NORCCAP (Norwegian colorectal cancer prevention): a randomised trial to assess the safety and efficacy of carbon dioxide versus air insufflation in colonoscopy

M Bretthauer, E This-Evensen, G Huppertz-Hauss, L Gisselsson, T Grotmol, E Skovlund, G Hoff

Background: To eliminate the risk of combustion during electrosurgical procedures and to reduce patient discomfort, carbon dioxide (CO₂) insufflation has been recommended during colonoscopy. However, air insufflation is still the standard method, perhaps due to the lack of suitable equipment and shortage of randomised studies.

Aims: This randomised controlled trial was conducted to assess patient tolerance and safety when using CO₂ insufflation during colonoscopy.

Patients: Over an eight month period a successive series of patients referred for a baseline colonoscopy due to findings in a flexible sigmoidoscopy screening trial were randomly assigned to the use of either air or CO₂ insufflation during colonoscopy.

Methods: End tidal CO₂ (ETCO₂), a non-invasive parameter of arterial pCO₂, was registered before and repeatedly during and after the examination. The patient’s experience of pain during and after the examination was registered using a visual analogue scale (VAS). Sedation was not used routinely.

Results: CO₂ insufflation was used in 121 patients (51%) and air in 119 patients (49%). The groups were similar in age, sex, and caecal intubation rate. No rise in ETCO₂ was registered. There were statistically significant differences in VAS scores between the groups with less pain reported when using CO₂.

Conclusions: This randomised study of unsedated patients shows that CO₂ insufflation is safe during colonoscopy with no rise in ETCO₂ level. CO₂ was found to be superior to air in terms of pain experienced after the examination.

This trial compared CO₂ with air insufflation in colonoscopy with regard to patient pain during and after the examination and investigated whether CO₂ insufflation leads to a rise in body CO₂ level.

METHODS

Attendees

The NORwegian Colorectal CAncer Prevention (NORCCAP) study is an ongoing screening trial for the prevention of colorectal cancer. Fourteen thousand presumptively healthy men and women, aged 55–64 years, living in two separate areas in Norway are randomly drawn from the population registry and invited to undergo a screening flexible sigmoidoscopy (FS) for colorectal adenomas and cancer. Patients with former colonic resections, severe heart or lung disease (New York Heart Association (NYHA) III-IV), or ongoing treatment for malignant disease are excluded. Biopitically verified adenoma at screening FS, irrespective of size, qualifies for a baseline colonoscopy with polypectomy, to be performed within 6–8 weeks after FS.

NORCCAP participants referred for colonoscopy between October 1999 and April 2000 (267 patients) were included in the present study. All examinations were performed by one of three experienced endoscopists. According to recently published guidelines, colonoscopies were performed without any

Abbreviations: ETCO₂, end-tidal carbon dioxide; FS, flexible sigmoidoscopy; NORCCAP, NORwegian Colorectal CAncer Prevention; NYHA, New York Heart Association; VAS, visual analogue scale.
routine use of sedation. However, on demand medication with intravenous midazolam was given if indicated, as judged by the endoscopist. For bowel cleansing, a 4 litre polyethylene glycol solution was used, taken orally on the day before the examination.

Randomisation

Participants referred for colonoscopy were successively assigned to appointments as referrals were received and no sessions were available other than for participants in the NORCCAP screening study. Whole day sessions for colonoscopy were randomised for CO2 or air insufflation, using sealed envelopes. Randomisation of whole sessions rather than individual patients was done to avoid unblinding by change of gas couplings between patients. Both participants and endoscopists were blinded with regard to which gas was being used.

Endoscopic examination

Procedures were performed using Olympus video colonoscopes (Olympus, Hamburg, Germany). However, the standard gas/water valves of the endoscopes which only redirect a continuous gas flow into the gut lumen or atmospheric air, were replaced with another type of valve (MIJ-521, Olympus) preventing gas leakage into the environment. To administer gas into the colon, the valve button had to be pushed halfway down. CO2 or air was administered using two different pressure and flow controlled devices connected to the CO2 and air reservoirs provided (Endoscopic CO2 Regulator; Key Med Ltd (Southend-on-Sea, Essex, UK) for CO2, Norsk Hydro Ltd (Oslo, Norway) for air). The endoscopy assistant was responsible for switching on and off the CO2 and air devices, respectively. To prevent unblinding, the devices were placed behind the endoscopy rack and hidden from the view of the endoscopist.

End-tidal CO2 measurements

End-tidal (ET) CO2 has been shown to give adequate approximations of arterial pCO2 in spontaneously breathing adults and is therefore a good and commonly used non-invasive method of expressing arterial pCO2.

How was the CO2 insufflation conducted in the study? Before starting the trial we conducted ETCO2 measurements and arterial pCO2 samples simultaneously on five consecutive patients. The two methods were comparable, with a maximum deviation of the ETCO2 value of only 0.2 kPa. Additionally, we performed repeated measurements within the group of investigators to test the validity of the method, with no evidence of method failure.

ETCO2 was measured successively: (1) at the start of each examination, (2) when the endoscope had reached the caecum, (3) when the rectum was passed during withdrawal, and (4) 10 minutes after finishing the examination. At these measuring points, patients were asked to take a deep breath and to expire deeply and slowly through the mouthpiece of the provided mainstream infrared capnograph (Novametrix Ltd, Wallingford, Connecticut, USA). The endoscopy assistant performed the measurements and registered the readings. Both the participant and endoscopist were blinded to the results.

Measurement of pain

Pain was registered on a questionnaire given to participants immediately after the examination. A 100 mm visual analogue scale (VAS) was used, ranging from “no pain” on the left to “pain as bad as it could be” on the right end. Participants were asked to score the amount of pain experienced at one, three, six, and 24 hours after the examination. In addition, the questionnaire contained a similar VAS scale for the amount of pain experienced during the examination. Once completed, questionnaires were returned by mail to the screening centre.

Statistical analysis

A pilot study was conducted to estimate the SDs of the pain and ETCO2 measurements, respectively. Regarding the pain measurements, with an assumed SD of 30 mm, we estimated that 240 patients were needed to achieve at least 95% power to detect a 15 mm difference in VAS between the two groups, which was considered to be clinically important. Assuming that the difference in ETCO2 would have to be >0.5 kPa to be clinically important (SD 0.6), the power to detect this with 150 patients was 95%.

For statistical analysis of repeated measurements of pain and ETCO2, ANOVA for repeated measures was used. Some variables were not normally distributed and thus the Wilcoxon rank sum test was used as a supplementary analysis to compare groups at each time point. The proportion of individuals reporting no pain on the VAS was compared at each time point using the χ2 test. Statistical significance was defined as p<0.05. Only two sided tests were used. Statistical analyses were performed using SPSS 9.0.

Ethics

The regional ethics committee approved the study protocol. Informed consent was obtained from all participants before entering the trial.

RESULTS

A total of 267 patients were randomised and examined; 249 patients (93%) completed the questionnaire. Ten patients (seven in the air group and three in the CO2 group; p<0.01) received sedation and were excluded from further analysis. Thus 240 patients were included in the study; 121 (51%) were receiving CO2 insufflation and air was used in 119 patients (49%). There were no differences in baseline characteristics between the two groups (table 1). No statistically significant differences were observed between endoscopists. There was a trend towards more rapid caecal intubation in patients in the CO2 group (table 1).

The only complication registered was one perforation requiring colonic resection (air group). This occurred after snare polypectomy of a large sessile adenoma with severe dysplasia in the sigmoid colon.

End-tidal CO2

According to power estimates it was considered sufficient to measure ETCO2 in a limited number of patients and hence ETCO2 measurements were restricted to the first 156 examinations performed without sedation, including 81 patients in the air group and 75 patients in the CO2 group. As shown in fig 1, there was no rise in ETCO2 during or after the examination in any group. On the contrary, we observed a significant reduction in ETCO2 levels during examination in both groups (p<0.001). This reduction was more pronounced when CO2 was used. The time point differences between the groups reached significance only for ETCO2 readings registered towards the end of the examination (p=0.01) (fig 1). However, this difference was estimated to be 0.28 kPa (95% confidence interval (CI) 0.06–0.49) and thus far below the

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Participant and examination characteristics in the CO2 and air groups</th>
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<tbody>
<tr>
<td></td>
<td>CO2 group (n=121)</td>
</tr>
<tr>
<td>Sex (M/F)</td>
<td>77/44</td>
</tr>
<tr>
<td>Mean age (y)</td>
<td>59.5</td>
</tr>
<tr>
<td>Caecum reached [No (%)]</td>
<td>109 [90]</td>
</tr>
<tr>
<td>Time to caecum [min (mean SD)]</td>
<td>13.1 [7.6]</td>
</tr>
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</table>

*p<0.09.
Pain during and after examination

Figure 1 Mean (SEM) end-tidal CO₂ values at the various observation points in the CO₂ (n=75) and air (n=81) groups. **p=0.01 compared with the CO₂ group at the corresponding time point by repeated measures ANOVA with multiple comparisons.

Figure 2 Mean (SEM) visual analogue scale (VAS) scores at the various observation points during and after examination in the CO₂ (n=121) and air (n=119) groups. *p<0.05, ***p<0.001 compared with the CO₂ group at corresponding time points by the Wilcoxon rank sum test.

Figure 3 Percentage of patients in the CO₂ (n=121) and air (n=119) groups who scored 0 (no pain) on the visual analogue scale (VAS) at the observation points during and after examination. Values are mean (SD). ***p<0.001 compared with the CO₂ group at corresponding time points by the χ² test.

There was a statistically significant difference in pain scores, favouring CO₂ insufflation at all observation points after examination. The overall mean difference was 7.8 mm (95% CI 4.4–11.2) (p<0.001). The pain reduction after examination was significantly more rapid in the CO₂ group (p=0.003). The maximum difference (14 mm (95% CI 9–19); p<0.001) was observed one hour after the examination. Comparison of the two groups by non-parametric analysis at each separate time point gave results similar to the overall analysis.

An alternative visualisation of the pain score results is to compare the proportion of patients with score zero (no pain) on the VAS. Figure 3 shows these proportions at each measurement. The finding of a clinically relevant difference in pain, favouring the use of CO₂ was supported.

DISCUSSION

This randomised double blind trial in unsedated patients showed that CO₂ insufflation during colonoscopy reduced the amount of pain during and after the examination. No rise in ETCO₂ during or after the examination was registered.

The current study the use of sedation differed between the two groups, with more individuals in the air group requiring sedation. If these patients had not been excluded from the present analyses, the observed differences would have been somewhat larger. Although the need for on demand administration of sedation in this study was judged subjectively by the endoscopist, the endoscopist’s observation of the patient’s need was consistent with the differences between the groups in VAS scores given by the patients.

End-tidal CO₂

The ideal gas for insufflation during colonoscopy should be inert. However, this gas has yet to be found. CO₂ is not ideal as it interferes with normal metabolic processes. It has been
known for a long time that intraperitoneal CO₂ insufflation during laparoscopic surgery causes a rise in arterial pCO₂ levels but this side effect is probably less marked in retroperitoneal compared with intraperitoneal insufflation. Furthermore, a laparoscopic procedure is quite different from colonoscopy (mechanical ventilation, Trendelenburg position, CO₂ kept under a positive pressure). Therefore, there was a need to investigate the effect on pCO₂ during insufflation of CO₂ in colonoscopy. To measure arterial pCO₂, arterial blood samples are needed. This was considered impractical in the present study. We used ETCO₂ measurement as an approximation of arterial pCO₂. Continuous measurement of ETCO₂, often through a nasal or nasopharyngeal canula, has been the preferred method in other studies. In the present study patients were awake, non-sedated, and would probably not have tolerated such canulae. We therefore measured ETCO₂ repeatedly using a mouthpiece connected to a capnograph. This method may not be as accurate as those published previously but it is an easily performed approximation to arterial pCO₂ in this setting. The purpose of our measurements was to exclude a clinically significant rise in body CO₂ level, not to evaluate exact values within the reference area.

We did not detect any rise in ETCO₂ levels. On the contrary, we observed an overall decrease in ETCO₂ during and after the procedure, slightly more marked in the air group. A possible explanation for this is that patients may have hyperventilated during the procedure, and that insufflation of CO₂ to some extent outweighed this reduction in ETCO₂. Rogers reported a rise, although not statistically significant, in arterial pCO₂ during colonoscopy. In that study, all patients were sedated and sedation is known to cause changes in pCO₂ and other metabolic parameters. Hence these results are difficult to compare with ours.

No patient with severe heart or lung diseases (NYHA III-IV) was included in this study. Therefore, our results cannot be generalised to this patient categories. The safety for these patients and for sedated patients has to be investigated further before extending recommendations to other than our described patient population.

CONCLUSIONS

This randomised trial of unsedated patients showed that CO₂ insufflation during colonoscopy is safe with no rise in ETCO₂ levels. CO₂ was found to be superior to air regarding pain after the examination. In this study, CO₂ insufflation led to an almost complete absence of post examination pain. We recommend CO₂ insufflation in colonoscopy.

ACKNOWLEDGEMENTS

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