To the Editor:—In their very interesting study, Loop et al.\(^1\) demonstrate an inhibitory effect of thiopental on nuclear factor \(\kappa B\) (NF-\(\kappa B\)) activation in T cells.

However, we missed some essential background information in the discussion. First, the concentration of thiopental used in this study is much higher than plasma concentrations\(^2\) noted in clinical practice or in other models presenting inhibitory effects of thiopental on interferon-\(\gamma\) (IFN-\(\gamma\)) production.\(^3\) Second, the fact that thiopental suppresses NF-\(\kappa B\) translocation in T cells may not directly reflect general immune suppression. Regulation of cytokines in T cells is simplified in this article and may lead to false interpretation.

Therefore, it would be helpful to give some more information beyond that provided by the authors in their statement that “other transcription factors may be involved.”\(^1\)

Cytokine expression is regulated in a cell-type and stimuli-specific manner. This might explain why Loop et al.\(^1\) were not able to demonstrate any effect of propofol on cytokine production or NF-\(\kappa B\) activation, whereas Takaono et al.\(^5\) describe inhibition of interleukin 6 (IL-6) production in lipopolysaccharide-stimulated peripheral blood mononuclear cells after propofol treatment. In the same study, thiopental (up to 200 \(\mu g/\mu l\)) had no significant effect on IL-6 production.

Furthermore, the ability of transcription factors to bind DNA and modulate gene transcriptions is tightly regulated in normal cells. There are four transcription factors that play a major role in the regulation of inflammatory gene expression: activator protein 1, activating transcription factor 2, signal transducers and activators of transcription, and NF-\(\kappa B\) (and, in T cells, nuclear factor of activated T cells [NFAT]). The pattern of their activation regulates expression of inflammatory mediators. Inhibition of one transcription factor may include enhanced activation of another factor.\(^5\) Loop et al.\(^1\) show that thiopental reduces the production of IL-2, IL-6, and IFN-\(\gamma\) in phorbol-12-myristate-13-acetate-stimulated peripheral blood mononuclear cells. The authors conclude that this is due to the inhibitory effect of thiopental on NF-\(\kappa B\) activation. However, the transcription of IL-2 and IFN-\(\gamma\) requires the activation of NFAT and activator protein 1 more than that of NF-\(\kappa B\).\(^6\)

This is of importance since NFAT is also required for transcription of IL-10, an antiinflammatory cytokine induced by thiopental.\(^8\) In this regard it also must be mentioned that the regulation of IFN-\(\gamma\) seems to be different in Jurkat cells than in “normal” T cells.\(^7\) In addition, the presentation of NFAT binding in the presence of other anesthetics would have been very interesting since it has been shown that ketamine decreases cytokine production in whole blood preparations at concentrations comparable to those used by Loop et al.\(^1\) Further-more, it has been shown that ketamine suppresses endotoxin-induced NF-\(\kappa B\) activation in other models.\(^9\) Therefore, it is surprising that Loop et al.\(^1\) do not see any effect of ketamine on NF-\(\kappa B\) activation in T cells or on cytokine production in peripheral blood mononuclear cells.

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Fourth, Drs. Haerleber and Kiefer state, “cytokine expression is regulated in a cell-type and stimuli-specific manner.” We fully agree with this statement. To briefly recapitulate, we examined the effects of anesthetics in T lymphocytes on the tumor necrosis factor–dependent activation of NF-kB, as well as the cytokine expression induced by phorbol-12-myristate-13-acetate (125 ng/ml) in mononuclear cells. Therefore, the cited study is not comparable because cytokine expression in mononuclear cells was induced by lipopolysaccharide. Likewise, in this study, the effects of thiopental were investigated at concentrations of less than 200 μg/ml, whereas we used 400-μg/ml thiopental. At this lower concentration, Takaono et al. did not note a significant effect; however, a trend toward inhibition could be observed.

Later in their letter, Haerleber and Kiefer state that “ketamine decreases cytokine production...and suppresses endotoxin-induced NF-kB activation.” These studies used whole blood, a human glioma cell line, or extracts of mice brain. Kawasaki et al. observed the inhibition of tumor necrosis factor–induced interleukin 6 and interleukin 8 production by ketamine at concentrations higher than 100 μg/ml. In contrast, we used 60 μg/ml or less ketamine in our study. Sakai et al., while observing an inhibition of lipopolysaccharide-induced NF-kB activation at 50 μg/ml, used a glioma cell line and brain extracts in these experiments. Therefore, we believe that the comparability of these studies with ours is impaired because of the different experimental settings and the fact that cytokine expression is, indeed, “regulated in a cell-type and stimuli-specific manner.”

To the Editor:—We read with interest the article published by Keller et al., who tested the Combitube 37 F SA (Kendall-Sheridan Catheter Corp., Argyle, NY). Our main observations about the study are as follows.

First, contrary to their ethical consideration of not using the recommended volumes in elective surgery patients because of the high risk of trauma, the authors used much greater volumes than those they wanted to avoid. The manufacturer recommends 85 ml (not 100 ml) of air for the pharyngeal balloon and 12 ml (not 20 ml) for the distal cuff of the 37 F SA model, which are inflation volumes for only emergency intubation. In elective cases, the minimal leakage technique should be used to minimize pressure exerted on the pharyngeal mucosa, which means that from 40 up to 85 ml and less than 12 ml, respectively, are usually sufficient to achieve a tight seal.

Furthermore, use of the Combitube is contraindicated when the gag reflexes are intact, as occurs in awake, nonanesthetized patients, a fact that the authors completely overlooked. This seems unethical to us because without sufficient knowledge of the basics of what they intended to test, the investigators exposed the volunteers to an increased risk of complications. In fact, all of them reported sore throat.

Second, the main implications of the study relate to the safety of using the Combitube in elective surgery patients. Other investigators have tested the device in such a population, concluding that elective use of the Combitube is not only safe but also reliable and feasible in most patients. Thus, the results of a study involving only four patients must be interpreted cautiously.

Third, concerning the cause of pharyngeal–esophageal lesions, although it has not been fully investigated, our impression is that, with other procedures in anesthesia, they relate much more to the performing hands than to the device itself. According to the American Society of Anesthesiologists closed claims database, most claims for esophageal injuries were for esophageal perforation (43 of 48; 90%), which involved difficult tracheal intubation in 67% of cases (n = 29). In addition, the intubating laryngeal mask airway has been reported to cause fatal esophageal perforation during elective general anesthesia. Nevertheless, the authors do not label the laryngeal mask airway as risky as they so superficially did with the Combitube. The cited reports of esophageal lesions associated with Combitube use are incidental cases involving paramedics in an out-of-hospital emergency setting who, exactly like the authors themselves, did not follow the recommendations. According to the criteria of evidence-based medicine, such reports should be classified as class C evidence, and they represent a poor basis for analysis.

Fourth, we missed in this study some essential qualities of good trial design, such as randomization or blind assessment of outcome. In conclusion, we are alarmed that colleagues who are not very experienced with these aspects of clinical investigation might falsely interpret this article. Finally, we are worried that biased studies could overshadow serious attempts to investigate in clinical practice.

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References


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Is the Combitube Traumatic?

To the Editor:—We read with interest the article published by Keller et al., who tested the Combitube 37 F SA (Kendall-Sheridan Catheter Corp., Argyle, NY). Our main observations about the study are as follows.

First, contrary to their ethical consideration of not using the recommended volumes in elective surgery patients because of the high risk of trauma, the authors used much greater volumes than those they wanted to avoid. The manufacturer recommends 85 ml (not 100 ml) of air for the pharyngeal balloon and 12 ml (not 20 ml) for the distal cuff of the 37 F SA model, which are inflation volumes for only emergency intubation. In elective cases, the minimal leakage technique should be used to minimize pressure exerted on the pharyngeal mucosa, which means that from 40 up to 85 ml and less than 12 ml, respectively, are usually sufficient to achieve a tight seal.

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Second, the main implications of the study relate to the safety of using the Combitube in elective surgery patients. Other investigators have tested the device in such a population, concluding that elective use of the Combitube is not only safe but also reliable and feasible in most patients. Thus, the results of a study involving only four patients must be interpreted cautiously.

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Fourth, we missed in this study some essential qualities of good trial design, such as randomization or blind assessment of outcome. In conclusion, we are alarmed that colleagues who are not very experienced with these aspects of clinical investigation might falsely interpret this article. Finally, we are worried that biased studies could overshadow serious attempts to investigate in clinical practice.

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To the Editor:—We read with interest the article by Keller et al. 
entitled "The Influence of Cuff Volume and Anatomic Location on 
Pharyngeal, Esophageal, and Tracheal Mucosal Pressures with the 
Esophageal Tracheal Combitube." The authors used a cadaver model and 
healthy volunteers to measure the pressures exerted by the esophageal-tracheal Combitube 37 F SA (ETC; Kendall-Sheridan Catheter Corp., Argyle, NY) on the pharyngeal, esophageal, and tracheal mucosa. To our knowledge, this is the first description of the use of the Combitube in awake volunteers. The very low amount and concentration of local anesthetic used (10 puffs of 1% lidocaine) demonstrate the ease of esophageal insertion of the device, even in awake volunteers. We appreciate the work of the authors but want to comment on several other aspects of their paper. The authors inflated the oropharyngeal balloon and distal cuff to a maximum volume of 100 and 20 ml, respectively, which is far above the maximum volume recommended by the manufacturer (85 and 12 ml, respectively). The authors observed relatively high mucosal pressures, potentially exceeding mucosal perfusion pressures, and do not recommend the ETC for routine anesthesia cases. However, several recent publications have clearly shown that neither the oropharyngeal balloon nor the distal cuff has to be inflated to the maximum volume recommended, and that much lower volumes are sufficient in the majority of patients. Hartmann et al. 

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(Accepted for publication November 4, 2002.)

Is It Unethical to Use the Combitube in Elective Surgery Patients?

In Reply:—We would like to thank Drs. Krafft, Hartmann, Agro, Gaitini, and Vaida and Drs. Urtubia and Gazmuri for their interest in our study that demonstrated high pharyngeal and esophageal mucosal pres-

ures with the esophageal-tracheal Combitube (ETC; Kendall-Sheridan Catheter Corp., Argyle, NY) in fresh cadavers and awake volunteers. We will respond to each point in turn, dealing with the former group first. We were aware that our maximum cuff volumes exceeded those for the small-adult ETC. In clinical practice, recommended volumes are frequently exceeded, either by initial overinflation (both accidental and deliberate) or through nitrous oxide diffusion (and subsequent failure to limit intracuff pressure increases), and we wished to test these conditions. The maximum volumes we chose were those recom-

recommended for the normal-adult ETC. Our subjects, both living and dead, were adults of normal—not small—size.

Obviously, and as we showed in our study, an effective pharyngeal seal can be obtained at lower cuff volumes, but many patients still...
require the maximum volume. Krafft et al. seem to ignore the fact that we measured mucosal pressures over the full inflation range and found that even when cuff volume was reduced to the minimum required to form a pharyngeal seal of 30 cm H₂O, mucosal pressures were still two or three times higher than mucosal perfusion pressure. A meta-analysis of data from similar studies suggests that when cuff volumes are reduced so that the pharyngeal seal is no greater than 30 cm H₂O, pharyngeal mucosal pressures for the ETC are the highest among modern extraglottic airway devices (table 1).

Krafft et al. suggest that a pharyngeal seal of 30 cm H₂O can never be reached with use of an LMA-Classic™ (Laryngeal Mask Company, San Diego, CA) or an LMA-ProSeal™ (Laryngeal Mask Company) and quote a study by our group in which the mean pharyngeal seals were 16 and 27 cm H₂O, respectively.² However, if the authors had read the studies more carefully they would have realized that these results were from a mixed male and female population in which the size 4 laryngeal masks were used. The mean ± SD maximum pharyngeal seal for the size 4 LMA-Classic™ and LMA-ProSeal™ in women is 21 ± 3 cm and 36 ± 6 cm H₂O,³ respectively, and the maximum pharyngeal seal for the size 5 LMA-Classic™ and LMA-ProSeal™ in men is 24 ± 5 cm and 32 ± 7 cm H₂O,³ respectively. These values are by no means the highest reported in the literature, and both values for the LMA-Classic™ exceed 30 cm H₂O.

Citing two studies,⁵,⁶ Krafft et al. state that the incidence of bleeding, dysphagia, and sore throat for routine anesthesia can be reduced by experienced users to 10–27%, 16%, and 8%, respectively. However, another study by experienced users showed that the incidence of bleeding, dysphagia, and sore throat was 36%, 68%, and 48%, respectively. Interestingly, in the former two studies the ETC was inserted with laryngoscopic guidance,⁵,⁶ and in the latter study it was inserted blindly.⁷ Perhaps it is the insertion technique rather than the experience level that reduces morbidity.

The authors imply that inflation of the distal cuff to the maximum recommended volume is safe because it ‘almost never blindly enters the trachea.’ This statement is astonishing given that one of the authors (Dr. Agro) recently wrote a review on the ETC.⁸ A more careful analysis of the literature reveals that the mean incidence of tracheal placement when it is inserted blindly is 9% (range, 3–17%).⁹–¹² Furthermore, accidental tracheal placement can occur even in an attempt to insert the distal cuff into the esophagus with laryngoscopic guidance, as demonstrated by yet another one of the authors (Dr. Gaitini).⁵ Krafft et al. state that the ETC provides aspiration prophylaxis when used properly in the esophageal position, but they cite no evidence. Two studies have addressed this issue. One revealed no evidence of gastroesophageal reflux, as determined by swallowed dye, but the other showed that the incidence of aspiration was 12% with the ETC in the esophageal position, as determined by tracheal pH changes. Other evidence that the esophageal cuff does not always seal off the esophagus is that gastric insufflation can still occur. One recent study showed an incidence of 2.5%.¹³ In addition, gastric rupture¹³–¹⁵ has been reported to occur with the esophageal obturator airway, which also uses an esophageal cuff. The most likely explanation for failure of the esophageal seal is that the recommended volumes are too low.

We were unable to find any data about the cuff volume required to seal off the human esophagus. The only data we could find come from a 1974 study in which a canine model was used, which showed that inflation of a Foley catheter cuff with 30 ml prevented pharyngeal regurgitation of fluid.¹⁶ The implication is that the manufacturer’s recommended cuff volume of 12–20 ml may be below that which is needed to provide protection. Interestingly, our study suggested that esophageal mucosal blood flow might be impeded at cuff volumes as low as 6 ml.¹³

Drs. Urtubia and Gazmuri imply that it was unethical to insert the ETC into awake volunteers because the manufacturer considers it contraindicated for patients with intact gag reflexes. However, these recommendations relate to its use in semicomatose patients and not awake volunteers. We believe that using the ETC was entirely ethical for two reasons. First, numerous studies have been conducted on other airway devices in awake volunteers with potentially intact gag reflexes; for example, Bemumof²⁷ used 60 such volunteers to test a new airway device. Second, all volunteers understood the risks involved, topical anesthesia was applied, the efficacy of topicalization was tested with a spaulata, and none of the subjects (as it happened) gagged with the ETC. We used only four volunteers to minimize risk.

Urtubia and Gazmuri are somewhat contradictory in their comments, claiming on the one hand that it was unethical to use awake volunteers and claiming on the other that we should have studied more. We agree that data from only four patients should be interpreted cautiously, but the main findings of our study were based on data from 20 fresh cadavers. In our study we presented evidence that a fresh cadaver is a reasonable model of the anesthetized, paralyzed patient.¹³ Urtubia and Gazmuri state that 12 ml is usually sufficient to achieve an airtight seal with the esophagus. However, as mentioned earlier, little is known about the volumes required to form an airtight seal in the esophagus. Furthermore, the distal cuff volume cannot be reduced to the minimum required to form an effective seal since there is no easy way of measuring the esophageal seal, unlike the pharyngeal seal.

Urtubia and Gazmuri state that the ETC is safe for elective surgery. To date there have been only three studies in which the ETC was used for elective surgery, and, indeed, there have been no major complications in a metapopulation of 275 patients.⁶–⁷ However, these numbers are too small to claim that a technique is safe. In addition, Klein et al.¹⁸ reported that in a patient undergoing elective surgery with the ETC as part of a clinical trial, an esophageal tear occurred after difficult blind placement. Direct trauma or increased intraluminal pressure distal to the cuff may have caused the tear. The patient underwent a thoracotomy for esophageal repair and fortunately survived. The patient was only the ninth enrolled in the study. The incidence of esophageal tearing according to data collected so far is therefore 0.4% (1 of 284), a figure that could hardly be considered to confirm the safety of use of the ETC in patients undergoing elective surgery.

Urtubia and Gazmuri state that esophageal lesions associated with the ETC are incidental cases involving paramedics in out-of-hospital emergency settings and that according to the criteria of evidence-based medicine, these reports form a poor base for analysis. However, the case of Klein et al.¹⁸ occurred in a prospective study performed by

**Table 1. Composite Data for Directly Measured Mucosal Pressures in the Anterior (Base of Tongue), Lateral, and Posterior Pharynx for Six Modern Extraglottic Airway Devices at a Pharyngeal Seal Pressure of 30 cm H₂O or at Maximum Seal**

<table>
<thead>
<tr>
<th>LMA-Classic™</th>
<th>LMA-ProSeal™</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laryngeal Tube Airway (Cadaver)</td>
<td>Cuffed Oropharyngeal Airway†</td>
</tr>
<tr>
<td>Lateral pharynx</td>
<td>3 (1–4)</td>
</tr>
<tr>
<td>Base of tongue</td>
<td>11 (8–14)</td>
</tr>
</tbody>
</table>

Units are cm H₂O. Data from Keller et al.¹,²,¹ⁱ,²¹,²³ and Brimacombe et al.²²


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anesthetists presumably adhering to the guidelines. The out-of-hospital evidence of major trauma is not just incidental. One such study showed that 0.7% of patients (8 of 1,139) developed subcutaneous emphysema, and another showed that major tissue trauma occurred in 0.6% (10 of 1,563).

Finally, we take offense to the authors’ suggestion that our study lacked good design in terms of randomization and blinding. There was nothing to randomize and nothing to blind other than perhaps the volume of air in the cuff. Blinding with respect to whether the subject was a cadaver or an awake volunteer would have been difficult.

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2. S, Schwarz S, Fitzgerald RD: Complications following the use of the Combitube, CPB and acute myocardial ischemia of those models is that no human undergoes CPB alone, and therefore, ”as it contributes further evidence of the cardioprotective properties of infusion in a model of myocardial ischemia improved left-ventricular function resulting from intravenous esmolol we beg to differ with the authors’ statement that ”a paucity of studies exist on effectiveness, rationale, and/or mechanisms underlying the use of βAR [β-adrenergic receptor] antagonists in this setting” (i.e., CPB surgery during acute myocardial ischemia).

We would like to remark that the intraoperative use of esmolol is now a well established technique of myocardial protection that was clinically introduced 10 yr ago. A number of clinical studies have investigated the impact of intraoperative esmolol administration on outcome. We also take issue with the authors’ perception that “most animal models to date have focused on CPB alone. The criticism of those models is that no human undergoes CPB alone, and therefore, the model reffers not reflect CPB surgery.”

A number of experimental studies, some of which were conducted by our group, have investigated the impact of esmolol in models of CPB and acute myocardial ischemia-reperfusion injury and showed that esmolol improved myocardial function and reduced infarct size. We believe that the discussion in this otherwise excellent paper by Booth et al. suffers significantly from the failure to consider this previous work.

Another detail of concern is the combination of esmolol and cold crystalloid cardioplegia in the treatment group, which makes no sense from a cardiac surgeon’s point of view. Intraoperative myocardial protection with esmolol is considered an alternative rather than an adjunct to cardioplegic arrest. In fact, combining both principles sacrifices the major advantages of the esmolol technique, such as the avoidance of additional global myocardial ischemia and prevention of crystalloid perfusion-induced myocardial edema.

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References


In Reply.—We thank Drs. Geissler, Melhior, Laine, and Allen for commenting positively on our recent article.1 We agree that the authors have contributed significantly over the past 10 yr to demonstrating the safety and efficacy of intracoronary β-adrenergic receptor (β-AR) antagonists as an alternative to cardioplegia in the setting of coronary artery bypass grafting surgery. In contrast, our study does not compare or contrast methods for administration of β-ARs during coronary artery bypass grafting surgery, but, rather, the mechanisms by which protection might occur. As such, our comments on “the paucity of data” in the area of β-AR antagonists during cardiac surgery refers to both a lack of mechanistic data demonstrating the rationale for beneficial effects and a lack of large-scale, randomized, clinical trials.

To support this claim, we cited recent American College of Cardiology and American Heart Association concerns,2 as well as American College of Cardiology and American Heart Association guidelines that state that, at this time, “there is no universally applicable myocardial protection technique” for reducing the risk of perioperative myocardial dysfunction.3 Geissler et al. further comment that the use of intravenous esmolol as an adjunct to cold crystalloid cardioplegia (e.g., rather than intracoronary esmolol) makes “no sense from a cardiac surgeon’s point of view.” However, the model used in our study was based on the common practice in the United States of intravenous β-blockade during cardiopulmonary bypass, a strategy that has recently been shown to have benefit in terms of neuroprotection in humans.4

Our study also clearly demonstrates prevention of acute myocardial β-AR desensitization in a canine model with use of this approach. Crystalloid cardioplegia has been supplemented with many agents over the years in the quest for better myocardial protection; Geissler et al. have contributed in very positive ways to alternative thinking in this regard in their use of esmolol as a sole “cardioplegic” agent. Our study does not purport to answer this controversial question. Rather, our quest for understanding the mechanisms by which stress affects the myocardium, in this case via β-ARs, is based on the hope that such insight may lead to novel approaches for protecting the heart during cardiac surgery in the future.

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Use of Recombinant Activated Factor VII in Patients with Severe Coagulopathy and Bleeding

To the Editor—The emergence of recombinant activated factor VII (rFVIIa) as potentially a panhemostatic agent that will “save the day” for patients with severe hemorrhage is truly exciting. The positive experiences reported by Tobias,1 Slappendel et al.,2 and Svartholm et al.3 add valuable data to help define the role of this agent in clinical practice. With time, however, it is likely that we will find out that, in addition to cost advantages, there are problems and limitations associated with the use of rFVIIa. Just as antibiotics have not eliminated the need for infection control and for expert management of infections in patients, rFVIIa will not replace the need for aggressive prevention and reversal of coagulopathy in patients.

By definition, dilutional coagulopathy suggests that insufficient coagulation factors have been given during the course of severe hemorrhage and blood product replacement. It is therefore theoretically preventable and reversible, mainly with the use of adequate amounts of fresh frozen plasma (FFP). In the first case presented by Tobias,1 the patient was clearly having significant ongoing bleeding, as indicated by chest tube drainage, deterioration of the coagulation indices and hematocrit, and other evidence. When the international normalized ratio reached 2.0, plasma factor concentrations had probably decreased to just slightly above 50% of normal.4 At that point, the inadequate use of

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In the case presented by Slappendel et al. 2 we are surprised that the patient’s hemostatic parameters were not at least partially corrected before the elective surgery, especially in light of the type of surgery being performed and the use of spinal anesthesia. Intraoperatively, the use of cell saver does not prevent further dilution of coagulation factors, as is clearly reflected in the abnormal laboratory results. The appropriate action should have been the transfusion of FFP before surgery or, at the very least, as soon as bleeding started.

In the case presented by Svartholm et al., 3 the patient had received 191 (equivalent to 65 units) of packed erythrocytes and 4.5 L (18 units) of FFP. If we assume that some crystalloid or colloid solution had also been given (such that the patient’s hematocrit was not excessively high due to the large amount of packed erythrocytes and the relatively small volume of FFP), then using this amount of FFP equaled transfusing whole blood with a plasma coagulation factor concentration of 30% or less. In the face of severe hemorrhage, such a low level of FFP dosing is inadequate. Prothrombin complex concentrate does not contain all of the factors. Exacerbating the situation was that another 8 L of packed erythrocytes was given over the next 11 h, with no apparent supplementation with FFP. With coagulation factor concentrations low and continuing to dwindle, the reliance on pharmacologic supplements rather than on coagulation factor replenishment seemed inappropriate.

We respectfully suggest that in all of the aforementioned cases 1–3 the patients might have suffered from an underdosing of FFP. This likely resulted in the transfusion of increased amounts of blood products. While we are excited by the impressive evidence accumulating on the use of rFVIIa in “refractory” bleeding, we wish to caution against an excessive and premature reliance on the use of this (or any new) technology to bail us out of difficult situations that we could have avoided getting into in the first place.

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In Reply:—Drs. Ho, Dion, and Karmakar suggest that hemostasis in the patients reported by Tobias, 1 Svartholm et al., 2 and Slappendel et al. 3 might have been achieved with administration of additional fresh frozen plasma. Ho et al. point out, as did I in the editorial 1 that accompanied their reports, that recombinant activated coagulation factor VII (rFVIIa) has not been shown to be efficacious for states of dilutional coagulopathy. Parenthetically, it should be noted that when massive transfusion (one blood volume or more) is expected or achieved, use of whole blood rather than packed erythrocytes will provide the required coagulation factors and obviate or decrease the need for administration of plasma. 3 The only currently approved use for rFVIIa is for patients who have hemophilia with inhibitors of coagulation factors VIII or IX. Part of the value of the case reports detailing “off-label” uses of rFVIIa is that the reports have provided a stimulus for the initiation of prospective, controlled, randomized, blinded studies examining the efficacy and safety of rFVIIa administered for bleeding secondary to causes other than hemophilia.

While there are sound theoretical rationales and laboratory data to suggest that rFVIIa might be efficacious for treating a variety of coagulation defects, demonstration of clinical efficacy awaits the results of these ongoing clinical trials. These trials should also produce data regarding the incidence (if any) of the possible theoretical adverse responses: vascular thrombosis and embolism, myocardial infarction, and coagulopathy resulting from intravascular coagulation. My previous comments continue to pertain: “diagnosis of the specific [hemo-

Dr. Weiskopf consults for Novo Nordisk A/S (Copenhagen, Denmark) regarding their product, recombinant activated coagulation factor VII.

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In Reply:—The reaction of Drs. Ho, Dion, and Karmakar in their letter to the editor is clear and simple: the best way to act on a dilution coagulopathy is to recognize it early and to treat it using adequate amounts of fresh frozen plasma. In our view, the main reason to avoid fresh frozen plasma or any homolog blood products in cases of orthopedic surgery is also simple. Since the first human blood transfusion there have been unnecessary related risks: administration and handling mistakes, transmission viruses, transfusion reactions, and immunosuppression. Transfusion-related immunosuppression is considered to favor postoperative infections, to perturb postoperative wound healing, and thereby to result in a protracted hospital stay. The availability of new recombinant DNA medicine, like recombinant activated factor VII, is very promising because of the nearly complete lack of side effects. We agree with Ho et al. that treatment with fresh frozen plasma for dilution coagulopathy is, at present, the first choice. We expect that it’s only a matter of time (for double-blinded, randomized trials) and money before recombinant DNA drugs such as recombinant activated factor VII replace (classic) coagulant drugs such as aprotinin and desmopressin.

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(Accepted for publication November 17, 2002.)
**Propofol Preservation of Myocardial Function in Patients Undergoing Coronary Surgery Using Cardiopulmonary Bypass is Dose Dependent**

To the Editor:—We read with great interest the article recently published by De Hert et al.1 entitled "Sevoflurane but Not Propofol Preserves Myocardial Function in Coronary Surgery Patients." In this study, the authors based their conclusion primarily on myocardial mechanics measured before and 15 min after cardiopulmonary bypass. Hearts were paced at 90 beats/min during the measurements. This approach makes the cardiac mechanics measured more comparable between groups.

It would be very helpful for us to know if the plasma concentration of propofol is available and/or if the hemodynamic data are comparable at 12 h or more postoperatively in this study. An experimental study has shown that the protective effect of propofol on myocardial function following global myocardial ischemia and reperfusion is dose dependent, being effective at concentrations of 50 μM or more and not effective at concentrations of 10 μM or less.2 This may explain why propofol is not cardiac protective at "clinically relevant" concentrations of 1 μg/ml (5.6 μM) or up to 10 μM.3 Our preliminary clinical study indicated that propofol is cardiac protective in a dose-dependent manner in coronary surgery patients when applied at "clinically achievable" concentrations of 4 and 11.8 μg/ml during surgery. This was evident when the cardiac index beyond 12 h post-cardiopulmonary bypass was compared.4

To understand this we conducted an experiment in an isolated heart model, applying 12 μg/ml (67 μM) propofol during global myocardial ischemia and during the early phase of reperfusion and then 5 μg/ml after 15 min of reperfusion (to avoid the depressant effect of high-dose propofol on myocyte contraction). We have observed that this approach provides better long-term myocardial functional recovery than 5 μg/ml propofol applied throughout ischemia and reperfusion.5

De Hert et al should be congratulated for their efforts. They provided evidence that cardiac troponin I was significantly lower in the sevoflurane group than in the propofol group up to 36 h postoperatively, a cardiac protection probably related to the preconditioning effects of sevoflurane. It should be noted, however, that from 24 to 36 h postoperatively, cardiac troponin I levels decreased in all 10 patients in the propofol group, while they increased in 2 or 3 patients (2 of 10 or 5 of 10) in the sevoflurane group. This is a significant difference between groups.

We must point out that a decrease in heart rate and/or the depression of cardiac contraction during the early phase of reperfusion might be an integral part of the cardiac-protective effect of propofol in the long run. Given that cardiac output increased from post-cardiopulmonary bypass to the end of the operation only in the propofol group and that there was no difference in clinical outcome between groups, we feel that the statement "propofol is not myocardial protective," as implied in the title of the article by De Hert et al.,1 is somewhat misleading.

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In Reply:—We appreciate the interest of Drs. Ansley and Xia regarding our study on the preservation of myocardial function by sevoflurane in coronary surgery patients.1 Transient myocardial dysfunction after coronary bypass surgery is a well known phenomenon,2 but its pathophysiologic mechanisms still are not fully elucidated. Although its causes are multifactorial, reperfusion injury is thought to play an important role.

Recently, increasing evidence has indicated that anesthetic agents may have a potential beneficial role in the prevention and/or treatment of reperfusion injury. Volatile anesthetics have been shown to directly precondition or indirectly enhance ischemic preconditioning, resulting in protection of the myocardium, with the adenosine triphosphate-regulated potassium channel playing an important role. Central to the pathogenesis of ischemia-reperfusion injury is the production of free radicals. Several reports have demonstrated that propofol may enhance antioxidant capacity, and this property has been claimed to protect the myocardium. However, data on this subject are essentially experimental, and, to date, few studies have demonstrated a potential beneficial effect of anesthetic agents in myocardial stunning in the clinical situation.

For the volatile anesthetics, the beneficial effects were demonstrated on both functional hemodynamic parameters and biochemical markers of myocardial damage. For propofol, the evidence of a myocardial protective effect is mainly circumstantial. Several studies have, indeed, demonstrated that propofol increases the antioxidant capacity of erythrocytes and tissue. However, the possible implications of this increase in antioxidant capacity for preservation of tissue function remain to be demonstrated.

Ebel et al.3 found no protective effect of propofol (at clinically relevant concentrations) against myocardial reperfusion injury, but Ko et al.4 demonstrated that propofol at higher concentrations (100 μM) attenuated mechanical, biochemical, and histologic changes caused by myocardial ischemia and reperfusion. In a recent publication, Ansley et al.5 observed that high-dose propofol (a 2–2.5 mg/kg bolus followed by a continuous infusion of 200 μg·kg⁻¹·min⁻¹) enhances erythrocyte antioxidant capacity during cardiopulmonary bypass in humans. Erythro-
Correspondence

Effect of Nitrous Oxide on Sevoflurane Vaporizer Setting

To the Editor:—I read the article by Hendrickx et al.1 with great interest. In their study, the authors found that during minimal-flow anesthesia (MFA; 0.5 l/min), the vaporizer dial setting required to maintain the end-tidal sevoflurane concentration constant at 1.3% is lower when sevoflurane is delivered in an oxygen-nitrous oxide mixture than in oxygen alone because less gas and vapor are wasted through the pop-off valve with the oxygen–nitrous oxide mixture. During low-flow anesthesia (LFA; 1 l/min), however, vaporizer dial settings are similar with oxygen-nitrous oxide or oxygen, presumably because the proportion of excess gas leaving the pop-off valve relative to the amount taken up by the patient increases. However, I carefully examined the authors’ experimental design and found that the vaporizer setting during minimal-flow anesthesia (MFA) was set to keep the end-tidal sevoflurane concentration constant at 1.3% lower when sevoflurane is delivered in an oxygen–nitrous oxide mixture than in oxygen alone during MFA, simply because of the differences in the initial flow setup and not because less gas and vapor are wasted through the pop-off valve. We must recognize that the very large space of the anesthesia circuit (4.3 l) and the patient’s functional residual capacity (2.5–3 l) existed before the patient’s alveolar membrane; therefore, the use of MFA with oxygen alone certainly requires a much higher vaporizer dial setting to maintain the end-tidal sevoflurane concentration constant at 1.3%. On the other hand, MFA with oxygen–nitrous oxide mixture and a high fresh gas flow of 6 l/min was used for 10 min; therefore, the circuit and functional residual capacity were pre-filled to keep the end-tidal sevoflurane concentration constant at 1.3%. Certainly lower sevoflurane vaporizer dial settings are required afterwards, even at MFA.

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Reference


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In Reply.—We appreciate the comments made by Dr. Lin. He correctly points out that the anesthesia circuit and the patient’s functional residual capacity can be considered to be a reservoir of vapor (in this study, sevoflurane). However, because the desired end-expired sevoflurane concentration was obtained within 5 min in all groups, and because the sevoflurane concentration was kept constant throughout the study period (10–60 min), the anesthesia circuit and functional residual capacity are not a “source” of sevoflurane thereafter nor do they contribute to sevoflurane consumption during the observation period. In other words, the anesthesia circuit and the patient’s functional residual capacity, once saturated at the desired sevoflurane concentration, do not affect the vaporizer setting required to keep end-expired sevoflurane concentration constant. This concept is not influenced by the initial fresh gas flow settings.

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To the Editor:—I read the article by Hendrickx et al.1 with great interest. In their study, the authors found that during minimal-flow anesthesia (MFA; 0.5 l/min), the vaporizer dial setting required to maintain the end-tidal sevoflurane concentration constant at 1.3% is lower when sevoflurane is delivered in an oxygen-nitrous oxide mixture than in oxygen alone because less gas and vapor are wasted through the pop-off valve with the oxygen–nitrous oxide mixture. During low-flow anesthesia (LFA; 1 l/min), however, vaporizer dial settings are similar with oxygen-nitrous oxide or oxygen, presumably because the proportion of excess gas leaving the pop-off valve relative to the amount taken up by the patient increases. However, I carefully examined the authors’ experimental design and found that the vaporizer setting during minimal-flow anesthesia (MFA) was set to keep the end-tidal sevoflurane concentration constant at 1.3% lower when sevoflurane is delivered in an oxygen–nitrous oxide mixture than in oxygen alone during MFA, simply because of the differences in the initial flow setup and not because less gas and vapor are wasted through the pop-off valve. We must recognize that the very large space of the anesthesia circuit (4.3 l) and the patient’s functional residual capacity (2.5–3 l) existed before the patient’s alveolar membrane; therefore, the use of MFA with oxygen alone certainly requires a much higher vaporizer dial setting to maintain the end-tidal sevoflurane concentration constant at 1.3%. On the other hand, MFA with oxygen–nitrous oxide mixture and a high fresh gas flow of 6 l/min was used for 10 min; therefore, the circuit and functional residual capacity were pre-filled to keep the end-tidal sevoflurane concentration constant at 1.3%. Certainly lower sevoflurane vaporizer dial settings are required afterwards, even at MFA.

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In Reply.—We appreciate the comments made by Dr. Lin. He correctly points out that the anesthesia circuit and the patient’s functional residual capacity can be considered to be a reservoir of vapor (in this study, sevoflurane). However, because the desired end-expired sevoflurane concentration was obtained within 5 min in all groups, and because the sevoflurane concentration was kept constant throughout the study period (10–60 min), the anesthesia circuit and functional residual capacity are not a “source” of sevoflurane thereafter nor do they contribute to sevoflurane consumption during the observation period. In other words, the anesthesia circuit and the patient’s functional residual capacity, once saturated at the desired sevoflurane concentration, do not affect the vaporizer setting required to keep end-expired sevoflurane concentration constant. This concept is not influenced by the initial fresh gas flow settings.

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( Accepted for publication November 20, 2002.)
To the Editor:—We read with interest the article by Gamperl et al., which describes studies evaluating effects of isoflurane on isolated porcine coronary microvessels. The main finding from this well-performed study was that isoflurane caused dilation, which confirmed our results in a similar model.

The data from Gamperl et al. add to the existing body of compelling evidence that isoflurane is a dilator in the coronary circulation and not a constrictor, as was previously argued. We hope that these findings will finally put to rest the controversy relating to the vasomotor effect of isoflurane in the coronary circulation.

Over the past decade, studies in our laboratory and others have sought to clarify the effects of the volatile anesthetics in the coronary circulation. From these studies, several definitive findings have emerged. First, the volatile anesthetics, as a class of drugs, are dose-dependent coronary vasodilators. The coronary vasodilating effect of isoflurane is, by far, greater than that of the other volatile anesthetics (halothane, enflurane, sevoflurane, and desflurane). Second, volatile anesthetic-induced coronary vasodilation is mediated by the adenosine triphosphate–sensitive potassium channels and is endothelium dependent, although nitric oxide does not appear to be involved. Third, coronary vascular smooth muscle adapts to the relaxing effect of volatile anesthetics; thus, the coronary vasodilation caused by these anesthetics is minimized by slow or extended administrations.

It is worth recognizing that these findings in the coronary circulation served as a springboard for studies demonstrating that the volatile anesthetics may precondition the myocardium against reperfusion injury, an action that may have enormous therapeutic value in patients undergoing surgical and nonsurgical coronary revascularization procedures.

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Unusual Cause of Intraoperative Urinary Retention

To the Editor:—While intraoperative oliguria in not a reliable predictor of postoperative renal dysfunction, vigilant effort is made by the anesthesiologist to keep the patient’s urine output at an acceptable level. The total absence of urine in the collection bag suggests either a renal or bladder problem. While intraoperative oliguria is not a reliable predictor of postoperative renal dysfunction, vigilant effort is made by the anesthesiologist to keep the patient’s urine output at an acceptable level. The total absence of urine in the collection bag suggests either a renal or bladder problem. An 85-yr-old woman underwent coronary artery bypass surgery with aortic valve replacement. A Foley catheter was placed (CRITICORE

Fig. 1. Two Foley catheters: the top catheter is normal, and the lower catheter is defective, with lack of an orifice.

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Foley tray; Bard Medical) following induction of anesthesia. The lack of urine draining into the catheter prompted a call to the urology service. After establishing correct placement (transabdominal palpation of inflation and deflation of the Foley catheter balloon), irrigation was attempted without success. The catheter was then replaced. The replaced Foley catheter was examined and found to be lacking an orifice (same defect as depicted in fig. 1).

While intraoperative oliguria is indicative of renal hypoperfusion, anuria suggests the possibility of either catheter occlusion or the detachment of the tubing from the collection bag. Irrigation of the Foley catheter always should be the first step in the evaluation of the anuric patient. In these two cases, it was not possible to irrigate the bladder due to a lack of communication between the catheter lumen and the patient’s bladder. As the first patient voided immediately prior to his arrival in the operating room, the initial lack of return of urine into the Foley collection bag, even with suprapubic compression, was not surprising. Since anesthesiologists rely on urine output as an indicator of renal perfusion, confirming proper placement and function of the catheter by demonstrating the flow of urine should be a routine procedure prior to the beginning of surgery.

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The Use of Three-dimensional Computed Tomography to Visualize Thoracic Epidural Catheters

To the Editor:—Thoracic epidural catheterization was implemented in a 62-yr-old woman scheduled for subtotal gastrectomy. A FlexTip Plus® epidural catheter (single end hole; Arrow International, Reading, PA) was inserted at the T9–T10 intervertebral space and directed cephalad for 5 cm. With the patient’s consent, a 10-cm slab of a thin-cut spiral computed tomography (CT) scan was obtained through T6–T12 with 1-mm collimation at a pitch of 2.140 kVp and 160 mA and with a 13-cm field of view (HiSpeed Advantage scanner; GE Medical Systems, Milwaukee, WI).

The raw data were processed with a 0.5-mm overlap using the GE Advantage workstation (DentaScan, GE Medical Systems). The axial transverse section of the CT image demonstrated the existence of a catheter in the epidural space (fig. 1A). With an oblique view, the path of the catheter in the epidural space was displayed with multiplanar reconstruction (fig. 1B) or a three-dimensional surface-shaded display (fig. 2). There was no kinking or knotting of the catheter despite the curved path in the epidural space.

A substantial incidence of failed or unilateral epidural block may be due to the complexity of the epidural space.1 When looping, kinking, entrapment, or knotting of epidural catheters occurs, it is not easy to visualize the path of the radiopaque catheter within the epidural space.2,3 Although conventional radiography,4,5 ultrasonographic imaging,6 epidurography with contrast medium,7 CT,8–10 or magnetic resonance imaging11 might be useful, a potential allergic reaction to contrast medium, interference with metal coils, or blockade or occlusion of the epidural catheter could still be problematic. Thin-cut volumetric CT scanning, coupled with two-dimensional multiplanar or three-dimensional model reconstruction, has been used for evaluation of the upper airway anatomy.12

Given the exquisite tissue contrast of CT in displaying the epidural space, its potential in the evaluation of a problematic epidural catheter has never been explored. An axial scan shows the exact location of the

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Fig. 1. (A) An axial computed tomography image shows the epidural catheter between the spinal lamina and the dural sac. The epidural catheter contains radiopaque metal spring coils. (B) A multiplanar reconstructed image of the lower thoracic spine in an oblique coronal section shows the complex but smooth, looping course of the epidural catheter (arrow), without kinking.

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catheter in the epidural space and thus is useful in detecting malposition. With two-dimensional multiplanar and three-dimensional surface display models, the course of the catheter in the epidural space is depicted. In conclusion, three-dimensional CT imaging appears to be a novel tool to visualize the position and path of an epidural catheter.

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To the Editor—Epidural hematoma formation following lumbar regional anesthesia is a rare complication, with incidences of 1 in 220,000 after spinal anesthesia and 1 in 150,000 after epidural anesthesia.1 While severe lower back pain that is enhanced by percussion or movements of the spine typically represents the initial clinical symptom of a large epidural hematoma,2 cauda equina syndrome with paraparesis and dysfunctions of the bladder and bowel may develop after a delay of several hours.

We report the case of an 83-yr-old woman who underwent spinal anesthesia for minor gynecologic surgery. Preoperative evaluation revealed an ASA physical status of III with atrial fibrillation, discrete pretibial edema, and chronic rheumatoid pain in both legs. Blood chemistry values and coagulation status were normal. Daily medication, including 1,000 mg naproxen, was discontinued on the evening before surgery.

Due to degeneration of the spine, puncture of the lumbar subarachnoid space required multiple attempts at intervertebral spaces L2–L3 and L3–L4, with use of a 27-gauge Quincke needle. After appearance of clear cerebrospinal fluid without shooting pain or paresthesia, 75 mg hyperbaric lidocaine was injected.

Unilateral Presentation of a Large Epidural Hematoma

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Fig. 2. Surface display images (three-dimensional) of the epidural catheter in the thoracic epidural space. The arrows depict the path of the catheter. The laminae were removed during the reconstruction process.
In the early postoperative period, the patient reported her typical rheumatoid pain in the legs, and diclofenac was administered topically. Seventeen hours after surgery, a neurologic examination revealed monoparesis of the right leg in the absence of back pain. Sensory functions were reduced in radicular segments L4–L5 and S1–S4 on the right side. The right patellar tendon reflex was reduced, and ankle tendon reflexes were absent on both sides. There were no signs of bladder or bowel dysfunction. Primarily, a pressure palsy of the lumbar plexus or the femoral and sciatic nerves was assumed.

Lumbar computed tomographic images were obtained and revealed high-grade compression of the dural sac by a large, hyperdense mass. Subsequent magnetic resonance imaging (MRI) of the lumbar spine (fig. 1) demonstrated a subtotal obstruction of the spinal canal at the L2 and L5 levels. As imaging findings were highly suggestive of a large epidural hematoma, the patient underwent urgent decompressive laminectomy of L2–L4, and the hematoma was removed.

Postoperatively, the patient initially showed no significant benefit from decompressive therapy, although MRI examinations 3 weeks after the procedure confirmed complete removal of the hematoma. Following a rehabilitation process of 3 months, neurologic function was recovered and the patient was able to walk independently.

Major risk factors for bleeding complications following spinal or epidural anesthesia include impaired coagulation, difficult or multiple punctures, and insertion of a catheter. Naproxen, a nonsteroidal antiinflammatory drug, may also have played a causal role in the development of the epidural hematoma, because platelet aggregation is normal in only 50% of patients 2 days after withdrawal of naproxen, and in our patient naproxen was withdrawn 12 h before surgery.

The strictly unilateral sensorimotor dysfunction of a radicular pattern without lower back pain in our patient was certainly an atypical clinical presentation of a large epidural hematoma expanding from the lower thoracic to the lower lumbar spine. Furthermore, in view of the age of the patient, the absence of ankle tendon reflexes on both sides might, instead, have been attributed to mild polyneuropathy than to an impairment of the first sacral root on both sides.

Magnetic resonance imaging is the diagnostic procedure of choice to verify an epidural hematoma. Therapy consists of urgent decompressive surgery with removal of the hematoma, usually in combination with hemilaminectomy. While outcome is influenced mainly by the severity of neurologic impairment at the time of surgical intervention, data and clinical experience suggest that the length of the delay before surgery also is important to recovery. Minor neurologic disturbances and treatment within 12 h and up to 36 h are correlated with a better prognosis for recovery, but patients may continue to benefit from surgical therapy up to 3 weeks after damage.

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