Outcomes in video laryngoscopy studies from 2007 to 2017: systematic review and analysis of primary and secondary endpoints for a core set of outcomes in video laryngoscopy research

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Abstract

Background: Airway management is crucial and, probably, even the most important key competence in anaesthesiology, which directly influences patient safety and outcome. However, high-quality research is rarely published and studies usually have different primary or secondary endpoints which impedes clear unbiased comparisons between studies. The aim of the present study was to gather and analyse primary and secondary endpoints in video laryngoscopy studies being published over the last ten years and to create a core set of uniform or homogeneous outcomes (COS).

Methods: Retrospective analysis. Data were identified by using MEDLINE® database and the terms “video laryngoscopy” and “video laryngoscope” limited to the years 2007 to 2017. A total of 3351 studies were identified by the applied search strategy in PubMed. Papers were screened by two anaesthesiologists independently to identify study endpoints. The DELPHI method was used for consensus finding.

Results: In the 372 studies analysed and included, 49 different outcome categories/columns were reported. The items “time to intubation” (65.86%), “laryngeal view grade” (44.89%), “successful intubation rate” (36.56%), “number of intubation attempts” (23.39%), “complications” (21.24%), and “successful first-pass intubation rate” (19.09%) were reported most frequently. A total of 19 specific parameters is recommended.

Conclusions: In recent video laryngoscopy studies, many different and inhomogeneous parameters were used as outcome descriptors/endpoints. Based on these findings, we recommend that 19 specific parameters (e.g., “time to intubation” (inserting the laryngoscope to first ventilation), “laryngeal view grade” (C&L and POGO), “successful intubation rate”, etc.) should be used in coming research to facilitate future comparisons of video laryngoscopy studies.

Keywords: Airway management, Video laryngoscopy, Primary outcome, Primary endpoint
Background
Airway management is at least one crucial but probably even the most important key competence in anaesthesiology, which directly influences the safety and outcome of anaesthetised patients [1, 2]. Fortunately, anaesthesia-specific mortality has been significantly decreasing over the last decades and is now estimated to be approximately 1 per 100,000 cases [3–5]. Airway-related problems were reported to cause approximately 40% of anaesthesia-related deaths [6].

Mortality rate is approximately 5.6 per million general anaesthetics or one per 180,000 patients anaesthetised [7]. Taking these numbers into account, it is not surprising that airway management is a major research focus and each year thousands of studies are published analysing many specific problems during airway management [1]. Several specific patient groups have even a higher risk of problems [2].

Endotracheal intubation by using video laryngoscopy has significantly increased over the last decade in both pre- and in-hospital airway management [8]. Today, it is considered standard for difficult airway management and specific emergencies. It is even questioned whether it should be the first choice method.

However, high-quality research is rarely published [1] and studies usually have different primary or secondary endpoints, which impedes high quality comparisons between studies and hampers the possibility to draw meaningful conclusions to significantly and systematically improve safety and quality of clinical care. Different definitions and an inconsistent outcome reporting in studies which investigate comparable clinical problems will, therefore, limit results of research [9–11].

Insufficient attention has been paid on the choice of outcomes used for clinical trials in recent years [12]. To describe and analyse the same intubation performance, some studies use different “time” definitions and intervals (e.g., time to intubation, time to visualize glottis, time to place the endotracheal tube, etc.) and others use anatomical parameters (e.g., Cormack & Lehane grade [13], POGO score [14], etc.). Hence, no standard has been established to facilitate comparisons of results among different studies.

Consensus and consistency when using appropriate outcome measures in clinical trials should enhance the interpretation of research [9]. So far, no conclusive analysis of primary and secondary endpoints being used in studies has been published.

The aim of the present study was to gather and analyse primary and secondary endpoints in video laryngoscopy studies published over the last ten years, i.e., during 2007 to 2017. This data is used to create a basis for development of a core set of outcomes items to be used to facilitate comparisons in future trials. Besides parameters found in published literature, the list would be amended by parameters considered essential in airway management studies.

Methods
Systematic PubMed search
Data gathering was performed using MEDLINE® database (http://www.ncbi.nlm.nih.gov/pubmed). To identify relevant literature, the search terms “video laryngoscopy” and “video laryngoscope” were used.

A total of 3351 studies were identified by the applied search strategy in PubMed (Fig. 1). First, a filter restricting the time period of the search (10 years range; going from 22/June/2007 to 22/June/2017) was applied. The PubMed® article categories selected were “Clinical Study”, “Clinical Trial”, “Comparative Study”, “Controlled Clinical Trial”, “Evaluation Studies”, “Multicenter Study”, “Observational Study” and “Randomized Controlled Trial”. The final raw dataset consisted of 582 papers (the number of results is referred to a search made on 11/July/2017).

The complete list of items, including whole article names, authors and PubMed® URLs as well as the table of the results sorted by year was downloaded directly
from PubMed in a CSV format. Manuscripts presenting scientific data on video laryngoscopy as well as outcome parameters were included for analysis. If outcome parameters were not presented, the specific manuscript was excluded for analysis.

Data analysis
Papers were screened manually by two anaesthesiologists to identify study endpoints. For each study, “primary outcome”/”primary endpoint” or “secondary outcome”/”secondary endpoint” were collected if clearly stated in the abstract and/or in the full text (when available in the University of Naples Federico II or University of Cologne digital libraries).

If a study did not contain parameters described as “primary outcome/endpoint” or “secondary outcome/endpoint”, alternative measurements were included in the analysis. Alternatively, the “aim/goal/objective/target/purpose/null hypothesis” information was used from the abstract. Also, all studies evaluating directly a video laryngoscopy system or evaluating how the effectiveness of these laryngoscopes could be improved by other ancillary devices (e.g., different endotracheal tubes, stylets, gum elastic boogie, etc.) were included.

A chess-like table was hence built with the several outcomes as columns and 372 papers included as rows (Fig. 2). Outcomes from different studies, having different names but concerning the same variable were fused in a single column (e.g., “dental compression” and “number of audible dental click sounds”). Outcomes concerning a group of variables, slightly different among the studies but mostly overlapping were fused, as well.

Data processing
The number of row-column matches for each column/category was reported in a different table and the ratio between it and the total number of included articles was calculated. Derived from the percentages of each parameter, a suggestion was provided of which parameters should be reported in future video laryngoscopy studies to facilitate study comparisons.

Generation of recommendations
The chosen outcomes outcomes need to be relevant to both health care providers and healthcare users on one hand, and also to those involved in making decisions and choices about health care, on the other hand [12]. However, a lack of attention for using clinical outcomes in studies has led to avoidable losses in both the production and reporting of research. Moreover, the outcomes which have been included in studies have not always been those being most important or relevant for patients [15]. To develop a consensus between the authors concerning use of different parameters, the Delphi system was used [16, 17].

To develop relevant recommendations for video laryngoscopy studies, the four-step Core Outcome Set (COS) process was used. First, the scope was defined, followed by checking if a set exists as a second step. Third, a procedure for the development of the COS was defined and last, it was defined what specific parameters should be measured in future studies [12].

In a COS framework, the method is used for achieving convergence of opinion from experts on the importance of different outcomes in sequential questionnaires (or rounds) sent either by post or electronically [12]. In the present study, all authors (n = 4) participated in the process. Three rounds were planned. The answers for each of the outcomes were summarised and fed back anonymously until a consensus was reached with at least 75%. After considering the views of others before re-rating each item, participants were able to change their initial responses based on the feedback from the previous rounds. Direct communication concerning the specific parameters was not possible. Therefore, the feedback provides a mechanism for reconciling different opinions of participants and is essential to achieving a consensus [12]. In terms of the overall validity for the final consensus, this approach has significant advantages as compared to round-table discussions [18].

Results
Number of studies
A total of 3351 studies were identified by the applied search strategy in PubMed (Fig. 1). Using the filters for date and article type, a reduction of 2769 papers was obtained. The final raw dataset consisted of 582 papers. During the detailed analysis, 210 papers were excluded because they were considered off topic or belonged to an article type different from the ones chosen during the search filters setting (e.g. meta-analysis, review, etc.). Of the n = 210 articles excluded, n = 169 (80.48% of 210) were off the topic, n = 23 (10.95% of 210) did not evaluate directly a video laryngoscopy system or the effectiveness of the association with ancillary devices, n = 13 (6.19% of 210) belonged to a category of articles not included in the search strategy and n = 5 (2.38% of 210) did not provide an endpoint.

After exclusion of not relevant papers, N = 372 were considered eligible for final analysis (Fig. 1).

Analysis
The analysis of each item, excluding the off topic articles or the not chosen article types, led to a table consisting of 49 outcome categories depicted horizontally and 372 publications depicted vertically (Fig. 2). The ratio between the number of row-column matches for each
outcome category (numerator) and the total number of included articles (denominator) was calculated (the total number of outcome categories was not chosen as denominator since it could be more susceptible to subjective evaluation).

Among the total of 49 collected parameters, the items “time to intubation” (65.86%), “laryngeal view grade” (44.89%), “successful intubation rate” (36.56%), “number of intubation attempts” (23.39%), “complications” (21.24%), “successful first-pass intubation rate” (19.09%) were reported most frequently in the investigated studies (Table 1). Furthermore, these items were grouped in eight parameter categories (Table 1). Besides these six parameters having the highest reporting rate in previous studies and found with the purely computationally approach, additional six parameters were identified in the Delphi round. The Delphi method was used to find a consensus what parameters are of utmost importance. These parameters were “time to glottis view”, “ease of intubation (subjective scoring)”, “dental compression AND number of audible dental click sounds”, “optimization manoeuvres AND use of airway back-up devices”, “haemodynamic parameters”, and “lowest arterial oxygen saturation”. We, therefore, highlighted top twelve of outcomes with a prevalence range from 65.86 to 3.49% (Table 1).

Besides the previously reported parameters, seven additional parameters were identified with the Delphi round which should be reported in future airway management studies. Two of them should be reported in any study (patient and manikin study): “exact specifications of the device used” and “exact specifications of the patient group”. For patient studies, the parameters “death”,

Fig. 2 Primary and secondary endpoints of the analysed studies. The parameter with the highest incidence is depicted on the left, with the lowest incidence on the right. This figure shows graphically how often parameters were used in the studies analysed. On the x-axis (columns): from P1 to P49 the different parameters analysed (for their definitions see Table 1). On the y-axis (lines): the specific studies (i.e., n = 372). For the percentages of each specific parameters see Table 1.
Table 1 Frequency of primary and secondary endpoints used in the studies analysed. The “Top-12” of the used parameters are indicated in Bold-type

| Outcome Name | Number of Matches | Percent | Column Name |
|--------------|-------------------|---------|-------------|
| Time         |                   |         |             |
| Time to intubation (#1) | 245 | 65,86% | P1 |
| Time to glottic view (#2) | 36 | 9,68% | P9 |
| Endotracheal tube insertion time | 18 | 4,84% | P15 |
| TTSt on the first attempt | 1 | 0,27% | P18 |
| Time to ventilation after intubation | 7 | 1,88% | P23 |
| Total time of chest compression interruption during ETIb | 2 | 0,54% | P36 |
| Time to supraglottic ventilation | 1 | 0,27% | P38 |
| Views |                   |         |             |
| Laryngeal view grade (CL AND/OR POGOj) (#3) | 167 | 44,89% | P2 |
| Intubation Success |                   |         |             |
| Successful intubation rate (#4) | 136 | 36,56% | P3 |
| Successful first-pass intubation rate (#5) | 71 | 19,09% | P6 |
| Ease of intubation (subjective scoring) (#6) | 48 | 12,90% | P7 |
| Failed intubation | 23 | 6,18% | P11 |
| Intubation difficulty score (IDS) | 22 | 5,91% | P12 |
| Factors complicating intubationa | 11 | 2,96% | P19 |
| Proportion of difficult intubation | 6 | 1,61% | P25 |
| DoubleLumenTube position | 4 | 1,08% | P27 |
| Successful tracheal intubation rate after failed initial laryngoscopy | 4 | 1,08% | P28 |
| Intubation success rate in patients with difficult laryngoscopy predictors | 2 | 0,54% | P32 |
| Reason for intubation failure | 2 | 0,54% | P34 |
| Likelihood of successful intubation | 2 | 0,54% | P35 |
| Adequate ETTf position | 1 | 0,27% | P41 |
| Factors that affect FPSg in trauma patients | 1 | 0,27% | P43 |
| Proportion of successful to failed intubations | 1 | 0,27% | P45 |
| Accuracy of correct unilateral placement | 1 | 0,27% | P48 |
| Number of attempts |                   |         |             |
| Number of intubation attempts (#7) | 87 | 23,39% | P4 |
| Number of tube insertions | 2 | 0,54% | P37 |
| Complications |                   |         |             |
| Complicationsh (#8) | 79 | 21,24% | P5 |
| Dental compression AND number of audible dental click sounds (#9) | 24 | 6,45% | P14 |
| Severity of force applied to the upper airway | 10 | 2,69% | P20 |
| Variables reflecting morbidityi | 6 | 1,61% | P24 |
| Potential laryngeal trauma | 2 | 0,54% | P33 |
| Gagging severity score at the time of best laryngeal visualization | 1 | 0,27% | P44 |
| Device use & operator variables |                   |         |             |
| Optimization manoeuvres AND use of airway back-up devices (#10) | 48 | 12,90% | P8 |
| Device difficult score | 20 | 5,38% | P13 |
| Device preference | 9 | 2,42% | P21 |
| Overall participant satisfaction | 7 | 1,88% | P22 |
| Ergonomicsj | 3 | 0,81% | P29 |
| Postural analysis | 3 | 0,81% | P30 |
| Learning process | 3 | 0,81% | P31 |
| Reasons for using methods other than McGrath MAC video laryngoscope | 1 | 0,27% | P39 |
Table 1 Frequency of primary and secondary endpoints used in the studies analysed. The ‘Top-12’ of the used parameters are indicated in Bold-type (Continued)

| Outcome Name                                                      | Number of Matches | Percent | Column Name |
|------------------------------------------------------------------|-------------------|---------|-------------|
| Practitioner experience                                          | 1                 | 0.27%   | P49         |
| Monitoring                                                       |                   |         |             |
| Haemodynamic parameters (#11)                                    | 30                | 8.06%   | P10         |
| Lowest arterial oxygen saturation (#12)                          | 13                | 3.49%   | P16         |
| Cervical vertebral angle                                         | 12                | 3.23%   | P17         |
| SpO2 immediately after removing the blade from the patient       | 1                 | 0.27%   | P40         |
| Bispectral index score                                           | 1                 | 0.27%   | P46         |
| Intraocular pressure                                             | 1                 | 0.27%   | P47         |
| Other                                                            |                   |         |             |
| Airway grade<sup>6</sup>                                         | 6                 | 1.61%   | P26         |
| Intubation conditions<sup>7</sup>                                | 1                 | 0.27%   | P42         |

<sup>1</sup> Time to successful intubation
<sup>2</sup> Endotracheal intubation
<sup>3</sup> Cormack-Lehane score and Percentage Of Glottic Opening
<sup>4</sup> mainly a visual analogue scale score ranging from 1 (extremely easy) to 10 (extremely difficult) with several exceptions (e.g., numerical rating scale 1 = the easiest, 5 = the most difficult)
<sup>5</sup> e.g., visualization difficulty related to obscured view from fogging, secretions or blood in the airway; difficulty passing the tracheal tube past the vocal cords; inappropriate endotracheal tube size for the patient; or difficulty controlling the direction of the tracheal tube using the video display
<sup>6</sup> Endotracheal tube
<sup>7</sup> First-pass success
<sup>8</sup> Pre- and post-intubation correlated complications (e.g., upper airway morbidity, swallowing difficulties or any dental injuries)
<sup>9</sup> e.g., in-hospital mortality, hospital length of stay, duration of mechanical ventilation, duration of ICU stay, ICU mortality, etc
<sup>10</sup> Biomechanical performance of doctors during the ETI (e.g., assessed using surface electromyography and inertial measurement units)
<sup>11</sup> Airway assessment predictors: Mallampati test, mouth opening, thyromental distance, cervical flexion-extension, and neck thickness, snoring, retrognathia, and other types of anomalies also considered as predictors of a difficult airway
<sup>12</sup> Ease of Laryngoscopy, Vocal cords position, Reaction to insertion of the tracheal tube and cuff inflation (Diaphragmatic movement/coughing), direction of the ETT by the forceps and advancement of the ETT by the forceps

Table 2 Suggested minimal endpoints categories for reporting of video laryngoscopy studies. The table consists of 12 previously reported parameters plus seven additional parameters from the Delphi round

| Category                          | Parameter                                                                 |
|-----------------------------------|---------------------------------------------------------------------------|
| Time                              | Time to intubation (taking the laryngoscope to first successful ventilation) |
|                                   | Time to glottis view                                                      |
| View                              | Laryngeal view grade (CL and POGO)                                        |
| Intubation Success                 | Successful first-pass intubation rate                                     |
|                                   | Successful intubation rate                                                |
|                                   | Ease of intubation                                                        |
| Number                            | Number of intubation attempts                                             |
| Complications                     | Any clinically significant complication                                   |
| Devices                           | Dental compression AND number of audible dental click sounds              |
| Monitoring                        | Optimization manoeuvres AND use of airway back-up devices                 |
| Patients                          | Hemodynamic parameters                                                   |
| Additional parameters (not covered by the studies before)          | Lowest arterial oxygen saturation                                         |
| Patients outcome                  | Death                                                                     |
|                                   | ICU admission                                                             |
|                                   | hospital length of stay                                                   |
|                                   | dysphagia                                                                 |
|                                   | reduced quality of life                                                   |
| Patients                          | Exact specifications of the patient group                                 |
| Devices                           | Exact specifications of the device used                                   |
“ICU admission”, “hospital length of stay”, “dysphagia”, and “reduced quality of life” should be reported.

Discussion
Over the last ten years, many different and inhomogeneous parameters were used as outcome descriptors/endpoints in video laryngoscopy studies. In order to facilitate literature comparison, taking into account the percentages of items used in previous publications, we suggest that 12 parameters should be used in future video laryngoscopy studies (Table 2). Additionally, the seven patient outcome parameters not covered by the studies before should be reported.

Video laryngoscopy studies
The use of video laryngoscopes has increased significantly over the last years for many pre-hospital and in-hospital situations [8]. Today, it is considered standard for difficult airway management and emergencies and in some scenarios, it is even questioned whether it should be the method of first choice. Whereas the use of video laryngoscopes was limited to elective intubations several years ago, especially for the anticipated difficult airway, these devices are used today for a broad spectrum of indications such as anticipated difficult airway, teaching and training, or even awake intubation. As airway management is a topic of major research interest and each year thousands of studies that probe all the specific problems of airway management are published, even video laryngoscopy studies seek to compare a variety of devices, arbitrarily chosen, in a variety of settings [1].

In previous studies, a great multiplicity of measured outcomes has been subsequently used to assess the capability of video laryngoscopes to modify the intubation-related variables in comparison to the classic direct laryngoscopy or within the category itself among the different devices [1]. However, none of these outcomes are present in all studies and in some cases, like in the case of the “time to intubation” endpoint, divergence subsists in single definitions, making it difficult to perform any comparison between the outcomes obtained in different articles.

The aim of the study was, therefore, to provide a simple analysis of a part of the scientific literature on the subject so that it would be possible to derive a common basis on which future studies on video laryngoscopy can be built. Standardizing endpoints will also improve the validity of pooled analysis of clinical trials and assist those wanting to replicate trial results [9]. Besides parameters found in published literature, the list will be amended by parameters considered essential in airway management studies.

Set of parameters
Nearly no analysed study used the same set of parameters to quantify and qualify performance of intubation with a video laryngoscope. Furthermore, parameters used were often non-specific and not clearly defined, since so far, in this field of research, no consolidated minimal reporting dataset does exist, unlike in other fields of research [19].

This problem of definition, even for the meaning of single parameters, is well represented in the main category for prevalence, i.e. “time to intubation”. Three examples could well reflect the high variability in definition since time to intubation is described as (i) the time “from the passage of the tip of the laryngoscope past the patient’s teeth to the appearance of CO 2 on the capnograph trace” [20], (ii) the moment “from when the facemask was removed from the patient’s face to when end-tidal CO 2 of at least 20 mm Hg was measured on
the end-tidal gas monitor” [21] or when (iii) “the time started to run when a participant took a laryngoscope into his/her hand and stopped when the appropriate position of the tube was confirmed by the fact that it was possible to ventilate with a bag valve mask and by the movements of the chest and the abdomen” (Fig. 3) [22].

Future aspects
From a practical point of view, several (comparable) parameters and categories should be reported in video laryngoscopy studies (Table 2). This may enhance comparisons of parameters in different studies and facilitate meta-analyses as well as systematic reviews in video laryngoscopy studies. Comparison between studies is made easier, and other investigators will have a stronger foundation on which to design future, definitive trials [9].

Usefulness of parameters
In the present study, a set of 19 parameters for video laryngoscopy studies is presented. Whereas using many different parameters increases the possibility to compare different studies, it may be cumbersome to record all parameters. Besides this practical point, the feasibility of the different parameters itself varies significantly. Whereas, e.g., “time to intubation”, “number of intubation attempts”, or “successful intubation rate” are quite clear and objective to assess, “ease of intubation” and “hemodynamic parameters monitoring” are far more subjective. Moreover, “laryngeal view grade” or “optimization manoeuvres AND use of airway back-up devices” clearly depends on the skills and expertise as well as anatomical factors of the patient. Therefore, it is essential to keep these limitations also in mind when comparing different studies.

Limitations
The present study provides an overview on parameters used in previous studies on video laryngoscopy. However, it has also some limitations which should be mentioned. From a total of \( n = 582 \) articles identified, only \( n = 372 \) (63.92%) could be included due to the inclusion-/exclusion-criteria mentioned.

Furthermore, not all studies provided information in which model video laryngoscopes were investigated: if in patients, in cadavers, or in manikins. Finally, the definition of video laryngoscopes is quite broad and comparison between the different available models is often impossible.

Conclusions
Over the years, many different and inhomogeneous parameters were used as outcome endpoints in video laryngoscopy studies.

The final result of parameters offers several recommendations for choosing the endpoints, but the fact remains that these endpoints are still numerous, which reflects the literature on this field although we have demanded a consensus limiting the endpoints to those most relevant and most clinical based from the experts. The example of “Laryngeal view grade” (CL and/or POGO), even if widely cited, does not reflect the difficulty of intubation in video laryngoscopy [23].

The standardization of endpoints for video laryngoscopy studies could lead to improve the effectiveness of literature review and facilitate a more valid comparison between outcomes obtained in future studies.

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Authors’ contributions
JH and PK conceived the study. JH and II performed data analysis and wrote the manuscript. PK and EDR significantly improved the manuscript and checked analyses. All authors read and approved the final manuscript.

Ethics approval and consent to participate
Not required since retrospective data analysis.

Consent for publication
Granted. Not applicable.

Competing interests
J.H. is chairman of the Working group ”Standards, Recommendations, and Guidelines” from the German Society of Aerospace Medicine (DGARM) as well as of the European Society of Anaesthesiology (ESA) Guideline Task Force on “Intraoperative Cardiac Arrest”. P.K. is member of the ESA guidelines committee and E.D.R. is past-chairman of the ESA guidelines committee. For all other authors: NONE.

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