Quality Indicators for Barrett’s EndoTherapy (QBET): UK consensus statements for patients undergoing Endoscopic therapy for Barrett’s neoplasia

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- Endoscopy
- Endoscopic Mucosal Resection (EMR)
- Endoscopic Submucosal Dissection (ESD)
- Radio Frequency Ablation (RFA)

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Competing Interest:
None declared

- This Project received funding from the BSG to cover travel expenses endured by the investigators in order to attend the face to face meeting in London for round 2 voting.
- The current main text is now 2960 words (excluding abstract, tables and references)
ABSTRACT:

Introduction:
Endoscopic therapy for the management of patients with BE neoplasia has significantly developed in the past decade; however, significant variation in clinical practice exits. The aim of this project was to develop expert physician-lead Quality Indicators for Barrett’s Endoscopic Therapy.

Method:
The RAND/UCLA Appropriateness Method was utilised to combine the best available scientific evidence with the collective judgment of experts to develop QBET in 4 sub-groups: Pre-endoscopy, intra-procedure (resection & ablation) and post-endoscopy. International experts including gastroenterologists, surgeons, BE pathologist, clinical nurse specialist, and patient representative participated in a 3-round process to develop 15 QIs that fulfilled the RAND/UCLA definition of appropriateness.

Results:
17 experts participated in Round 1 and 20 in Round 2. Of the 24 proposed QIs in round 1, 20 were ranked as appropriate (put through to round 2) and 4 as uncertain (discarded). At the end of round 2 a final list of 15 QIs were scored as appropriate.

Conclusions:
This UK national consensus project has successfully developed QIs for patients undergoing BET. These QIs can be used by service providers to ensure that all patients with BE neoplasia receive uniform and high quality care.
Key Summary:

Summarise the established knowledge on this subject:

Endoscopic eradication therapy for barrett’s neoplasia has revolutionised the management of patients with barrett’s neoplasia; however, despite various societal guidelines there still exist a great variation in clinical practice that results in variable patient outcomes.

What are the significant and/or new findings of this study:

Quality indicators in barrett’s endotherapy have been published recently in the United States. Adherence to these quality indicators have shown improvement in dysplasia detection rate; however quality indicators for barrett’s endotherapy in United Kingdom and Europe are lacking. These quality indicators identify important steps for providing a unified high quality care based on the best available evidence and expert opinion. These quality indicators may also be used for the training of the new generation of advanced endoscopists and adherence to these measures would ultimately result in improving patient outcomes.
Introduction:

The past decade has seen significant advancement in minimally invasive endoscopic treatment modalities for Barrett’s oesophagus (BE) neoplasia. Short and long term data report high eradication rates, acceptable disease durability and good safety profile that are comparable to the outcomes of surgical treatment (1). There has been great emphasis on targeting patients at earlier disease stages amenable to endoscopic eradication therapy (EET). EET for early neoplastic BE has been recommended by various major international guidelines (2).

EET for BE neoplasia has revolutionised the management of patients with BE neoplasia and is increasingly used at high volume tertiary referral centres and smaller district general hospitals. Adherence to Quality Indicators (QIs) introduced by the American Gastroenterological Association for the endoscopic management of patients with BE has been shown to improve dysplasia detection rate. Despite various societal guidelines (2), there still exist a great variation in clinical practice that results in variable patient outcomes.

It is important to note that the management of patients with BE neoplasia is just not confined to the endoscopic procedure only. It requires case discussion in a dedicated Multidisciplinary team (MDT) meeting with careful explanations to patients of their disease status and available therapies prior to and after endotherapy.

The current endoscopic management of BE neoplasia consists of endoscopic resection (ER) of visible lesions for accurate staging and risk stratification of patients followed by field ablation of remaining areas of flat BE to prevent the development of metachronous neoplasia. It is therefore important that cases are carefully selected for endoscopic therapy following discussion in MDTs with appropriate choice of therapy (after discussion with the patient) with strict follow up of these cases to ensure high quality service provision and better patient outcomes.

It is essential that medical resources are used appropriately and that health provision is shaped and maintained at the highest standard in order to ensure the best possible patient outcomes. Healthcare systems and providers will therefore need to be aligned to ensure a streamlined, efficient and high quality service provision to all patients. QI for Barrett’s endotherapy (BET) in the United Kingdom (UK) and Europe are lacking and have led to variable outcomes in the past (3). The aim of this project was to develop physician-lead Quality Indicators in BE Endotherapy (QBET) to define standardised clinical practice and achieve optimal clinical outcomes for all patients with BE neoplasia.
The aim from this project is not to replace existing guidelines but to create an adjunct so that clinicians can measure performance in a systematic way.

**Method:**

**This project was NOT a clinical trial and there was NO search conducted on humans.**

The RAND/UCLA Appropriateness Method (RAM):

The RAND/UCLA Appropriateness Method (RAM) was developed in the 1980s as part of the RAND Corporation/UCLA Health Services Utilisation Study. It is a tool used to measure the overuse and underuse of resources. In the RAM an appropriate measure refers to one in which the expected health benefit exceeds the expected negative consequences by a wide margin such that the procedure is worth performing without considering the cost (4). This methodology is used in situations where there is no adequate high quality research (e.g. randomised controlled trials) to guide clinical practice and therefore the best available evidence is combined with expert opinion, in order to develop quality indicators. RAM is a modified Delphi method that gives experts the opportunity to have a face to face discussion. RAM has been utilised in various clinical specialties including gastroenterology (5). This methodology was successfully utilised in establishing similar quality measures in EET in the US endorsed by the ASGE and ACG (5).

We utilised RAM to combine the best available scientific evidence with the collective judgment of experts to develop QBET in 4 sub-groups that are integral to patient selection, treatment and follow up in BET (Figure 1). The expert panel was selected based on membership in the UK RFA registry and publication history in the field of BE and BET. In addition, geographical variation was considered to ensure expert representation from all regions in the UK, which could be representative of the European variation in practice. The experts consisted of gastroenterologists and therapeutic endoscopists (n=20), including 2 surgeons performing surgery for advanced oesophageal adenocarcinoma (OAC) and providing BET and 1 BE expert pathologist. We also had participation from a BE clinical nurse specialist, a medical statistician and a patient representative. We developed QIs in 4 subgroups, as follows:

- Pre-endoscopy
- Intra-procedure (resection)
- Intra-procedure (ablation)
- Post endoscopy

Round Zero:

RAND/UCLA utilises 3 rounds as shown in figure 1. In round zero, experts were introduced to the project methodology and objectives (via teleconference on the 18th September 2017 by RJH, DA and KR) and familiarised with the RAM process. In addition, one expert was
allocated as lead for each subgroup to facilitate the discussions during the face to face meeting (round 2). After round zero, the core group leading the project (RJH, DA, KR, PS, OP) met to collate a list of potential QI’s. These were then reviewed with the project leads and the project leads (consisting of national and international experts) then proposed potential QIs for each of the 4 subgroups, which were put forward for ranking at round 1 (24 QIs in total).

Round One:
In round 1, 17 experts had the opportunity to rank each of the 24 QIs electronically in an independent fashion. This was done without interaction with other colleagues. The proposed QIs were sent to all the participating experts via a REDCap database.

Study data were collected and managed using REDCap electronic data capture tools hosted at University College London Hospital (6)(7). REDCap (Research Electronic Data Capture) is a secure, web-based software platform designed to support data capture for research studies, providing 1) an intuitive interface for validated data capture; 2) audit trails for tracking data manipulation and export procedures; 3) automated export procedures for seamless data downloads to common statistical packages; and 4) procedures for data integration and interoperability with external sources.

Instructions were also sent to the panel indicating that each QI should be scored by each expert based on their current expertise and knowledge on the topic. The experts were advised to score each QI as it would be applied to an average patient presenting to an average medical facility and to an average physician without the consideration for cost or feasibility of applying the QI in clinical practice. Each QI was ranked from 1 to 9 as per RAM protocol.

- Score of 1, 2, 3 = Inappropriate QI
- Score of 4, 5, 6 = Uncertain QI
- Score of 7, 8, 9 = Appropriate QI

Following round 1 voting, all the scores were collected and analysed using 4 statistical methods by an expert statistician with knowledge of the RAM process.

In addition, an extensive literature search on PubMed on the topic of BE and BET was performed around the proposed QIs. The literature search was limited to publications from 1st January 1990 to 23rd January 2018.

Prior to the round 2 face to face interaction and voting, the following was sent to all the investigators:
- A summary copy of the literature search for each QI
- A document showing the distribution of all the responses from round 1, including the
investigator’s personal response.

Round Two:
Only QIs that were deemed appropriate at round 1 (based on round 1 voting and statistical analysis), were put forward for discussion at round 2. The round 2 meeting (face to face meeting) took place on the 14th of March 2018 in London. At this meeting 20 investigators were provided with individual iPads containing all the overall results of the round 1 voting, a summary of all the literature searches around the QI’s, and full text copies of all manuscripts and references for reference and discussion. The lead for each subgroup led the discussion for each QI in that subgroup during this meeting. Each QI was discussed in detail taking into account the opinion from all those present and the available scientific literature. QIs were reworded, deleted and new QIs were developed (where necessary) for each of the 4 subgroups.

At the end of round 2 meeting, a set of 15 QIs were finalised and scored by each investigator [pre-endoscopy 2 QIs; intra-procedure (resection) 5 QIs; intra-procedure (ablation) 6 QIs; and post-procedure 2 QIs]. The experts also agreed on setting performance thresholds for each QI (if indicated) in order to set aspirational targets for all service providers. The median score (and range) of suggested performance thresholds are included with each QI. There were no set aspirational targets for QIs with pre-defined performance target in the text [e.g. Intra-procedural (ablation) QI number 4]. The expert panel recognised that some performance targets had to be set cautiously in order to avoid undermining established efficient practices and therefore aspirational targets were set to encourage centres to work towards enhancing their practice and performance. There were no attempts to force the expert panel to reach a consensus and each expert had the opportunity to score the finalised QIs independently.

Statistical Method:
Firstly, summaries of the number of responses in three categories were produced. Each response was categorised into one of the following categories:

- Inappropriate: Score 1-3
- Uncertain: Score 4-6
- Appropriate: Score 7-9

In addition to the categorisation, the median score for each QI was calculated and summarised.

The deviation in the responses between the panel members was assessed using a number of different methods. Firstly, deviation was assessed by the MAD-M statistics. This is the mean
absolute deviation from the median. Higher values of MAD-M indicate more spread in responses between the panel. A second measure was based on the BIOMED Concerted Action on Appropriateness definition. This method calculates the number of raters outside of the response category (i.e. inappropriate uncertain, appropriate) containing the median response. Disagreement was assumed if the number of raters outside this category meets a pre-defined threshold. In the RAND/UCLA handbook guidance is given for panel sizes up to 16 raters, but none is provided for 20 raters, as per this panel. Although there were no set guidelines for this number of raters, the decision was based on the same criteria as for a 16 rater panel (agreement if \( \leq 4 \) raters outside the category). The third measure used the RAND method that tests hypotheses about the distribution of ratings in a hypothetical population of repeated ratings. It is hypothesised that 90% of the hypothetical population of repeated ratings are within one of two extra wide regions (1-6 or 4-9). The binomial test was used to calculate the probability (p-value) that that ‘true’ value is below 90%. If the calculated probability is below the pre-determined level of 0.10, the conclusion will be reached that there is disagreement amongst raters. The final measure of deviation uses the IPRAS methods. This method is based on the inter-percentile range (IPR) between the 30th to 70th percentiles. The IPRAS is a statistic based on the IPR which is adjusted for symmetry. Disagreement was assumed if the IPRAS was larger than the IPR.

An additional set of analyses examined the threshold values for questions where these were appropriate. Median values and ranges were calculated for the thresholds.

The measures of spread included:
- The count of responses in each 3-point region (1,2,3 – 4,5,6 – 7,8,9)
- The mean absolute deviation from the median (MAD-M)

Appropriateness was measured using:
- Median rating
- BIOMED Concerted Action on Appropriateness definition
- P-value
- Interpercentile range adjusted for symmetry (IPRAS)

A QI was deemed appropriate if it met the definition of appropriateness, using ALL defined statistical methods.

**Results:**
Summary of responses from round 2 for each individual QI are shown in table 1. At round 2, 20 investigators ranked 15 QIs that were all deemed appropriate and shown in tables 2 to 5 with corresponding aspirational performance target (if indicated) and evidence summary. During round 1, 17 investigators ranked 24 QIs of which 20 were deemed appropriate and 4 uncertain (Table 6).
Discussion:

Endoscopic treatment for dysplastic BE and early OAC has been recommended by various major societal guidelines; however QIs for the management of patients with BE neoplasia have been lacking. This piece of work delivers a UK-based collection of QIs that will allow streamlined and accountable delivery of best clinical practice to patients undergoing BET. This nationwide project combined the best available evidence with the collective judgment of national and international experts in order to develop a set of formally validated QIs for the management of patient with BE neoplasia using a rigorous and validated methodology (RAM). The RAM, unlike the original Delphi, provides the expert panel with the opportunity to have a face-face discussion in round 2. Unlike guidelines which use a consensus methodology, the RAM reduces the possibility of results being influenced by the opinion of the most senior or most vocal member of the panel.

These UK-based QIs reflect those recently published QIs in BET in the United States (5); We were able to develop QIs for the intra-procedure component of patient care and for the management of patients at the pre-endoscopy and post endoscopy stage. In addition this UK-based project covered various aspects of patient care including the importance of formal training of endoscopists prior to service provision, the use of high quality endoscopic imaging modalities for lesion recognition in BE surveillance and the need for individual patient discussion at dedicated MDTs.

Adherence to Prague classification is known to result in improved dysplasia detection in patient with BE. This may be influenced by data from tertiary centres where diagnosis was obtained by expert BE endoscopist that are more likely to adhere to Prague classification with access to better endoscopic equipment including high definition endoscopy and virtual chromoendoscopy.

Our expert panel acknowledged the importance of endoscopic resection modalities (EMR and ESD) for the management of visible lesions in BE neoplasia. ESD is a feasible treatment option that allows en-bloc resection for histological staging and treatment of patients with early BE neoplasia. ESD is likely to expand in the near future and these QIs may need to evolve in order to cater for that in due course.

It is important that the clinical community recognises the balance between performing BET and the rate of success and stenosis. Therefore the expert group emphasized the importance of minimising stricture rates (not exceeding 10-15%) post BET and the need for discussion of patients’ care in MDTs prior to BET and when BET fails to achieve successful outcomes.
The current published evidence in BET (3) provides data that is confined to a limited time period (less than 10 years); however BET is expanding rapidly and therefore we need to continue long-term follow up in these patients and monitor outcomes, which will provide us with essential information that will shape our future practice.

In this project we were also able to set aspirational performance thresholds to ensure that patient care is of highest standard. Regulatory and accrediting agencies as well as hospitals and clinicians may use these QIs to measure performance and highlight areas for improvement. The regular audit of outcomes and adverse events will ensure the efficacy and safety of endoscopic therapy for patients with early BE neoplasia. Auditing results may be used to implement changes in routine practice nationally, allowing comparison of local practices to national standards. These QIs may also be used for teaching, service development and standardisation of care at all hospitals performing BET. Future studies will need to investigate the positive and the negative impact of these QI on patient outcomes.

There were some limitations to this study. First, high-quality evidence such as randomised controlled trials (RCTs) in the literature was not available for some QIs; however, this situation is common in many aspects of health care, and it was the very reason that the expert panel methodology such as RAM was developed (4). Second, some health care centres in the country may not be equipped with high quality endoscopic modalities and therefore these QIs may have a negative impact on their practice. Third, there was lack of validation of these QIs by an external committee and our expert panel voted on QIs that they developed themselves hence all the QIs in round 2 voting performed very well. Finally, the expert panel failed to determine the number of procedure needed to be performed by a centre to qualify as high volume centre and also failed to determine the adequate number of procedures needed by an endoscopist prior to performing independent BET.

In conclusion, this is the first UK national consensus project that has utilised a validated methodology to successfully develop process-based QIs for patients undergoing endoscopic treatment for early BE neoplasia. These indicators identify meaningful and important steps for providing a unified high quality care based on the best available evidence and expert opinion. These QIs may also be used for the training of the new generation of advanced endoscopists and adherence to these measures would ultimately result in improving patient outcomes.

**Acknowledgements:**
This study was previously presented as an abstract at UEGW 2018.
References:


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Tables:

**Table 1: Summary of responses from round 2 to individual QI**

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<th>Group</th>
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<th>Uncertain n (%)</th>
<th>Appropriate n (%)</th>
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<th>Median interpretation</th>
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<td>Pre-Endoscopy</td>
<td>1</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>20 (100%)</td>
<td>9</td>
<td>Appropriate</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>20 (100%)</td>
<td>9</td>
<td>Appropriate</td>
</tr>
<tr>
<td>Intra-Proced</td>
<td>1</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>20 (100%)</td>
<td>9</td>
<td>Appropriate</td>
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<tr>
<td>(EMR)</td>
<td>2</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>20 (100%)</td>
<td>9</td>
<td>Appropriate</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>20 (100%)</td>
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<tr>
<td></td>
<td>4</td>
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<td>0 (0%)</td>
<td>20 (100%)</td>
<td>8.5</td>
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</tr>
<tr>
<td></td>
<td>5</td>
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<td>0 (0%)</td>
<td>20 (100%)</td>
<td>8.5</td>
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<tr>
<td>Intra-Proced</td>
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<td>(RFA)</td>
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<td>9</td>
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<tr>
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<td>20 (100%)</td>
<td>8</td>
<td>Appropriate</td>
</tr>
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**Table 2: Appropriate Pre-endoscopy quality indicators after Round 2 voting with the median score, MAD-M, BIOMED Analysis, p-value, IPRAS analysis and the performance threshold**

<table>
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<tr>
<th>Pre-endoscopy QIs</th>
<th>Median Score</th>
<th>MAD-M</th>
<th>BIOMED Analysis</th>
<th>P-Value</th>
<th>IPRAS Analysis</th>
<th>Performance Threshold Median % (Range)</th>
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<tr>
<td>BET should be performed in high volume centres within a local cancer network to</td>
<td>9</td>
<td>0.2</td>
<td>No disagreement</td>
<td>1</td>
<td>No disagreement</td>
<td>100 (90, 100)</td>
</tr>
</tbody>
</table>
Aspirational performance target: 100% (range: 90-100)

Evidence summary:
Endoscopic training should start with knowledge acquisition, followed by resection and ablation in animal models, before training in human subjects. Endoscopist proficiency increases with the numbers of treatment sessions performed (8). Adherence to BE surveillance biopsy protocol in non-tertiary centres are poor, resulting in reduced dysplasia detection rate. Adherence to this protocol is further reduced with an increasing length of BE segment. Advanced imaging with HD-WLE and NBI have been shown to improve the detection rate of early neoplasia in patients with BE. The majority of gastroenterologists from academic centres use HD-WLE to classify BE as per guidelines and perform significantly more EET procedures per month, in comparison to those in district general hospitals. These factors favour the referral of patients with BE neoplasia to dedicated high volume centres.

In addition, data from the UK RFA registry has shown that increasing experience in performing EET is associated with significantly improved Complete Remission of Dysplasia (CR-D) and Complete Remission of Intestinal Metaplasia (CR-IM) rates, less number of rescue EMRs and faster protocol completion. At the start of the registry and at a time when only less than 20 patients were enrolled, the documented CR-D and CR-IM after completing EET were 79.8% and 71.3% respectively; however with increasing experience (i.e. once > 40 patients enrolled), the study was able to show significantly better CR-D (91%) and CR-IM (83.9%) (p<005) (9). This data supports improvement in experience and outcomes with increase in the number of procedures performed. The expert panel has therefore suggested that endoscopic therapy should be performed in high volume referral centres to optimise outcomes. Hospitals performing > 40 EET cases per year, may therefore be suitable centres for preforming BE endoscopic eradication therapy.

| Patients considered for BET, should be discussed in an Oesophago-Gastric MDT | 9 | 0.3 | No disagreement | 1 | No disagreement | 93 (85, 100) |

Aspirational performance target: 93% (range: 85-100)

Evidence summary:
The National Institute for Health and Clinical Excellence (NICE) (August 2010) guidelines on ablative therapy for the treatment of BE, recommends to discuss the MDT’s views on the range of appropriate treatments with the patient. It also recommends giving patients verbal and written information about their diagnosis, available treatments, patient support groups, and the uncertainty of the long-term outcomes of ablative therapies (10). In addition the BSG recommends that the treatment of patients with BE neoplasia should be discussed in a dedicated GI specialist MDT taking into account patient comorbidities, nutritional status, patient preferences and staging (2). Patients should be provided with information on all treatment options and offered verbal and written information on support groups available to them (2) including clinical nurse specialists. Despite little evidence, the expert panel advocates a MDT approach (consisting of an expert BE pathologist) for these patients in order to safeguard against incorrect use of BET in patients with more advanced disease and to ensure that the case management provided is directed to best patient interest.

Table 3: Appropriate Intra-procedure (Resection) quality indicators after Round 2 voting with the median score, MAD-M, BIOMED Analysis, p-value, IPRAS analysis and the performance threshold

<table>
<thead>
<tr>
<th>Intra-Procedural QIs (Resection)</th>
<th>Median Score</th>
<th>MAD-M</th>
<th>BIOMED Analysis</th>
<th>P-Value</th>
<th>IPRAS Analysis</th>
<th>Performance Threshold Median % (Range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adherence to the Prague and Paris classification is mandatory</td>
<td>9</td>
<td>0.1</td>
<td>No disagreement</td>
<td>1</td>
<td>No disagreement</td>
<td>95 (80, 100)</td>
</tr>
</tbody>
</table>

Aspirational performance target: 95% (range: 80-100)

Evidence summary:
Several studies have investigated the validity of the Prague circumferential and maximum length (C & M) classification showing high overall validity for the endoscopic assessment of visualised BE lengths amongst expert endoscopists, community hospital endoscopists and trainees. The BSG guidelines recommend endoscopic reporting be performed using the Prague criteria (2). Description of lesion morphology using the Paris classification is based on the Japanese system used to classify early gastric cancer. This provides information on the likelihood of invasion of cancer and helps communication between endoscopists. Description of lesion morphology using the Paris classification improves lesion recognition at the time of endoscopic therapy. It gives an indication of the likelihood of invasive cancer and aids communication between clinicians. The BSG recommends the use of Paris classification for all visible lesions; therefore adherence to the Prague and Paris classification is recommended.
All patients undergoing BET and follow up, should have assessment with High-definition white light (WL) endoscopy with (virtual) chromoendoscopy

<table>
<thead>
<tr>
<th></th>
<th></th>
<th>No disagreement</th>
<th>No disagreement</th>
<th>93 (80, 100)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Aspirational performance target: 93% (range: 80-100)**

**Evidence summary:**

Endoscopy in BE patients should be performed with careful inspection of the columnar-lined oesophagus using HD-WLE, with biopsy of any suspicious areas followed by 4-quadrant biopsies of the BE metaplasia. The use of the HD-WLE is associated with improved detection of dysplasia during routine BE surveillance. In addition, chromoendoscopy allows for detailed imaging of the mucosal and vascular surface patterns in BE. Recent studies have shown that imaging techniques such as chromoendoscopy or virtual chromoendoscopy increase the diagnostic yield for identification of dysplasia or cancer in patients with BE; however the evidence for advanced endoscopy boosting dysplasia detection rate on a per-patient basis is slim (11).

The application of a dilute acetic acid (AA) solution to the BE mucosa results in mucosal colour change and highlights mucosal patterns more clearly, facilitating sensitive and specific identification of potentially neoplastic areas. Furthermore, the premature loss of acetowhiteness in areas of the mucosa and the speed at which it disappears is also associated with the presence of early neoplasia. The efficacy of AA chromoendoscopy has been demonstrated in few studies showing a sensitivity and specificity of up to 98% and 96%, respectively.

Three main virtual chromoendoscopy modalities are currently available: narrow band imaging (NBI - Olympus), the i-Scan imaging system (Pentax), and blue laser imaging (BLI – Fujifilm). Recent studies have indicated the potential of NBI as a replacement for AA chromoendoscopy with an accuracy of 92%, and sensitivity and specificity of 91% and 93%, respectively, in the identification of early dysplastic lesions on still images (12). Other studies have also shown that i-Scan can improve neoplasia detection in patients with BE with an impressive accuracy and sensitivity, of up to 94% and 83%, respectively. The use of i-Scan can improve dysplasia detection rate. A recent study by Subramaniam et al. validated a classification system for Blue laser imaging (BLI) which identifies dysplastic BE tissue with sensitivity and specificity of 96%, based on both increased pit pattern irregularity and the presence of disordered and dilated microvessels (13). Currently, only AA and NBI have reached the ASGE PIVI requirement.

The current data on advanced imaging modalities in improving dysplasia yield is encouraging, but the data does not provide evidence on how these modalities can impact EET. Most studies to date have either been performed using still images or have been limited to high volume BE referral centres. The expert panel has therefore suggested that all patients undergoing BET and follow up should have assessment with HD WLE with chromoendoscopy or virtual chromoendoscopy.

All visible lesions should be entirely resected with EMR or ESD

<table>
<thead>
<tr>
<th></th>
<th></th>
<th>No disagreement</th>
<th>No disagreement</th>
<th>93 (80, 100)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

**Aspirational performance target: 93% (range: 80-100)**

**Evidence summary:**

ER is the cornerstone of endoscopic therapy of early oesophageal neoplasia, which aims to provide accurate histological staging with therapeutic intent. ER of early BE neoplasia with Multiband Mucosectomy (MBM) is effective and safe. Large number of studies have shown long-term complete remission rate of 85% to 96% with bleeding rates ranging from 0.7-7.9% and perforation rates ranging from 0.2-2.3% (14). EMR of all visible lesions has been shown to upgrade the pathological diagnosis in 39% of all patients. Most of the change was associated with upgrading of grade of dysplasia and neoplasia. EMR for all visible lesions have been recommended by the ASGE. In addition the provision of EMR specimens to the pathology department results in an improvement in interobserver agreement among pathologists compared with biopsy specimens only.

ESD for early stage BE neoplasia is also a feasible treatment option as it allows en bloc resection and accurate histopathologic analysis of lateral resection margins in BE neoplasia. Multiple studies have shown high en bloc resection rates ranging from 89-98.6% and R0 resection rates ranging from 72.4-87% with acceptable perforation (0-8.3%), bleeding (1.4-1.7%) and stricture rates (2.1-11.6%). When curative resections are achieved, good oncologic outcomes are likely in the management of early stage BE neoplasia by ESD.

The ESGE recommendations (2015) state that EMR is acceptable for resecting lesions confined to the mucosa, regardless of the size, but ESD may be considered for lesions larger than 15 mm, poorly lifting tumours, and lesions at risk for SM invasion (15). These data show that EMR and ESD are effective treatment modalities in the staging and treatment of early BE neoplasia with acceptable side effect profiles. It is however important to mention that operator skill and experience will have significant effect on
The use of EUS is not routinely recommended for patients undergoing BET

<table>
<thead>
<tr>
<th>Intra-Procedure QIs (Ablation)</th>
<th>Median Score</th>
<th>MAD-M</th>
<th>BIOMED Analysis</th>
<th>P-value</th>
<th>IPRAS Analysis</th>
<th>Performance Threshold Median % (Range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low and High grade dysplasia without visible lesions should undergo endoscopic ablation</td>
<td>9</td>
<td>0.4</td>
<td>No disagreement</td>
<td>1</td>
<td>No disagreement</td>
<td>95 (80, 100)</td>
</tr>
</tbody>
</table>

Aspirational performance target: 95% (range: 80-100)

Evidence summary:
The multicentre EURO II study showed that RFA can achieve a CR-D and CR-IM rates of 92% and 87%, respectively (21), in patients with early BE neoplasia. A systematic review by Desai et al also showed that ET of BE neoplasia with resection of visible lesions followed by ablation of the remaining segment of BE can achieve a CR-D rate of 93.4% and CR-IM of 73.1% (22). ET for early BE neoplasia should therefore be offered after appropriate discussion with the patient as ET is associated with high rate of CR-D and CR-IM and reduction in patient outcome and therefore good training is paramount.
in disease progression and development of cancer. The efficacy and safety profile of RFA suggests that it is the best ablative modality currently available for patients with LGD and HGD without visible lesions. The diagnosis of dysplasia should be reproduced and confirmed by expert BE pathologists prior to consideration for EET. Recent meta-analysis by Qumseya et al, studied the progression rates in LGD patients based on review by an expert GI pathologist. The group was able to show that the rate of progression from LGD to HGD/OAC was significantly higher among studies where expert GI pathologist confirmed the diagnosis of LGD compared with studies that did not use a GI pathologist (23).

<table>
<thead>
<tr>
<th>Following endoscopic resection, patients undergo ablative therapy, every 2-4 months in order to achieve CR-IM</th>
<th>9</th>
<th>0.3</th>
<th>No disagreement</th>
<th>1</th>
<th>No disagreement</th>
<th>90 (80, 100)</th>
</tr>
</thead>
</table>

**Aspirational performance target: 90% (range: 80-100)**

**Evidence summary:**
The initial UK RFA registry of 335 patients with BE and neoplasia that received ER for visible lesion followed by RFA every 3 months until all areas of BE were ablated or cancer developed showed that by 12 months after initial RFA treatment CR-D was achieved in 81% and CR-IM in 62% of patients (24). The registry’s later report in 2015 (consisting of 508 patients) showed a CR-D and CR-IM rates of 92% and 83%, respectively (3). There is increasing evidence to support the use of RFA (25) post-ER of any visible lesion in order to achieve CR-IM in the first 12-18 months post initial endoscopic ablation. Data is lacking on how often and at which interval RFA should be provided to these patients; however, our panel of experts suggest that an interval of 2-4 months would be acceptable practice.

<table>
<thead>
<tr>
<th>For patients undergoing RFA with a focal device, the dosimetry and treatment regimen is 12 J/cm² X 3, without interval cleaning and for patients undergoing RFA with a circumferential device the dosimetry and treatment regimen is 10J/cm², clean, 10J/cm²</th>
<th>8</th>
<th>0.4</th>
<th>No disagreement</th>
<th>1</th>
<th>No disagreement</th>
<th>N/A</th>
</tr>
</thead>
</table>

**Aspirational performance target: N/A**

**Evidence summary:**
Focal application of RFA without cleaning in between each ablation has been shown to be effective with 94% CR-D and 87% CR-IM with a stenosis rate of 11% (26). A multicentre randomised trial by Vilsteren et al showed that a simplified ablative regimen (3 X 15 J/cm²–no clean) is highly effective and can achieve higher complete remission of residual BE islands (73% vs 67%) than the standard method (2 X 15 J/cm²–clean–2 X 15 J/cm²) at 2 months (27). The same group was also able to show that the simplified regimen without cleaning was able to achieve higher BE surface regression (88% vs 83%) in comparison to the standard regimen in circumferential balloon based RFA with significantly shorter ablation time with the simplified technique (P < 0.01) (28). Furthermore, a multicentre RCT on focal RFA for dysplastic BE showed that the simplified RFA regimen (3 x 12 J/cm², without cleaning) is non-inferior to the standard regimen (2 x 15 J/cm², followed by cleaning, followed by 2 x 15 J/cm²) and therefore is the preferred RFA regimen for the management of patients with BE dysplasia (29).

The volume of evidence supporting the use of the circumferential RFA device in published literature is increasing. Recent data have shown a regression of 78% of BE segment at 3 months post ablation with the circumferential device using a dose of 12J and 85% regression with 10J (30). Furthermore a randomised trial in the Netherlands assessed treatment regimens for the 360 Express RFA balloon catheter (360 Express) using standard (1x10J/cm²-clean- 1x10J/cm²), simple-double (2x10J/cm²-no clean) and simple-single ablation regimen (1x10J/cm²-no clean). The simple-double arm of the study was terminated early as the result of significant severe stenosis; however, the study was able to show higher median BE regression in the standard arm compared to the simple-single group: 85% (IQR 75-94), 95% CI:78-92% versus 73% (IQR 48-90), 95% CI:59-85% (p=0.009) (30). It would therefore be appropriate to consider standard regimen (1x10J/cm²-clean- 1x10J/cm²) for the use of the circumferential RFA device.

<table>
<thead>
<tr>
<th>Centres undertaking BET should achieve CR-D ≥ 90% and CR-IM ≥ 80% within 18 months after the first treatment</th>
<th>8</th>
<th>0.4</th>
<th>No disagreement</th>
<th>1</th>
<th>No disagreement</th>
<th>N/A</th>
</tr>
</thead>
</table>

**Aspirational performance target: N/A**

**Evidence summary:**
The Ablation of Intestinal Metaplasia Containing Dysplasia (AIM) trial included a 5-year follow up analysis of patients with BE and dysplasia managed by RFA in a randomized controlled trial. Data showed BE recurrence after CR-IM by RFA in almost one-third of
patients with baseline dysplastic BE. Most recurrences occurred during the first year after CR-IM. However, patients that achieved CR-IM and remained BE free at 1 year after RFA had a low risk of BE recurrence (31). In addition, data from the UK RFA registry, the multicentre community practice registry, and the multicentre interventional EURO II study have all shown that ET is capable of achieving CR-D in 81-92% and CR-IM of 72-87% in patients with BE neoplasia at 12 months (3). Recent systematic reviews and a meta-analysis have also shown that EMR followed by RFA in patients with early BE neoplasia can achieve CR-D of 91-93% and CR-IM of 73-78% with 5-10% stricture rate, 1% bleeding rate and 0.2% perforation rate (22). Based on recent studies the expert panel suggests that centres undertaking BET should aim for CR-D > 90 % and CR-IM > 80 % at 18 months after the first treatment and end of treatment should be confirmed by 2 successive negative endoscopies after which patients should receive follow up endoscopies at appropriate intervals stratified according to risk of recurrence. The expert panel agreed that 18 months-time point is appropriate as standard clinical practice cannot always ensure timely visits and a 12month time point would be too restrictive.

Patients with residual dysplasia after 18 months, are to be re-discussed at a Oesophago-Gastric MDT

<table>
<thead>
<tr>
<th>Post-Procedure QIs</th>
<th>Median Score</th>
<th>MAD-M</th>
<th>BIOMED Analysis</th>
<th>P-value</th>
<th>IPRAS Analysis</th>
<th>Performance Threshold Median % (Range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Following successful BET, patients undergo follow up endoscopies at appropriate intervals stratified according to risk of recurrence</td>
<td>9</td>
<td>0.6</td>
<td>No disagreement</td>
<td>1</td>
<td>No disagreement</td>
<td>90 (80, 100)</td>
</tr>
</tbody>
</table>

Aspirational performance target: 90% (range: 80-100)

Evidence summary:

ET does not eliminate the need for continued endoscopic surveillance or completely eliminate the risk of synchronous or metachronous disease. Particular concern remains over IM, which is buried under neo-squamous epithelium after ET. This is a rare but recognized finding. The identification of these cases indicates the need for continued surveillance following RFA therapy, even after
CR-IM. Increasing age and length of BE segment are associated with a longer time to achieve CR-IM. It is therefore essential to continue surveillance after RFA. By dividing patients into simple categories, clinicians may stratify risk to choose the appropriate surveillance regimen. A large prospective study by Shaheen et al has shown impressive CR-D and CR-IM rates at 2 years (CR-D 95% and CR-IM 93%) and 3 years (CR-D 98% and CR-IM 91%) post initial BET with an annual rate of neoplastic progression of 1.37% per patient-years (25). Phoa et al also showed a 90% remission at 5 years post BET (35). The UK RFA registry has demonstrated a risk of neoplasia recurrence of 19% at 5 years with the predicted risk of IM recurrence at 13% at 26 months with a 32% risk of IM recurrence at 5 years (3).

The literature supports an IM/neoplasia recurrence rate between 10-32% at 5 years. Therefore, follow up post endoscopic therapy of BE neoplasia is needed to exclude recurrence and to deliver further therapy as needed (2). A recent study by Cotton et al provided evidence-based surveillance intervals after completion of ET in patients with BE neoplasia. For patients with LGD the group proposed surveillance endoscopy at 1 and 3 years after achieving CR-IM with ET. For patients with HGD or IMC, the proposed surveillance endoscopy was at 3 months, 6 months and 1 year and then annually (for 5 years) after achieving CR-IM with ET (36). Based on recent evidence, our expert panel felt that it would be reasonable to consider endoscopic follow-up proposed by Cotton et al (36).

At follow up endoscopy, biopsies should be taken from the Squamo-columnar junction and within the extent of the original BE length, for the first 2 years; thereafter biopsies should be taken from the Squamo-columnar junction and any visible lesion

| At follow up endoscopy, biopsies should be taken from the Squamo-columnar junction and within the extent of the original BE length, for the first 2 years; thereafter biopsies should be taken from the Squamo-columnar junction and any visible lesion | 8 | 0.6 | No disagreement | 1 | No disagreement | 90 (80, 100) |

Aspirational performance target: 90% (range: 80-100)

Evidence summary:

Adherence to biopsy protocol will significantly increase the detection rate of dysplasia in patients with BE. IM can reoccur at the gastro-oesophageal junction in the absence of visible BE following the successful eradication of BE neoplasia. Recent studies have suggested evidence of buried glands post BET in 5.5-7% of patients, but the majority of these were not detectable at subsequent endoscopies (37). Our expert panel suggests that endoscopic follow-up should include biopsies at the GOJ and within the previous extent of the BE epithelium (2). This should include a high resolution gastroscope to assess the treated and remaining area of BE (5). In order to exclude synchronous neoplastic lesions, 4 quadrant biopsies should be performed at 1–2-cm intervals throughout the entire BE segment (5).

Table 6: Quality indicators ranked as uncertain after Round 1 voting with the median score, MAD-M, BIOMED Analysis, p-value, IPRAS analysis

<table>
<thead>
<tr>
<th>Quality indicators</th>
<th>Median Score</th>
<th>MAD-M</th>
<th>BIOMED Analysis</th>
<th>P-Value</th>
<th>IPRAS Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-Endoscopy</td>
<td>Before undertaking EET, endoscopists need to have attended BET academia platforms.</td>
<td>7</td>
<td>1.2</td>
<td>Disagreement</td>
<td>0.83</td>
</tr>
<tr>
<td>Intra-procedure (Ablation)</td>
<td>It is recommended that prior to starting BET, a minimum of 30 supervised cases of endoscopic resection and 30 cases of endoscopic ablation should be performed to acquire competence in technical skills, management pathways and complications.</td>
<td>7</td>
<td>1.2</td>
<td>Disagreement</td>
<td>0.83</td>
</tr>
<tr>
<td>Post Procedure</td>
<td>For patients undergoing RFA with a circumferential device, the recommended dose is 10 J/cm2 CLEAN 10J/cm2 (EXPRESSION)</td>
<td>7</td>
<td>1.5</td>
<td>Disagreement</td>
<td>0.83</td>
</tr>
<tr>
<td></td>
<td>Following successful eradication after BET, patients should undergo follow up surveillance endoscopies at 3, 6, 9, 12 months and then annually (if fit for endoscopy)</td>
<td>8</td>
<td>1.7</td>
<td>Disagreement</td>
<td>0.51</td>
</tr>
</tbody>
</table>
Figure Legend:

Figure 1: RAND/UCLA Appropriateness Method (RAM) - Summary
Figure 1: RAND/UCLA Appropriateness Method (RAM) - Summary

- Recruitment of International Experts
- Round 0 Meeting (Familiarity with RAM)
- Generation of List of Potential Quality Indicators (QI) for 4 groups: pre-endoscopy, intra-procedure (resection), intra-procedure (ablation) and post-procedure
- QIs Proposed (n=24)
- Round 1: Independent electronic voting
- Analysis of Round 1 Voting and Literature Search
- QIs Appropriate (n=20), Inappropriate (n=0), Uncertain (n=4)
- Round 2: Panel Meeting (in-person discussion, re-wording, re-ranking)
- QIs Proposed (n=15)
- Determine Overall Appropriateness (Based on RAM Scoring Guide)
- Formally Validated Quality Indicators for EET:
  - Pre-procedure QIs: 2
  - Intra-procedure QIs (Resection): 5
  - Intra-procedure QIs (Ablation): 6
  - Post-procedure QIs: 2