Methods Data on consecutive patients who underwent endoscopic full thickness resection (eFTTR) at two UK teaching hospitals in November - December 2019 were analysed. The procedure was undertaken using the endoscope mounted gastroduodenal FTRD®. Main outcome measures were technical success (target lesion resection with FTRD®), total procedural time, specimen size, R0 resection, and adverse events. Need for dilatation to facilitate passage of device past cricopharyngeus or the pylorus was also documented.

Results All cases were undertaken under general anaesthetic. It was possible to insert the device to the lesion in all cases; in two, dilatation of the pylorus with a 20 mm through the scope balloon was required to facilitate passage of the device to the duodenum. Technical success and histological diagnosis were achieved in 5/5 (100%) cases. Median total procedural time was 23 minutes (range 18–65). Baseline and outcome data of the cases can be seen in table 1.

Two patients were kept for overnight observation and three were discharge on same day as the procedure. One patient reported shivering post procedure, which was thought to be general anaesthesia related, otherwise there were no immediate or delayed complications.

Conclusions eFTTR of SELs or heavily scarred lesions in the stomach and duodenum is feasible and safe with the gastroduodenal FTRD®. It facilitates acquisition of definite histology aiding diagnosis and R0 resection is possible, providing treatment or avoiding need for ongoing surveillance in selected patients. The device can be challenging to insert and in particular, pre-dilatation of the pylorus to facilitate insertion into the duodenum may be required.


REFERENCES

P15

SHOULD POST COLONOSCOPY COLORECTAL CANCER BE ADDED TO INDIVIDUAL’S KEY PERFORMANCE INDICATORS?

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Introduction Post colonoscopy colorectal cancer (PCCRC) is associated with a number of contributory factors, including poor endoscopist performance metrics. The JAG Global Rating Scale (GRS) tool currently only includes 8-day unplanned admission rates and 30-day mortality rates as a late outcome quality Key Performance Indicator (KPI) to feedback to individual endoscopists. Since 2016, units have been encouraged to subject each PCCRC to a root cause analysis (Rees CJ, et al. Gut 2016;65:1923–1929). We aimed to review PCCRC in our institution with a focus on KPI and specialty.

Methods This was a retrospective review of PCCRC over a 3-year period (Jan 2017-Dec 2019) at a 2-site hospital in North London serving a population of 500,000. All patients with an endoscopic diagnosis of colorectal cancer (CRC) were identified from the Unisoft GI reporting tool. Patients who had a prior colonoscopy within 3 and 5-years were identified (index colonoscopy). A definite missed cancer was considered as a PCCRC within 3 years of the index scope, and a probable missed cancer was within 5 years.

Results CRC was diagnosed in 618 patients, of whom 3.7% (23) were identified as having a PCCRC [Female 48%; mean age 75 years; 56–90 years]. The ‘definite miss rate’ was 2.1% (13/618) and the ‘probable miss rate’ was 1.6% (10/618). The mean time lag from index scope to CRC was 34.4 months. A definite missed cancer was identified within 3 years (Rees CJ, et al. Gut 2016;65:1923–1929).

Conclusions Our study demonstrates that our department meets quality standard for units which includes a target of <5% PCCRC at 3 years (Rees CJ, et al. Gut 2016;65:1923–1929). However, review of individual cases identified higher rates of PCCRC amongst endoscopists with lower volumes of procedures and sub-optimal Adenoma Detection Rates (ADR).

P16

ACCEPTABILITY OF KEY PERFORMANCE INDICATORS (KPI) IN THE NATIONAL ENDOSCOPY DATABASE (NED), A DELPHI PROCESS

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Introduction Automated Performance Reports to Improve Quality Outcomes Trial (APRIQOT) uses NED to provide endoscopists feedback on colonic detection KPI. Traditional adenoma detection rate is dependent on unavailable histological data. Our aim was to gain expert consensus on which available KPI are acceptable to endoscopists.

Method A Delphi panel of UK expert endoscopists was recruited online, purposively for professional background. Panellists interacted using an online form. In round one we provided a summary and acceptability statement for each KPI, participants rated agreement with a 5 point Likert scale and free-text comments. Responses were analysed anonymously. In subsequent rounds participants reviewed all graded consensus statements and comments. Statements were accepted with ≥80% consensus or redrafted. Rounds ran January to April 2019.

Results We recruited 21 UK expert endoscopists. 12 were female, 48% gastroenterology background, 29% nursing, 14% surgical and 9% trainees. All statements reached consensus by round 3 (Table 1). The panel agreed KPI adjusted for age, sex and indication were ‘more acceptable’. Polyp measure had risks of ‘gaming’ and distal hyperplastic polyp over reporting, but encompass significant non-adenomatous polyps. Mean number of polyps (MNP) reached consensus after discussing reduction of the ‘one and done’ phenomenon and using a cap of 5 polyps/colon to mitigate skew from polyposis. Proximal