Research Article

How to Improve Reprocessing of Flexible Endoscopes Nationwide? Data from the German Colorectal Cancer Screening Program

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Background and Aims. International studies revealed prevalences of around 50% of microbiological contaminations in reprocessed flexible endoscopes. In Germany a system was installed where the qualification for refund for colonoscopies was made conditional on successfully passing twice annually a microbiological surveillance test of reprocessed endoscopes. This study is an implementation and outcome evaluation as well as a general discussion of the quality assessment assurance in colonoscopy in Germany.

Methods. German data from 2003–2008 were analysed: number of endoscopic units performing therapeutic and/or screening colonoscopies; result of all microbiological surveillance tests of reprocessing quality; number of failed surveillance tests and retests; number of qualifications for refund from the public health system cancelled due to repeated failure of microbiological surveillance tests.

Results. After the introduction of the quality assessment assurance, the percentage of failed microbiological surveillance tests dropped significantly and steadily from close to 17% to below 5%.

Conclusions. This study evidences (1st) the successful implementation of the quality assessment assurance in Germany and (2nd) a substantial improvement in the quality of reprocessing flexible endoscopes achieved by these measures.

1. Introduction

Flexible endoscopes invariably become contaminated with microorganisms during clinical use. With the more widespread use of flexible endoscopy 30 years ago, nosocomial infections associated with oesophago-gastro-duodeno-pancreaticoscopy (ERCP), or bronchoscopies were observed and described in the literature [1–9]. A survey in the 70ies yielded an estimated risk of 1 per 10,000 endoscopic examinations (oesophagogastroduodenoscopies und colonoscopies) [10]. More recently, optimistic estimates suggest a lower risk of 1 in 1.8 million [11] or 1 in 5 million procedures [12] given millions of endoscopic examinations worldwide. Data from the late nineteen-seventies report an infection rate of 0.74% following endoscopic retrograde cholecysto-pancreaticoscopy (ERCP) [7, 8, 13]. The risk for transmission of, for example, Helicobacter pylori has been considered to be likely around of 2-3 per 1000 oesophagogastroduodenoscopies [14–16]. The problem of possible infections originating from endoscopic examinations has been controversially discussed by the media, that is, in Germany [17] and the US [18]. In Germany, this culminated in the repeated statement of a notable hygienist that he himself refrained from endoscopic examinations due to the risk of infection [19].

To minimize the risk of infection transmission via endoscopes many gastroenterological and hygienic societies worldwide developed national guidelines for reprocessing flexible endoscopes and endoscopic accessories [20–35]. However, implementation of these guidelines in clinical practice and outcome are not evaluated at all. Only small surveys based on questionnaires [36–44] are available, hard data on the effect of national reprocessing guidelines on reprocessing quality are still lacking.
The situation in Germany is different from other countries. In 2002, based on the results of the HYGEA study [45] confirming similar studies in other countries [22, 46] and a comparison of international guidelines for reprocessing flexible endoscopes [27] the commission for hospital hygiene and disease prevention at the Robert Koch institute, the German analogue to the Centers for Disease Control (CDC) published an evidence-based guideline for reprocessing flexible endoscopes and endoscopic accessories [47]. In addition, a national colorectal cancer screening program via colonoscopy was initiated in autumn 2002. Before nationwide implementation of this screening program a quality assurance agreement was established by the Kassenärztliche Bundesvereinigung (KBV) and the stakeholder of the public health insurances to ensure that screening colonoscopy will be performed safely and without risk for the screened patient. According to this quality assurance agreement [48] both qualification indicators for the performing gastroenterologist were defined (see Section 4) and monitoring of colonoscopy reprocessing by microbiological cultures was implemented. Certification to participation in the colorectal cancer screening program was given by the local KV only after documented fulfilling of these qualification requirements and financial funding for screening colonoscopy was coupled to compliance with the quality indicators and negative surveillance cultures of reprocessed colonoscopies. Under coordination by the central KBV and supervision and control by local Kassenärztliche Vereinigungen (KV) in the different states of the Federal Republic of Germany a nationwide system was established with certified microbiological laboratories performing two times per year microbiological surveillance cultures from swabs from the tip of the colonoscope, from flushing solution of disinfected channels of the instrument, and from water bottle solution of the optic flushing system.

The present study is an implementation and outcome evaluation of the German quality assurance agreement with the specific aims to analyse surveillance culture data from 2002 to 2008 in detail and to assess whether linking reprocessing quality indicators to funding of colonoscopy will improve colonoscopy reprocessing quality in the long run.

2. Materials and Methods

The following nationwide data, per year and per association of statutory health insurance physicians (in German: Kassenärztliche Vereinigung (KV)), were made available through the framework of the quality assurance agreement for the time period of 2003 to 2008: the number of endoscopic units performing therapeutic and/or screening colonoscopies within each KV, the results of all microbiological surveillance tests of reprocessed endoscopic units (two per year per unit), the number of failed surveillance tests and failed retests, and the number of qualifications for refund from the public health system cancelled due to repeated failure of microbiological surveillance tests. All surveillance tests were conducted by independent, accredited microbiological institutes/laboratories.

The frequencies of actually performed hygiene surveillance tests per year were expressed as a percentage of the tests prescribed by the assurance system, that is, 100% are reached when two tests per year were conducted per endoscopic unit. The frequencies of failed microbiological surveillance tests per year (i.e., the number of necessary retests) were expressed as a percentage of the number of actually conducted routine surveillance tests. The frequencies of repeated failure of microbiological surveillance tests per year were also expressed as a percentage of the number of actually conducted initial routine surveillance tests. The frequencies of qualifications for refund from the public health system cancelled due to repeated failure of microbiological surveillance tests were listed as raw numbers.

Analyses of variance for repeated measurements were used to statistically assess the time trends of the above detailed percentages. An overall alpha level of 5% was used. Pairwise comparisons between consecutive years were analysed by means of paired t-tests where a Bonferroni correction for multiple testing was applied.

3. Results

The time trends from 2003 to 2008 of the following key microbiological performance parameters are detailed in Table 1:

1) The number of endoscopy units accredited in Germany for therapeutic and/or preventive colonoscopies.

2) Percentages of actually performed surveillance tests (out of those prescribed by the quality assurance agreement). Since some associations of statutory health insurance physicians did not provide data for all years, two rows of data are presented: all available information (here the number of reporting associations may differ over the years) and complete information only (only those associations included that reported data for all years of the study period). The latter formed the basis for statistical tests.

3) Percentages of failed microbiological surveillance tests out of all conducted tests. Again two rows are displayed: for all information available and for complete series only.

4) Percentages of repeated failures of microbiological tests out of all conducted initial tests. Again two rows are displayed: for all information available and for complete series only.

5) The number of qualifications for refund from the public health system cancelled due to repeated failure of microbiological tests.

The time trend of percentages of conducted surveillance tests of all KV’s with complete data over the whole study period (n = 11) are depicted in Figure 1. The overall trend is statistically significant (P < 0.001). Consecutive pairwise comparisons only reveal a significant change between 2003 and 2004. The time trend of percentages of failed tests (out of conducted tests) of all KV’s with complete data (n = 10) is shown in Figure 2. The overall trend is statistically significant (P < 0.001). Consecutive pairwise comparisons reveal...
ISRN Endoscopy

Table 1: Time trends of key microbiological performance data in endoscopy in Germany from 2003 to 2008.

<table>
<thead>
<tr>
<th></th>
<th>2003</th>
<th>2004</th>
<th>2005</th>
<th>2006</th>
<th>2007</th>
<th>2008</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accredited endoscopic units</td>
<td>1695</td>
<td>2176</td>
<td>2645</td>
<td>2567</td>
<td>2583</td>
<td>2476</td>
</tr>
</tbody>
</table>

Percentages of conducted surveillance tests: all information used
- 2003: 56.0% (13)
- 2004: 94.9% (14)
- 2005: 97.4% (17)
- 2006: 93.1% (16)
- 2007: 97.6% (17)
- 2008: 96.4% (16)

Percentages of conducted surveillance tests: only complete information used
- 2003: 58.5% (11)
- 2004: 94.2% (11)
- 2005: 96.8% (11)
- 2006: 98.5% (11)
- 2007: 96.3% (11)
- 2008: 94.8% (11)

Percentages of failed tests: all information used
- 2003: 16.6% (12)
- 2004: 9.7% (14)
- 2005: 5.1% (16)
- 2006: 5.1% (16)
- 2007: 3.7% (17)
- 2008: 4.2% (16)

Percentages of failed tests: only complete information used
- 2003: 16.5% (10)
- 2004: 9.9% (10)
- 2005: 6.7% (10)
- 2006: 5.8% (10)
- 2007: 3.4% (10)
- 2008: 5.6% (10)

Percentages of repeated failures: all information used
- 2003: 0.9% (11)
- 2004: 1.4% (14)
- 2005: 0.8% (14)
- 2006: 0.2% (16)
- 2007: 0.3% (17)
- 2008: 0.2% (16)

Percentages of repeated failures: only complete information used
- 2003: 0.8% (9)
- 2004: 0.9% (9)
- 2005: 0.4% (9)
- 2006: 0.1% (9)
- 2007: 0.1% (9)
- 2008: 0.1% (9)

Number of accreditations cancelled due to repeated failure
- 2003: 5
- 2004: 3
- 2005: 17
- 2006: 2
- 2007: 9
- 2008: 6

Stated are the raw numbers and percentages as indicated. In brackets the numbers of the reporting associations of statutory health insurance physicians are given.

significant changes between 2003 and 2004 and between 2004 and 2005. The overall time trend of repeated failures (n = 9) is significant (P < 0.01). Consecutive pairwise comparisons revealed only between 2004 and 2005 a significant result.

4. Discussion

Since colorectal cancer is one of the main cancers in Western societies accounting for more than 50,000 deaths each year in the United States [49] and more than 30,000 deaths each year in Germany [50], colonoscopy as the gold standard for early detection of adenomas and carcinomas in symptom-free individuals is considered as a mass screening tool to reduce incidence and mortality from colorectal cancer by detecting and removing potential precursor lesions.

4.1. General Remarks: Quality Management of Screening Colonoscopy. Several studies have shown that colonoscopic polypectomy reduces colorectal cancer incidence [51–60]. Furthermore, screening with colonoscopy is costeffective compared to other screening strategies [61,62].

However, colonoscopy is an invasive procedure and not without risk. Therefore, those performing this test have to be well trained and experienced. In addition, in order to remove public concerns regarding the risk of infection transmission, reprocessing of the screening instrument, the flexible colonoscope has to be in accordance with recent evidence-based guidelines [20–35].

Data on the process quality and the incidence of acute complications and data on the diagnostic yield of screening colonoscopy of the German colorectal cancer screening program were recently published [63,64].

In addition, Germany was the first country worldwide having realized continuous monitoring of colonoscope reprocessing by microbiological surveillance cultures of reprocessed endoscopes.
4.2. The German System: Implementation of a Quality Control Program for Endoscope Reprocessing. In order to establish and maintain high-quality standards for screening colonoscopy the public health insurances and the KBV, the main players of the ambulance health system in Germany [65–67], arranged a quality assurance agreement before starting the German colorectal screening program by colonoscopy. According to this agreement [48] the following quality indicators for the performing gastroenterologist were defined as a marker for experience—a minimal number of more than 200 colonoscopies and 20 polypectomies per year—as a marker for performance—(electronic or videoprint) documentation of the ileocecal valve or the terminal ileum and—as a marker for staff performance/equipment reproprocessing quality—negative surveillance cultures of disinfected colonoscopes two times per year. The central issue in this quality management program consists in payment for performance and continuous documentation of the quality indicators for colonoscopy mentioned above. Each gastroenterologist, performing endoscopies in his office and fulfilling the qualification requirements, can participate in the German colorectal cancer screening program.

For a nationwide colonoscopy screening program, availability of an experienced gastroenterologist as well as minimizing problems concerning infection risk are crucial steps for acceptance of the screening program. Thus, the German colorectal cancer screening program was coupled and coordinated with a quality assurance agreement implementing outcome monitoring of endoscope reproprocessing. The present analysis is based on the collected data of this quality control program. Completeness and quality of the data needs critical comments. In 2003, the first year of implementation of this control system, surveillance cultures were not performed twice a year but only once a year. For the years 2003 and 2004 in several federal states only limited data were available, collection was done in retrospective and completeness of the data was questionable. Thus, in general data quality and completeness are somewhat limited for the first two years, but become valid in 2005 and the following years.

4.3. German and International Guidelines in Endoscope Reprocessing: Do We Need Surveillance Cultures? The German guideline for reproprocessing flexible endoscopes published in 2002 [26] was the first evidence-based guideline for scope reproprocessing and was based on a profound literature review comparing international guidelines for endoscope reproprocessing prepared by a working group [27] and the results of a large study on hygiene in endoscopy performed in the area of Munich in 2000 [45]. One and a half year later, several gastroenterological, microbiological, and hygienic societies in the United States published a similar evidence-based multisociety guideline [68].

One of the most important differences of these two guidelines is the answer to the problem whether microbiological surveillance cultures of reproprocessed endoscopes should be recommended or not. It is absolutely correct, that no data exist demonstrating that regular microbiological surveillance cultures of endoscope reproprocessing decrease the risk for infection transmission during endoscopy. Thus, the interpretation of the multisociety guideline [68] and other authors [69] that the value of regular microbiological surveillance cultures of reproprocessing is an “unresolved issue” is principally correct. In accordance with this statement microbiological controls of endoscope reproprocessing quality are not performed in the United States [49]. In recent years, however, usefulness of surveillance cultures is mentioned more often in the United States [37, 40, 44, 69].

In contrast to the multisociety guideline [68] and considering the poor compliance with hygienic recommendations and variability in implementation [70, 71] the German guideline recommended regular surveillance cultures of endoscope reproprocessing [26]. The recent European guideline for endoscope reproprocessing, published in 2007 [24], also recommended microbiological controls of endoscope reproprocessing quality.

The experience of smaller studies from Germany [72, 73], the results of the former nationwide study [74], and the present update reinforce the hypothesis underlying the German RKI guideline that measurement and feedback of reproprocessing quality will help to implement reproprocessing procedures in accordance with present guidelines. In other words and from a quality management point of view: feedback of reproprocessing outcome quality enhances the process quality and compliance with reproprocessing guidelines which in turn vice versa affects outcome quality.

4.4. Process Quality or Outcome Quality of Reprocessing Complex Medical Devices? The question whether we should focus on process quality or outcome quality of reproprocessing flexible endoscopes is still a matter of debate. Although standardized reproprocessing is possible in special washer-disinfectors using automatic single channel cleaning with enzymatic cleaners and programmed disinfection with glutaraldehyde at higher temperature [75], only microbiological controls of reproprocessing quality are able to detect equipment defects of flexible endoscopes (such as perforation of internal channels) which affect the outcome of the reproprocessing process and which would be without surveillance cultures only recognized in working ups of nosocomial infection outbreaks [76].

4.5. Summary and Conclusions. The quality assurance agreement for screening colonoscopy in Germany in 2002 [58] improved colonoscopy reproprocessing quality significantly. After implementation a quality control system for colonoscopy reproprocessing in ambulance offices the rate of positive surveillance cultures from reproprocessed endoscopes dropped from 50% of failed tests observed before the introduction in 2000–2001 to 17.7% in 2003 and to 10.1% in 2004 and reached a plateau of under 5% in 2008. Thus, the ethical principle that a screening procedure, such as a screening colonoscopy should be without risk of infection transmission, seems nearly fulfilled.

The question whether monitoring endoscope reproprocessing by regular microbiological surveillance cultures will reduce the infection transmission during endoscopy cannot
yet be answered by the present study. However, our data strongly favor regular surveillance cultures as an important catalyst for implementation and adherence to national endoscope reprocessing guidelines and for improvement of reprocessing quality. So in order to reduce infection transmission by endoscopy, we recommend that other countries considering implementation of screening colonoscopy should also consider implementation of outcome monitoring of endoscope reprocessing.

Conflict of Interests

The authors declare that they have no conflict of interests.

Authors’ Contribution

E. Fröhlich: study concept; acquisition of data, O. Leiß: critical revision of the paper, and R. Muller: statistical analysis, interpretation of data, and drafting the paper.

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