Multicentre prospective study of COVID-19 transmission following outpatient GI endoscopy in the UK

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MESSAGE
The COVID-19 pandemic has severely curtailed the practice of endoscopy (as an exemplar for outpatient diagnostic procedures) worldwide. Restart and recovery processes will be influenced by the need to protect patients and staff from disease transmission, but data on the risk of COVID-19 transmission after an endoscopy are sparse. This is of particular importance in later pandemic phases when the risk of harm from delayed or missed significant diagnoses is likely to far outweigh the risk of infection. The British Society of Gastroenterology guidance for restarting endoscopy included the stratification of diagnostic procedures according to aerosol generation or assessment of infectious risk as well as pragmatic guidance on the use of personal protective equipment (PPE). We sought to document the risk of COVID-19 transmission after endoscopy in this ‘COVID-minimised’ environment. Prospective data were collected from 18 UK centres for n = 6208 procedures. Pre-endoscopy, 3 of 2611 (0.11%; 95% CI 0.00%–0.33%) asymptomatic patients tested positive for SARS-CoV-2 on nasopharyngeal swab. Based on follow-up telephone symptom screening of patients at 7 and 14 days, no cases of COVID-19 were detected by any centre after endoscopy in either patients or staff. Although these data cannot determine the relative contribution of each component of a COVID-minimised pathway, they provide clear support for such an approach. The rational use of PPE and infection control policies should be continued and will aid in planning for outpatient diagnostics in the COVID-19 recovery phase.

IN MORE DETAIL
The COVID-19 pandemic has had an extraordinary impact on the delivery of GI endoscopy, with an initial reduction to 12% of prepandemic levels in the UK. In the deceleration and early recovery phases (up to end July 2020), this had risen to 42% of prepandemic levels. Recovery has been influenced by multiple factors including availability of staff, restrictions caused by longer room cleansing, physical distancing and the use of personal protective equipment (PPE) slowing lists. There are grave implications of this contraction in activity, with the delayed diagnosis of significant conditions like GI cancer or inflammatory bowel disease of particular concern.

The risk around the inadvertent periprocedure transmission of COVID-19 infection to both patients and staff is a primary concern, but is not well described. Early data from northern Italy described low rates of infection in patients and staff, even during the peak phase of the pandemic.

Upper GI endoscopy is widely accepted to be an aerosol-generating procedure (AGP); however, the relevance of small-volume aerosols (ie, 0.3 μm; which appear to predominate) for virus transmission is unclear. Infectious potential appears to be confined to particles 0.5 μm or larger, but this is a complex subject with a number of variables. There are also direct data to support the effectiveness of masks, including surgical face masks most widely used in endoscopy units, in preventing viral transmission.

COVID-19 infection rates have been demonstrated to be lower-than-expected in endoscopy staff (compared with other healthcare workers) even when the so-called ‘high-risk’ PPE, particularly face masks, were not universally available or applied. Whether lower GI endoscopy is an AGP is also important but has been pragmatically regarded as having low infectious potential as per British Society of Gastroenterology (BSG) guidance, whereas staff are still advised to use appropriate (stratified) PPE for all procedures.

Significant patient anxiety regarding the potential for contracting COVID-19 infection also exists and this has also been demonstrated to influence the ability to provide effective diagnostic services.

We therefore sought to study prospectively the number of patient infections following GI endoscopy from multiple centres across the UK through the peak, deceleration and early recovery phases of the pandemic. Taking into account the complexities of infection control, aerosols, infectivity and abrogation of risk by PPE, it was felt that the ultimate determinant of risk would be whether any COVID-19 cases were actually detected after endoscopy (in either patients or staff). The purpose of this study was to enable the quantification of the potential risk to patients and to inform endoscopy departments regarding the likelihood of transmitting the infection to patients. These data could be used to help communicate with patients and staff as well as inform planning for future outbreaks.

This multicentre prospective study collected data for consecutive outpatients attending for elective diagnostic or therapeutic endoscopy from 18 UK centres. No patient identifiable data were collected,
no treatment decisions were affected and no identifiable data were used, analysed or transferred. As such ethical approval was deemed not to be required by the Research Governance committee at the lead author’s institution.

Centres were selected to reflect differently sized units, tertiary and local, covering a wide range of demographics including those serving mixed socioeconomic populations and a mix of ethnicities. All centres prospectively completed an anonymised database of patients including procedure type, responses to preprocedure SCOTS criteria, preprocedure nasopharyngeal swab test result (if performed in that unit), source of referral and dates for all activities. The SCOTS criteria comprise telephone screening questions around Symptoms, infectious Contacts, Occupational risk, Travel risk, Shielding status and are recommended in BSG guidance. These were developed as an update to a pre-existing screening tool (FTOCC; developed in Hong Kong during the first SARS outbreak and proposed for using during the COVID-19 pandemic), to take into account considerations specific to COVID-19.

All centres conducted patient follow-up by telephone consultation at 7 and 14 days after the procedure to check for symptoms of COVID-19. If symptoms were reported that could be consistent with COVID-19, then these patients were advised to undergo COVID-19 nasopharyngeal swab testing. Data were collected on any patients with positive COVID-19 swab undertaken for any reason in the 14 days postendoscopy. Where patients were COVID-19 swab tested preprocedure and found to be positive, those patients were excluded from the follow-up study.

Data were collected from 6208 patients undergoing endoscopy at 18 centres between 30 April and 30 June 2020 (mean±SD age 59.3±15.4 years and n=2973 (48%) were female individuals).

The characteristics of the endoscopy units are shown in table 1. Follow-up data on symptoms were collected up to 14 July 2020. There were no cases of COVID-19 detected in the 2 weeks following endoscopy (0/6208, 95% CI 0.0%–0.08% with continuity correction).

Figure 1 shows the procedures performed and overall % of total. There was an approximate 40:60 split between upper and lower GI procedures (where combined oesophagogastroduodenoscopy and colonoscopy counted as upper GI—given the AGP status of the former procedure and therefore potential for greater risk).

Most centres were prioritising urgent symptomatic patients during this period with less than 4% (n=233) of procedures classed as ‘routine or surveillance’ and these were performed at the discretion of the performing centre, mostly within the last 2 weeks of the data collection period. Of the remainder, n=3166 (51%), were patients referred for suspected cancer on the pre-existing UK ‘2-week wait’ pathways (again split 60:40 in favour of lower GI). There were 1193 (19.0%) in a ‘shielded’ category as determined by the preprocedure telephone questionnaire.

Preprocedure nasopharyngeal swab testing for SARS-CoV-2 was performed in 2611 patients from 13 sites, in line with BSG guidance, after the confirmation of the absence of COVID-19 symptoms or risks. Only 3 (0.11%, 95% CI 0.03%–0.36% with continuity correction) patients were positive. All had their investigation safely deferred with no complications.

Following their procedure, 12 patients reported symptoms that were potentially compatible with COVID-19 infection at either the 7-day or 14-day telephone contact. All then underwent nasopharyngeal swab testing and were found to be negative. All symptoms settled with none deemed to have COVID-19. There

### Table 1 Characteristics of units submitting data

| Site | Centre type | Annual procedure count (pre-covid) | Cases submitted | Pre-endoscopy actions | Stratified use* of |
|------|-------------|-----------------------------------|-----------------|-----------------------|-------------------|
|      |             |                                   |                 | NP swab? | Mandatory self-isolation for patients (≥7 days)? | PPE | IPCPs |
| A    | T           | 20000                             | 198             | Yes       | Yes               | Yes | Yes   |
| B    | T           | 17000                             | 111             | No        | No                | Yes | Yes   |
| C    | L           | 20000                             | 436             | No        | No                | Yes | Yes   |
| D    | L           | 9800                              | 365             | Yes       | Not†              | No  | No    |
| E    | T           | 15000                             | 374             | Yes       | No                | Yes | Yes   |
| F    | L           | 7500                              | 58              | Yes       | No                | No  | No    |
| G    | L           | 9800                              | 206             | Yes       | No                | Yes | Yes   |
| H    | T           | 17000                             | 218             | Yes       | Not†              | Yes | Yes   |
| I    | L           | 15200                             | 351             | Yes       | No                | Yes | Yes   |
| J    | L           | 17400                             | 175             | Yes       | Yes               | Yes | Yes   |
| K    | T           | 15000                             | 84              | Yes       | No                | No  | No    |
| L    | T           | 15600                             | 950             | No        | No                | Yes | Yes   |
| M    | T           | 20000                             | 363             | Yes       | No                | Yes | Yes   |
| N    | T           | 3025                              | 219             | Yes       | No                | No  | No    |
| O    | T           | 15000                             | 715             | No        | No                | No  | No    |
| P    | L           | 24000                             | 672             | No        | No                | No  | No    |
| Q    | T           | 14000                             | 660             | No        | No                | Yes | Yes   |
| R    | L           | 20000                             | 251             | Yes       | Not†              | Yes | Yes   |
| Total| 10T, 8L     | 302825                            | 6208            | 12Y       | 16N               | 12Y | 12Y   |

*Stratified use refers to the differential use of ‘low-risk’ and ‘high-risk’ principles according to upper vs lower GI endoscopy and COVID19-confirmed vs COVID19-excluded patients (ie, a COVID-minimised pathway as explained in references 4, 5 and 11).
†Sites suggested, but did not mandate, self-isolation of patients for 7 days preprocedure.
IPC Ps, infection prevention and control policies; L, local; NP, nasopharyngeal; PPE, personal protective equipment; T, tertiary.
deceleration/recovery phases of the COVID-19 pandemic in the UK and should therefore inform periods of activity where similar rates of infection are seen.

When a COVID-minimised pathway is in place, patients (including those in a high-risk ‘shielding’ category) can now be reassured that concerns over COVID-19 infection should not stop them attending for GI endoscopy.

Correction notice This article has been corrected since it published Online First. The collaborator names have been updated.

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REFERENCES

1 Rutter MD, Brookes M, Lee TJ, et al. Impact of the COVID-19 pandemic on UK endoscopic activity and cancer detection: a national endoscopy database analysis. Gut 2021;70:537–43.

2 National endoscopy database. Available: www.ned.jets.nhs.uk [Accessed 01 Aug 2020].

3 Lai A, Denaxas S, Dave C. Estimating excess mortality in people with cancer and multimorbidity in the COVID-19 emergency 2020.

4 Rees CJ, East JE, Oppong K, et al. Restarting gastrointestinal endoscopy in the deceleration and early recovery phases of COVID-19 pandemic: guidance from the British Society of gastroenterology. Clin Med 2020;20:352–8.

5 Hayee Bu’Hussain, Thoufeeq M, Rees CJ, et al. Safely restarting GI endoscopy in the era of COVID-19. Gut 2020;69:2063–70.
