Reliable and rapid smooth extubation after Ketofol for induction of general anesthesia in Laparoscopic Drilling of polycystic ovary: A randomized controlled trial

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Research article

Keywords: ketofol, airway response, hemodynamics, smoothness of extubation, general anesthesia

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Abstract

Background

The outcome of ketofol on the hemodynamics and the airway response during induction of general anesthesia has been studied before. Its effect on smoothness of extubation has not been studied before. So, we aimed to assess the effect of ketofol on the smoothness of extubation and compare it with propofol only for induction of general anesthesia.

Methods

This double-blind, randomized, and controlled study was conducted on one hundred and six American Society of Anesthesiologists Physical status "ASA PS" class I and II female patients aged 18–40 years old and scheduled for laparoscopic drilling for polycystic ovary disease under general anesthesia. The patients were assigned into one of two groups (53) patients each; group KP = ketofol and group P = propofol.

Results

There was good sedation score during suction and extubation in the ketofol group. Airway response and smoothness of extubation were better in the ketofol group better than the propofol group.

Conclusion

Ketofol as an induction anesthetic agent was effective in attenuating the airway response during extubation more than propofol only.

Trial registration:

This trial was retrospectively registered at the Clinical Trial.gov with the Identification Number: NCT04365686.

Background

Tracheal extubation is a critical and stressful moment during general anesthesia at which many hazardous and unwanted hemodynamics and airway responses may occur such as tachycardia, hypertension, dysrhythmias, coughing, laryngospasm, and bronchospasm\(^1,2\).

These hazardous responses are ought to the sudden release of catecholamines during tracheal extubation\(^3\)
Many medications have been studied to reduce the stress response during extubation aiming for the achievement of a state of smooth extubation such as including intravenous lignocaine, short acting opioids as fentanyl and remifentanyl, esmolol, endotracheal local anesthetic instillation and dexametomidine\(^{(4-9)}\)

Propofol (2, 6-diisopropylphenol) is a sedative and hypnotic drug with antiemetic property used for induction and maintenance of general anesthesia and in the intensive care units (ICU) for sedation. It produces its effect by facilitation of inhibitory neurotransmission mediated by GABA (Gama Amino Butyric Acid) depression\(^{(10-12)}\)

Ketamine is a phencyclidine derivative used for induction of general anesthesia. It has favorable analgesic and amnestic properties but increases sympathetic activity, nausea, vomiting, and undesirable psychomimetic disorders.\(^{(13,14)}\)

Ketofol (ketamine-propofol mixture) as an induction agent drug and its effect on hemodynamics has been studied previously.\(^{(15,16)}\) Its effect on the airway responses during induction of anesthesia was studied by Aberra B et al.\(^{(17)}\)

We hypothesized that giving ketofol as an induction agent getting the beneficial effects of both medications would favorably affect both the hemodynamics and the airway response during extubation. The aim of this study was to compare the effect of injecting a single dose of ketofol versus propofol on the smoothness of extubation as regards the airway response and the hemodynamics when given for the induction of general anesthesia in laparoscopic drilling of polycystic ovary disease.

**Methods**

Following ethical committee approval and research is coded (R83) of Anesthesia department, Fayoum University and obtaining a written informed consent, this study was conducted on one hundred and six American Society of Anesthesiologists Physical Status "ASA PS" class I and II female patients aged above 18 years old up to 40 years old and scheduled for laparoscopic drilling for polycystic ovary under general anesthesia. This trial protocol was registered at the Clinical Trial.gov with the Identification Number: NCT04365686. This study was conducted in El Fayoum University Hospitals and took 12 months between April 2019 and April 2020. Patients suffering from cardiac, hepatic, renal diseases, history of epilepsy were excluded from this study. The patients were randomly allocated by a computer-generated table into one of two study groups. The randomization sequence was concealed in opaque sealed envelopes. The envelopes were opened by the study investigators just after recruitments and admission to the operation room. Only assessors and data collectors were blinded to the group's allocations. Our study adheres to CONSORT guidelines.

The two study groups of this randomized, double-blind and parallel clinical trial are: Group K (Ketofol group): includes fifty three female patients who received propofol (1 mg/kg) plus ketamine (0.5 mg/kg)
at induction of general anesthesia and Group P (propofol group): includes fifty three female patients who received propofol (2 mg/kg) only at induction of general anesthesia.

After securing an intravenous access by a 20 g i.v. cannula, intravenous premedication (midazolam 2 mg and 4 mg ondansetron) was administered to all patients. Standard ASA monitoring (5-lead ECG, noninvasive blood pressure (NIBP), and pulse oximetry) was applied to all the patients for recording of heart rate (HR), NIBP and oxygen saturation by a mutli-parameter monitor. Induction of general anesthesia was achieved as follows: in Group K (Ketofol group), fifty three female patients received propofol (1 mg/kg) plus ketamine (0.5 mg/kg) at induction of general anesthesia while in Group P (Propofol group), fifty three female patients received propofol (2 mg/kg) only at induction of general anesthesia.

Patients in both groups received intravenous fentanyl 2 µg/kg, and atracurium 0.5 mg/kg. After tracheal intubation, general anesthesia was maintained by isoflurane 1.5% in 2 L/min oxygen-air mixture 50%:50% and atracurium 0.1 mg/kg every 30 minutes, if needed. At the end of surgery, inhalational anesthesia was stopped and reversal of the neuromuscular blockade was done by intravenous neostigmine 0.05mg/kg and atropine 0.01mg/kg.

Hemodynamics [HR and mean arterial blood pressure (MAP)] were assessed at 5 minutes interval from the time of reversal of muscle relaxant up to 30 minutes after extubation. The level of sedation during suction and extubation was assessed using observer assessment sedation score (Table: 1) and the airway response under direct laryngoscopy to suction was noted by five point scale (Table 2). (18) After 5 minute interval the level of sedation and smoothness of extubation was noted by four point scale (Table: 3). (18)

<table>
<thead>
<tr>
<th>Observation</th>
<th>Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Responds readily to name spoken in normal tone</td>
<td>5</td>
</tr>
<tr>
<td>Lethargic response to name spoken in normal tone</td>
<td>4</td>
</tr>
<tr>
<td>Responds only after name is called loudly and/or repeatedly</td>
<td>3</td>
</tr>
<tr>
<td>Responds only after mild podding or shaking</td>
<td>2</td>
</tr>
<tr>
<td>Does not responds to mild podding or shaking</td>
<td>1</td>
</tr>
</tbody>
</table>
### Table 2
**Grading of air way reflexes.**

<table>
<thead>
<tr>
<th>Grade</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Excellent (breathing well, no response to laryngoscopy &amp; suctioning)</td>
</tr>
<tr>
<td>2</td>
<td>Good (breathing well, minimal grimacing response to laryngoscope &amp; suctioning)</td>
</tr>
<tr>
<td>3</td>
<td>Satisfactory (breathing well coughing attempt to laryngoscopy &amp; suctioning)</td>
</tr>
<tr>
<td>4</td>
<td>Poor (breathing well, coughing on tube laryngoscopy)</td>
</tr>
<tr>
<td>5</td>
<td>Very poor (breathing well, coughing on tube laryngoscopy)</td>
</tr>
</tbody>
</table>

### Table 3
**Smoothness of extubation.**

<table>
<thead>
<tr>
<th>Grade</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>No coughing on endotracheal tube</td>
</tr>
<tr>
<td>2</td>
<td>Coughing on the tube</td>
</tr>
<tr>
<td>3</td>
<td>Vomiting</td>
</tr>
<tr>
<td>4</td>
<td>Laryngospasm</td>
</tr>
</tbody>
</table>

### Table 4
**Demographic data.**

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Group K (Ketofol), N = 53 Mean ± SD</th>
<th>Group P (Propofol), N = 53 Mean ± SD</th>
<th>P-Value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>27.68 ± 3.67</td>
<td>26.49 ± 3.06</td>
<td>0.073</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>164.13 ± 5.20</td>
<td>166.25 ± 5.05</td>
<td>0.036**</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>74.42 ± 10.19</td>
<td>73.36 ± 7.92</td>
<td>0.552</td>
</tr>
<tr>
<td>BMI</td>
<td>27.70 ± 4.18</td>
<td>26.56 ± 2.80</td>
<td>0.103</td>
</tr>
</tbody>
</table>

*Using independent t test, **p-value is significant at or below 0.05.
Table 5
Hemodynamic variables at different study time points.

<table>
<thead>
<tr>
<th>Group</th>
<th>Ketofol group (Ketofol), N = 53</th>
<th>Propofol group (Propofol), N = 53</th>
<th>P-value*</th>
<th>95% CI of the difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAP at (5) min before extubation.</td>
<td>107.19 ± 10.34</td>
<td>115.64 ± 11.01</td>
<td>&lt; 0.001**</td>
<td>-12.6, -3.4</td>
</tr>
<tr>
<td>MAP at the time of extubation.</td>
<td>107.49 ± 12.52</td>
<td>112.96 ± 11.10</td>
<td>0.019**</td>
<td>-10.0, -0.9</td>
</tr>
<tr>
<td>MAP at (5) min after extubation.</td>
<td>103.04 ± 13.00</td>
<td>104.36 ± 7.86</td>
<td>0.528</td>
<td>-5.4, 2.8</td>
</tr>
<tr>
<td>MAP at (10) min after extubation.</td>
<td>94.92 ± 5.59</td>
<td>97.66 ± 4.61</td>
<td>0.007**</td>
<td>-4.7, -0.8</td>
</tr>
<tr>
<td>MAP at (15) min after extubation.</td>
<td>90.43 ± 5.49</td>
<td>93.75 ± 5.28</td>
<td>0.002**</td>
<td>-6.7, -1.8</td>
</tr>
<tr>
<td>MAP at (20) min after extubation.</td>
<td>83.30 ± 5.99</td>
<td>90.15 ± 3.86</td>
<td>&lt; 0.001**</td>
<td>-8.8, -4.9</td>
</tr>
<tr>
<td>HR/min at (5) min before extubation.</td>
<td>85.47 ± 13.93</td>
<td>104.55 ± 10.83</td>
<td>&lt; 0.001**</td>
<td>-23.8, -12.4</td>
</tr>
<tr>
<td>HR/min at the time of extubation.</td>
<td>87.75 ± 1.22</td>
<td>103.72 ± 6.48</td>
<td>&lt; 0.001**</td>
<td>-19.5, -12.4</td>
</tr>
<tr>
<td>HR/min at (5) min after extubation.</td>
<td>82.26 ± 8.01</td>
<td>97.34 ± 7.03</td>
<td>&lt; 0.001**</td>
<td>-118, -12.2</td>
</tr>
<tr>
<td>HR/min at (10) min after extubation.</td>
<td>79.25 ± 7.36</td>
<td>92.81 ± 7.00</td>
<td>&lt; 0.001**</td>
<td>-16.3, -10.8</td>
</tr>
<tr>
<td>HR/min at (15) min after extubation.</td>
<td>75.36 ± 6.51</td>
<td>90.23 ± 6.08</td>
<td>&lt; 0.001**</td>
<td>-17.3, -12.4</td>
</tr>
<tr>
<td>HR/min at (20) min after extubation.</td>
<td>72.98 ± 5.40</td>
<td>86.38 ± 5.07</td>
<td>&lt; 0.001**</td>
<td>-15.4, 11.4</td>
</tr>
</tbody>
</table>

* using independent t test, ** p-value is significant at or below 0.05.
Our primary outcome was assessment of smoothness of extubation while, our secondary outcomes were assessment of airway response to laryngoscopy and suction, sedation score, and hemodynamics as mentioned before.

### Sample size calculation

Based on our pilot study, that was done on thirty patients who were randomly allocated by blind envelope methods into three groups ten patients each. These patients were not included in the final study. Sample size was calculated (G* power version 3.0.10). Minimal sample size was calculated as 48 patients in both groups needed to get the power level of 0.80, alpha level of 0.05 (two tailed), and effect size of 0.58 for grading of smoothness of extubation grade 1 (mean ± SD) in the study group and in the control group only is (1.5 ± 0.8) and (2.0 ± 1.2) respectively. Based on the results of the pilot study to overcome problem of loss of follow up, calculated sample size was increased by 10% to reach 53 in each group.
Statistical analysis

Statistical analysis was done SPSS for windows software version 23. Descriptive statistics was done for categorical variables and presented as numbers and percentages. While for numerical data, descriptive statistics were done the mean and the standard deviation being normally distributed. Independent t test was used to compare numeric variables in the two groups being either normally distributed data or based on the central limit theorem. Chi-Square test to analyze the difference in the extubation quality, the sedation score and the adverse events. P value < 0.05 was considered as statistically significant.

Results

A total of 106 patients participated in this study. They were randomized into the K group (53 patients) and the P group (53 patients). There were no statistically significant differences between the two study groups as regard the demographic data except for the height which is clinically insignificant (Table: 4).

Hemodynamic parameters (MAP and HR) 5 min before extubation, at extubation, and every 5 min later till 20 minutes post-extubation are detailed in (Table:5).

The MAP was significantly higher in the propofol group than in the ketofol group at all the readings except at 5 min after extubation where the difference was non-significant (Table: 5).

HR was lower in the ketofol group than in the propofol group at all reading times and that difference was statistically significant (Table: 5).

The grade of smoothness of extubation, our primary outcome, was 1 in all cases of the ketofol group and the grade was 2 to 4 for the cases of the propofol group. The relationship is highly significant and the Somers' D value = 1 meaning that there is a perfect trend association (Table: 6) and (Figure: 1).

The level of sedation during suction was 1 in 83% of the ketofol cases with lower numbers at levels 2 and 3. While in the propofol group, it was at level of 1 in19% of the cases whereas, the rest of the cases were at levels of 2 and 3. This relationship is highly significant and the Somers' D value = 0.67 meaning that there is a strong trend association (Table: 6).

The grade of airway reflex was 2 in 11 cases (20.8%) and 3 in 33 cases (62.3%) of the ketofol cases with lower numbers at grades 4 and 5. While in the propofol group, 12 cases (22.6%) of the cases were at grade 2 and 33 cases (62.3%) at grade 4. Also, this relationship is highly significant and the Somers' D value = 0.55 meaning that there is a strong trend association (Table: 6).

The level of sedation at extubation was 5 in all cases of the propofol group whereas; it was at the level of 2 and 3 for the cases of ketofol group. The relationship is highly significant and the Somers' D value = 1 meaning that there is a perfect trend association (Table: 6).
Discussion

Stress response during extubation is an unwanted and unpredictable response that makes anesthesiologists vigilant and attentive for minimizing its effect on hemodynamics and airway reflexes.

In our study we compared the effect of injecting a ketofol compared to propofol at induction of general anesthesia on the smoothness of extubation as regards airway response and hemodynamics in laparoscopic drilling of polycystic ovary disease.

Aboeldahab et al studied the effect of ketofol compared to its two constituents as an induction agent in 60 patients undergoing hernia repair under general anesthesia. They examined its effect as an induction agent clinically by assessing the hemodynamics and by using the BIS index where, 20 of them were given ketofol, 20 were given propofol and the last 20 were given ketamine during induction of anesthesia. During extubation HR in ketofol group (81.65 ± 2.60) was lower than in the propofol group (81.73 ± 3.93) but, with no statistical significance and MAP in the ketofol group (83.90 ± 3.30) was lower than in the propofol group (85.66 ± 3.43) with also, no statistical significance. This might be due to the lengthy procedure in their study and/or due to the small sample size of the groups. Our results came in agreement with Aboeldahab et al, as ketofol was associated with more stable hemodynamics than propofol during extubation. We attributed this to the good level of sedation of ketofol during suction and extubation due to the additive sedative effect of both ketamine and propofol.

We concluded that the effect of ketofol on HR was more important and more significant in stabilizing the hemodynamics than its effect on MAP when compared to popofol.

On the other hands, Jalili et al, who compared the effect of propofol and ketofol on emergence delerium in 87 ASA I and II children aged from 3–12 years and underwent adenotonsillectomy, reported no statistically significant difference between the two groups regarding HR in the recovery room at 0, 10, and 20 minutes postoperatively.

The smoothness of extubation without coughing, laryngospasm and vomiting on the tube was examined in both groups and it was in favor of ketofol group.

Recently, Aberra et al, in a study done on 120 pediatric patients aged from 2 to 15 years undergoing elective ophthalmic surgical procedures under general anesthesia using laryngeal mask airway (LMA) to compare the ketamine–propofol mixture (ketofol) with propofol on the ease of laryngeal mask airway insertion conditions and hemodynamic effects during induction of general anesthesia, found that 54 patients in the ketofol group compared to 52 in the propofol group developed no cough, 6 patients in ketofol group compared to 7 patients in propofol group developed slight cough (coughing which can occur immediately after LMA and subside by itself) and 1 patient in propofol group developed gross cough (coughing which needs deepening of anesthesia to be relieved) with no significant difference between them.
The sedation scores during suction and extubation were significantly lower in the ketofol group compared to the propofol group. We attributed this good level of sedation during suction and extubation to the analgesic effect of ketamine.

In our study, we reported that the ketofol group showed better airway response than the propofol group. One of the most unwanted adverse effects of stress response on airway is cough. Kim and Bishop reported an incidence rate of 75% of cough in patients during emergence and extubation. Hypertension, tachycardia, myocardial ischemia, and bronchospasm are adverse effects related to cough.

In our study, the majority of patients in the ketofol group [44 patients (83.1%)] developed better airway reflexes (grade 2,3) during suction better than the majority of patients in the propofol group [39 patients (73.6%)] developed higher degree of airway reflexes (grade 4 or 5). From all of the previous, we gave another explanation for the better hemodynamic stability during extubation in the ketofol group.

Aberra et al reported that no patient in the ketofol group developed laryngospasm while, 2 patients in the propofol group developed partial laryngospasm with no statistical significance. They concluded that Ketofol provided equivalent laryngeal mask airway insertion conditions and it can be used as an alternative propofol for LMA insertion.

**Conclusion**

Giving ketofol during induction of general anesthesia in laparoscopic drilling of polycystic ovary provided a good sedation score during suction and extubation, better airway response during suction, smoothness of tracheal extubation and more stable hemodynamics than propofol only group.

**Limitation**

We did this study in a relatively short surgical procedure that could be a limitation of our study.

**Abbreviations**

ASA PS  
Society of Anesthesiologists Physical Status.

HR  
Heart Rate.

GABA  
Gama Amino Butyric Acid.

ICU  
Intensive Care Unit.

LMA  
Laryngeal Mask Airway.

MAP
Declarations

• **Ethics approval and consent to participate:**
  
  • Fayoum University Ethical Committee approval.

• **Availability of data and material:**
  
  • All data are available.

• **Competing interests:**
  
  • No competing interest.

• **Funding:**
  
  Fayoum University Hospital resources and our own resources.

• **Authors Contribution:**
  
  All authors contributed equally in this work. **AM** invited the idea, shared in the statistical analysis and shared in writing the manuscript. **SG** shared in writing the manuscript and language editing. **JB** edited shared in writing, revised the manuscript, language editing and the corresponding author. All authors red and approved the manuscript.

• **Acknowledgements:**
  
  • our families.

References


**Figures**

**Figure 1**

comparing the smoothness of extubation between the two groups

**Supplementary Files**

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- [CONSORT2010Checklist12.doc](#)