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| **ITU-T Focus Group Deliverable** | |
| **(09/2023)** | |
|  | Focus Group on Artificial Intelligence for Health  (FG-AI4H) | |
|  | **DEL10.20 – FG-AI4H Topic Description Document for the Topic Group on AI for Endoscopy (TG‑Endoscopy)** | |
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| ITU-T FG-AI4H Deliverable  DEL10.20 – FG-AI4H Topic Description Document for the Topic Group on AI for Endoscopy (TG-Endoscopy) |
| Summary  Endoscopy is the core technical means for early diagnosis and screening of digestive cancer, while AI solutions for endoscopy are expected to help clinicians improve the quality of their examinations and reduce the number of missed diagnoses. This document describes the application of artificial intelligence (AI) in endoscopic procedures, specifically focusing on two subtopics: colonoscopy and endoscopic ultrasound (EUS). In addition to a general description of AI for endoscopy, this document defines a framework for standardized benchmarking of AI systems designed to improve the early diagnosis and screening of digestive cancers. The document details AI tasks, existing solutions, data annotation processes, benchmarking considerations, and regulatory aspects within these medical domains. |
| Keywords  Artificial intelligence; health; topic groups; overview; ethics; regulations; data quality; data audit; clinical relevance; topic description; endoscopy; colonoscopy; endoscopic ultrasound; data annotation; benchmarking; gold standard. |
| Note  This is an informative ITU-T publication. Mandatory provisions, such as those found in ITU-T Recommendations, are outside the scope of this publication. This publication should only be referenced bibliographically in ITU-T Recommendations. |
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| --- | --- | --- |
| Editors: | Jianrong Wu Tencent Healthcare (Shenzhen), China | E-mail: [edwinjrwu@tencent.com](mailto:edwinjrwu@tencent.com) |
|  | Shan Xu CAICT, China | E-mail: [xushan@caict.ac.cn](mailto:xushan@caict.ac.cn) |
| Contributors: | Yajun Zhang  Tencent Technology (Shenzhen), China | E-mail: [yajunzhang@tencent.com](mailto:yajunzhang@tencent.com) |
|  | Yi Cai  Olympus Medical Systems Corp, Japan | E-mail: [i.sai@olympus.com](mailto:i.sai@olympus.com) |
|  | Junbo Li Suzhou Institute of Biomedical Engineering and Technology, Chinese Academy of Sciences, China | E-mail: [lijb@sibet.ac.cn](mailto:Lijb@sibet.ac.cn) |
|  | Yanchun Zhu China Unicom (Guangdong) Industrial Internet, China | E-mail: [zhuyc82@chinaunicom.cn](mailto:zhuyc82@chinaunicom.cn) |

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**ITU-T FG-AI4H Deliverable**

**DEL10.20 – FG-AI4H Topic Description Document for the Topic Group on AI   
for Endoscopy (TG-Endoscopy)**

# 1 Introduction

Endoscopy is the core technical means for early diagnosis and screening of digestive cancer. Implementing endoscopic screening for digestive cancer can detect and treat precancerous lesions, which can drastically reduce the incidence and mortality of digestive cancer. Due to factors such as the endoscopic doctor operating, the doctor’s ability to identify lesions, and visual fatigue, a considerable proportion of lesions in clinical diagnosis, including even advanced and precancerous lesions, may be missed by the endoscopic doctor.

In recent years, with the breakthrough of the new generation of artificial intelligence (AI) technology represented by deep learning, revolutionary progress has been made in the field of automatic recognition of medical images. The real-time assistance of AI for detecting and classifying gastrointestinal (GI) lesions is expected to help clinicians improve their examination quality and reduce the number of missed diagnoses.

This topic description document (TDD) specifies the standardized benchmarking for AI for endoscopy systems. It serves as deliverable DEL 10.20 of the ITU/WHO Focus Group on AI for Health (FG‑AI4H).

# 2 About the FG-AI4H topic group on AI for endoscopy (TG-Endoscopy)

The introduction highlights the potential of a standardized benchmarking of AI systems for AI for endoscopy to help solve important health issues and provide decision-makers with the necessary insight to successfully address challenges.

To develop this benchmarking framework, FG-AI4H decided, at meeting I held online on 7 and 8 May 2020, to create TG-Endoscopy.

FG-AI4H assigns a topic driver to each topic group (similar to a moderator) who coordinates the collaboration of all topic group members on the TDD. During FG-AI4H meeting I, e-meeting, 7‑8 May 2020, Dr. Jianrong Wu from Tencent Healthcare was nominated as topic driver for TG‑Endoscopy.

## 2.1 Documentation

This document is the TDD for TG-Endoscopy. It introduces the health topic, including the AI tasks, outlines its relevance and the potential impact that the benchmarking will have on the health system and patient outcome, and provides an overview of the existing AI solutions for endoscopy. It describes the existing approaches for assessing the quality of AI for endoscopy systems and provides the details that are likely relevant for setting up new standardized benchmarking. It specifies the actual benchmarking methods for all subtopics at a level of detail that includes technological and operational implementation. There are individual subsections for all versions of the benchmarking. Finally, the TDD summarizes the results of the topic group's benchmarking initiative and benchmarking runs, and also addresses ethical and regulatory aspects.

The TDD will be developed cooperatively over time by all members of the topic group and updated TDD iterations are expected to be presented at each FG-AI4H meeting.

The final version of this TDD will be released as deliverable "DEL 10.20 AI for Endoscopy (TG‑Endoscopy)". The topic group is expected to submit, at each FG-AI4H meeting, input documents (see Table 1) reflecting updates to the work on this deliverable.

Table 1 – Topic group output documents

| Number | Title |
| --- | --- |
| FGAI4H-S-025-A01 | Latest update of the Topic Description Document of the TG-Endoscopy |
| FGAI4H-J-025-A02 | Latest update of the Call for Topic Group Participation (CfTGP) |
| N/A | The presentation summarizing the latest update of the Topic Description Document of the TG-Endoscopy |

## 2.2 Status of this topic group

The following subsections describe the updates on the collaboration within TG-Endoscopy for the official focus group meetings.

### 2.2.1 Status update for meeting S

• Rearranged the TDD following FG-AI4H-J-105

• Updated the TDD for AI for endoscopy

### figure2.2.2 Status update for meeting P

• Updated the TDD for AI for endoscopy

### 2.2.3 Status update for meeting N

• Updated the TDD for AI for endoscopy

• Modified the structure of sections

• Invited new participants

### 2.2.4 Status update for meeting M

• Updated the TDD for AI for endoscopy

• Added new subtopic as endoscopic ultrasound

• Invited new participants

### 2.2.5 Status update for meeting L

• Converted the initial document of TG-Endoscopy into the TDD template format

• Invited new participants

### 2.2.6 Status update for meeting K

• Updated the initial AI for endoscopy documents

• Invited new participants

### 2.2.7 Status update for meeting J

• Started the draft of the TDD

• Stared the draft of the call for participation

• Presented the initial documents on AI for endoscopy (TG-Endoscopy)

### 2.2.8 Status update for meeting I

• Discussed the proposal from Tencent Healthcare

• Approved AI for endoscopy as a use case for FG-AI4H

• Established the topic group at meeting I (online, 7-8 May 2020)

• Nominated the topic group driver

## 2.3 Topic group participation

The participation in both FG-AI4H and in a topic group is generally open to anyone (with a free ITU account). For this topic group, the corresponding CfTGP can be found here:

• <https://www.itu.int/en/ITU-T/focusgroups/ai4h/Documents/tg/CfP-TG-Endoscopy.pdf>

Each topic group also has a corresponding subpage on the ITU collaboration site. The subpage for this topic group can be found here:

• <https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/tg/SitePages/TG-Endoscopy.aspx>

For participation in this topic group, interested parties can also join the regular online meetings. For all topic groups, the link will be the standard ITU-TG zoom link:

• <https://itu.zoom.us/my/fgai4h>

All relevant administrative information about FG-AI4H, such as upcoming meetings or document deadlines, will be announced via the general FG-AI4H mailing list [fgai4h@lists.itu.int](mailto:fgai4h@lists.itu.int).

All topic group members should subscribe to this mailing list as part of the registration process for their ITU user account, by following the instructions in the CfTGP at this link:

• <https://itu.int/go/fgai4h/join>

Regular FG-AI4H workshops and meetings proceed about every two months at changing locations around the globe or remotely. More information can be found on the official FG-AI4H website:

• <https://itu.int/go/fgai4h>

# 3 Topic description

This clause contains a detailed description and background information of the specific health topic for the benchmarking of AI in AI for endoscopy and how this can help to solve a relevant “real-world” problem.

Topic groups summarize related benchmarking AI subjects to reduce redundancy, leverage synergies, and streamline FG-AI4H meetings. However, in some cases different subtopic groups can be established within one topic group to pursue different topic-specific fields of expertise. TG‑Endoscopy has two subtopics, namely, colonoscopy and endoscopic ultrasound (EUS), discussed in the following subsections.

## 3.1 Subtopic: colonoscopy

### 3.1.1 Definition of the AI task

This clause provides a detailed description of the specific task that the AI systems of this topic group are expected to solve. This clause corresponds to [DEL03](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/_layouts/15/WopiFrame.aspx?sourcedoc=%7B7997F2C1-5A1D-4409-B2A0-CBC4E9CE8CDA%7D&file=DEL03.docx&action=default) "*AI requirements specifications*", which describes the functional, behavioural, and operational aspects of an AI system.

The application of AI in the field of colonoscopy varies according to different clinical goals. In general, it is mainly divided into the following three categories: classification, detection, and segmentation.

#### 3.1.1.1 Classification

Classification is a machine learning (ML) task for determining which classes are in an image, video or other types of data. It refers to training ML models with the intent of finding out which classes are present.

In clinical applications, it is possible to classify colorectal polyps in endoscopic images from a patient into different categories, such as non-adenomatous polyps, adenomatous polyps and cancerous. Different categories would need specific treatments. With the help of the classification results, clinicians can make more accurate diagnoses. It is also possible to evaluate the image quality of all the endoscopic images of the patient, like categorization of bowel cleanliness, for which the quality of the image should meet the diagnostic quality requirements.

#### 3.1.1.2 Detection

Object detection combines classification and localization to determine what objects are in the image or video and specify where they are in the image. Generally, bounding boxes are used to distinguish objects in video frames or images.

In clinical applications, it is possible to detect different findings in colonoscopy for different purposes, such as polyps, angiectasia, bleeding, inflammations, esophagitis, ulcerative colitis, pylorus, cecum, dyed polyp, dyed resection margins and stool. Specifically, polyp detection is the most usual AI application in colonoscopy. Detection for polyps can effectively reduce the polyp miss rate in colorectal screening, which would further reduce the adenomatous miss rate.

#### 3.1.1.3 Segmentation

Image segmentation separates an image into regions at the pixel level, with a particular shape and border, delineating potentially meaningful areas for further processing, such as measurement, classification and object detection. The regions may not take up the entire image, but the goal of segmentation is to highlight foreground elements and make it easier to be evaluated. Image segmentation provides pixel-by-pixel details of an object, distinguishing it from classification and object detection.

For example, in endoscopy, the polyp size can be automatically calculated based on the segmentation results, while the polyp size is one of the key factors for polyp diagnosis by clinicians.

### 3.1.2 Current gold standard

This clause provides a description of the established gold standard for the addressed health topic.

Colorectal cancers (CRCs) are the third most prevalent cancer and the second-highest cause of cancer deaths worldwide. Colonoscopy is considered the gold standard for CRC screening to detect and remove the polyps and adenomas in the colorectum [1]. In clinical practice, colonoscopy requires both manipulation and observation at the same time, and it cannot detect all colonic polyps, some of which may be neoplasms. Colonoscopy has been reported to miss 17% to 48% of adenomas which are considered to be the cause of 50% to 60% of interval cancers.

Over the last two decades, computer-assisted polyp detection has been actively explored to improve inspecting quality and reduce adenoma miss rates (AMR). Recently, AI has made remarkable breakthroughs in medical fields with deep learning and convolutional neural networks (CNNs). With enough qualified learning materials, CNNs can reach even higher real-time detecting accuracy than human experts, which demonstrates that computer-aided detection (CADe) systems might have the potential to serve as real-time “experts” to improve the quality of colonoscopies.

To build an AI solution for colonoscopy, such as CADe and computer-aided diagnosis (CADx) systems, it is necessary to have a gold standard and colonoscopy images that could be annotated by objective or subjective methods. By objective methods, the gold standard would be annotated by information from clinical diagnosis reports. For example, to train a CADx classifying the nature of polyps by image, the nature of polyps in pathology report could be used for the gold standard [2][3][4]. By subjective methods, the gold standard would be made by colonoscopists manually [5]. Usually, subjective methods are used for polyp detection [6][7] and segmentation tasks [8][9][10], whose annotation results are bounding box and mask, respectively. The subjective method might involve one or multiple colonoscopists in single-step or multi-step procedures of annotation.

### 3.1.3 Relevance and impact of an AI solution

This clause addresses the relevance and impact of the AI solution and describes how solving the task with AI improves a health issue.

The significance of screening colonoscopy lies largely in the detection and removal of colorectal polyps. In clinical practice, colonoscopy is a highly operator-dependent procedure and requires both manipulation and observation at the same time, which may lead to significant variation of adenoma miss rates between individual endoscopists [11]. CADe with real-time automatic polyp detection or classification powered by AI algorithms has been proposed to help endoscopists improve the polyp detection rate (PDR) and adenoma miss rate [12].

A prospective study including 1058 patients was designed as a randomized controlled trial (RCT) to investigate the impact of an automatic polyp detection CADe acting as an assistant to the endoscopist on PDR and adenoma detection rate (ADR), in Endoscopy Center of the Sichuan Provincial People's Hospital, China. The colonoscopy with CADe showed increased ADR (29.1% vs 20.3%, p < 0.001), the mean number of adenomas per patient (0.53 vs 0.31, p < 0.001) and hyperplastic polyps (114 vs 52, p < 0.001) [13].

Over 71000 images from 20 centres were used to train and test a deep learning-based CADe in Changhai Hospital in Shanghai, China. The CADe was able to identify polyps in the test dataset with 95.0% sensitivity and 99.1% specificity. Colonoscopists can detect more polyps (0.90 vs 0.82, P < 0.001) and adenomas (0.32 vs 0.30, P = 0.045) with the aid of CADe, particularly polyps < 5 mm and flat polyps (0.65 vs 0.57, P < 0.001; 0.74 vs 0.67, P = 0.001, respectively) [14].

Besides detection, there has been an attempt at classification. A deep convolutional neural network model was trained to predict the histology of polyps using only narrow band imaging. The accuracy of the model was 94% (95% CI 86% to 97%); the sensitivity for identification of adenomas was 98% (95% CI 92% to 100%); specificity was 83% (95% CI 67% to 93%); negative predictive value 97%; and positive predictive value 90% [2].

### 3.1.4 Existing AI solutions

This clause provides an overview of existing AI solutions for colonoscopy that are already in operation. AI solutions for colonoscopy are moving towards commercialization and clinical practice, while there are more and more CADe products approved by the Chinese National Medical Products Administration (NMPA), the U.S. Food and Drug Administration (FDA) and European CE.

• Tencent Healthcare and Changhai Hospital developed a CADe system in 2021 that was built based on the "You Only Look Once" v2 deep learning framework [15]. The system detects potential polyps and presents an alert rectangle surrounding polyps on a second monitor for colonoscopists. Colonoscopists detect more polyps and adenomas with the aid of CADe systems, particularly polyps<5 mm and flat polyps[14]. The real-time polyp detection system by Tencent Healthcare was certificated by the NMPA in June 2023.

• In 2017, National Cancer Center Hospital and NEC Japan successfully developed a system that immediately detects colorectal cancer and ulcerative colon polyps, a precursor to cancer, during an endoscopic examination using AI. It automatically detects colorectal cancer and polyps from images and videos taken during an endoscopic examination of the colon and aids in the discovery of lesions by endoscopists. It improves polyp detection, which was an issue during such exams, and increases the detection rate. In this way, it contributes greatly to the prevention and early detection of colorectal cancer [16].

• Wision AI and Sichuan Provincial People's Hospital developed a real-time automatic polyp detection system in 2018 [8] that detects colorectal polyps during an endoscopic examination using deep learning. The detection algorithm is a deep CNN based on SegNet architecture. If any polyp is detected by the system, a hollow tracking box around it would be shown on the monitor. As a conclusion, in a low-prevalence ADR population, an automatic polyp detection system during colonoscopy resulted in a significant increase in the number of diminutive adenomas detected, as well as an increase in the rate of hyperplastic polyps. The cost-benefit ratio of such effects has to be determined further [13][17]. The product for real-time colorectal polyps' detection, named EndoScreener, has been certificated by the NMPA, the FDA and European CE-MDR (Medical Device Regulation).

• In 2018, National Chiao Tung University and Tri-Service General Hospital developed a CADx with a deep neural network to analyze narrow-band images of diminutive colorectal polyps. The system could classify the polyps in narrow-band images as neoplastic or hyperplastic [18].

• In 2018, Sun Yat-sen University developed a CADe system called Gastrointestinal Artificial Intelligence Diagnostic System (GRAIDS) with deep learning to detect upper GI cancers by endoscopy). It is the first real-time AI-aided image recognition system that has been implemented in clinical practice for detecting upper GI cancers during endoscopy [19].

• Zhongshan Hospital and University of California developed an AI-based CNN-CAD system through transfer learning, leveraging a state-of-the-art pretrained CNN architecture, ResNet50. The system is used to determine the invasion depth of the gastric cancer and screen patients for endoscopic resection. This system distinguished early gastric cancer from deeper submucosal invasion and minimized overestimation of invasion depth, which could reduce unnecessary gastrectomy [20].

• Cancer Institute Hospital Ariake, AI Medical Service and Tada Tomohiro Institute of Gastroenterology and Proctology developed a CNN-based diagnostic system based on Single Shot MultiBox Detector architecture to detect gastric cancer in endoscopic images. This constructed CNN system for detecting gastric cancer can process numerous stored endoscopic images in a very short time with a clinically relevant diagnostic ability [21].

• Renmin Hospital of Wuhan University developed a system using a novel deep convolution neural network (DCNN) to detect early gastric cancer (EGC) without blind spots during esophagogastroduodenoscopy (EGD). This system could identify EGC from non-malignancy and classify gastric location into 10 or 26 parts with high accuracy [22].

• Kindai University developed a system in 2017 that could diagnose colon polyps as adenomatous or non-adenomatous using a simple CNN [23].

• Wuhan ENDOANGEL Medical Technology Co., LTD developed an AI system called EndoAngel, consisting of polyp detection and quality monitoring functions. The polyp detection function can provide a reference to endoscopists regarding the location of the polyp. The quality monitoring function can monitor the velocity of insertion of the endoscope, record the time of insertion and withdrawal of the endoscope, and assist endoscopists with any blind areas caused by intestinal segment slipping [24]. It was certificated by NMPA in May 2023.

## 3.2 Subtopic: endoscopic ultrasound

### 3.2.1 Definition of the AI task

Endoscopic ultrasound (EUS) is a minimally invasive procedure in which endoscopy is combined with ultrasound to obtain images of the internal organs. Compared to colonoscopy, EUS is a multi-modality procedure capturing ultrasound and image at the same time. In general, the AI task with EUS is mainly divided into classification, detection, and segmentation.

#### 3.2.1.1 Classification

Classification is an ML task for determining which classes are in an image, video, or other types of data. It refers to training ML models with the intent of finding out which classes are present.

As EUS is frequently used in the assessment of digestive disease, the clinical applications of AI for EUS largely involve their use in classifying suspicious lesions in the upper and lower digestive tract and surrounding tissues from endoscopic images and ultrasound data (RF, B-mode, colour, flow, contrast enhanced ultrasound, elastography, etc.) In addition, AIassisted EUS could be used for respiratory and urinary systems to distinguish malignant from benign lesions. With the classification result, clinicians could make more accurate diagnoses, reduce unnecessary EUS-guided biopsies and provide more suitable treatment. It is also possible to evaluate the quality of EUS images of the patient, like station classification and quality assessment for pancreatis EUS scans.

#### 3.2.1.2 Detection

Object detection combines classification and localization to determine what objects are in the image or video and specify where they are in the image. Generally, bounding boxes are used to distinguish objects in video frames or images.

Unlike conventional endoscopy, where AI-assisted detection is possibly used to avoid missing blind spots during the procedure, EUS can hardly be used as the first-line screening choice for digestive or respiratory tract due to its limited image quality for endoscopic imaging. Instead, it can be used to identify issues beneath or surrounding the digestive and respiratory tract in EUS, such as lymph nodes, bleeding and inflammation.

#### 3.2.1.3 Segmentation

Image segmentation separates an image into regions at the pixel level, with a particular shape and border, delineating potentially meaningful areas for further processing, such as measurement, classification and object detection. The regions may not take up the entire image, but the goal of segmentation is to highlight foreground elements and make them easier to be evaluated. Image segmentation provides pixel-by-pixel details of an object, distinguishing it from classification and object detection.

In EUS, the size and extension of the suspicious lesion can be automatically calculated based on the segmentation results, which can help clinicians provide more suitable treatment.

### 3.2.2 Current gold standard

Clinical evidence has shown the benefits of EUS over potential adverse events (AEs) and clinical guidelines have been published and continuously updated to ensure the safe use of the procedures [25][26]. EUS has emerged as an important imaging modality for the diagnosing and staging of benign and malignant lesions in the upper digestive tract and the respiratory system, and it is most commonly used for the staging of GI malignancies, evaluating pancreaticobiliary disease, evaluating subepithelial abnormalities, evaluating extraluminal abnormalities, the staging of lung cancer, and image guidance for therapeutic procedures [27]. The European Society of Gastrointestinal Endoscopy (ESGE) has suggested EUS for pancreatic cancer screening in selected high-risk patients, recommended EUS-guided sampling for pancreatic solid lesions as first line procedure and EUS-guided sampling for biochemical analysis plus cytopathologic examination for pancreatic cystic lesions, etc., and recommended EUS as therapeutic procedures for various types of diseases, including percutaneous transhepatic biliary drainage (PTBD), and pancreatic duct (PD) drainage. Specifically, EUS is capable of identifying small pancreatic tumours with a staging sensitivity greater than 90% [28], and endobronchial ultrasound (EBUS) has been used for lung cancer staging with a diagnostic accuracy of 90% to 100% [29].

Research on AI in EUS is still limited [30][31][32][33][34]. Only a handful of reports were published based on limited clinical data through retrospective or prospective studies, with a main focus on pancreatic diseases. Currently, there is no commercial AI product for EUS on the market.

### 3.2.3 Relevance and impact of an AI solution

EUS has been proven to be an effective imaging modality for local and regional staging of GI tumours. The diagnostic ability of EUS is higher than that of computed tomography (CT), transabdominal ultrasonography, and magnetic resonance imaging (MRI) [35][36]. It has also proved to be a useful alternative therapeutic modality in surgery. However, EUS may be less accurate for early staging of oesophageal cancer. According to a meta-analysis by Puli et al., the diagnostic accuracy of EUS was higher for T3-T4 lesions (>90%) than T1-T2 (65%) [36]. Using EUS has also shown low accuracy in differentiating benign and malignant rectal cancer after treatment. Another limitation for EUS (as well as other ultrasonography procedures) is its operator dependency. The performance of EUS improves with experience. High inter-observer variability (61% to 77%) has been reported and a wide range of overall accuracy for tumour staging could be found among different studies (63% to 95%).

AI is believed to play an important role in endoscopic procedures, not only for detecting anatomical features, differentiating benign and malignant lesions, and delineating lesion contours but, more importantly, for reducing learning time for junior endoscopists, decreasing workload and standardizing the overall quality of endoscopic procedures.

### 3.2.4 Existing AI solutions

This clause provides an overview of existing AI solutions for EUS that are already in operation. It should be noted that currently there is no well-accepted AI solution for EUS. The solutions listed below are mainly premature prototype systems or even models from academic or industrial research.

• Researchers from Pusan National University Hospital, Silla University, Asan Medical Center and Yonsei University College of Medicine developed a CNN-CAD system to analyse gastric mesenchymal tumours on EUS images. The CNN-CAD system can differentiate gastrointestinal tumours (GISTs) from non-GIST tumours within a short amount of time and with high sensitivity and specificity. However, the datasets used in the study were relatively small and only high-quality EUS images were selected for the training and test datasets [37].

• Researchers from Changhai Hospital reported a single-centre retrospective study in 2010. Support vector machine (SVM)-based classification was implemented to differentiate pancreatic cancer from normal tissue, with high accuracy, sensitivity, and specificity [38]. A further study was reported in 2013 with more data from Changhai Hospital [39]. Results show the superiority of SVM-based CAD systems for pancreas EUS. However, a substantial decrease in classification performance can be found between the two studies, using data from the same clinical site and very similar technology.

• Tokyo Medical University developed a CNN-based EUS-CAD system and assessed its ability to detect pancreatic ductal carcinoma (PDAC), using control images from patients with chronic pancreatitis (CP) and those with a normal pancreas (NP). Results indicate EUS-CAD systems can work, not only in assisting the training of beginners of EUS instead of an instructor but also in supporting fatigued experts or reducing carelessness caused by performing a large number of screening examinations [40].

• The European EUS Elastography Multicentric Study Group performed a prospective multicentric study in 2012 and develop an artificial neural network (ANN)-based CAD to differentiate benign from malignant pancreatic lesions using real-time EUS elastography [41]. In 2015, another prospective multicentric study was conducted to access ANN-based CAD, to classify pancreatic cancer using dynamic contrast-enhanced EUS [42]. Results from two studies suggest that integration of clinical data into efficacious ANNs, in concordance with imaging enhancements (real-time sono-elastography, contrast-enhancement, hybrid imaging, 3‑dimensional imaging, and so forth) and cytologic parameters, would certainly be beneficial for improved clinical decision-making in patients with focal pancreatic lesions.

• Wuhan EndoAngel Medical Technology Company, Renmin Hospital of Wuhan University, Wuhan Union Hospital and Wuhan Puai Hospital developed a pancreaticobiliary master (BP MASTER) system for training in, and quality control of, pancreatis EUS scans. Results show the BP MASTER system has potential to play an important role in shortening the pancreatic EUS learning curve and improving EUS quality control in the future. [43]

• China Medical University Hospital and National Taiwan University developed a CNN-based CAD to classify lung lesions, using EBUS images. The results showed that the fusion of the fine-tuned CaffeNet and SVM system have the potential to assist lung cancer detection [44].

• Shimane University of Japan developed an EBUS-computer-aided diagnosis system using CNN to differentiate benign from malignant lesions based on EBUS findings. The developed EBUS-computer-aided diagnosis system is capable of reading EBUS findings that are difficult for clinicians to judge with precision, and helps differentiate between benign lesions and lung cancers [45].

• NeuralSeg Ltd, St. Joseph's Healthcare Hamilton and McMaster University performed a clinical study to access an AI algorithm (NeuralSeg) for identifying and predicting lymph node malignancy based on EBUS images. Results suggest that NeuralSeg is able to accurately rule out nodal metastasis and can possibly be used as an adjunct to EBUS when nodal biopsy is not possible or is inconclusive [46].

• Olympus, Chiba University and Dokkyo Medical University developed CNN-based CAD for the detection and classification of nodal metastasis from EBUS images. The prediction of LN metastasis by CAD using EBUS images showed high diagnostic accuracy with high specificity. CAD during transbronchial needle aspiration endobronchial ultrasound (EBUS-TBNA) may help improve the diagnostic efficiency and reduce the invasiveness of the procedure [47].

# 4 Ethical considerations

The rapidly evolving field of AI and digital technology in the fields of medicine and public health raises a number of ethical, legal, and social concerns that have to be considered in this context. These are discussed in deliverable DEL01 "*AI4H ethics considerations*", which was developed by the working group on "Ethical considerations on AI4H" (WG-Ethics). This clause refers to DEL01 and should reflect the ethical considerations of TG-Endoscopy.

Collecting massive data is necessary for the development of AI solutions. However, ethical considerations, such as patient privacy concerns, should be taken into careful consideration and relevant regulations should be followed. The privacy of patients must be protected in the process of data collection, transmission, and use. If the data contains patients’ private information or identified codes, data desensitization must be performed. Generally, it is better for data sources, such as hospitals and other clinical institutions, to be responsible for handling the ethical, legal and privacy aspects of the relevant data.

The following procedures are executed in our practice and recommended to other practices of AI for endoscopy.

• Patients consent procedure at each individual institution.

• Review of the data collection plan by a local medical ethics committee or an institutional review board.

• Anonymization of the video or image frames (including demographic information) by clinical institution prior to sending to AI developer.

• Anonymization of the video or image frames (including demographic information) by AI developer prior to use (optional).

# 5 Existing work on benchmarking

This clause focuses on the existing benchmarking processes in the context of AI and AI for endoscopy for quality assessment. The goal is to collect all relevant learnings from previous benchmarking that could help to implement the benchmarking process in this topic group.

## 5.1 Subtopic: colonoscopy

### 5.1.1 Publications on benchmarking systems

Some work has been done in the scientific community assessing the performance of AI for endoscopy. This clause summarizes insights from the most relevant publications on this topic. It covers parts of the deliverable DEL7 "*AI for health evaluation considerations*",DEL7.1 "*AI4H evaluation process description*", DEL7.2"*AI technical test specification*", DEL7.3 "*Data and artificial intelligence assessment methods (DAISAM)*", and DEL7.4 "*Clinical Evaluation of AI for health*".

#### 5.1.1.1 EndoCV

The International Workshop and Challenge on Computer Vision in Endoscopy (EndoCV) was held from 2019 to 2022 annually [48][49][50][51][52][53], aimed at benchmarking methods on larger test sets, comprising mostly video sequences as in the real-world clinical scenario, for endoscopy artefact detection, endoscopy disease detection and polyp generalization [54][55][56][57]. The latest 4th International Workshop and Challenge on Computer Vision in Endoscopy (EndoCV2022) was held in conjunction with IEEE International Symposium on Biomedical Imaging (ISBI2022).

#### 5.1.1.2 EndoVis

Endoscopic Vision Challenge (EndoVis) [58] organizes high-profile international challenges for the comparative benchmarking and validation of endoscopic vision algorithms that focus on different problems each year at International Conference on Medical Image Computing and Computer Assisted Intervention (MICCAI), from 2015 until now, except for 2016, while there were several sub-challenges in each year.

##### 5.1.1.2.1 GIANA

Gastrointestinal Image ANAlysis (GIANA) was one of the sub-challenges in EndoVis, which was held in 2017, 2018 [59] and 2021 [60].

##### 5.1.1.2.2 CATARACTS

In 2017, the Challenge on Automatic Tool Annotation for cataRACT Surgery (CATARACTS) [61] released 50 cataract surgery videos accompanied by instrument usage annotations including frame-level instrument presence information.

##### 5.1.1.2.3 Detection of Abnormalities in Gastroscopic Images (DAGI)

The challenge of detection of abnormalities in gastroscopic images (DAGI) [62] was one of the sub-challenges in EndoVis 2015, focusing on comparing different abnormal detection methods for recognizing abnormal regions from GI images. The abnormal detection for GI images was addressed with different abnormal patterns, such as gastritis, cancer, ulcers and bleeding.

##### 5.1.1.2.4 Automatic polyp detection in colonoscopy videos (APDCV)

The challenge of automatic polyp detection in colonoscopy videos (APDCV) [63] was one of the sub-challenges in EndoVis 2015 and the first challenge about polyp detection. This challenge was to automatically detect polyps in colonoscopy videos, thereby reducing polyp miss rates and the subsequent mortality rate from colon cancer.

#### 5.1.1.3 EndoTect

The EndoTect challenge [64] at the International Conference on Pattern Recognition (ICPR) 2020 aimed to motivate the development of algorithms that aid medical experts in finding anomalies that commonly occur in the GI tract [65].

### 5.1.2 Benchmarking by AI developers

All developers of AI solutions for endoscopy implemented internal benchmarking systems for assessing the performance. This clause outlines the insights and learnings from this work that are of relevance for benchmarking in this topic group.

#### 5.1.2.1 EndoCV

To achieve high diversity, the dataset of EndoCV was built with data from multiple centres in different countries that include Egypt, France, Italy, Norway, Sweden and UK. The resolution of data included standard definition (SD), high definition (HD) and ultra HD. The data was collected by different endoscopy manufacturers, includes Olympus (mostly), Fujifilm and Karl Storz.

There were two sub-challenges in EndoCV2022: endoscopy artefact detection (EAD 2.0) and polyp generalization (PolypGen 2.0). The aim of the sub-challenge EAD 2.0 was to localize bounding boxes, and predict class labels and pixel-wise segmentation of eight artefact classes for given clinical endoscopy video clips, namely, specularity, bubbles, saturation, contrast, blood, instrument, blur and imaging artefacts. PolypGen 2.0 aimed to benchmark methods on the basis of generalization capabilities to unseen colonoscopy video sequence data for both detection and segmentation deep learning methods.

• **AI task**

o **Detection task**: The aim of this task was to test the performance of participants' methods for the detection and localization tasks on comprehensive and sorted multicentre datasets. The participants were tested on both detection-based metric and localization metric. A weighted final metric was used to evaluate the best-performing method.

o **Segmentation task**: Each participant’s method was evaluated on multicentre curated and sorted datasets.

• **Dataset**

o **EAD 2.0**: A total of 280 patient videos from multiple organs and institutions, with 45,478 annotations on both single frame and sequence video data. Training data for the detection task consisted of a total of 2531 frames with 31,069 bounding boxes and 643 frames with 7511 binary masks for the segmentation task. Additionally, there was a new set of test data were curated that included unique video sequences consisting of more than 500 frames.

o **PolypGen 2.0**: This dataset was composed of a total of 6282 frames, including both single and sequence frames from six centres, incorporating more than 300 patients. It consisted of 3762 positive sample frames and 2520 negative sample frames with 3446 annotated polyp labels with precise delineation of polyp boundaries (pixel level for segmentation task and bounding boxes for detection task) verified by six senior gastroenterologists.

o In addition to this dataset, an additional 23 unique patient video clips (> 100 frames per video), were collected, making a total of 46 sequences for PoypGen2.0 and 24 sequences for EAD2.0.

• **Annotation**

o First, a small subset of the dataset was annotated by all clinical experts and a joint consensus was made available.

o Then, the remaining subset of the dataset was annotated by post-doctoral researchers (working on endoscopy) and validated by clinicians at two different centres (10-fold cross-validation).

o Finally, through a joint conference call, all annotation validation was achieved.

• **Metrics**

• **Detection** **task**

o Standard computer vision metric: mean average precision (mAP).

o Standard intersection over union (IoU).

o Final detection score (trade-off between mAP and IoU): 0.6\*mAP + 0.4\*IoU.

o Generalization gap (Gerror): defined as the difference between the detection score and the generalization score (on unseen data).

o Centroid localisation error (Lerror): defined as the distance between centroids’ positions of detected boxes between the consecutive frames in a video (new).

o Clinical applicability metrics: runtime (to be used post challenge only).

• **Segmentation** **task**

o Standard segmentation metrics that include Dice coefficient (DICE), F2-error, positive predictive value (PPV), Hausdorff distance (HD) and sensitivity (recall) were used.

o The ranking on leaderboard was based on the highest mean value between Dice similarity coefficient (DSC), PPV and sensitivity, and the least HD value.

o Generalizability difference (Gerror): the difference between DSC on mixed sample data and DSC on unseen data will be key in deciding the winner of this task.

o Clinical applicability metrics: runtime (to be used post challenge only).

#### 5.1.2.2 EndoVis

##### 5.1.2.2.1 GIANA

In general, GIANA includes polyp detection, segmentation and classification in colonoscopy images, and polyp segmentation, angiodysplasia detection and localization in wireless capsule images. See Table 2.

| Table 2 – GIANA | | | | | | |
| --- | --- | --- | --- | --- | --- | --- |
| Year | Task | Modality | Definition | Clinical use | Database content | Ground truth |
| GIANA 2018 | Polyp detection | Colonoscopy | Ability to detect presence/absence of polyps in each frame and, in case of polyp presence, locate it within the image | Prevention of colorectal cancer | 18 short videos for training, more than 20 short and long videos for testing | Polyp masks, Paris classification  Clinical partner: Hospital Clinic, Barcelona, Spain |
| Polyp segmentation | Colonoscopy | Delimit the region the polyp occupies in the image | Preliminary stage for lesion classification through analysis of polyp region content | Two sets: SD images (300 images for training, 612 for testing) and High-Definition images (more than 150 images) | Polyp masks, Paris classification Clinical partner: Hospital Clinic, Barcelona, Spain |
| Angiodysplasia detection | Wireless capsule endoscopy | Label each of the frames into angiodysplasia containing or not | Automatic detection of small bowel lesions related to bleeding | 600 images for training (same number of positive and negative examples) and 600 images for testing. | Angiodysplasia mask Clinical partner: Saint Antoine Hospital, Paris, France |
| Angiodysplasia localization | Wireless capsule endoscopy | Label each of the frames into angiodysplasia containing or not and, in case angiodysplasia is detected, localize the region it occupies in the image | Automatic detection of small bowel lesions related to bleeding | 600 images for training (same number of positive and negative examples) and 600 images for testing. | Angiodysplasia mask Clinical partner: Saint Antoine Hospital, Paris, France |
| GIANA 2021 | Polyp detection | Colonoscopy | Ability to detect presence/absence of polyps in each frame and, in case of polyp presence, locate it within the image | Prevention of colorectal cancer | 18 short videos for training, more than 20 short and long videos for testing | Polyp masks, Paris classification  Clinical partner: Hospital Clinic, Barcelona, Spain |
| Polyp segmentation | Colonoscopy | Delimit the region the polyp occupies in the image | Preliminary stage for lesion classification through analysis of polyp region content | Two sets: S D images (300 images for training, 612 for testing) and HD images (more than 150 images) | Polyp masks, Paris classification Clinical partner: Hospital Clinic, Barcelona, Spain |
| Polyp classification (frames) | Colonoscopy | Label each of the frames with one of the following categories: 1) adenoma, 2) non-adenoma | In-vivo diagnosis, advance patient treatment | 1000 images for training and validation and 200 images for testing | Label of each frame, polyp region Clinical partner: Hospital Clinic, Barcelona, Spain |

##### 5.1.2.2.2 CATARACTS

Pixel-wise semantic annotations for anatomy and instruments of 36 classes for 4670 images sampled from 25 videos of the CATARACTS training set were released in 2020, including 4 anatomical classes, 29 instruments and 3 classes of other objects appearing in the scene. As one of the sub-challenges in EndoVis 2020, there were three sub-tasks to assess participating solutions on anatomical structure and instrument segmentation: 1) anatomy and instruments, 2) anatomy and grouped instruments, and 3) anatomy, instrument tips and handles [66].

Their performance was assessed on a hidden test set of 531 images from 10 videos of the CATARACTS test set. There were 25 classes in the test set, including 4 anatomical classes, 18 instruments and 3 other objects in the scene in particular. The mean intersection over union (mIoU) was used to assess model performance.

##### 5.1.2.2.3 DAGI

In total, 800 gastroscopic images from 137 volunteers were involved, while three senior experts were invited to annotate the lesion/abnormal regions independently, and the pixel-level ground truth was defined as the average.

For benchmark and evaluation, the area under curve (AUC) of the receiver operating characteristic (ROC) curve was used. The performance was based on the image-level predictions in particular. For positive images, an image can be considered as true positive if at least 40% of the truly abnormal pixels are detected; otherwise, it will be considered as false negative. For negative images, an image can be considered as true negative only when no abnormal pixel is detected; otherwise, it will be considered as false positive.

##### 5.1.2.2.4APDCV

There were two tasks, namely, frame classification of polyp existence and polyp detection in image, which were defined in the challenge as polyp detection and polyp localization, respectively.

Three public databases were used in the context of the benchmark: CVC-CLINIC [67], ETIS-LARIB and ASU-Mayo Clinic Colonoscopy Video Database [68].

• CVC-CLINIC contains 612 SD frames and comprises 31 different polyps from 31 sequences.

• ETIS-LARIB database contains 196 HD frames and comprises 44 different polyps from 34 sequences. Ground truth of each polyp consists of a polyp mask for both databases, which was generated by the annotated boundary of polyp by expert video endoscopists from the corresponding associated clinical institution. In the challenge, these two datasets were used for scenario of still frame analysis.

• The ASU-Mayo Clinic Colonoscopy Video Database comprises a set of short and long colonoscopy videos, collected at the Department of Gastroenterology at Mayo Clinic, Arizona. This database consists of 38 different, fully annotated videos. Ground truth consisting of binary masks (polyp frames) and black frames (non-polyp frames) were created by volunteer students at Arizona State University and have been reviewed and corrected by a trained expert. This dataset was used for scenario of video analysis in the challenge.

For benchmarking and evaluation, general metrics of precision, recall, specificity, F1-measure and F2-meature were used. An additional video analysis performance metric was introduced to assess how fast a polyp detection method was introduced. That was detection latency, representing the delay in frames between the first appearance of the polyp in the video sequence and the first actual detection of the polyp by a method.

#### 5.1.2.3 EndoTect

A large dataset named as HyperKvasir [69], containing images taken from several endoscopies, was used. In total, the dataset contains 110,079 images and 374 videos where it captures anatomical landmarks, pathological findings, and normal findings, while the dataset can be split into four distinct parts.

• **Labelled images**. In total, the dataset contains 10,662 labelled images stored using the JPEG format. The labeled images represent 23 different classes of findings.

• **Unlabelled images**. In total, the dataset contains 99,417 unlabelled images.

• **Segmentedimages**. The original image, a segmentation mask and a bounding box for 1,000 images from the polyp class were provided.

• **Annotated videos**. The dataset contains a total of 373 videos containing different findings and landmarks. Each video was manually assessed by a medical professional working in the field of gastroenterology and resulted in a total of 171 annotated findings.

There were three tasks.

• **Detection task**: classifying images from the GI tract into 23 distinct classes. Metrics of precision, recall/sensitivity, specificity, F1-measure and Matthews correlation coefficient (MCC) for multi-classification were used and the MCC was used for benchmarking to rank the submission.

• **Efficient detection task**: efficient classification measured by the amount of time spent processing each image. Metric of a combination of the MCC classification score and the number of frames processed per second was used for benchmarking to rank the submission.

• **Segmentation task**: automatically segmenting polyps. Metrics of precision, recall, DICE, and the IoU, also known as the Jaccard index, were used and the IoU was used for benchmarking to rank the submission.

### 5.1.3 Relevant existing benchmarking frameworks

Triggered by the hype around AI, recent years have seen the development of a variety of benchmarking platforms where AIs can compete for the best performance on a determined dataset (see Table 3).

**•** **EvalAI**: EvalAI is an open source platform for evaluating and comparing ML and AI algorithms at scale [70].

• **AIcrowd**: AIcrowd enables data science experts and enthusiasts to collaboratively solve real-world problems, through challenges [71].

•**Kaggle**: Kaggle offers a no-setup, customizable, Jupyter Notebooks environment. Access to graphics processing units (GPUs) at no cost and a huge repository of community published data and code [72].

• **CodaLab**: CodaLab is an open source platform to learn, create and collaborate through challenges [73].

• **Grand challenge**: A platform for end-to-end development of ML solutions in biomedical imaging.

Table 3 – Relevant existing benchmarking frameworks

| Benchmarking frameworks | Challenges | Users | Submissions | Organizations | Papers published | Awarded in prizes | Datasets/ stored data | Notebooks/ algorithms |
| --- | --- | --- | --- | --- | --- | --- | --- | --- |
| EvalAI | 200+ | 18k+ | 180k+ | 30+ |  |  |  |  |
| AIcrowd | 246+ | 59k+ |  |  | 60+ | 823+ USD | 13 TB+ |  |
| Kaggle |  | 13.6m+ |  |  |  |  | 50k+ | 400k+ |
| CodaLab | 1946 | 36k+ | 409k+ |  |  |  |  |  |
| Grand challenge | 356 | 82k+ | 88k+ |  |  |  |  | 27k+ |

There are several public datasets related to AI for endoscopy, as shown in Table 4.

| Table 4 – Available public dataset | | | |
| --- | --- | --- | --- |
| Dataset | Findings | Size | Availability |
| CVC-612 (CVC-ClinicDB) [67] | Polyp,  with mask | 612 images | Open academic |
| ASU-Mayo polyp database [9] | Polyp,  with mask | 18,781 images | By request  (not available anymore) |
| ETIS-Larib Polyp DB [74] | Polyp,  with mask | 196 images | Open academic |
| KID [75] | Angiectasia, bleeding, inflammations, polyp | 2371 images, 47 videos | Open academic  (not available anymore) |
| GIANA'17 [59] | Angiectasia, with mask | 600 images | By request |
| GASTROLAB [76] | GI lesions | Some 100s of images + few videos | Open academic  (video capsule endoscopy) |
| WEO Clinical Endoscopy Atlas [77] | GI lesions | 152 images | By request  (video capsule endoscopy) |
| GI Lesions in Regular Colonoscopy Data Set [78] | GI lesions, with mask | 76 images | By request |
| Atlas of Gastrointestinal Endoscope [79] | GI lesions | 1295 images | Unknown  (not available anymore) |
| GastroAtlas [80] | GI lesions | 5,071 video clips | Open academic  (video capsule endoscopy) |
| Kvasir [81] | Polyps, esophagitis, ulcerative colitis, Z-line, pylorus, cecum, dyed polyp, dyed resection margins, stool | 8,000 images | Open academic |
| Nerthus [82] | Stool - categorization of bowel cleanliness | 21 videos | Open academic |
| Kvasir-SEG [83] | Polyps, with mask | 1000 images | Open academic |
| HyperKvasir [69] | GI findings including polyps | 110,079 images and 374 videos | Open academic |
| Kvasir-Capsule [84] | GI findings including polyps (video capsule endoscopy) | 4,741,504 images | Open academic |
| CVC-ColonDB [85] | Polyps, with mask | 380 images | Open academic  (not available anymore) |
| EDD2020 [56] | GI lesions including polyps | 386 images | Open academic |

## 5.2 Subtopic endoscopic ultrasound (EUS)

### 5.2.1 Publications on benchmarking systems

Although research on AI for EUS application has increased rapidly in the last few years, publicly accessible datasets and benchmarking systems do not exist. Several review papers have been published to summarize latest research in the field, but none of those can provide comparable benchmarking for different studies [86][87][88][89][90]. It is extremely important for the scientific community to establish a public accessible EUS image database and benchmarking system to push forward AI-assisted EUS research.

### 5.2.2 Benchmarking by AI developers

All developers of AI solutions for EUS implemented internal benchmarking systems for assessing the performance. Depending on the tasks of the AI solutions (detection, classification, segmentation etc.), different metrics will be used in order to enable performance comparison. These metrics are not much different from those used in medical image analysis and computer vision, specifically colonoscopy, such as mAP, IoU, DICE, PPV, precision, recall, specificity, F1-measure, MCC and AUC of ROC curve.

### 5.2.3 Relevant existing benchmarking frameworks

Relevant existing benchmarking frameworks of AI EUS are the same as colonoscopy, including EvalAI, AIcrowd, Kaggle, CodaLab and Grand challenge. For more details, section 5.1.3 could be referred to. Currently, there seems to be no publicly available dataset for AI EUS.

# 6 Benchmarking by the topic group "For further study"

This clause describes all technical and operational details regarding the benchmarking process for the AI for endoscopy AI task, including subsections for each version of the benchmarking that is iteratively improved over time.

It reflects the considerations of various deliverables: DEL5 *"Data specification"* (introduction to deliverables 5.1-5.6), DEL5.1 *"Data requirements"* (which lists acceptance criteria for data submitted to FG-AI4H and states the governing principles and rules), DEL5.2 *"Data acquisition"*, DEL5.3 *"Data annotation specification"*, DEL5.4 *"Training and test data specification"* (which provides a systematic way of preparing technical requirement specifications for datasets used in the training and testing of AI models), DEL5.5 *"Data handling"* (which outlines how data will be handled once they are accepted), DEL5.6 *"Data sharing practices"* (which provides an overview of the existing best practices for sharing health-related data based on distributed and federated environments, including the requirement to enable secure data sharing and addressing issues of data governance), DEL06 *"AI training best practices specification"* (which reviews best practices for proper AI model training and guidelines for model reporting), DEL7 *"AI for health evaluation considerations"* (which discusses the validation and evaluation of AI for health models, and considers requirements for a benchmarking platform), DEL7.1 *"AI4H evaluation process description"* (which provides an overview of the state of the art of AI evaluation principles and methods and serves as an initiator for the evaluation process of AI for health), DEL7.2 *"AI technical test specification"* (which specifies how an AI can and should be tested *in silico*), DEL7.3 *"Data and artificial intelligence assessment methods (DAISAM)"* (which provides the reference collection of WG-DAISAM on assessment methods of data and AI quality evaluation), DEL7.4 *"Clinical Evaluation of AI for health"* (which outlines the current best practices and outstanding issues related to the clinical evaluation of AI models for health), DEL7.5 *"FG-AI4H assessment platform"* (which explores assessment platform options that can be used to evaluate AI for health for the different topic groups), DEL9 *"AI for health applications and platforms"* (which introduces specific considerations for the benchmarking of mobile- and cloud-based AI applications in health), and DEL9.1 *"Mobile based AI applications,"* and DEL9.2 *"Cloud-based AI applications"* (which describe specific requirements for the development, testing and benchmarking of mobile- and cloud-based AI applications).

## 6.1 Subtopic colonoscopy

The benchmarking of AI for endoscopy is being developed and improved continuously to reflect new features of AI systems or changed requirements for benchmarking. It should be noted that the benchmarking by TG-Endoscopy is not mature and needs further study.

### 6.1.1 Benchmarking version V1.0

This clause includes all technological and operational details of the benchmarking process for the benchmarking of version V1.0.

#### 6.1.1.1 Overview

This clause provides an overview of the key aspects of this benchmarking iteration, version V1.0. Besides recommended testing metrologies and scoring matrices, and data format requirements of input data and output data, the training and testing of data annotation quality control are also involved in the method for AI benchmarking.

#### 6.1.1.2 Benchmarking methods

This clause provides details about the methods of the benchmarking, version V1.0. All developers of AI solutions for endoscopy implemented internal benchmarking systems for assessing performance.

##### 6.1.1.2.1 Benchmarking system architecture

Referring to the benchmarking and evaluation of the APDCV challenge , the benchmarking system of colonoscopy should consist of AI tasks, benchmarking metrics and task-based metrics calculation, see Figure 1. Benchmarking version V1.0 is being built following this structure.

While the selection and calculation of metrics differs from AI tasks and applications, task-based metrics calculation is decided by the type of AI tasks and benchmarking metrics used. For example, PDR is applicable for polyp detection but not polyp classification.

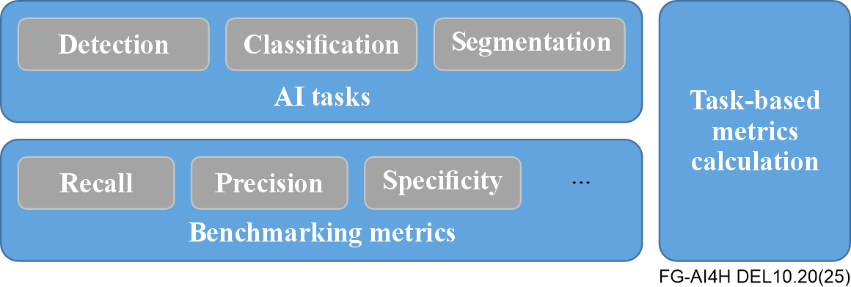


Figure 1 – Architecture of benchmarking version

##### 6.1.1.2.2 Benchmarking system dataflow

Initially, benchmarking version V1.0 will be a standalone and internal system for assessing performance. The type of AI tasks and the benchmarking metrics are defined manually, and the prediction by AI system is needed to be done before benchmarking.

The dataflow is illustrated in Figure 2. First, the test dataset was predicted by AI system to generate the prediction of test dataset, whose data structure need to applicable to the benchmarking system. Then, the AI task and benchmarking metrics need to be set based on the feature of algorithm, so as to calculate the task-based metrics. Finally, the prediction of test dataset by AI system was evaluated with ground truth of test dataset by the task-based metrics.

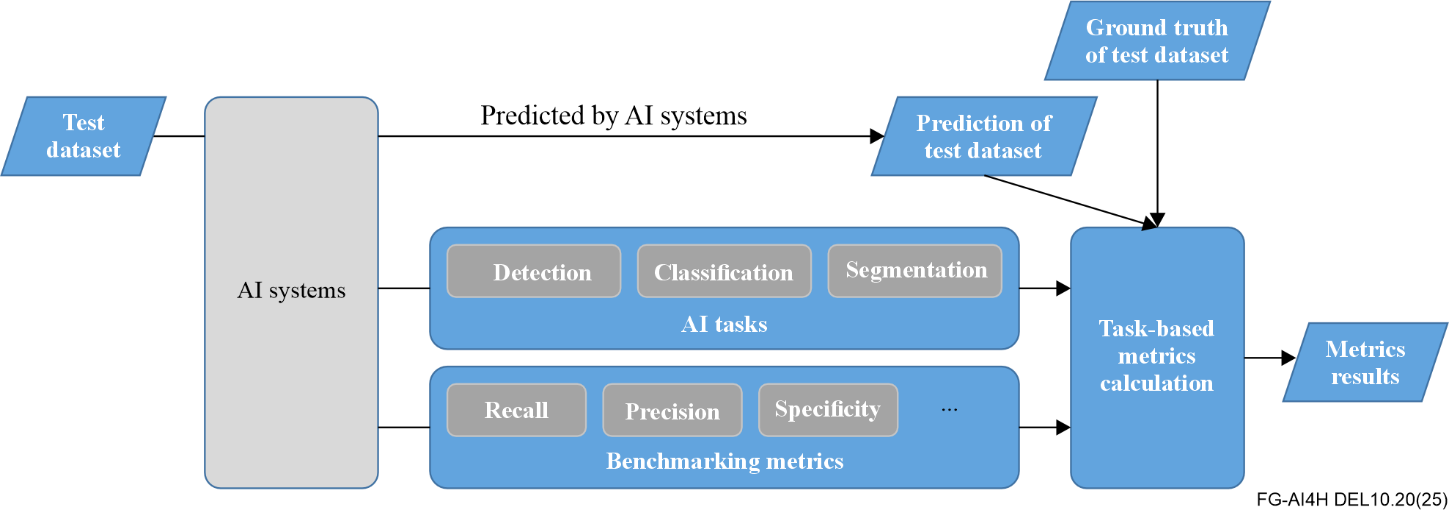


Figure 2 – Dataflow of benchmarking version

##### 6.1.1.2.3 Safe and secure system operation and hosting

The access to the system will only be authorized inside the corporation. More details of safety and security will be considered in the following version.

##### 6.1.1.2.4 Benchmarking process

The current version of the benchmarking system will be a standalone system and not for open access. The prediction of test dataset by AI systems, definition of AI tasks and benchmarking metrics in benchmarking, and execution of benchmarking calculation will be handled and done by authorized AI developers.

#### 6.1.1.3 AI input data structure for the benchmarking

This clause describes the recommended structure of input data provided to the AI solutions as part of the benchmarking of AI for endoscopy.

Endoscopic images or videos captured with colonoscope should be submitted as separate files in the following format:

• Image file format: JPEG format, PNG format or BMP format.

• Image file names: be unique in the dataset and anonymize the personal information of the patient.

• Image resolution: original resolution as captured with endoscopic device.

• Video file format: AVI format or MPEG-4 format.

• Video file names: be unique in the dataset and anonymize the personal information of the patient.

• Video resolution: original resolution as captured with endoscopic device.

#### 6.1.1.4 AI output data structure

Similar to the input data structure for the benchmarking, this clause describes the recommended structure of output data the AI systems are expected to generate in response to the input data.

The output should be documented in an arranged and clear way, like a CSV, XML or JSON file, with the following information.

• Information of data (name, format, etc).

• Result of the data. It would depend upon the specific condition and the type of task that is being benchmarked.

##### 6.1.1.4.1 Detection

• Data information: data name, data format, etc.

• Result information

• Category information: the types would depend on the task.

• Location information: coordinates of a specific point (left-top or centre of the bounding box) in the image. For video data, the slice index should be recorded.

• Size information: height and width in pixels.

• Task info (optional): task ID, task name, task type, etc.

##### 6.1.1.4.2 Classification

• Data information: data name, data format, etc.

• Result information

• Category information: the types would depend on the task.

• Task information (optional): task ID, task name, task type, etc.

##### 6.1.1.4.3 Segmentation

• Data information: data name, data format, etc.

• Result information

• Category information: the types would depend on the task.

• Path of segmentation file: the stored path of the segmentation file.

• Segmentation border information (optional): coordinates of points of the segmentation mask.

• Task information (optional): task ID, task name, task type, etc.

#### 6.1.1.5 Test data label/annotation structure

While the AI systems can only receive the input data described in the previous sections, the benchmarking system needs to know the expected correct answer (sometimes called 'labels', 'ground truth' or 'gold standard') for each element of the input data, so that it can compare the expected AI output with the actual one. Since this is only needed for benchmarking, it is encoded separately.

To guarantee the quality of data annotation and reduce individual differences among doctors, it is recommended that the annotation process should involve multiple steps by multiple doctors, such as independent annotation, cross-annotation, arbitration, and review. Specially, arbitration and review may be combined as one step by one doctor.

If appropriate, a corresponding clinical diagnosis report or pathological report would be recommended for reference, even the gold standard in data annotation.

Before annotation, the data needs preliminary filtering and laundering to eliminate worthless data, such as missing data, image parameter mismatch, non-inspection site data, foreign matter in the data, image artefacts, and images whose quality cannot satisfy the diagnostic requirements.

##### 6.1.1.5.1 Annotation of detection

The annotation of detection includes localizing the object inside the data and categorizing it, see Figure 3. The bounding box is usually used to localize the object with a rectangular box which is called a bounding box.

• Independent annotation: Independent annotation by two doctors to confirm whether the endoscopic image/video contains lesions or intended objects and, if so, mark the location and size of the lesion or intended objects with a bounding box. All the marked bounding boxes should be documented in a clear way, like a CSV file. Independent annotation requirements include:

o Non-annotating information (optional): Image/video name, image/video identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information (mandatory): Annotated results (bounding box like [*x, y, w, h, s*], where *s* is the slice index in video and equal to 0 in image), annotator information, annotation procedure information, the date, annotation serial number.

• Cross-annotation: The independent annotations by different annotators are cross-evaluated to identify the relationship between each other by calculating the similarity, like IoU [91]. If the similarity satisfies pre-set requirements, the independent annotations would be merged to the gold standard candidate in a specific manner, like average. Cross-annotation requirements include:

o Non-annotating information (optional): Image/video name, image/video identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information (mandatory): Cross-annotated results (bounding box like [x, y, w, h, s], if the pre-set requirements are not satisfied, the bounding box should be [0, 0, 0, 0, 0]), annotation serial numbers for merging, merge manner, annotation procedure information, the date, annotation serial number.

• Arbitration:If the similarity calculated in the cross-annotation step does not satisfy the pre-set requirements, the corresponding data will be transferred to the arbitration doctor to review and re-annotate as a gold standard candidate. Otherwise, the gold standard candidate in step cross-annotation will be transferred to the review doctor. Arbitration requirements include:

o Non-annotating information(optional): Image/video name, image/video identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information(mandatory): The arbitrated results (bounding box like [*x, y, w, h, s*]), annotation serial numbers for arbitration, arbitration doctor information, annotation procedure information, the date, annotation serial number.

• Review: The gold standard candidates would be confirmed by the review doctor one by one. The data approved by the review doctor would be marked as the gold standard. Otherwise, the data without review approval would be sent back to the arbitration procedure or modified by the review doctor to generate the gold standard. Review requirements include:

o Non-annotating information(optional): Image/video name, image/video identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information(mandatory): The review results (gold standard or sent back to arbitration), serial number for review, review doctor information, annotation procedure information, the date, annotation serial number.

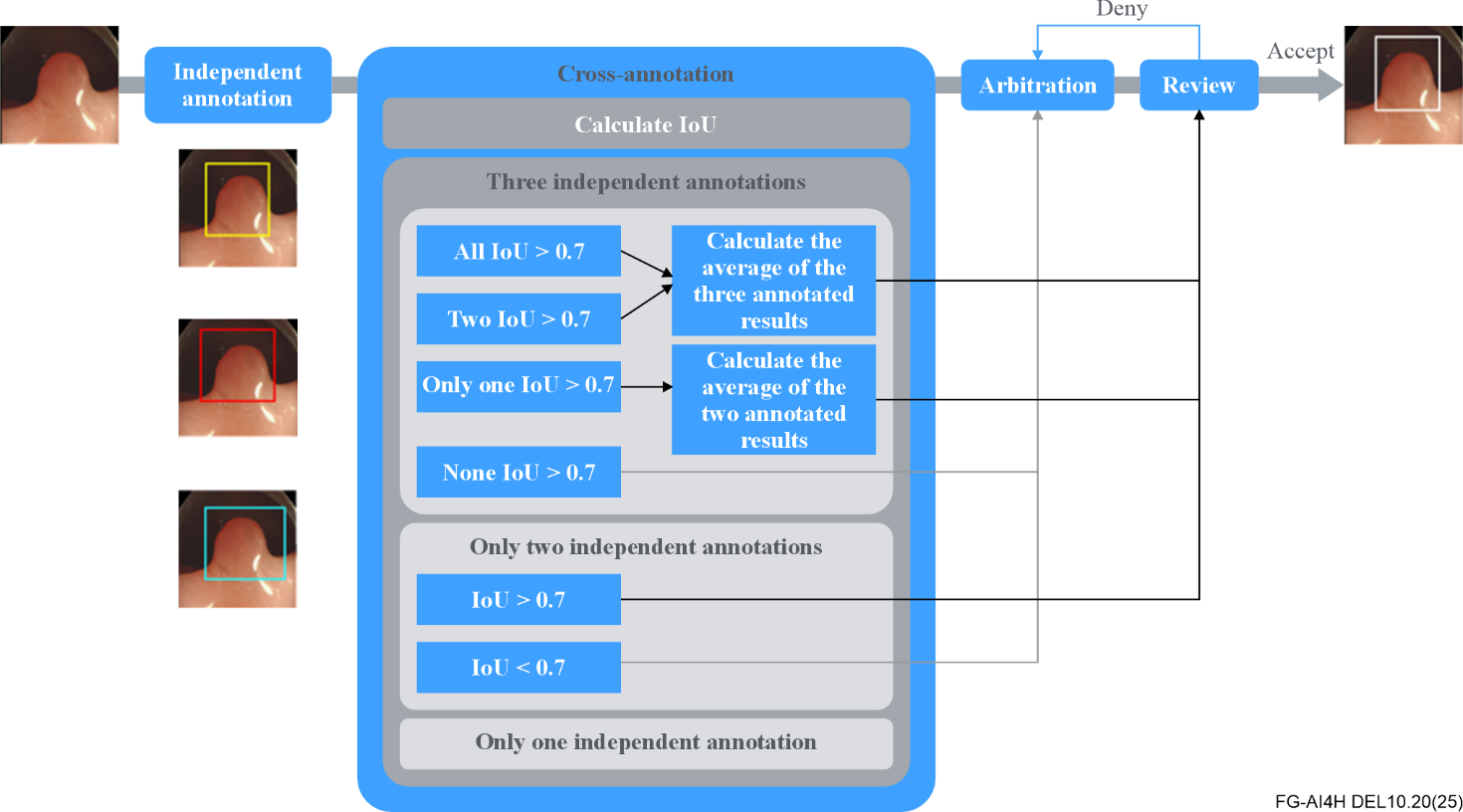


Figure 3 ؘ– Illustration of annotation procedure for detection

##### 6.1.1.5.2 Annotation of classification

Annotation of classification means arranging a category for the data. For example, the decision of the category might be made subjectively, based on the manual observing of features in the entire or part of data. Also, the category might be made objectively, based on the corresponding objective evidence, like pathological results.

In the subjective annotation procedure, the annotation would be made manually without objective evidence.

• Independent annotation: Independent annotation of classification by two doctors to confirm which category the data should be arranged. All the annotated results should be documented in a clear way, like a CSV file. Independent annotation requirements include:

o Non-annotating information (optional): Image/video name, image/video identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information(mandatory): Annotated results, annotator information, annotation procedure information, the date, annotation serial number.

• Cross-annotation: The independent annotations by different annotators are cross-evaluated to identify the relationship between each other by calculating the level of consistency. If the level of consistency satisfies pre-set requirements, the independent annotations will be merged to the gold standard candidate in a specific manner, like majority rule. Cross- annotation requirements include:

o Non-annotating information (optional): Image/video name, image/video identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information (mandatory): Cross-annotated results, annotation serial numbers for merging, merge manner, annotation procedure information, the date, annotation serial number.

• Arbitration: If the level of consistency of independent annotations does not satisfy the pre-set requirements, the corresponding data will be transferred to the arbitration doctor to review and re-annotate as a gold standard candidate. Otherwise, the gold standard candidate in step cross-annotation will be transferred to the review doctor. Arbitration requirements include:

o Non-annotating information (optional): Image/video name, image/video identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information (mandatory): The arbitrated results, annotation serial numbers for arbitration, arbitration doctor information, annotation procedure information, the date, annotation serial number.

• Review: The gold standard candidates will be confirmed by the review doctor one by one. The data approved by the review doctor will be marked as the gold standard. Otherwise, the data without review approval will be sent back to the arbitration procedure or modified by the review doctor to generate the gold standard. Review requirements include:

o Non-annotating information(optional): Image/video name, image/video identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race)

o Annotating information(mandatory): The review results (gold standard or sent back to arbitration), serial number for review, review doctor information, annotation procedure information, the date, annotation serial number.

The process is illustrated in Figure 4.

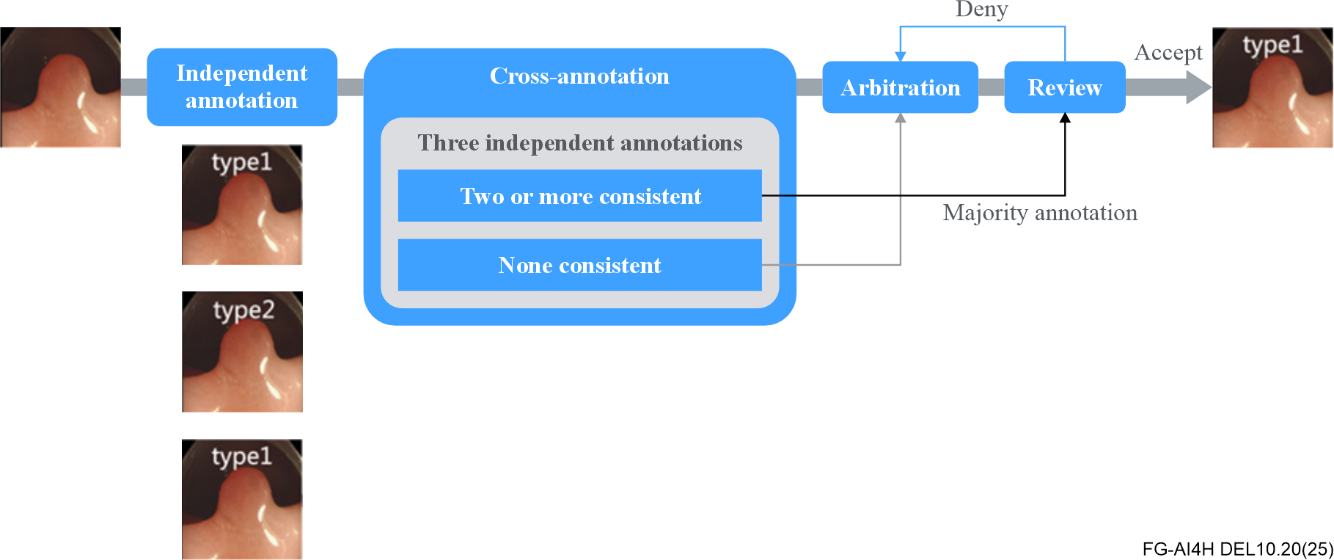


Figure 4 – Illustration of annotation procedure for classification

##### 6.1.1.5.3 Annotation of segmentation

Annotation of segmentation means the annotation of every pixel in an object within a data. Practically, there are two methods for annotation of segmentation: annotating the contour of the object with a polygon and annotating the region of the object with a mask.

• Initial annotation: Initial annotation to sketch the contour or mask of the object by one doctor. All the annotated results should be well recorded and linked to corresponding images in a clear way. Initial annotation requirements include:

o Non-annotating information (optional): Image name, image identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information (mandatory): Annotated results, annotator information, annotation procedure information, the date, annotation serial number.

• Review: The initial annotation will be confirmed and modified by the review doctor. The data approved by the review doctor will be marked as the gold standard. Review annotation requirements include:

o Non-annotating information (optional): Image name, image identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information (mandatory): The gold standard, serial number for review, review doctor information, annotation procedure information, the date, annotation serial number.

#### 6.1.1.6 Scores and metrics

Scores and metrics are at the core of the benchmarking. This clause describes the scores and metrics applicable to measure the performance, robustness, and general characteristics of AI systems. Table 5 presents a list of applicable scores and metrics that could be used in demands.

| Table 5 – Benchmarking metrics | | |
| --- | --- | --- |
| Methodology | Description | AI Task |
| True positive (TP) | The number of correctly identified positive samples. The number of frames with endoscopic findings which are correctly identified as frames with an endoscopic finding. | Detection  classification |
| True negative (TN) | The number of correctly identified negative samples, i.e. the number of frames without an endoscopic finding which are correctly is identified as frames without an endoscopic finding. | Detection  classification |
| False positive (FP) | The number of wrongly identified positive samples, i.e. commonly called a "false alarm". The number of frames without an endoscopic finding which are erroneously identified as frames with an endoscopic finding. | Detection  classification |
| False negative (FN) | The number of wrongly identified negative samples. The number of frames with an endoscopic finding which erroneously are identified as frames without an endoscopic finding. | Detection  classification |
| Recall (REC) or sensitivity (SENS) | This metric is also frequently called sensitivity, probability of detection and true positive rate, and it is the ratio of samples that are correctly identified as positive among all existing positive samples. | Detection  classification |
| Precision (PREC) or positive predictive value (PPV) | This metric is also frequently called the positive predictive value. It shows the ratio of samples that are correctly identified as positive among the returned positive samples (the fraction of retrieved samples that are relevant). | Detection  classification |
| Negative predictive value (NPV) | This shows the ratio of samples that are correctly identified as negative among the predicted negative samples (the fraction of retrieved samples that are relevant). | Detection  classification |
| Specificity (SPEC) | This metric is frequently called the true negative rate. It shows the ratio of negatives that are correctly identified as such (e.g. the fraction of frames without an endoscopic finding are correctly identified as a negative result). | Detection  classification |
| Accuracy (ACC) | The percentage of correctly identified true and false samples. | Detection  classification |
| Matthews correlation coefficient (MCC) | MCC takes into account true and false positives and negatives. It is a balanced measure even if the classes are of very different sizes. | Classification |
| F1 score (F1) | A measure of a test's accuracy by calculating the harmonic mean of the precision and recall. | Classification |
| Dice coefficient (DICE) | This metric measures the similarity between two sets of data and is most broadly used in the validation of image [segmentation](https://radiopaedia.org/articles/segmentation?lang=us). It equals twice the number of elements common to both sets divided by the sum of the number of elements in each set. | Segmentation |
| Jaccard coefficient or Intersection over union (IoU) | This metric measures the similarity between two sets of data and is most broadly used in the validation of object detection and image segmentation. It equals the number of elements common to both sets divided by the sum of the number of unique elements in each set. | Segmentation |
| Polyp detection rate (PDR) | This metric is the percentage of patients undergoing screening endoscopy who have one or more polyp detected. | Detection |
| Adenoma detection rate (ADR) | This metric is the percentage of patients undergoing screening endoscopy who have one or more conventional adenomas detected. | Detection |
| Detection latency | This metric is the delay in frames between the first appearance of the polyp/lesion/object in the video sequence and the first actual detection of the polyp/lesion/object by a method. | Detection |
| Average precision (AP) | This metric is the average precision in the P-R curve. | Detection |
| Mean average precision (mAP) | This metric is the average value of AP of every class. | Detection |
| Runtime | This metric is the cost time of running one image or frame by a method. | Detection  Classification  Segmentation |

#### 6.1.1.7 Test dataset acquisition

The test dataset acquisition is in progress.

#### 6.1.1.8 Data sharing policies

After finishing the test dataset acquisition, the sharing of the dataset should be protected by special agreements or contracts that cover, for instance, the data sharing period, patient consent, and update procedure (see also [DEL5.5](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/_layouts/15/WopiFrame.aspx?sourcedoc=%7B71FE8B9D-ACB3-48CE-AA3F-136409B550A4%7D&file=DEL05_5.docx&action=default) on *data handling* and [DEL5.6](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/_layouts/15/WopiFrame.aspx?sourcedoc=%7B5C95327E-96A5-4175-999E-3EDB3ED147C3%7D&file=DEL05_6.docx&action=default) on *data sharing practices*).

#### 6.1.1.9 Baseline acquisition

The baseline will be acquired after the test dataset acquisition is finished.

#### 6.1.1.10 Reporting methodology

The results of benchmarking runs will be shared with AI developers internally. There is no public reporting methodology, except the publication of technical papers.

#### 6.1.1.11 Results

Results will be not available before the benchmarking version 1.0 and test dataset acquisition are finished.

#### 6.1.1.12 Discussion on benchmarking

This clause provides insights on benchmarking iterations, giving details about the 'outcome' of the benchmarking process (e.g. giving an overview of the benchmark results and process).

In the benchmarking of subtopic colonoscopy, recommended requirements for benchmarking methods, data structure of input and output, annotation structure and information, scores and metrics, test datasets and results are described. In the corresponding clauses, there are individual requirements for detection, segmentation, and classification.

Referring to the benchmarking and evaluation of APDCV, the benchmarking in this subtopic is being built as a standalone system initially, consisting of AI tasks, benchmarking metrics and task-based metrics calculation. As there are several existing benchmarking systems, the data structure of input/‌output and the annotation structure and information are considered first, so the progresses of data structure of input/output, annotation structure and information are more advanced than benchmarking system and test dataset acquisition. Data annotation is currently a spontaneous non-standard process. It is a challenging task to guarantee the accuracy and representativeness of learning materials without the standardized data annotation quality control measures which are widely recognized by the industry. Moreover, this lack of accuracy may also bring a greater risk of erroneous judgment for endoscopic-assisted diagnosis. The benchmarking version 1.0 tries to propose a general standardized solution of requirements for data structure and annotation.

#### 6.1.1.13 Retirement

Generally, the retirement of the AI system and dataset should follow the policy agreed with providers and users before the benchmarking activity. It might be desirable to keep the database for traceability and future use. Alternatively, there may be security or privacy reasons for deleting the data. Further details can be found in the reference document of This clause [DEL04](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/_layouts/15/WopiFrame.aspx?sourcedoc=%7BC68833D1-9B31-4E8E-8A4A-3939D7DEA56F%7D&file=DEL04.docx&action=default) *"AI software lifecycle specification"* (identification of standards and best practices that are relevant for the AI for health software life cycle).

## 6.2 Subtopic endoscopic ultrasound

### 6.2.1 Benchmarking version V1.0

This clause includes all technological and operational details of the benchmarking process for benchmarking version V1.0. It should be noted that, the benchmarking by TG-Endoscopy is not mature and needs further study.

#### 6.2.1.1 Overview

This clause provides an overview of the key aspects of this benchmarking iteration, version V1.0.

The method for AI benchmarking, including recommended requirement of data format for input data and output data, and training and testing data annotation quality control, as well as testing metrologies and scoring matrices, are described.

#### 6.2.1.2 Benchmarking methods

##### 6.2.1.2.1 Benchmarking system architecture

Compared to colonoscopy, EUS is a multi-modality procedure, capturing ultrasound and image at the same time, but the requirements of benchmarking are similar. EUS and colonoscopy share the same benchmarking system architecture described in clause 6.1.1.2.1.

##### 6.2.1.2.2 Benchmarking system dataflow

EUS and colonoscopy share the same benchmarking system dataflow described in clause 6.1.1.2.2.

##### 6.2.1.2.3 Safe and secure system operation and hosting

EUS and colonoscopy share the same safe and secure system operation and hosting described in clause 6.1.1.2.3.

##### 6.2.1.2.4 Benchmarking process

EUS and colonoscopy share the same benchmarking process described in clause 6.1.1.2.4.

#### 6.2.1.3 AI input data structure for the benchmarking

Ultrasound images or videos captured with EUS should be submitted as separate files in the following format:

• Image file format: JPEG format, PNG format, BMP format or DICOM format.

• Image file names: be unique in the dataset and anonymize the personal information of the patient. For the DICOM file, anonymizing should be performed to remove sensitive information in the DICOM tag.

• Image resolution: original resolution as captured with EUS.

• Video file format: AVI format or MPEG-4 format or DICOM format.

• Video file names: be unique in the dataset and anonymize the personal information of the patient. For the DICOM file, anonymizing should be performed to remove sensitive information in the DICOM tag.

• Video resolution: original resolution as captured with EUS.

#### 6.2.1.4 AI output data structure

The output should be documented in an arranged and clear way, like a CSV, XML or JSON file with the following information.

• Information of data (name, format, etc).

• Result of the data. It would depend on the specific condition and the type of task that is being benchmarked.

##### 6.2.1.4.1 Detection

• Data information: data name, data format, etc.

• Result information:

o Category information: the types would depend on the task.

o Location information: coordinates of a specific point (left-top or centre of the bounding box) in the image. For video data, the slice index should be recorded.

o Size information: height and width in pixels.

• Task info (optional): task ID, task name, task type, etc.

##### 6.2.1.4.2 Classification

• Data information: data name, data format, etc.

• Result information

o Category information: the types would depend on the task.

• Task information (optional): task ID, task name, task type, etc.

##### 6.2.1.4.3 Segmentation

• Data information: data name, data format, etc.

• Result information

o Category information: the types would depend on the task.

o Path of segmentation file: the stored path of the segmentation file.

o Segmentation border information (optional): coordinates of points of the segmentation mask.

• Task information (optional): task ID, task name, task type, etc.

#### 6.2.1.5 Test data label/annotation structure

While the AI systems can only receive the input data described in the previous sections, the benchmarking system needs to know the expected correct answer (sometimes called 'labels', 'ground truth' or 'gold standard') for each element of the input data so that it can compare the expected AI output with the actual one. Since this is only needed for benchmarking, it is encoded separately.

Comparing with colonoscopy, EUS is a multi-modality procedure capturing ultrasound and image at the same time. The test data label/annotation structure needs to consider of two modalities as ultrasound data and image/video.

Referring to the section on 'Test data label/annotation structure' of colonoscopy, it is recommended that the annotation process should involve multiple steps by multiple doctors, such as independent annotation, cross-annotation, arbitration, and review. Specially, arbitration and review may be combined as one step by one doctor.

##### 6.2.1.5.1 Annotation of detection

The annotation of detection includes localizing the object inside the data and categorizing it. The bounding box is usually used to localize the object, using a rectangular box called a bounding box.

• Independent annotation: Independent annotation by two doctors to confirm whether the image/video/ultrasound data contains lesions or intended objects and, if so, mark the location and size of the lesion or intended objects with a bounding box. All the marked bounding boxes should be documented in a clear way, like a CSV file. Independent annotation requirements include:

o Non-annotating information (optional): Image/video/ultrasound data name, image/video/ultrasound data identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information(mandatory): Annotated results (bounding box like [*x, y, w, h, s*], where *s* is the slice index in video and equal to 0 in image), annotator information, annotation procedure information, the date, annotation serial number.

• Cross-annotation: The independent annotations by different annotators are cross-evaluated to identify the relationship between each other by calculating the similarity, like IoU. If the similarity satisfies pre-set requirements, the independent annotations would be merged to the gold standard candidate in a specific manner, like average. Cross-annotation requirements include:

o Non-annotating information (optional): Image/video/ultrasound data name, image/video/ultrasound data identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information (mandatory): Cross-annotated results (bounding box like [x, y, w, h, s], if the pre-set requirements are not satisfied, the bounding box should be [0, 0, 0, 0, 0]), annotation serial numbers for merging, merge manner, annotation procedure information, the date, annotation serial number.

• Arbitration:If the similarity calculated in the cross-annotation step does not satisfy the pre-set requirements, the corresponding data will be transferred to the arbitration doctor to review and re-annotate as a gold standard candidate. Otherwise, the gold standard candidate in step cross-annotation would be transferred to the review doctor. Arbitration requirements include:

o Non-annotating information (optional): Image/video/ultrasound data name, image/video/ultrasound data identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information (mandatory): The arbitrated results (bounding box like [x, y, w, h, s]), annotation serial numbers for arbitration, arbitration doctor information, annotation procedure information, the date, annotation serial number.

• Review: The gold standard candidates would be confirmed by the review doctor one by one. The data approved by the review doctor would be marked as the gold standard. Otherwise, the data without review approval would be sent back to the arbitration procedure or modified by the review doctor to generate the gold standard. Review requirements include:

o Non-annotating information (optional): Image/video/ultrasound data name, image/video/ultrasound data identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race)

o Annotating information (mandatory): The review results (gold standard or sent back to arbitration), serial number for review, review doctor information, annotation procedure information, the date, annotation serial number.

##### 6.2.1.5.2 Annotation of classification

Annotation of classification means arranging a category for the data. For example, the decision of the category might be made subjectively, based on the manual observing of features in the entire or part of data. Also, the category might be made objectively, based on the corresponding objective evidence, like pathological results.

In the subjective annotation procedure, the annotation would be made manually without objective evidence.

• Independent annotation: Independent annotation of classification by two doctors to confirm in which category the data should be arranged. All the annotated results should be documented in a clear way, like a CSV file. Independent annotation requirements include:

o Non-annotating information (optional): Image/video/ultrasound data name, image/video/ultrasound data identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information (mandatory): Annotated results, annotator information, annotation procedure information, the date, annotation serial number.

• Cross-annotation: The independent annotations by different annotators are cross-evaluated to identify the relationship between each other by calculating the level of consistency. If the level of consistency satisfies pre-set requirements, the independent annotations would be merged to the gold standard candidate in a specific manner, like majority rule. Cross- annotation requirements include:

o Non-annotating information (optional): Image/video/ultrasound data name, image/video/ultrasound data identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information (mandatory): Cross-annotated results, annotation serial numbers for merging, merge manner, annotation procedure information, the date, annotation serial number.

• Arbitration: If the level of consistency of independent annotations does not satisfy the pre-set requirements, the corresponding data will be transferred to the arbitration doctor to review and re-annotate as a gold standard candidate. Otherwise, the gold standard candidate in step cross-annotation would be transferred to the review doctor. Arbitration requirements include:

o Non-annotating information (optional): Image/video/ultrasound data name, image/video/ultrasound data identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information (mandatory): The arbitrated results, annotation serial numbers for arbitration, arbitration doctor information, annotation procedure information, the date, annotation serial number.

• Review: The gold standard candidates would be confirmed by the review doctor one by one. The data approved by the review doctor would be marked as the gold standard. Otherwise, the data without review approval would be sent back to the arbitration procedure or modified by the review doctor to generate the gold standard. Review requirements include:

o Non-annotating information (optional): Image/video/ultrasound data name, image/video/ultrasound data identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information (mandatory): The review results (gold standard or sent back to arbitration), serial number for review, review doctor information, annotation procedure information, the date, annotation serial number.

##### 6.2.1.5.3 Annotation of segmentation

Annotation of segmentation means the annotation of every pixel in an object within a data. Practically, there are two methods for annotation of segmentation, including annotating the contour of the object with a polygon and annotating the region of the object with a mask.

• Initial annotation: Initial annotation to sketch the contour or mask of the object by one doctor. All the annotated results should be well recorded and linked to corresponding images in a clear way. Initial annotation requirements include:

o Non-annotating information(optional): Image/ultrasound data name, image/ultrasound data identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race).

o Annotating information(mandatory): Annotated results, annotator information, annotation procedure information, the date, annotation serial number.

• Review: The initial annotation would be confirmed and modified by the review doctor. The data approved by the review doctor would be marked as the gold standard. Review annotation requirements include:

o Non-annotating information (optional): Image/ultrasound data name, image/ultrasound data identification code, collection device model, collection date, image size, collection frame rate, hospital, patient information (age, gender, race)

o Annotating information (mandatory): The gold standard, serial number for review, review doctor information, annotation procedure information, the date, annotation serial number.

#### 6.2.1.6 Scores and metrics

EUS and colonoscopy share the same scores and metrics described in clause 6.1.1.6.

#### 6.2.1.7 Test dataset acquisition

The test dataset acquisition is in progress.

#### 6.2.1.8 Data sharing policies

After finishing the test dataset acquisition, the sharing of the dataset should be protected by special agreements or contracts that cover, for instance, the data sharing period, patient consent, and update procedure (see also [DEL5.5](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/_layouts/15/WopiFrame.aspx?sourcedoc=%7B71FE8B9D-ACB3-48CE-AA3F-136409B550A4%7D&file=DEL05_5.docx&action=default) on *data handling* and [DEL5.6](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/_layouts/15/WopiFrame.aspx?sourcedoc=%7B5C95327E-96A5-4175-999E-3EDB3ED147C3%7D&file=DEL05_6.docx&action=default) on *data sharing practices*).

#### 6.2.1.9 Baseline acquisition

The baseline will be acquired after finishing the test dataset acquisition.

#### 6.2.1.10 Reporting methodology

EUS and colonoscopy will share the same reporting methodology described in clause 6.1.1.10.

#### 6.2.1.11 Results

Currently, there is no publicly available EUS dataset and benchmarking system. It is impossible to perform comparable benchmarking for different AI solutions. Several review papers have been published to summarize the latest research in AI-assisted EUS in different clinical fields [86][87][88][89][90]. For instance, Dumitrescu. et. al. conducted meta-analysis for the diagnostic value of AI-assisted EUS for pancreatic cancer with 10 clinical studies and 1871 patients [88]. The overall diagnostic accuracy showed 0.92 (95% CI, 0.89–0.95) sensitivity and 0.9 (95% CI, 0.83–0.94) specificity.

#### 6.2.1.12 Discussion on benchmarking

In general, EUS produces ultrasound images of different kinds (B mode, contrast enhanced ultrasound, elastography), just like conventional endoscopy does (white light, narrow band imaging, dye-spray chromoendoscopy). So, the benchmarking methods, general requirements for data structure of input and output, annotation structure and information, scores and metrics, test dataset and result are not very different from those of endoscopic subgroups. General descriptions are given in above sections. With regards to the difference of different AI tasks, there are individual requirements for detection, segmentation, and classification in the corresponding clauses.

It should be noted that one data type, radiofrequency (RF) data, and its related AI task (for instance, beamforming, data compression, denoising, reconstruction, etc.) are discarded from the current version of the benchmarking process for mainly two reasons: (1) only a few EUS manufacturers and research facilities have the ability to access EUS RF data from EUS system. Research on RF data-based AI-EUS is rare at the moment; and (2) currently there is no standard for storing RF data for different EUS manufacturers. It can be added in the future if needed.

#### 6.2.1.13 Retirement

EUS and colonoscopy will share the same reporting methodology described in clause 6.1.1.13.

# 7 Overall discussion of the benchmarking

Endoscopy is the core technical means for early diagnosis and screening of digestive cancer, which can drastically reduce the incidence and mortality caused by digestive cancer. Furthermore, with the breakthrough of the new generation of AI technology represented by deep learning, revolutionary progress has been made, and the real-time assistance of AI for detecting and classifying GI lesions is expected to help clinicians improve their examination quality and reduce the number of missed diagnoses. This TDD specifies the standardized benchmarking for AI for endoscopy systems in two subtopics, namely, colonoscopy and EUS.

Colonoscopy is considered the gold standard for CRC screening for detecting and removing polyps and adenomas in the colorectum. By the effort of researchers, some work has been done by the scientific community in assessing the performance of such application, such as challenges and datasets. In each challenge, general elements were involved, including tasks, data, annotation and metrics. It should be noted that there might be different definitions and selection mechanisms for these elements. The fundamental element would be a determined dataset. A variety of datasets have been released and open accessed. There are datasets annotated with single type of lesions or multiple types of lesions and also datasets of images or videos. Based on these challenges and datasets, researchers have published a variety of high-quality publications, and there are already commercial AI products on the market. In this TDD, recommended requirements for the data structure of input and output, annotation structure and information, score and metrics, test datasets and results are described. There is more specific description of recommended requirements for data structure of input and output, annotation structure and information, which aims to guarantee the accuracy and representativeness of datasets and annotations. Referring to the benchmarking and evaluation of APDCV, the benchmarking in this subtopic is being built as a standalone system initially. The access to the system will be only authorized inside the corporation.

As a fresh technology, clinical evidence has shown the benefits of EUS over the potential adverse events, and clinical guidelines have been published and continuously updated to ensure the safe use of the procedures. In general, EUS produces ultrasound images of different kinds (B mode, contrast enhanced ultrasound, elastography), just like conventional endoscopy does(white light, narrow band imaging, dye-spray chromoendoscopy). So the benchmarking methods, general requirements for data structure of input and output, annotation structure and information, score and metrics, test dataset and result are no big different with other endoscopic subgroup. Although research on AI in EUS is still limited, it is believed that AI will play an important role in EUS procedures, not only in detecting anatomical features, differentiating benign and malignant lesions, and delineating lesion contours but, more importantly, by reducing the learning time for junior endoscopists, decreasing workloads and standardizing the overall quality of endoscopic procedures.

Generally, it should be noted that the benchmarking by TG-Endoscopy is not mature and needs further study.

# 8 Regulatory considerations

For AI-based technologies in healthcare, regulation is not only crucial to ensuring the safety of patients and users, but also to accomplishing market acceptance of these devices. This is challenging because there is a lack of universally accepted regulatory policies and guidelines for AI-based medical devices. To ensure that the benchmarking procedures and validation principles of FG-AI4H are secure and relevant for regulators and other stakeholders, the working group on "[Regulatory considerations on AI for health" (WG-RC)](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/wg/SitePages/WG-RC.aspx) compiled the requirements that consider these challenges.

The deliverables with relevance to regulatory considerations are [DEL2](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/_layouts/15/WopiFrame.aspx?sourcedoc=%7BF2F46A99-7457-4BC8-81A3-0E1E63D6072A%7D&file=DEL02.docx&action=default) *"AI4H regulatory considerations"* (which provides an educational overview of some key regulatory considerations), [DEL2.1](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/_layouts/15/WopiFrame.aspx?sourcedoc=%7B6AF7C004-8BCE-4151-9F44-45F041A1EB1D%7D&file=DEL02_1.docx&action=default) *"Mapping of IMDRF essential principles to AI for health software",* and[DEL2.2](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/_layouts/15/WopiFrame.aspx?sourcedoc=%7B1ED0D4D1-876C-4A0F-AEF7-06D3F445F5E6%7D&file=DEL02_2.docx&action=default) *"Guidelines for AI based medical device (AI-MD): Regulatory requirements"* (which provides a checklist for understanding the expectations of regulators, promotes step-by-step implementation of safety and effectiveness of AI-based medical devices, and compensates for the lack of a harmonized standard). [DEL04](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/_layouts/15/WopiFrame.aspx?sourcedoc=%7BC68833D1-9B31-4E8E-8A4A-3939D7DEA56F%7D&file=DEL04.docx&action=default) identifies standards and best practices that are relevant for the *"AI software lifecycle specification"*. The following sections discuss how the different regulatory aspects relate to TG‑Endoscopy.

## 8.1 Existing applicable regulatory frameworks

Most of the AI systems that are part of the FG-AI4H benchmarking process can be classified as software as a medical device(SaMD), and a multitude of regulatory frameworks that are already in place will be applicable to them. In addition, these AI systems often process sensitive personal health information that is controlled by another set of regulatory frameworks.

If the AI systems for endoscopy are for a clinical purpose and classified as SaMD, it would be covered by existing regulatory frameworks, such as the NMPA, MDR, FDA, General Data Protection Regulation (GDPR), and International Standardization Organization (ISO), and the AI manufacturers will need to address all the requirements of those regulatory frameworks.

## 8.2 Regulatory features to be reported by benchmarking participants

In most countries, benchmarked AI solutions can only be used legally if they comply with the respective regulatory frameworks for the application context.

The benchmarking participants need to provide compliance features and certifications as part of the metadata following the regulatory requirements in [DEL2](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/_layouts/15/WopiFrame.aspx?sourcedoc=%7BF2F46A99-7457-4BC8-81A3-0E1E63D6072A%7D&file=DEL02.docx&action=default) *"AI4H regulatory considerations"*.

## 8.3 Regulatory requirements for the benchmarking systems

The benchmarking system itself needs to comply with regulatory frameworks (e.g. some regulatory frameworks explicitly require that all tools in quality management are also implemented with a quality management system in place). This clause outlines the regulatory requirements for software used for benchmarking in this topic group.

Referring to the regulatory requirements in [DEL2](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/_layouts/15/WopiFrame.aspx?sourcedoc=%7BF2F46A99-7457-4BC8-81A3-0E1E63D6072A%7D&file=DEL02.docx&action=default) "*AI4H regulatory considerations*", if the benchmarking system is built for evaluating medical devices, it might need to comply with the following requirements.

| Table 6 – Regulatory requirements for the benchmarking systems | | |
| --- | --- | --- |
| Requirement(s) | Checklist item(s) | Applicable standards and regulations |
| The manufacturer should plan the model evaluation. | − There is an evaluation plan.  − The plan specifies the evaluation activities, the roles involved and the milestones at which these activities have to be performed.  − The plan foresees the evaluation with clinically relevant datasets independent from training datasets. | [ISO 13485] clauses 7.3.2, 7.3.6 and 7.3.7  [ISO 14971] clause 10.  GMLP guiding principle (8) (by FDA et al.) |
| The manufacturer should gain an understanding of how the machine makes a decision to evaluate the correctness and robustness of the model. | − There is a validation specification and validation results for the evaluation of the model with validation dataset.  − There is a test specification and test results for the final evaluation of the model with new test data.  − There are documented values for specified quality metrics.  − There may be an analysis of datasets that have exhibited good model performance versus datasets that have performed badly.  − For individual datasets there may be an evaluation of the feature that the model particularly determined in the decision.  − There may be an analysis/visualization of the dependency (strength, direction) of the prediction of the feature values.  − There may be a synthetization of datasets that activate the model particularly strongly.  − There may be an approximation of the model using a simplified surrogate model. | [EU-MDR (2017/745)] Annex I (17), Annex II (6.1).  [IEC 62304] clauses 5.5 ff.  [ISO 13485] clause 7.3.4 ff.  [b-XAVIER] "Perspectives and good practices for AI and continuously learning systems in healthcare"  [b-XAVIER University] "Building explainability and trust for AI in healthcare"  DIN SPEC 2  [b-ISO/IEC TR 24028] clauses 10.2 and 10.3  GMLP guiding principles (6) (e.g. overfitting) and (8) (confounding factors) (by FDA et al.) |

## 8.4 Regulatory approach for the topic group

Building on the outlined regulatory requirements, this clause describes how the topic group plans to address the relevant points in order to be compliant. The discussion here focuses on the guidance and best practices provided by the [DEL2](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/_layouts/15/WopiFrame.aspx?sourcedoc=%7BF2F46A99-7457-4BC8-81A3-0E1E63D6072A%7D&file=DEL02.docx&action=default) *"AI4H regulatory considerations"*.

To comply with applicable regulatory requirements, TG-Endoscopy will refer to the guidance and best practices provided by [DEL2](https://extranet.itu.int/sites/itu-t/focusgroups/ai4h/_layouts/15/WopiFrame.aspx?sourcedoc=%7BF2F46A99-7457-4BC8-81A3-0E1E63D6072A%7D&file=DEL02.docx&action=default) *"AI4H regulatory considerations"*.

Annex A  
  
Glossary

This clause lists all the relevant abbreviations, acronyms and uncommon terms used in the document.

| Acronym/Term | Expansion | Comment |
| --- | --- | --- |
| AE | Adverse Event |  |
| AI | Artificial Intelligence |  |
| AI4H | Artificial Intelligence for Health |  |
| AI-MD | AI based Medical Device |  |
| AMR | Adenoma Miss Rates |  |
| ANN | Artificial Neural Network |  |
| APDCV | Automatic Polyp Detection in Colonoscopy Videos |  |
| API | Application Programming Interface |  |
| AUC | Area Under Curve |  |
| CADe | Computer Assisted-Detection systems |  |
| CADx | Computer-Aided Diagnosis system |  |
| CATARACTS | The Challenge on Automatic Tool Annotation for cataRACT Surgery |  |
| CfTGP | Call for Topic Group Participation |  |
| CNN | Convolutional Neural Network |  |
| CP | Chronic pancreatitis |  |
| CRC | Colorectal Cancer |  |
| CSV | Comma-Separated Values |  |
| CT | Computed Tomography |  |
| DAGI | the challenge of Detection of Abnormalities in Gastroscopic Images |  |
| DCNN | Deep Convolution Neural Network |  |
| DEL | Deliverable |  |
| DSC | Dice Similarity Coefficient |  |
| EBUS | Endobronchial Ultrasound |  |
| EGC | Early Gastric Cancer |  |
| EUS | Endoscopic Ultrasound |  |
| FDA | Food and Drug Administration |  |
| FGAI4H | Focus Group on AI for Health |  |
| GDP | Gross Domestic Product |  |
| GDPR | General Data Protection Regulation |  |
| GIANA | Gastrointestinal Image Analysis |  |
| GIST | Gastrointestinal Tumour |  |
| GPU | Graphics Processing Unit |  |
| HD | Hausdorff Distance/High Definition |  |
| IMDRF | International Medical Device Regulators Forum |  |
| IoU | Intersection over Union |  |
| IP | Intellectual Property |  |
| ISBI | IEEE International Symposium on Biomedical Imaging |  |
| ISO | International Organization for Standardization |  |
| ITU | International Telecommunication Union |  |
| LMIC | Low- and middle-Income Countries |  |
| mAP | mean Average Precision |  |
| MCC | Matthews Correlation Coefficient |  |
| MDR | Medical Device Regulation |  |
| MICCAI | international conference on Medical Image Computing and Computer Assisted Intervention |  |
| ML | Machine Learning |  |
| MRI | Magnetic Resonance Imaging |  |
| NP | Normal Pancreas |  |
| NPV | Negative Predictive Value |  |
| PD | Pancreatic duct Drainage |  |
| PDAC | Pancreatic Ductal Carcinoma |  |
| PII | Personally Identifiable Information |  |
| PPV | Positive Predictive Value |  |
| PTBD | Percutaneous Transhepatic Biliary Drainage |  |
| ROC | Receiver Operating Characteristic curve |  |
| SaMD | Software as a Medical Device |  |
| SVM | Support Vector Machine |  |
| TDD | Topic Description Document |  |
| TBNA | Transbronchial Needle Aspiration |  |
| TG | Topic Group |  |
| WG | Working Group |  |
| WHO | World Health Organization |  |

Annex B  
  
Declaration of conflicts of interest

The contributors declare that they have no conflicts of interest.

Tencent Healthcare (Shenzhen) Co., Ltd

Harnessing the technical capabilities, Tencent Healthcare aims to promote innovation in technologies, applications and cooperation models in the healthcare sector. Through upstream and downstream partnerships, Tencent strives to strengthen the digital capabilities for the industry, resulting in improved medical services, enhanced diagnostic efficiency, and ultimately leading to a new digital healthcare ecosystem. Tencent Healthcare encompasses Medical AI diagnosis, Smart Hospital, and Tencent Medipedia, offering comprehensive, convenient, precise and efficient medical and healthcare services to the public.

The China Academy of Information and Communications Technology

Founded in 1957, the China Academy of Information and Communications Technology (hereinafter referred to as CAICT) is a scientific research institute directly under the Ministry of Industry and Information Technology (MIIT) of China. Committed to "the think-tank and enabler for innovation and development in an information society", CAICT has provided strong support for major strategy, plan, policy, test, and certification for the development of the national ICT sector and the IT application, thus proving itself an important facilitator in the leapfrog development and innovation of China's information and communications sector, playing an important role in international cooperation related to the ICT sector and the integration of industrialization and informatization.

Olympus Medical Systems Corp.

At Olympus Medical Systems, we focus on improving patient care quality every day. We do this through developing and designing world-leading, clinically advanced, precision technologies and services. Our products enable healthcare professionals, from a broad range of specialties, to 'peer' inside the body, using endoscopic procedures. This allows them to see more and do more. By focusing on early detection and minimally invasive treatment of a broad range of diseases, our mutual mission is to improve patient outcomes, minimize discomfort, and accelerate the recovery process. Our innovative technologies and services can also optimize workflow and maximize operational efficiency.

Suzhou Institute of Biomedical Engineering and Technology, Chinese Academy of Sciences

Suzhou Institute of Biomedical Engineering and Technology (SIBET), Chinese Academy of Sciences (CAS) is the only institute for research and development of biomedical instruments in CAS. To meet the significant needs in biomedical products, we focused on the basic, strategic, prospective researches in advanced biomedical instruments, reagents and biomedical materials, stimulated the development of biomedical engineering technology, established a platform for the innovation and transformation of medical instruments. Its main research fields cover medical optics, biomedical diagnostics, and rehabilitation technology.

China Unicom (Guangdong) Industrial Internet Co., Ltd

China Unicom (Guangdong) Industrial Internet Co., Ltd. is the first subsidiary with independent legal personality established by China Unicom in Guangdong Province. The company is positioned as an industrial Internet innovation service provider, with the mission of "industrial Internet expert", integrates innovative techniques such as big data, cloud computing, Internet of Things, artificial intelligence, data security, etc., and empowers thousands of industries. Up to now, the company has served more than 1,000 enterprises and more than 200 government units, promoting regional economic development.

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