The AirSeal® insufflation device can entrain room air during routine operation

R. P. Weenink1 · M. Kloosterman2 · R. Hompes3 · P. J. Zondervan4 · H. P. Beerlage4 · P. J. Tanis3 · R. A. van Hulst5

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Abstract
Background Surgical procedures that use insufflation carry a risk of gas embolism, which is considered relatively harmless because of the high solubility of carbon dioxide. However, an in vitro study suggested that valveless insufflation devices may entrain non-medical room air into the surgical cavity. Our aim was to verify if this occurs in actual surgical procedures.

Methods The oxygen percentage in the pneumoperitoneum or pneumorectum/pneumopelvis of eight patients operated with use of the AirSeal® was continuously measured, to determine the percentage of air in the total volume of the surgical cavity.

Results Basal air percentage in the surgical cavity was 0–5%. During suctioning from the operative field air percentage increased to 45–65%.

Conclusions The AirSeal® valveless insufflation device maintains optimal distension of the surgical cavity not only by insufflating carbon dioxide, but also by entraining room air, especially during suctioning from the operative field. This may theoretically lead to air embolism in patients operated on with this device.

Keywords Laparoscopy · Air embolism · Transanal endoscopic surgery · Insufflation

Introduction

Venous gas embolism (GE) can occur as a complication of surgical procedures that utilize insufflation. The reported incidence varies widely, and probably depends on the type of procedure and the method used to detect embolization. When surgeons are retrospectively asked to recall clinically significant GE during transanal total mesorectal excision (TaTME), incidence is 0.45% [1]. On the other hand, transesophageal echocardiography detects gas bubbles in up to 100% of laparoscopic hysterectomies [2]. Irrespective of the incidence of intraoperative GE, these bubbles are believed to be subclinical, unless they amount to such a volume that they cause massive pulmonary GE or cardiac air lock. This is based on the assumption that the bubbles consist of carbon dioxide (CO₂), which, due to its high solubility, will resorb before irreversible damage has occurred.

Recently, Huntington and colleagues published an in vitro study demonstrating that valveless insufflation devices may entrain room air into the surgical cavity [3]. This would put patients at risk of air embolism, which may lead to more severe symptoms because of slower speed of resorption. In this study, we investigated whether the findings of this in vitro study would be replicable during actual surgical procedures.

Materials and methods

Eight patients who underwent surgery using the AirSeal® (ConMed, Utica, New York, United States) were included. After induction of general anesthesia, a pneumoperitoneum
or pneumoretum/pneumopelvis was created. In cases that
only involved a pneumoperitoneum, the AirSeal® insuffla-
tion tubing was attached to an AirSeal® 8 mm access port
placed through the abdominal wall. In combined transanal
and abdominal procedures, the abdominal part of the pro-
cedure was initiated with a standard insufflation device. For
the transanal part of the procedure, the insufflation tubing
of the AirSeal® was attached to an AirSeal® 8 mm access
port inserted through the gelcap of a GelPOINT path trans-
al access platform (Applied Medical Resources Corpora-
tion, Rancho Santa Margarita, California, United States).
An insufflation pressure of 12 mmHg was used uniformly.

During use of the AirSeal®, a sampling line with an inner
diameter of 1.3 mm was placed 10–20 cm into the surgi-
cal cavity, either by passing it next to one of the trocars, or
through the gel cap of the access platform. Absence of leak-
age of gas alongside the sampling line was confirmed. If the
presence of the sampling line obstructed the surgeon’s view
of the operating field, it was attached to one of the luer lock
valves of the access platform (in transanal surgery) or the
luer lock connector on one of the trocars (in laparoscopic
surgery) (Fig. 1).

The sampling line was attached to a calibrated gas ana-
lyzer (Fluke Corporation, Everett, Washington, United
States). The difference between the pressure in the surgical
cavity and atmospheric pressure created a small flow from
the surgical field to the gas analyzer, which continuously
measured the oxygen concentration in the sample. During
the procedure, surgical manipulation such as insertion or
removal of an instrument, or suctioning from the oper-
ative field was noted. Data analysis was done in MATLAB
R2019b (MathWorks, Natick, Massachusetts, United
States). The amount of room air as a percentage of the
total volume of gas in the surgical cavity at any point in
time was calculated as the oxygen percentage multiplied by
4.78, based on the assumptions that (1) room air contains a
constant 20.9% of oxygen (100/20.9 = 4.78); (2) room air
was the only source of oxygen in the surgical cavity; (3)
the gas in the surgical cavity was of homogenous composi-
tion; and therefore (4) the gas sample as collected by the
analyzer was representative of the contents of the surgical
cavity.

Results

Details of the eight patients included in this study are pro-
vided in Table 1. All surgical procedures were uneventful;
specifically, no signs of GE were observed. During normal
surgical conditions, the amount of room air as a percentage
of total volume of the surgical cavity was 0–5%. However,
multiple increases in air percentage were seen in all proce-
dures. Some of these increases could be related to removal
and reinsertion of a surgical instrument through the laparo-
scopic ports. This lead to relatively small increases of air, up
to 10–25%, which immediately restored to baseline values
(Fig. 2). When suctioning with 40 l/min through a standard
5 mm laparoscopic aspirator was performed, the air percent-
age increased to higher values, around 45–65% (Figs. 3 and
4). Maximum observed flow rate of CO₂ as displayed on the
AirSeal® during suctioning was 9 l/min, while at the same
time no decrease in the distention of the surgical cavity was
notable. During suctioning, when the AirSeal® access port
was not occupied with a surgical instrument, flow of air from
the operating room into the surgical cavity could be felt by
placing a finger on the access port. Placement of the sup-
plied noise cancelling cap on the access port did not prevent
air entrainment. We observed no differences in measured

![Fig. 1 Experimental setup during laparoscopic surgery (left) and transanal surgery (right). (A) AirSeal® 8 mm access port with insuf-
flation tubing attached. (B) GelPOINT path transanal access platform. (C) Sampling line, passed next to the access port into the pneumoperitoneum. (D) Luer lock connection on the access platform to which the sampling line is attached (the sampling line itself cannot be seen)
values, whether the sampling line was placed in the surgical cavity, or attached to a luer lock connector on a trocar or on the access platform.

**Discussion**

Our data confirms the observation made in an in vitro study [3] that the valveless AirSeal® insufflation device can entrain room air into the surgical cavity. This occurs especially during suctioning from the operative field, but also with removal and insertion of instruments. Airflow into the surgical cavity could indeed be felt at the AirSeal® trocar during suctioning. The maximum air percentage of 65% corresponds with the value found in the in vitro study, when the same suctioning flow of 40 l/min was used [3].

CO₂ is the laparoscopic gas of choice because it is readily available, inert, non-combustible, and has a high solubility in blood. This latter property renders CO₂ relatively safe in cases of intravascular introduction of gas. Air is 20

Table 1 Details of the surgical procedures performed in 8 patients

| Patient | Sex | Age (years) | Indication | Procedure | Use of AirSeal |
|---------|-----|-------------|------------|-----------|---------------|
| 1       | M   | 65          | Renal cell carcinoma | Nephrectomy and adrenalectomy | Pneumoperitoneum |
| 2       | M   | 19          | Ulcerative colitis, status post subtotal colectomy | Completion proctectomy with ileoanal pouch | Pneumoperitoneum and pneumorectum/-pelvis |
| 3       | V   | 74          | Rectal fistula post rupture of sphincter with dynamic graciloplasty and sacrocolpopexy | Proctectomy, removal of mesh and pelvic omentoplasty | Pneumoperitoneum and pneumorectum/-pelvis |
| 4       | M   | 43          | Ulcerative colitis, status post subtotal colectomy | Completion proctectomy with ileoanal pouch | Pneumoperitoneum and pneumorectum/-pelvis |
| 5       | M   | 78          | Presacral sinus after low anterior resection | Intersphincteric resection of colorectal anastomosis and pelvic omentoplasty | Pneumoperitoneum and pneumorectum/-pelvis |
| 6       | M   | 49          | Ulcerative colitis, status post subtotal colectomy | Completion proctectomy with ileoanal pouch | Pneumoperitoneum and pneumorectum/-pelvis |
| 7       | F   | 72          | Rectal cancer | Transanal total mesorectal excision | Pneumoperitoneum and pneumorectum/-pelvis |
| 8       | F   | 35          | Space-occupying lesion | Partial nephrectomy | Pneumoperitoneum |

![Fig. 2 Example of air percentage in the pneumorectum of case 4 during normal use (no suctioning)](image-url)
times less soluble in blood than CO₂, mainly due to the low solubility of the nitrogen it contains, which makes air emboli potentially much more harmful [4]. Indeed, studies suggest that abrupt introduction of 200 ml of air into the venous vasculature of an adult is lethal, while this amount is approximately 1000 ml for CO₂ [4]. Other disadvantages of using air as insufflation gas are that it results in more postoperative shoulder pain, caused by intraperitoneal air retained for a prolonged duration, as well as an increased risk of combustion [4]. Additionally, when room air instead
of medical grade air is used, the introduction of contaminants from the operating room cannot be excluded.

Suctioning from the operative field is often performed in cases of bleeding, which is also a risk factor for gas embolization [5]. Potentially even more dangerous is direct vascular injury that is not accompanied by bleeding. This situation may occur during laparoscopic or endoscopic surgery in which the operating field is above the level of the heart, which results in venous pressure being lower than insufflation pressure, such as TaTME. Under these circumstances, vascular injury may lead to massive flow of insufflation gas into the vascular cavity. This would then lead to influx of not only CO₂ but potentially also of air. One could hypothesize that this mechanism may explain the relatively high incidence of clinically significant GE in TaTME [1].

Our study has a few drawbacks, which however do not weaken the validity of our findings. The small number of patients may limit the generalizability of our results. Our findings, however, were observed in all 8 patients, and we have no reason to assume that inclusion of more patients or other types of procedures would lead to different conclusions. We have tested only the AirSeal® device, so care should be taken when extrapolating our findings to other insufflation devices. Specifically, we did not compare our results to findings in traditional (non-valveless) devices, but a study in 14 laparoscopies using a non-valveless device showed a mean air percentage of only 3.2% [7]. In this study, which was published as an abstract, average maximum amount of oxygen percentage in 16 patients in whom the AirSeal® was used was 43%, which corresponds with our findings. Lastly, our study was not designed to actually determine the occurrence of GE. Because of the low incidence of clinically significant GE this would require a much larger study, ideally with retrieval of gas bubbles from the vessels to determine their composition.

## Conclusions

The AirSeal® device maintains optimal insufflation not only by insufflating CO₂, but also by entraining room air, particularly during sudden decrease of pressure as occurs during suctioning. We believe the surgical and anesthesiological community should be aware of this phenomenon, most importantly because gas emboli occurring during use of this device cannot be assumed to be ‘harmless’ CO₂ emboli, but may also contain air. We urge manufacturers of insufflation devices to optimize their design in order to prevent entrainment of air.

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