclic citrullinated peptide (Anti-CCP) antibody testing is more recent, is more sensitive than RF and may appear much earlier in the course of RA.

A positive RF or anti-CCP test does not mean that the patient has RA. Other features must be present such as symptoms of pain and swelling in the joints, involvement of many joints with inflammation, morning stiffness in the joints for longer than 45 minutes, x-ray evidence of the characteristic bone damage in the joints and extra-articular features of RA (meaning features that are outside the joints), such as nodules. Other blood tests often used prior to diagnosis include ESR and CRP, which measure the amount of inflammation in the joints.

**Common indications**

Common indications for the performance of a Rheumatology ultrasound scan are:

- joint pain and swelling. ?inflammatory arthritis
- RA on disease modifying anti-rheumatic drugs (DMARDS). ?active synovitis
- RA in remission. ?subclinical synovitis

**Common pathologies**

**Joint effusion:** A joint effusion is a collection of fluid within a joint or tendon sheath. It is typically anechoic, compressible and does not display any internal Doppler signal. Small effusions are commonly seen in normal joints but if they become large, or are associated with other inflammatory appearances, they may be a sign of acute disease.

**Synovial hypertrophy:** Synovial proliferation and resulting hypertrophy is the primary event in rheumatoid arthritis that is visible on imaging. The recognised definition of synovial hypertrophy on ultrasound is ‘Abnormal hypoechoic (relative to subdermal fat, but sometimes may be isoechoic or hyperechoic) intraarticular tissue that is non-displaceable and poorly compressible’.

However, the significance of the presence of synovial hypertrophy in joints without internal vascularity on Doppler is uncertain as it may be present in both inflammatory arthritis and osteoarthritis.

**Active synovitis:** When inflamed, synovial hypertrophy becomes active synovitis and is a sign of active inflammatory arthritis. It frequently affects the wrists, MCP and PIP joints of the hands and ankles and MTP joints of the feet.

The recognised definition of synovitis on ultrasound is:

‘Synovial hypertrophy which may exhibit Doppler signal’.

The degree of synovial hypertrophy and vascularity within joints or tendons may be estimated and documented either using an agreed grading system or ‘mild/moderate/severe’.
There are several grading systems for the degree of vascularity/grade of active synovitis. The most commonly used grades both synovial hypertrophy and active synovitis. Any pathology found should be documented in two planes. If a Doppler signal is seen within a joint in the longitudinal plane, its site must be confirmed within the joint in the transverse plane.

Comparison with the contra-lateral side (assuming it is asymptomatic) will help when determining the clinical significance of age/activity-related changes and should be imaged and documented in the report.

**Erosions:** RA is characterised by a chronic inflammatory process that targets the synovial lining of some joints. As the disease advances, there is evidence of progressive destruction of the structural components of the joints which targets the articular cartilage and bone at the joint margins.

The recognised definition of an erosion on ultrasound is: ‘An intra-articular discontinuity of the bone surface that is visible in two perpendicular planes.

The aim of modern treatments is to halt soft tissue inflammation and prevent or arrest the development of adjacent bone erosions and joint damage. Historically, radiographs have been used to detect and monitor erosions but ultrasound is now proven to detect them earlier but there are areas of most joints that are difficult to access with the transducer making erosions difficult to exclude. Research studies suggest that detection of flow within erosions on Doppler is suggestive of active bone destruction and should be highlighted to the referrer.

**Tenosynovitis:** Tenosynovitis is inflammation of the lining of the sheath that surrounds a tendon. Causes of tenosynovitis include infection, overuse or injury but tenosynovitis can also be associated with some kinds of inflammatory arthritis.

In the normal state, the sheath of the tendon can barely be detected and is seen as a thin, hypoechoic band around the tendon.

Once inflamed, the sheath becomes increasing hypoechoic, thickened and may display internal vascularity on Doppler.

The recognised definition of tenosynovitis on ultrasound is: ‘Hypoechoic or anechoic thickened tissue with or without fluid within the tendon sheath, which is seen in two perpendicular planes and which may exhibit a Doppler signal’.
Note: Not all tendons are surrounded by a sheath and the ultrasound appearances of inflammation vary, for example the extensor tendons of the fingers, over the MCP joints. Inflamed tendons without sheaths may have more diffuse peripheral oedema because inflammatory fluid is not contained. Instead of ‘tenosynovitis’, these tendons show evidence of ‘Para tendinitis’.

**Spondyloarthropathies:** Spondyloarthropathies are a group of inflammatory diseases in patients whose blood tests are negative for RA. These diseases include ankylosing spondylitis, reactive arthritis, enteropathic arthritis and psoriatic arthritis (PsA).

Common features of seronegative disease are listed in the table below.

<table>
<thead>
<tr>
<th>Features</th>
<th>Ankylosing spondylitis</th>
<th>Reactive arthritis</th>
<th>Psoriatic arthritis</th>
<th>Enteropathic arthritis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age of onset</td>
<td>20-30 years</td>
<td>20-30 years</td>
<td>35-45 years</td>
<td>Any age</td>
</tr>
<tr>
<td>Male: female ratio</td>
<td>3:1</td>
<td>5:1</td>
<td>1:1</td>
<td>1:1</td>
</tr>
<tr>
<td>Peripheral arthritis</td>
<td>Asymmetrical lower extremities</td>
<td>Asymmetrical lower extremities</td>
<td>Any joints</td>
<td>Asymmetrical lower extremities</td>
</tr>
<tr>
<td>Spine involvement</td>
<td>Symmetrical sacroiliitis (100%), delicate marginal syndesmophytes: lumbar spine and lower thoracic spine involved initially</td>
<td>Asymmetrical sacroiliitis, bulky marginal syndesmophytes</td>
<td>Asymmetrical sacroiliitis, bulky marginal syndesmophytes: cervical spine involvement most commonly</td>
<td>Symmetrical sacroiliitis, delicate marginal syndesmophytes</td>
</tr>
<tr>
<td>Enthesitis</td>
<td>Uncommon</td>
<td>Common</td>
<td>Common</td>
<td>Less common</td>
</tr>
<tr>
<td>Dactylitis</td>
<td>Uncommon</td>
<td>Common</td>
<td>Common</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Dermatological manifestations</td>
<td>Non-specific</td>
<td>Keratoderma blennorrhagica, circinate balanitis</td>
<td>Psoriasis</td>
<td>Erythema nodosum, pyoderma gangrenosum</td>
</tr>
<tr>
<td>Uveitis</td>
<td>Occasional</td>
<td>Common</td>
<td>Occasional</td>
<td>Occasional</td>
</tr>
<tr>
<td>Other extra-articular</td>
<td>Aortic regurgitation, conduction defects, upper lobe pulmonary fibrosis, IgA nephropathy</td>
<td>Aortic regurgitation</td>
<td>Aortic regurgitation</td>
<td>Aortic regurgitation</td>
</tr>
<tr>
<td>manifestations</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Familial aggregation</td>
<td>Common</td>
<td>Common</td>
<td>Common</td>
<td>Common</td>
</tr>
<tr>
<td>HLA B27</td>
<td>90%</td>
<td>80%</td>
<td>40%</td>
<td>30%</td>
</tr>
</tbody>
</table>

Psoriatic arthritis (PsA) is probably the most common of these arthritides and affects around 30% of patients who suffer from psoriasis. PsA may also affect people who have no signs of psoriasis themselves, but who have a family history of the disease.

In addition to joint synovitis, PsA commonly affects the tendons and entheses around joints. Common ultrasound features include Achilles tendinopathy /enthesopathy, plantar fasciitis and dactylitis. Unlike rheumatoid arthritis, the areas affected are often not symmetrical in PsA.

As these pathologies are not specific to PsA, the association between ultrasound appearances and a history (or family history) of psoriasis may be made during scanning and suggested in the report.
Crystal arthritis

Gout

Gout is characterised by acute attacks of inflammatory arthritis, often, but not only in the 1st metatarsophalangeal joints which become red, hot, tender and swollen. Gout is caused by elevated levels of uric acid in the blood which crystallises, and the crystals are deposited in joints and soft tissues.

Ultrasound appearances that are suggestive of gout include synovitis and erosions but also a ‘double contour’ sign. This is thought to be due to deposition of uric acid crystals on the surface of cartilage and is seen most commonly on the heads of the first metatarsals. Research into this is ongoing.

Tophi are aggregates of uric acid crystals in joint capsules or soft tissues. On ultrasound, they are seen as solid, hyperechoic but non-shadowing deposits within joint capsules, tendons or bursae.

Calcium pyrophosphate deposition

Calcium pyrophosphate deposition (CPPD) is an umbrella term for the various clinical subsets, whose naming reflects an emphasis on particular features. For example, pseudo gout refers to the acute symptoms of joint inflammation or synovitis: red, tender, and swollen joints that may resemble gouty arthritis.

Chondrocalcinosis, on the other hand, refers to the radiographic evidence of calcification in hyaline and/or fibrocartilage. Common sites are the triangular fibrocartilage (TFCC) of the wrist and the menisci of the knee.

Pitfalls, contraindications and limitations

Pitfalls in ultrasound scanning for inflammatory arthritis include the use of incorrect equipment settings and lack of appreciation of the effects of current treatments.

Scanning technique

Good technique for scanning is vital in accurately assessing joints for synovitis. Of particular importance is the lack of transducer pressure on the skin surface. Too much pressure may compress the small blood vessels present in joint synovitis or tenosynovitis and obliterate the Doppler signal. Gentle, meticulous scanning across the joints is essential to exclude inflammation.

Equipment settings

As Doppler is used frequently to detect the presence of active inflammation it is vital that the ultrasound system used is sensitive to detect low flow. Important settings include a low pulse repetition frequency (PRF), appropriate gain settings and a low level of wall filter.

Steroid and non-steroidal anti-inflammatory drug (NSAID) use

Steroids – oral, intramuscular or infusion - either for the treatment of joint disease or concurrent problems such as asthma, bronchiectasis and chronic obstructive pulmonary disease, will temporarilily reduce inflammation and hyperaemia. There is some evidence that NSAIDS have a similar effect.
In inflammatory arthritis, corticosteroids have been demonstrated to reduce synovial inflammation in imaging studies. It is therefore important to take steroid use into account when reporting on ultrasound studies to detect or grade active synovitis. The images below show the effects of steroid use on Doppler flow in a case of RA.

The use of oral, intra muscular or intra-articular steroids will affect the level of Doppler signal and may even remove the ultrasound features of inflammatory arthritis. It is therefore suggested that ultrasound examinations should be scheduled at least six weeks after any steroid intervention. If that is not possible, the sonographer should add a comment in the report to ensure that the referrer is aware.

For example: ‘The patient reports that they are taking oral steroids/has had a recent intra-articular or intramuscular steroid. This may reduce the ultrasound features of inflammatory arthritis and could affect true grading of synovitis. If there is ongoing clinical concern, a rescanning 6 weeks after any steroid use is suggested.’

The use of NSAIDS prior to a scan may be more difficult to manage as they are widely used and required for pain relief. It may not be practical to limit their use, but care should be taken when questioning the patient to ensure that it is known that they are being taken. In these cases, areas of synovial proliferation with no Doppler signal and no obvious cause should be taken into consideration when reporting and may require rescanning.

**Considerations when scanning**

As age-related changes are common in the musculoskeletal system and may not be the cause of the patient’s symptoms, ultrasound appearances must always be taken in clinical context and the referrer should be made aware of its limitations in the report. For example:

‘Ultrasound cannot exclude intra-articular pathology’

Diagnostic ultrasound is often used as a precursor to therapeutic injections and care should be taken to assist in the direction of that injection. Some structural changes may not be currently relevant and may not be associated with pain. For example, synovial hypertrophy without Doppler signal to
suggest activity is seen in many kinds of arthritis. The presence of a Doppler signal within the synovial hypertrophy is more indicative of active synovitis. The addition of the site of current symptoms on a report may help.

The pitfalls of ultrasound interpretation are widely documented and can be reduced by education and experience of the individual ultrasound practitioner.

Scan protocol

Hands and wrists

The structures that should be identified as a minimum in a hand ultrasound scan for rheumatology are:

- wrist/carpal joints – scan in longitudinal and transverse to interrogate joints and overlying extensor tendons
- extensor carpi ulnaris tendon (ECU) – scan in longitudinal and transverse
- all dorsal metacarpophalangeal (MCP) joints – scan in longitudinal, and if suspected pathology, confirm in transverse
- all dorsal proximal interphalangeal (PIP) joints – scan in longitudinal, and if suspected pathology, confirm in transverse
- flexor tendons – scan in longitudinal and transverse
- if querying seronegative inflammatory arthritis such as Psoriatic arthritis, it may be useful to include distal interphalangeal (DIP) joints.

Wrist

Begin your scan at the wrist crease. Scan the dorsal wrist/carpal joint in longitudinal from the medial to the lateral border and in transverse, to cover the joints.

**Look for:** Synovial hypertrophy, effusion, bone erosions, degenerative change.

Repeat the scan using Power Doppler

**Look for:** Active synovitis. If present, grade using grading guidelines.

Scan the extensor tendons in transverse from the musculo-tendinous junction to the distal insertion and then in longitudinal. If necessary, compare with the contralateral side.

**Look for:** Tendon sheath thickening, tendon thickening/thinning, tendon discontinuity, effusion.

Repeat the scan using Power Doppler.

**Look for:** Active tenosynovitis – mild, moderate or severe.

Scan the volar wrist/carpal joint in longitudinal from the medial to the lateral border and in transverse, to cover the joints. If necessary, compare with the contralateral side.

**Look for:** Synovial hypertrophy, effusion, bone erosions, degenerative change.

Repeat the scan using Power Doppler.

**Look for:** Active synovitis. If present, grade using grading guidelines.

Scan the flexor tendons in transverse from the musculo-tendinous junction to the distal insertion and then in longitudinal. If necessary, compare with the contralateral side.

**Look for:** Tendon sheath thickening, tendon thickening, tendon discontinuity, effusion.

Repeat the scan using Power Doppler.
Look for: Active tenosynovitis – mild, moderate or severe

Dorsal MCP/PIP/DIP joints

Begin your scan at the relevant joint. Repeat for each joint of each digit.

Scan the joint in longitudinal from the medial to the lateral border and in transverse, to cover the joint. Look for: Synovial hypertrophy, effusion, bone erosions, degenerative change. Repeat the scan using Power Doppler. Look for: Active synovitis. If present, grade using grading guidelines and image in longitudinal and transverse.

Imaging protocol

A standard hand and wrist series should include the following minimum images for a normal rheumatology scan:

- Dorsal wrist and carpal joints in longitudinal with colour box.
- ECU tendon with Colour box
- All MCPJs in longitudinal with Colour box
- All PIPJs in longitudinal with Colour box
- Flexor tendons with Colour box

Feet and ankles

Scan protocol

The structures that should be identified as a minimum in a foot ultrasound scan for rheumatology are:

- ankle – tibiotalar joint – scan in longitudinal and transverse to interrogate joints and overlying extensor tendons – anterior tibial, extensor hallucis longus and extensor digitorum longus.
- medial and lateral ankle tendons – posterior tibial, flexor digitorum longus and flexor hallucis longus tendons medially and peroneus longus and brevis laterally. Scan in longitudinal and transverse
- all dorsal metacarpophalangeal (MTP) joints – scan in longitudinal, and if suspected pathology, confirm in transverse if possible (this can be difficult due to the angle of the joints – a hockey stick transducer may help)
- MT heads on the plantar surface looking for erosions that may be difficult to see on the dorsal aspect, especially around the 5th MT head laterally
- flexor tendons – scan in longitudinal and transverse
- if querying seronegative inflammatory arthritis such as Psoriatic arthritis, it may be useful to include the Achilles and plantar fascia insertions.

Begin your scan at anterior ankle joint. Scan the dorsal tibiotalar joint in longitudinal from the medial to the lateral border and in transverse, to cover the joint. Look for: Synovial hypertrophy, effusion, bone erosions, degenerative change.

Repeat the scan using Power Doppler. Look for: Active joint synovitis – mild, moderate or severe
Scan the anterior, medial and lateral ankle tendons in transverse from the musculo-tendinous junction to the distal insertion and then in longitudinal. If necessary, compare with the contra lateral side.

**Look for:** Tendon sheath thickening, tendon thickening/thinning, tendon discontinuity, effusion

Repeat the scan using Power Doppler.

**Look for:** Active tenosynovitis – mild, moderate or severe

Scan the dorsal MTPJs in longitudinal from the medial to the lateral border and in transverse, to cover the joint.

**Look for:** Synovial hypertrophy, effusion, bone erosions, degenerative change.

Repeat the scan using Power Doppler.

**Look for:** Active joint synovitis – mild, moderate or severe

Scan the plantar surface of the MTPJs in longitudinal from the medial to the lateral border looking for erosions.

**Imaging protocol**

A standard foot/ankle series should include the following minimum images for a normal rheumatology scan:

- Dorsal ankle joint in longitudinal with colour box.
- Medial tendons with Colour box
- Lateral tendons with Colour box
- All MTPJs in longitudinal with Colour box

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**4.12.7.1 Reporting examples**

A report is intended to answer a clinical question and to assist with the patient journey so local opinion from radiologists and rheumatologists should be taken into account in the construction of local report templates.

The standard shoulder report should include:

- documentation of the normal anatomy;
- documentation of any pathology including measurements/ any increase in vascularity if appropriate;
- documentation of any limitation to range of movement, the site and degree of pain
- documentation of any difficulties with interpretation of the ultrasound appearances.

The sample reports below are intended as a guide only as reporting style may be specific to individuals/departments.

---

**Sample report for a rheumatology scan showing no evidence of inflammatory arthritis**

Ultrasound of the both hands with verbal consent
No evidence of active synovitis seen arising from the wrists or within the MCP or PIPJs. No tenosynovitis. No erosions seen.

Or
Ultrasound of the both hands with verbal consent
Degenerative changes noted but no erosions seen and no evidence of active synovitis seen arising from the wrists or within the MCP or PIPJs. No tenosynovitis.

Or

Ultrasound of the both hands with verbal consent
Degenerative changes noted with mild inactive synovial hypertrophy arising from the wrists. No evidence of active synovitis seen arising from the wrists or within the MCP or PIPJs. No tenosynovitis.

Referral for symptoms following recent intramuscular steroid injection
Ultrasound of the both hands with verbal consent

There is obvious synovial hypertrophy arising from the dorsal surfaces of both wrists but no evidence of internal Doppler signal. The patient reports an intramuscular injection of Depomedrone 2 weeks ago. This will reduce Doppler signal and may mask the ultrasound appearances of active synovitis. If symptoms return after 4-6 weeks and there is ongoing suspicion of inflammatory arthritis, a rescan is suggested before any further steroids are given.

Referral for a rheumatology scan with positive findings for inflammatory arthritis.
Ultrasound of the both hands with verbal consent

There is grade 2 (or moderate) active synovitis arising from the right wrist and within the right index and middle and left index finger MCPJs. No evidence of active synovitis arising from the left wrist or within the remainder of the MCPJs or the PIPJs. No tenosynovitis. No erosions seen.

Referral for suspected seronegative inflammatory arthritis
Ultrasound of the both hands with verbal consent

There is evidence of moderate synovitis arising from the left ankle joint but none seen on the right. The distal portion of both Achilles tendons are thickened and hypoechogenic with loss of the normal fibrillar pattern. There is a moderate degree of internal vascularity within the tendons at the insertion and erosions on the posterior aspect of the calcaneum. Appearances suggest erosive enthesitis. There is also effusion and hyperaemia of both retro calcaneal bursae suggesting bursitis. In the absence of an obvious biomechanical cause for these appearances, an inflammatory arthropathy should be considered and if there is clinical correlation, a rheumatology opinion is suggested.

References:


Further information can be found at:

4.13 Elastography

Advances in ultrasound technologies have expanded the clinical applications of ultrasound over recent years. Historically ultrasound has been a primary imaging modality, but it is now common for ultrasound technology to be utilised in other techniques such as needle placement, joint injections, guided biopsies, interventional procedures and more recently in the diagnosis and management of patients with chronic liver diseases.

Recent advances in elastography techniques such as acoustic radiation force impulsion (ARFI) and transient elastography (TE) have expanded the use of ultrasound into disease management. They can also enable lesion characterisation and the mapping of tissue stiffness. This reproduces / replaces the palpation performed by clinicians.

At present there are two main forms of elastography which are being used routinely in clinical practice.

Strain Elastography (SE)

Quasi-static or strain elastography (SE), where the tissue deformation is produced by external palpation with the ultrasound transducer. Most manufacturers overlay a speckle map on the ultrasound image, which is coded in a colour or greyscale to show a pattern of strain, which is inversely related to the tissue stiffness and can be assessed subjectively. These images are semi-quantitative and do not directly depict the elasticity which is derived from Young’s modulus, (stress/strain ratio).

Shear Wave Elastography (SWE)

Conventional ultrasound and shear waves are both pressure waves that are conducted through soft tissue. Unlike conventional ultrasound waves; shear waves are transverse in their directionality, they are rapidly attenuated by soft tissue and travel more slowly (between 1 and 10 m/s). Their speed is closely related to Young’s modulus of elasticity. Shear waves are created naturally from muscle movements or in cardiac activity and can be induced by ultrasound scanners which use them to measure their speed in a particular tissue or organ.

Liver Fibrosis Assessment

Most chronic liver diseases have alterations in the functional and structural appearance of the liver. The risk of developing cirrhosis and liver related complications in viral and non-viral chronic liver diseases is correlated to the amount of liver fibrosis. The detection and classification of liver fibrosis is pivotal for assessing progression and when to commence patients on antiviral therapies. Liver biopsy has long been the traditional approach for fibrosis assessment and disease classification. Liver biopsy by its nature is invasive and has potentially severe complications in up to 1% of cases with associated cost implications. NICE (2015) in their medical technologies guidance [MTG27] advocate elastography in the diagnosis and monitoring of fibrosis in chronic hepatitis. The economic benefits of using elastography is explored in the guideline, a saving of around £434 per patient is quoted when using Virtual Touch Quantification (VTq) over conventional liver biopsy. The safety implications to patients should also be considered as well as its tolerability and its ability to be undertaken in an outpatient setting. Giving the economic challenges facing healthcare today, this technique could have the potential to deliver large savings.

One difficulty with liver elastography is the different methods and multitude of ultrasound manufacturers providing a method of stiffness assessment. This results in the varying cut-off values being ultrasound scanner specific and not transferable across different makes of equipment.
Transient Elastography (TE)

One method of SWE is transient elastography (TE) which is being performed using the Fibroscan® (Echosens, Paris, France). This is a mechanical system using a single element ultrasound transducer which is used to apply light pressure on the patient’s skin, usually intercostally to assess the liver parenchyma. The resultant shear wave travels through liver tissue along the direction of the ultrasound beam and is used to measure its speed in a manner similar to M-mode. In patients with chronic hepatitis C, liver stiffness (LS) values are greater than 6.8–7.6 kPa. The cut off for predicting cirrhosis range between 11.0 – 13.6 kPa. Evidence suggests that TE has limitations in differentiating between mild and significant fibrosis. Other limitations include the requirement of dedicated equipment with specific probes and an intercostal scan approach. The technique is standardised and can be employed in various settings. It is currently being used mainly outside the radiology department in outpatient settings.

Acoustic Radiation Force Impulse (ARFI)

ARFI generates shear waves by causing small displacements in the soft tissue, causing sideways pressure waves away from the ultrasound beam which allow ultrasound systems to measure speed with good spatial resolution.

ARFI accuracy for the assessment of liver fibrosis has been similar to TE, with some more promising results in patients with non-alcoholic fatty liver disease (NALFD), non-alcoholic steatohepatitis (NASH) and in post transplantation. It has the advantage of allowing a radiological assessment of the liver parenchyma and more accurate localisation of the sample site. Therefore, fatty infiltration can be avoided so as not to not skew fibrosis scoring.

The European Federation of Societies of Ultrasound in Medicine and Biology (EFSUMB) guidelines “EFSUMB Guidelines and Recommendations on the Clinical Use of Ultrasound Elastography” (2013) and “EFSUMB Guidelines and Recommendations on the Clinical Use of Liver Ultrasound Elastography, Update 2017 (Long Version)” (2017) recommend the use of shear wave elastography to assess the severity of patients with liver disease with viral hepatitis. The guidelines also state that shear wave elastography is promising in NAFLD and post-transplantation.

Clinical Application

Whilst differing manufacturers have different shear wave velocities for stages of liver disease it is noted that the following (based on the VTq imaging as described in the NICE publication [MTG27]) are
given as examples of disease staging

- Normal < 1.2 m/s
- Fibrosis ≥ 1.21 – 1.34 m/s
- Cirrhosis ≥ 1.55 – 2.00 m/s

Extreme caution is required when quoting shear wave velocities in ultrasound reports. It is useful to report the shear wave velocity and quote the relevant reference values for the machine used to minimise confusion between normal and abnormal readings compared to the stage of liver disease. Practitioners are advised to refer to individual manufacturers’ reference ranges when reporting shear wave studies in liver disease.

Practical points on acquisition

- The patient should be fasted. Shear wave speed measurements are performed intercostally in the right liver with conventional curved arrays. The probe is aligned along an intercostal space.
• 10 measurements should be acquired and the mean value and standard deviation stated.
• The recommended depth is between 3 and 7 cm. Shear wave methods can be utilised in patients with ascites.

References:

Further information can be found at:
• EFSUMB Guidelines and recommendations

4.14 Contrast enhanced ultrasound (CEUS)

Ultrasound contrast agents are formulated into a microbubble structure which performs as a blood pool tracer when used in conjunction with contrast specific imaging modes using a low mechanical index (MI) technique. An example is ‘Sonovue’ (sulphur hexafluoride). These agents are intravenously injected in order to demonstrate the vasculature and microvasculature of organs and potential lesions.

CEUS is primarily used for hepatic applications, as recommended by NICE (2012) in the ‘diagnostics guidance 5 [DG5]’ but according to ‘The EFSUMB Guidelines and Recommendations for the Clinical Practice of Contrast-Enhanced Ultrasound (CEUS) in Non-Hepatic Applications: Update 2017 (Long Version)’ are also useful in a variety of other applications.

**Administration of ultrasound contrast agent**

Administration is performed via an intravenous cannula.

Use of contrast agents by non-medical staff, usually sonographers, should come within the remit of a patient group direction (PGD) or other legal mechanism which allows the prescribing and safe administration of the drug (ref: section 4.16).

**IMPORTANT**

A sonographer must be statutorily registered to follow a PGD.

Voluntary registration is not sufficient.

Any contraindication to the agent used should be carefully observed, and documentation of the dose administered recorded according to local rules.
Safety

The microbubble agents currently available are not nephrotoxic, making them a very useful alternative in patients with renal compromise. They have a very low incidence of adverse reactions although there is a small risk of anaphylactoid reactions to CEUS (rate estimated at 1:10,000) and resuscitation facilities with emergency equipment and personnel trained in its use should be available. It is recommended to keep the patient under close medical supervision during, and for at least 30 minutes following the administration of sulphur hexafluoride (Sonovue). A programme of annual basic life support training should be in place for staff, according to the RCR and SCoR ‘standards for the provision of an ultrasound service’.

Microbubble agents are currently not licensed in pregnancy or in children but, in children, may be used either ‘off label’ i.e. outside of the manufacturers marketing authorisation or ‘unlicensed’ i.e. no UK license or being made up in a way that it wasn’t intended to be used. The use of ‘off label’ is allowed when considered ‘best clinical practice’ according to the Specialist Pharmacy Service (2018) document “Medicines Matters: A guide to mechanisms for the prescribing, supply and administration of medicines (in England)”.

The FDA gave approval in April 2016 for the use of CEUS in the USA in children with focal liver lesions. EFSUMB guidelines (2017) “Role of Contrast Enhanced Ultrasound (CEUS) in Paediatric Practice: An EFSUMB Position Statement” assess the current status of CEUS applications in children and make suggestions for further development of this technique.

Equipment

Contrast-specific software is required to perform CEUS.

A low MI technique helps to preserve the microbubbles, enabling a longer available survey time.

The facility to record a short video clip, of all phases for at least three minutes, is essential when using CEUS to characterise lesions. The temporal resolution of CEUS is superior to any other current imaging modality, and the ability to review the arterial phase, frame by frame, provides a valuable contribution to the diagnostic process. Portal washout is a hallmark of malignancy.

Main applications

- Characterisation of focal liver lesions in non-cirrhotic patients.
- Characterisation of benign lesions.
- Detection of liver metastases in patients at high risk of liver metastases (e.g. those with a primary cancer who have not undergone CT staging, patients with sudden unintentional weight loss, abnormal LFTs and otherwise equivocal livers on a baseline scan). Patients for pre-treatment staging should ideally undergo CT.
- Doppler ‘rescue’ to identify flow in technically difficult examinations e.g. hepatic artery demonstration in a difficult post-transplant scan.
- Lesion localisation pre-ablation and for US guided biopsy.
- Post ablation checks.
- Trauma, to identify soft organ damage, if CT cannot be performed.
- Trauma follow-up looking for pseudo-aneurysms and resolution of lacerations.
Cirrhotic patients with focal lesions should ideally be referred to MRI for characterisation and further exclusion of hepatocellular carcinoma (HCC). However, if the patient is unable to undergo MRI, then CEUS may provide valuable additional information in lesion characterisation.

**Extrahepatic applications**

- Characterisation of complex renal cysts\(^46\) (e.g. with septa or containing debris/possible solid material). CEUS may be offered as a first line characterisation. Complex, obviously suspicious cysts and solid masses (Bosniak 4) should be referred urgently to CT.
- Vascular patency/ detection of infarcts in difficult renal transplant scans.
- Sentinel node detection in patients with breast cancer.
- Follow-up of endovascular aneurysm repair (EVAR) stents to establish patency and look for leaks.
- Hysterosalpingo-Contrast-Sonography (HyCoSy) for tubal patency testing.

It is the practitioner’s responsibility to be aware of the licensed use of Sonovue. Where Sonovue is used “off label”, local agreement to its use, and by whom within the department, should be documented. The patient’s verbal consent for the use of contrast “off label” should be sought and documented.

**EFSUMB (2018)** highlight further applications, with less supportive evidence at this stage, include the spleen, endoscopic US including pancreas, GI tract, scrotal and prostate. CEUS can also be used to facilitate ultrasound guided biopsy in difficult-to-see lesions.

**References:**


**4.15 Interventional and ‘extended scope’ examinations for sonographers [2018]**

As part of Health Education England’s “Multi-professional framework for advanced clinical practice” (2018) many sonographers are extending their scope of clinical practice, in addition to undertaking work in the other advanced practice areas of audit and research, education and leadership. There are a wide range of examinations that come within the overall scope of practice of ultrasound practitioners. For those who are not medically qualified or registered with the GMC (e.g. sonographers) they are sometimes termed ‘extended scope’ examinations. They include diagnostic procedures such as biopsy, fine needle aspiration, hystero-salpingo contrast sonography (HyCoSy), drainage of body cavities and therapeutic procedures such as joint injections.

Like all ultrasound examinations, the sonographer must be trained, competent and authorised to undertake them.
Examples of the type of ultrasound examinations undertaken by sonographers and to which this section refers are as follows:

i) Biopsy / fine needle aspiration e.g. breast, prostate, liver, thyroid.
ii) Drainage e.g. drainage of ascites
iii) Therapeutic e.g. joint injections
iv) Diagnosis e.g. HyCoSy, contrast enhanced ultrasound (CEUS).

When setting up such ‘extended scope’ services the approval of the clinical lead, service manager and Trust, Health Board or provider organisation should be sought.

The following will all need to be considered:

- support from a lead medical practitioner
- appropriate training
- professional indemnity Insurance
- quality assurance and audit programmes
- consent procedures and acceptance that a sonographer is the person who should obtain this
- protocols available for each individual type of procedure that the sonographer will undertake
- Patient Group Directions (PGD) set up as necessary. Some sonographers, depending on professional background, may be supplementary or independent prescribers. (ref: section 4.16). Sonographers must be statutorily registered to use a PGD, voluntary registration is not sufficient.

### 4.16 Patient group directions

When considering using pharmacy products in a clinical setting due reference has to be made to legislation and relevant publications. Reference is made to “Medicines Matters: A guide to mechanisms for the prescribing, supply and administration of medicines (in England)” (2018).

This document provides a comprehensive overview of which professionals can do what and under which legal framework.

In order to deliver safe and effective healthcare that also provides a good experience for patients, providers need to ensure that they are maximising the full potential of their entire clinical workforce. The aim is to enable quality improvement, innovation and greater productivity in service delivery.

Sonographers are a diverse group of healthcare professionals who deliver high quality care to patients across a wide range of care pathways in a variety of clinical settings. Ultrasound services are having increasing demands on their services, not least with compliance with referral to treatment targets (RTT), seven day working, and greater access for patients. A national shortage of sonographers leading to recruitment and retention issues further intensifies the challenge of providing a robust ultrasound service.

The existing arrangements by which sonographers prescribe and supply medicines to their patients are complex and for most pathways statutory registration is required. Supplementary prescribing training is available to some professional groups as is training to be an independent prescriber. Patient Group Directions are available to many sonographers who are also statutorily registered as (for example) a radiographer, physiotherapist, nurse or midwife.
Traditionally, the preferred way for patients to receive medicines they need is for a prescriber to provide care for a patient on a one-to-one basis. This method changed following publication of the final Crown report review on prescribing, supply and administration of medicines (1999)\textsuperscript{47}. Legal frameworks were developed that have allowed services to be re-designed and for healthcare professionals to work more flexibly for the benefits of patients. As a result, there are now several legal options for supplying and/or administering medicines including the commonly used Patient Group Directions (PGDs).

There are an increasing number of PGDs that will allow medicines to be administered without the need for a doctor to be present.

Greater flexibility of prescribing and medicines supply has the potential to reduce treatment delays, improve specificity and responsiveness of prescribing and thereby reduce patients’ exposure to safety risks. Safety considerations relate to training arrangements, communication of prescribing and governance arrangements.

The current situation and definitions

The law states that some groups of statutorily registered healthcare professionals are allowed to supply and/or administer medicines using Patient Group Directions. Some professional groups can train to become supplementary prescribers or independent prescribers.

**Independent Prescribing**

Independent Prescribing means that the prescriber takes responsibility for the clinical assessment of the patient, establishing a diagnosis and the clinical management required, as well as prescribing where necessary and the appropriateness of any prescription.

**Supplementary Prescribing**

Supplementary Prescribing is defined as a voluntary partnership between an independent prescriber (a doctor or dentist) and a supplementary prescriber to implement an agreed patient-specific Clinical Management Plan with the patient’s agreement.

**Patient Specific Directions (supply or administration)**

A Patient Specific Direction (PSD) is a direct written instruction for a named patient and does not require assessment of the patient by the individual instructed before administering, unlike a Patient Group Direction. In primary care, this might be a simple instruction in the patient’s notes. Examples in secondary care include instructions on a patient’s ward drug chart. If a radiologist writes the type, strength and amount of contrast agent to be given to a named patient then that is a Patient Specific Direction. No variation is allowed.

Where a Patient Specific Direction exists, there is no need for a Patient Group Direction. Anyone can follow a Patient Specific Direction as long as they are acting in accordance with the directions of an appropriate practitioner (i.e. the radiologist) following patient assessment, and delegation of supply and/or administration.

A Patient Specific Direction can also be used where a Patient Group Direction cannot be used, e.g. where the administration is carried out by a registered professional not covered by a Patient Group Direction or an unregistered healthcare professional such as an assistant practitioner or a sonographer who is not statutorily registered.
Patient Group Directions (supply or administration)

A Patient Group Direction (PGD) is a written instruction for the supply or administration of medicines to groups of patients who may not be individually identified before presentation for treatment. It is not a form of prescribing and there is no specific training that health professionals must undertake before they are able to work under a Patient Group Direction. They are a way for some groups of statutorily registered healthcare professionals to be able to supply and administer most medicines and contrast agents used in imaging and radiotherapy departments. However, certain requirements apply to the use of Patient Group Directions.

Appropriate training, competency assessment and accountability is required and due consideration is given to the following NICE guidance “Patient group directions. Medicines practice guideline [MPG2]”, section 1.7:

Training and competency

- Identify the senior person in each profession who is responsible for ensuring that only fully competent, qualified and trained health professionals use PGDs.
- Identify gaps in competency and establish a comprehensive and appropriate training programme for all people involved in considering the need for, developing, authorising, using and updating PGDs.
- Ensure that adequate educational materials are available to enable individual people and organisations to deliver safe and effective services in which PGDs are used.
- Consider collaborating with other organisations and sharing existing educational materials to ensure a comprehensive approach.
- Ensure that training and re-training of health professionals using PGDs incorporates a post-training assessment of competency.

Patient Group Directions are a way for some groups of statutorily registered healthcare professionals to be able to supply and administer most medicines and contrast agents used in imaging and radiotherapy departments.

N.B. Only some (not all) groups of statutorily registered health care professionals can use Patient Group Directions. Assistant practitioners are not a registered and regulated workforce and therefore are not allowed to supply or administer under Patient Group Directions. Some sonographers cannot achieve statutory registration and are instead voluntary registered. Voluntary registration is not sufficient for the use of PGDs; statutory registration is required.

Further information can be found at:

- Royal Pharmaceutical Society and Royal College of Nursing (2019) “Professional guidance on the administration of medicine in healthcare settings”

- SCoR. “Prescribing” at https://www.sor.org/practice/other-groups/prescribing

- Specialist Pharmacy Service. “The first stop for professional medicines advice”.

- Specialist Pharmacy Service. “Contrast agent PGD templates”
  - There are templates available for PGDs for iodinated contrast agents and for gadolinium.
• There are no templates currently available for ultrasound contrast agents but some of the information within the templates may be relevant when developing PGDs for these.

PGD flow diagrams:

Reference:

Citation: