

RadGenome-Chest CT: A Grounded Vision-Language Dataset for Chest CT Analysis.

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ABSTRACT

Developing generalist foundation model has recently attracted tremendous attention among researchers in the field of AI for Medicine (AI4Medicine). A pivotal insight in developing these models is their reliance on dataset scaling, which emphasizes the requirements on developing open-source medical image datasets that incorporate diverse supervision signals across various imaging modalities. In this paper, we introduce **RadGenome-Chest CT**, a comprehensive, large-scale, region-guided 3D chest CT interpretation dataset based on CT-RATE. Specifically, we leverage the latest powerful universal segmentation and large language models, to extend the original datasets (over **25,692** non-contrast 3D chest CT volume and reports from **20,000** patients) from the following aspects: (i) organ-level segmentation masks covering **197** categories, which provide intermediate reasoning visual clues for interpretation; (ii) **665 K** multi-granularity grounded reports, where each sentence of the report is linked to the corresponding anatomical region of CT volume in the form of a segmentation mask; (iii) **1.3 M** grounded VQA pairs, where questions and answers are all linked with reference segmentation masks, enabling models to associate visual evidence with textual explanations. All grounded reports and VQA pairs in the validation set have gone through manual verification to ensure dataset quality. We believe that **RadGenome-Chest CT** can significantly advance the development of multimodal medical foundation models, by training to generate texts based on given segmentation regions, which is unattainable with previous relevant datasets. We will release all segmentation masks, grounded reports, and VQA pairs to facilitate further research and development in this field.

Background & Summary

In the recent literature, the evolution of large-scale foundation models^{1–5} has sparked significant interest in the development of generalist medical AI (GMAI) systems^{6–10}, particularly within the realm of radiology—a crucial component of medical diagnostics. By training on large-scale visual-language medical datasets, *i.e.*, medical scans paired with global clinical reports, for example, MIMIC-CXR¹¹ has chest X-ray scans from 227,835 studies, and CT-RATE¹² contains chest CT scans from 20,000 patients. These medical models have demonstrated the preliminary ability for writing clinical reports, aiming to support radiologists throughout their workflow and markedly reducing workloads.

However, existing datasets only provide global reports for the medical scan, which has posed limitations on training models that enables grounded report generation, grounded question answering, *i.e.*, describing regional abnormalities and relevant normal findings, or answer questions corresponding to certain regions. To further push forward the training of more capable generalist models, we propose to extend the existing image-reports datasets with region-wise description, *i.e.*, linking the descriptive labels or findings from diagnostic reports to their corresponding anatomical regions in the images, in the form of segmentation masks for explainability.

In this paper, we introduce **RadGenome-Chest CT**, a comprehensive, large-scale and fine-grained annotated dataset for 3D chest CT interpretation, built upon the publicly available CT-RATE¹². *Initially*, we employ the latest powerful text-prompted universal segmentation model, SAT¹³, to segment primary anatomical targets in the image. *Subsequently*, utilizing large language models and NER models, we break all reports into an anatomically hierarchical structured format, and link the reports’ sentences to visual regions in CT volume. *Finally*, we further generate visual question-answering pairs closely related to the structured report and segmented image, from both region level and case level. In summary, we have extended the original image-report datasets from the following aspects:

- i) Organ-level segmentation masks that covers **197** categories, *i.e.*, all the critical regions existing in clinical CT reports;

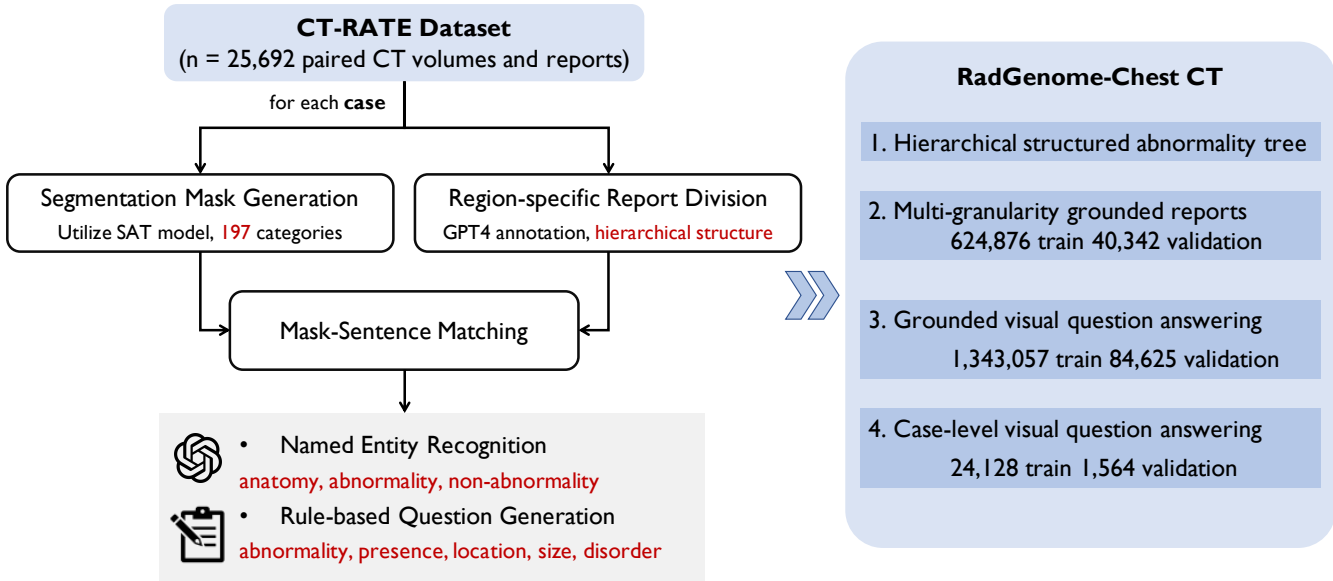


Figure 1. Data construction pipeline of **RadGenome-Chest CT**.

- ii) **665k** multi-granularity grounded reports, with each sentence grounded to the corresponding anatomical region.
- iii) **1.3 M** grounded VQA pairs, concerning both critical region-wise findings and comprehensive case-wise impressions. All the questions and answers are linked to segmentation masks for reference.

We believe that **RadGenome-Chest CT**, with provided region-to-report associations, can significantly advance the development of agent-based multimodal medical foundation models, that enables to generate texts, grounded on the corresponding visual regions, which is unattainable with previous relevant datasets.

Methods

In this section, we start by introducing the source dataset that **RadGenome-Chest CT** is built on. Next, we provide a detailed description of the collection procedure for obtaining segmentation masks, and region-wise reports, as illustrated in Fig. 1. The outcomes from each step are presented in Fig. 2.

Data Source

We initiate our study with CT-RATE¹² (<https://huggingface.co/datasets/ibrahimhamamci/CT-RATE>), it is a dataset of 25,692 non-contrast 3D chest CT volumes derived from 21,304 unique patients, each volume is accompanied by a radiology text report and annotated with 18 distinct types of abnormalities. These 25,692 non-contrast 3D chest CT volumes have been reconstructed with various methods to accommodate different window settings, totaling 50,188 images. For consistency in this paper, we have standardized all CT volumes to a uniform voxel spacing of $1 \times 1 \times 3\text{mm}$, resulting in only 25,692 paired CT volumes and reports. We follow the official division: 20,000 patients (24,128 volumes) were allocated to training and 1,304 (1,564 volumes) for validation.

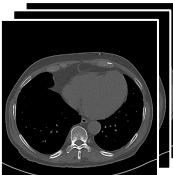
Constructing RadGenome-Chest CT

The pipeline consists of three major stages, as shown in Fig. 1: (i) **segmentation mask generation**, where detailed masks for each anatomical region in the chest CT volumes are created; (ii) **region-specific report division**, that involves the annotation and categorization of radiology text reports by the anatomical regions they refer; (iii) **rule-based question generation**, which entails extracting entities from the sentence, and formulating visual question answering (VQA) pairs linked to specific segmentation masks.

Segmentation Mask Generation

To segment as many anatomical regions as possible, we employ the recent SAT¹³ model. It is a knowledge-enhanced segmentation model, that employs natural language as prompts to effectively segment 3D medical volumes. The model has been trained on 72 diverse segmentation datasets, covering 498 classes across various anatomical regions including the

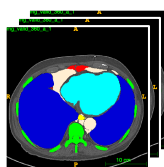
Case from CT-RATE Dataset



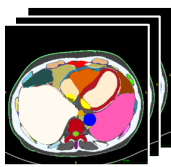
Findings: As far as can be observed: Heart contour and size are normal. There are atheromatous plaques in the aorta and coronary arteries. The widths of the mediastinal main vascular structures are normal. There are no pathologically enlarged lymph nodes in the mediastinum and hilar regions. No pleural or pericardial effusion was detected. There is no pathological wall thickness increase in the esophagus within the sections. Trachea and both main bronchi are open. No occlusive pathology was detected in the trachea and ...

Impression: Hypodense lesions found to be prostate ca, liver metastases during follow-up. Atherosclerotic changes in the aorta and coronary arteries. Emphysematous changes and atelectasis in both lungs. Millimetric nodules in both lungs.

Segmentation Mask Generation



9 regions



197 anatomies

Region-specific Report Division

Lung: There are minimal emphysematous changes and occasional linear atelectasis in both lungs...

Trachea and bronchie: Trachea and both main bronchi are open.

Pleura: No pleural or pericardial effusion was detected.

Mediastinum: The widths of the mediastinal main vascular structures are normal.

Heart: As far as can be observed: Heart contour and size are normal.

Esophagus: There is no pathological wall thickness increase in the esophagus...

Bone: "Vertebral corpus heights, alignments and densities within the sections are normal.

Thyroid: Thyroid appears normal in size, shape, and echotexture.

Abdomen: Hypodense lesions were observed in both lobes of the liver. The...

Name Entity Recognition

Finding sentence: Hypodense lesions were observed in both lobes of the liver.

Region: Abdomen -> Liver

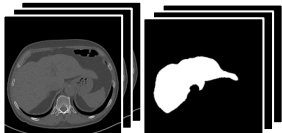
Entities: Anatomy: lobes of the liver. Abnormality: hypodense lesions

Impression sentence: Hypodense lesions found to be prostate ca, liver metastases during follow-up. Atherosclerotic changes in the aorta and coronary arteries. Emphysematous changes and atelectasis in both lungs. Millimetric nodules in both lungs.

Entities: prostate cancer, liver metastases, atherosclerosis, coronary artery disease, atelectasis, pulmonary emphysema, pulmonary nodules.

Rule-based Question Generation

I. Grounded visual question answering pairs



Input: <image> + <mask> What abnormality is seen in the liver?

Output: Hypodense lesions were observed in both lobes of the liver.

2. Case-level visual question answering pairs

Input: <image> What disorder can be identified from this volume?

Output: Prostate cancer, liver metastases, atherosclerosis, coronary artery disease, atelectasis,....

Figure 2. Overview of results obtained from each step of the data construction pipeline.

brain, head and neck, thorax, spine, abdomen, and limbs. For our research, SAT is adopted to execute detailed segmentation across all volumes of the CT-RATE dataset. Specifically, we focus on segmenting 197 regions pertinent to chest CT scans, enabling precise anatomical analysis. The list of segmented anatomies was organized into a hierarchical tree as shown in Supplementary B. This includes several major regions such as the lungs, trachea and bronchi, mediastinum, heart, pleura, bones, thyroid, breasts, abdomen, and other areas.

Region-wise Reports Generation

The goal here is to break the entire reports into region-wise descriptions, we observe that the OpenAI GPT-4² model can complete such task in very high accuracy^{14,15}. However, employing GPT-4 on the entire set would be prohibitively expensive, thus, we train a model for such report division. Specifically, we first employ GPT-4² to annotate the anatomical regions of each sentence in the “**FINDINGS**” section of 2,500 radiology reports, which comprise all reports in the validation set. The prompt

used is as follows.

You are a radiologist tasked with extracting anatomical regions from the “**FINDINGS**” section of radiology reports. For each sentence provided, identify the corresponding anatomical regions. Ensure each identified region is an entry from a predefined list: [region_list]. If a sentence mentions “left” or “right”, these qualifiers should precede the anatomical region (e.g., left kidney). Given input in the format: <Input><findings><\Input>. Please reply in the following JSON format: {<sentence>: [region₁,region₂,...], <sentence>: [region₁]}.

This process results in 15,926 annotated sentences. Subsequently, we divide these sentences into training and validation subsets in an 8:2 ratio and train a GPT-2 model using the annotated sentence along with the two preceding and following sentences from the report as input, if available. The model is designed to output the list of anatomical regions associated with each sentence. For instance, for the target sentence “No pleural effusion was detected on the left.”, the input is “There is minimal pleural effusion on the right. No pleural effusion was detected on the left. Atelectasis is observed in the middle lobe and lower lobe of the right lung. A malignant mass is observed around the lower lobe bronchi of the left lung.”, and the expected output is “left lung”. The model achieves an accuracy of **94.56%** on the validation set. Consequently, we employ this model to perform inference on all sentences across the entire dataset of reports. More examples of segmentation results and structured reports can be seen in Supplementary Section A.

Named Entity Recognition

Through the abovementioned process, each “**FINDINGS**” is divided into multiple sentences, and each is associated with one or multiple segmented regions. To facilitate the generation of detailed question-answer pairs from these sentences, we initially employ an in-house Named-Entity Recognition (NER) model to analyze all sentences. This process involves extracting entities that can be categorized into “anatomy”, “abnormality”, and “non-abnormality”. Here, “anatomy” pertains to the anatomical regions, “abnormality” refers to findings or diseases identified as present, and “non-abnormality” indicates findings or diseases that are reported as absent. Subsequently, all extracted “abnormality” and “non-abnormality” entities undergo quality evaluation using GPT-4, allowing us to filter out and revise any inaccuracies. For instance, in cases where the NER model extracts abnormalities such as “structural distortion and volume loss”, GPT-4 will segment it into “structural distortion” and “volume loss” for more accurate categorization. We filter out abnormalities with a GPT-4 output of “no” and update them to the revised versions provided by GPT-4. The prompt used is as follows.

You are a radiologist. You will receive an input that consists of a sentence from the findings section of a chest CT report, followed by a phrase extracted from that sentence. Assess whether the extracted phrase is an abnormality. The abnormality should not be a size or a location. If the extracted phrase does not accurately represent an abnormality, your output should provide the correct abnormalities. The input format will be <Input><sentence><sep><phrase></Input>. Your response should be in JSON format, structured as follows: {abnormality: [abnormality1, abnormality2,...], phrase: “yes” or “no”}. Note that the output should be in JSON format.

In addition, for the “**IMPRESSION**” section, we directly utilize GPT-4 to extract all disorders mentioned, and detailed information regarding the presence of any abnormalities in specific anatomical regions. The prompt used is as follows.

You are a radiologist performing disorders summary from the “**IMPRESSION**” section in the patient’s chest CT scan report. Given input in the format: <Input><impression><\Input> Please reply with the JSON format following template: {disorders:[disorder₁,disorder₂,...]}

Rule-based Question Generation

Here, we describe the procedure for generating grounded visual question-answering (VQA) data based on “**FINDINGS**” extracted from the report section, while case-level visual question-answering data is generated from the “**IMPRESSION**” section. Tab. 1 presents the 5 question types in the proposed dataset. The detailed rules will be introduced in the following sections.

First, after the region-wise report generation, the sentences in the findings section can be classified as follows:

- **Normal Findings:** Sentences that report no significant changes from normal health conditions.
 - **No abnormality entities** in the sentence: Sentences that mention specific anatomical regions without noting any abnormalities. For example, “Thoracic aorta diameter is normal.”

Level	Question type	Answer type	Example	Train	Validation
Region	Abnormality	Open	What abnormality is seen in the {region}?	483,507	30,831
Region	Presence	Close	Is there evidence of {abnormality} in the {region}?	556,050	34,785
Region	Location	Close & Open	Where in the image is the {abnormality} seen?	285,111	17,750
Region	Size	Open	What is the size of the {abnormality} in the {region}?	18,389	1,259
Case	Disorder	Open	What disorder can be identified from this volume?	24,128	1,564

Table 1. Question and answer types in the proposed datasets. The placeholders {region} and {abnormality} in the question templates are dynamically replaced with entities extracted from sentences.

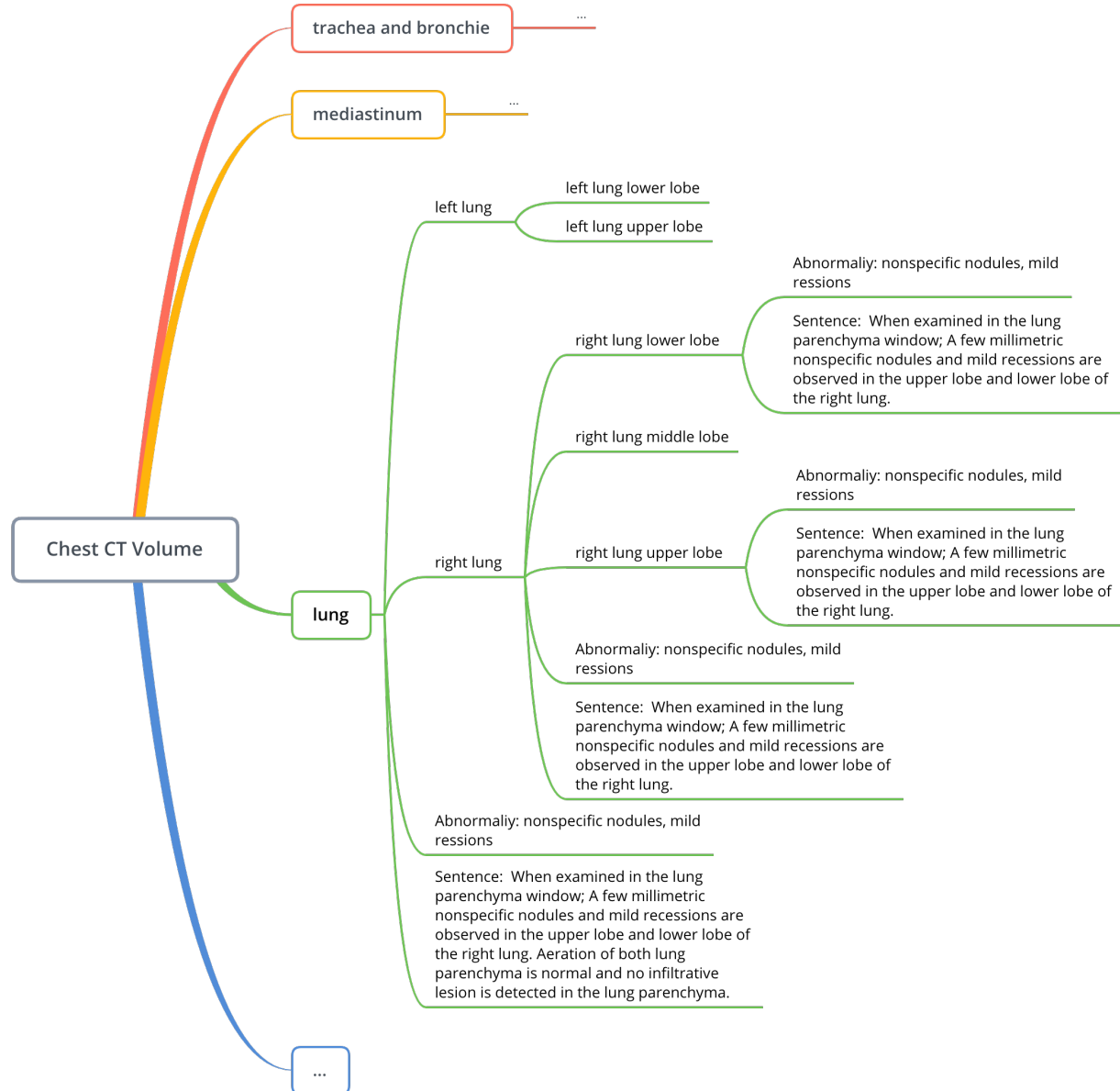


Figure 3. Partial visualization of the anatomical hierarchical tree with abnormalities annotated.

- **No anatomy entities** in the sentence: Sentences that solely note the absence of specific abnormalities without referring to any anatomical regions. For example, “No pleural effusion was detected.”
- **With anatomy and abnormality entities** in the sentence: Sentences that explicitly state the absence of abnormalities in specific anatomical regions. For example, “Bilateral adrenal glands were normal and no space-occupying lesion was detected.”
- **Abnormal Findings:** Sentences that report differences from normal anatomical conditions.
 - **No anatomy entities** in the sentence: Sentences that report an abnormal finding, but do not specify an anatomical region. For example, “Mild hiatal hernia is observed.”
 - **With anatomy entities** in the sentence: Sentences that include both an anatomical reference and describe an abnormality. For example, “There is narrowing of the spinal canal at the dorso-lumbar level.”

We then construct an anatomical disorder tree for each report, based on the anatomical hierarchical tree introduced in Supplementary Section B. This involves marking any abnormalities on the tree for all nodes within the hierarchy if they are present. As shown in Fig. 3 This comprehensive data enables us to transit to generating questions. Taking inspiration from the previous research^{16,17}, we categorize the questions into four types: 1) abnormality, 2) presence, 3) location, 4) size. Tab. 1 shows examples of the different question types. Note that we have designed 50 templates for each question type. The details of all templates are provided in the supplementary materials. For instance, when analyzing a sentence indicating normal findings, such as “Bilateral adrenal glands were normal and no space-occupying lesion was detected.”, questions can include “Is there any evidence of abnormality in adrenal glands?” and “What abnormality is seen in the adrenal glands?”. Conversely, for a sentence in abnormal findings, “There is a narrowing of the spinal canal at the dorso-lumbar level.”, the question can be “Is there any evidence of narrowing in the spinal canal?” “What abnormality is seen at the dorso-lumbar level of the spinal canal?” and “Where in the spinal canal is the narrowing located?”. For impression sentences, since we have already extracted disorders, we can generate case-level questions such as “What disorder can be identified from this volume?” for each case.

In summary, we have successfully generated 1.3M grounded Visual Question Answering (VQA) pairs for training and 85k for validation, along with 24,128 case-level visual question answering pairs for training and 1,564 for validation. Detailed counts for each type of VQA pair in both the training and validation sets are provided in Table 1.

Dataset Analysis

In this section, we analyze the abnormalities of the proposed dataset. The proposed hierarchically structured abnormality tree for each case enables us to systematically extract and analyze the abnormal findings. First, we calculate the normal-to-abnormal case numbers for both the training set and the validation set, as shown in Fig. 4. Second, based on the identified and extracted abnormalities from all anatomical regions, we proceed to visualize these findings using a word cloud. The word cloud of abnormalities is presented in Fig. 4.

Conclusion

In this paper, we develop an automated pipeline for generating grounded datasets and introduce **RadGenome-Chest CT**, a comprehensive, large-scale, region-guided 3D chest CT interpretation dataset based on CT-RATE. **RadGenome-Chest CT** include **197** organ-level segmentation masks, **665 K** multi-granularity grounded reports, and **1.3 M** grounded VQA pairs. We anticipate that **RadGenome-Chest CT** will significantly advance multimodal medical AI models, enabling them to generate texts based on segmentation regions, thus enhancing interpretability and patient care. We will release all segmentation masks, grounded reports, and VQA pairs to support future research in this field.

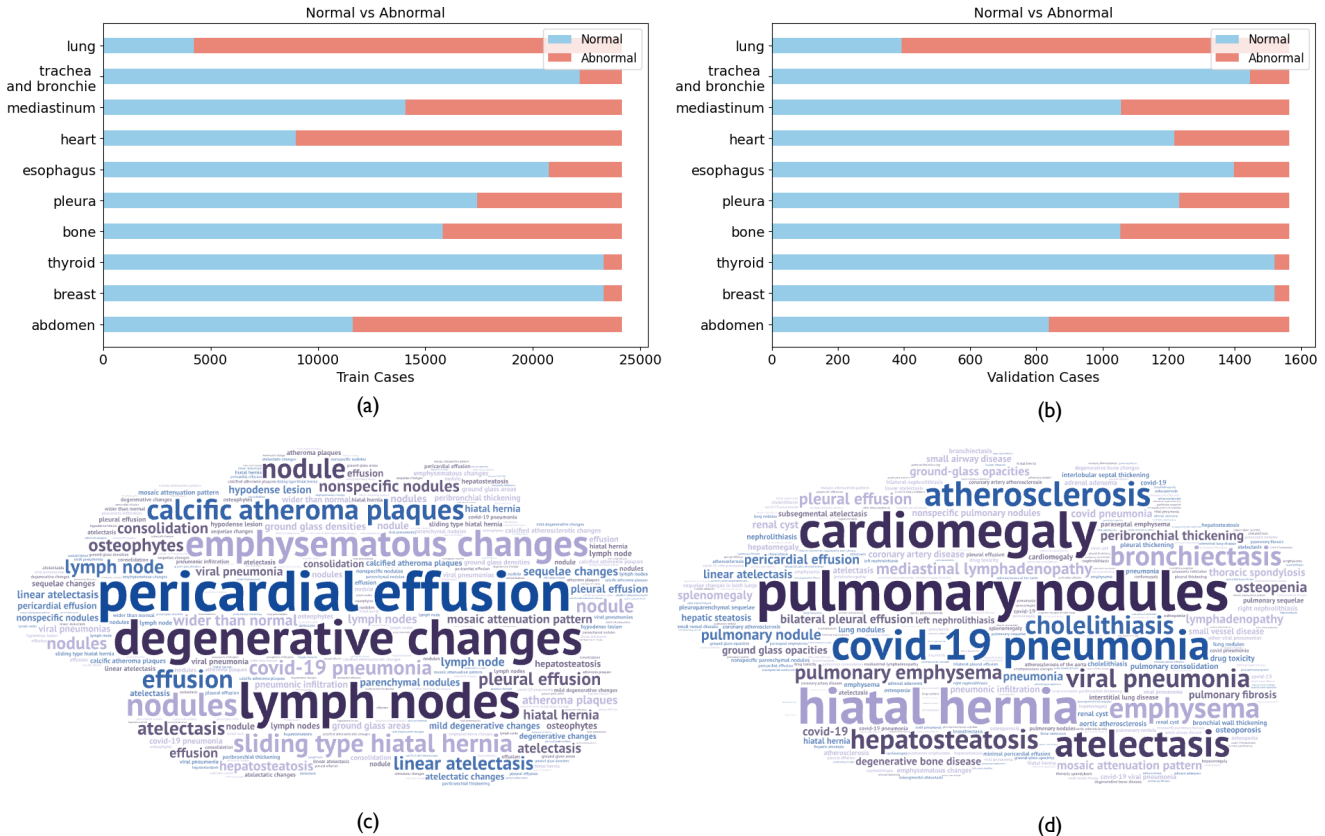
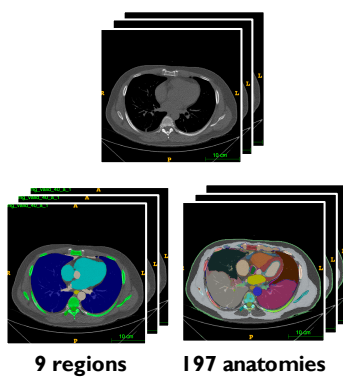


Figure 4. (a) Distribution of Normal vs. Abnormal Cases: Training set analysis. (b) Distribution of Normal vs. Abnormal Cases: Validation set analysis. (c) Abnormalities Word Cloud: Visual summary of frequent abnormalities from findings. (d) Disorders Word Cloud: Visual summary of frequent disorders from findings.

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A Case Examples



Lung: When examined in the lung parenchyma window; Ground-glass densities, which are more prominent in the lower lobes of both lungs, are common and scattered. No nodular lesions were detected in both lung parenchyma.

Trachea and bronchie: Trachea and both main bronchi are open.

Pleura: In addition, subpleural consolidation areas are observed in the right lung lower lobe laterobasal segment and left lung upper lobe apicoposterior segment.

Mediastinum: Mediastinal main vascular structures, heart contour, size are normal.

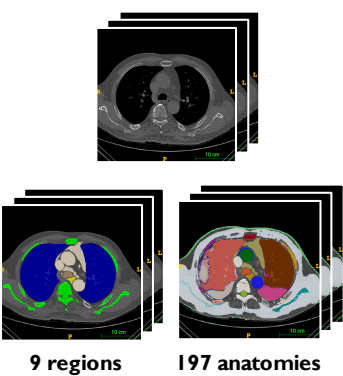
Heart: Pericardial effusion-thickening was not observed.

Esophagus: Thoracic esophageal calibration was normal and no significant tumoral wall thickening was detected.

Bone: Bone structures in the study area are natural. Vertebral corpus heights are preserved.

Thyroid: Thyroid appears normal in size, shape, and echotexture.

Abdomen: Thoracic aorta diameter is normal. Upper abdominal organs included in the sections are normal. No space-occupying lesion was detected in the liver that entered the cross-sectional area. Bilateral adrenal glands were normal and no space-occupying lesion was detected.



Lung: No lymph node was detected in the mediastinum and in both hilar levels in pathological size and configuration. When examined in the lung parenchyma window; Both hemithorax ...

Trachea and bronchie: The calibration of the trachea and main bronchi is normal and their lumens are clear.

Pleura: Bilateral pleural effusion pneumothorax was not detected.

Mediastinum: Calibration of other major vascular structures in the mediastinal is natural.

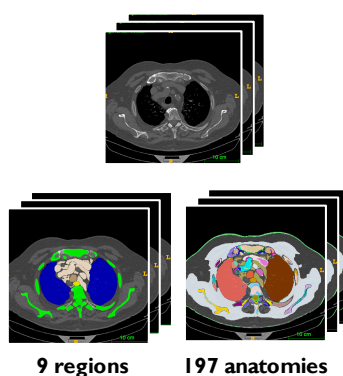
Heart: Heart contour and size are normal.

Esophagus: Thoracic esophagus calibration was normal and no significant tumoral wall thickening was detected. Mild hiatal hernia is observed.

Bone: Degenerative changes are observed in the bone structures in the study area. There is narrowing of the spinal canal at the dorso- lumbar level.

Thyroid: Thyroid appears normal in size, shape, and echotexture.

Abdomen: Calibration of the aortic arch is at the maximal physiological limit. There are bilateral irregular density increases in the perinephric areas. A decrease in density is observed in the liver, ...



Lung: When examined in the lung parenchyma window; Parenchymal fibrosis areas causing structural distortion in both lungs, emphysematous changes, prominence in interlobular septa and honey...

Trachea and bronchie: Trachea and lumen of both main bronchi are open. No occlusive pathology was detected in the trachea and lumen of both main bronchi.

Pleura: Bilateral pleural thickening-effusion was not detected.

Mediastinum: Mediastinal structures were evaluated as suboptimal since the examination was unenhanced. The diameter of the main pulmonary artery was 43 mm, the diameter of ...

Heart: Heart size increased.

Esophagus: Thoracic esophagus calibration was normal and no significant pathological wall thickening was detected.

Bone: Thoracic kyphosis has increased. Degenerative changes were observed in bone structures.

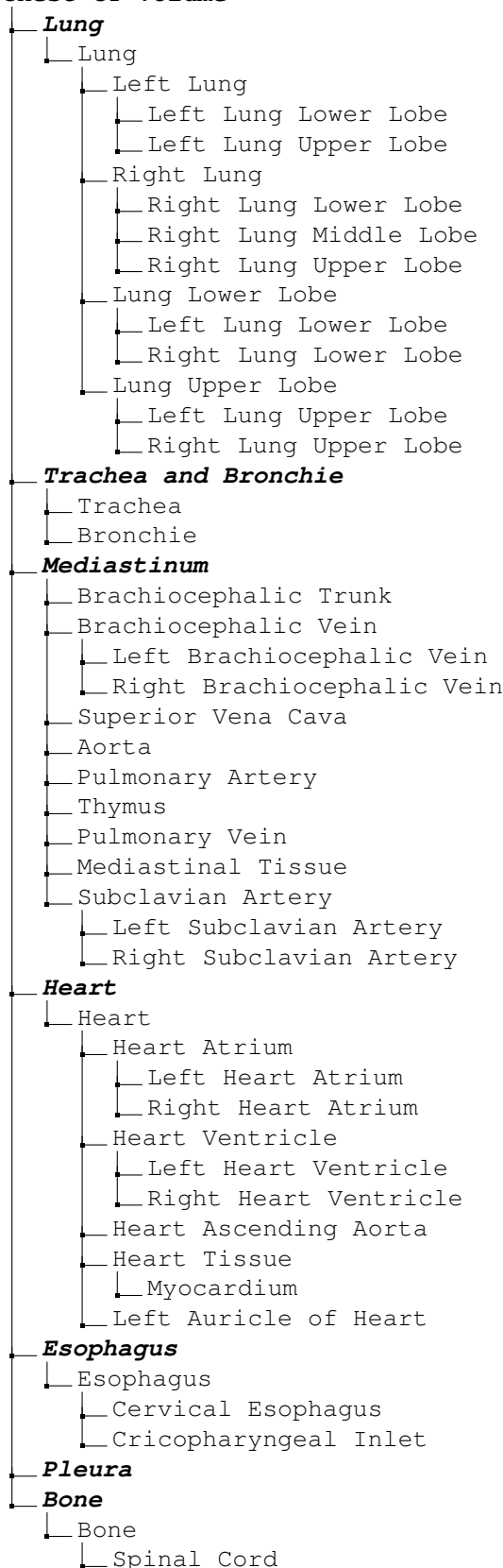
Thyroid: As far as can be seen; A 32 mm diameter hypodense nodule was observed in the right thyroid lobe.

Abdomen: Calcific atherosclerotic changes were observed in the thoracic aorta and ...

Figure 5. Case examples of segmentation masks and structured reports.

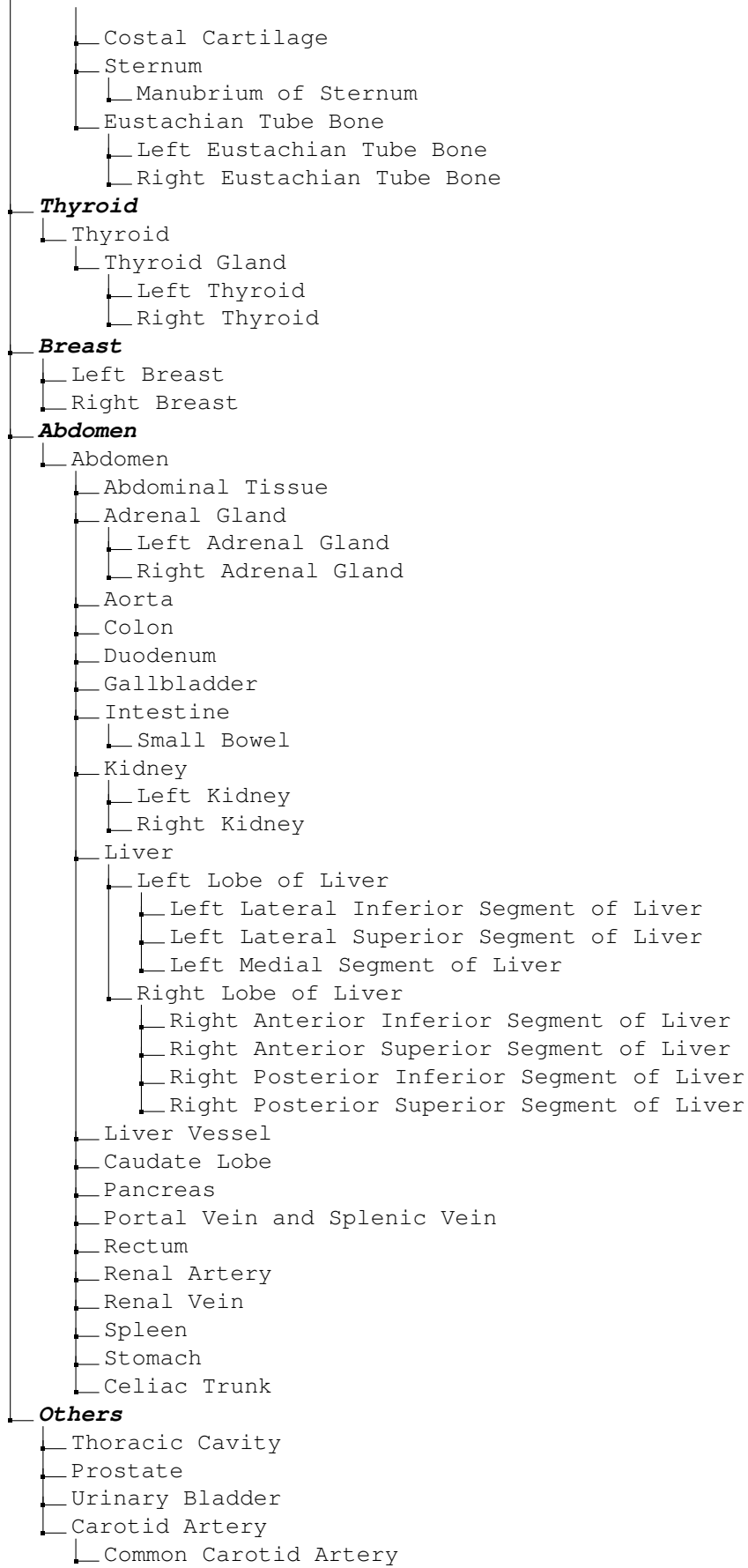
B Hierarchy of anatomical regions

Chest CT Volume



- Spinal Canal
- Vertebrae
 - Cervical Vertebrae
 - Cervical Vertebrae 1 (C1)
 - Cervical Vertebrae 2 (C2)
 - Cervical Vertebrae 3 (C3)
 - Cervical Vertebrae 4 (C4)
 - Cervical Vertebrae 5 (C5)
 - Cervical Vertebrae 6 (C6)
 - Cervical Vertebrae 7 (C7)
 - Thoracic Vertebrae
 - Thoracic Vertebrae 1 (T1)
 - Thoracic Vertebrae 2 (T2)
 - Thoracic Vertebrae 3 (T3)
 - Thoracic Vertebrae 4 (T4)
 - Thoracic Vertebrae 5 (T5)
 - Thoracic Vertebrae 6 (T6)
 - Thoracic Vertebrae 7 (T7)
 - Thoracic Vertebrae 8 (T8)
 - Thoracic Vertebrae 9 (T9)
 - Thoracic Vertebrae 10 (T10)
 - Thoracic Vertebrae 11 (T11)
 - Thoracic Vertebrae 12 (T12)
 - Lumbar Vertebrae
 - Lumbar Vertebrae 1 (L1)
 - Lumbar Vertebrae 2 (L2)
 - Lumbar Vertebrae 3 (L3)
 - Lumbar Vertebrae 4 (L4)
 - Lumbar Vertebrae 5 (L5)
 - Lumbar Vertebrae 6 (L6)
 - Sacral Vertebrae 1 (S1)
- Clavicle
 - Left Clavicle
 - Right Clavicle
- Scapula
 - Left Scapula
 - Right Scapula
- Humerus
 - Left Humerus
 - Right Humerus
- Femur
 - Left Femur
 - Right Femur
- Head of Femur
 - Left Head of Femur
 - Right Head of Femur
- Rib
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 - Left Rib 2
 - Left Rib 3
 - Left Rib 4
 - Left Rib 5
 - Left Rib 6
 - Left Rib 7

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    |       Right Rib 8
    |
    |   Rib 9
    |       Left Rib 9
    |       Right Rib 9
    |
    |   Rib 10
    |       Left Rib 10
    |       Right Rib 10
    |
    |   Rib 11
    |       Left Rib 11
    |       Right Rib 11
    |
    |   Rib 12
    |       Left Rib 12
    |       Right Rib 12
    |
    |   Rib Cartilage
```



- └─ Internal Carotid Artery
- └─ Left Carotid Artery
 - └─ Left Common Carotid Artery
 - └─ Left Internal Carotid Artery
- └─ Right Carotid Artery
 - └─ Right Common Carotid Artery
 - └─ Right Internal Carotid Artery
- └─ Iliac Artery
- └─ Iliac Vena
- └─ Iliac Vein
- └─ Left Iliac Artery
- └─ Left Iliac Vena
- └─ Right Iliac Artery
- └─ Right Iliac Vena
- └─ Inferior Vena Cava
- └─ Internal Jugular Vein
- └─ Larynx
 - └─ Larynx Glottis
 - └─ Larynx Supraglottis
- └─ Muscle

C Question template

Abnormality

1. What are the abnormalities in the {region}?
2. What abnormalities are present in the {region}?
3. What types of abnormality are visible in the {region}?
4. What type of abnormality is visible in the {region}?
5. What kind of abnormalities are observed in the {region}?
6. What kinds of abnormalities can be identified in the {region}?
7. What specific abnormalities are detected in the {region}?
8. What specific types of abnormalities are evident in the {region}?
9. What types of abnormalities are evident upon examination of the {region}?
10. What types of abnormalities can be seen in the {region}?
11. What are the anomalies in the {region}?
12. What anomalies are present in the {region}?
13. What types of anomalies are visible in the {region}?
14. What type of anomalies is visible in the {region}?
15. What kind of anomalies are observed in the {region}?
16. What kinds of anomalies can be identified in the {region}?
17. What specific anomalies are detected in the {region}?
18. What specific types of anomalies are evident in the {region}?
19. What types of anomalies are evident upon examination of the {region}?
20. What types of anomalies can be seen in the {region}?
21. What are the abnormal findings in the {region}?
22. What abnormal findings are present in the {region}?
23. What types of abnormal findings are visible in the {region}?
24. What type of abnormal findings is visible in the {region}?
25. What kind of abnormal findings are observed in the {region}?
26. What kinds of abnormal findings can be identified in the {region}?
27. What specific abnormal findings are detected in the {region}?
28. What specific types of abnormal findings are evident in the {region}?
29. What types of abnormal findings are evident upon examination of the {region}?
30. What types of abnormal findings can be seen in the {region}?
31. What are the irregular findings in the {region}?
32. What irregular findings are present in the {region}?

33. What types of irregular findings are visible in the {region}?
34. What type of irregular findings is visible in the {region}?
35. What kind of irregular findings are observed in the {region}?
36. What kinds of irregular findings can be identified in the {region}?
37. What specific irregular findings are detected in the {region}?
38. What specific types of irregular findings are evident in the {region}?
39. What types of irregular findings are evident upon examination of the {region}?
40. What types of irregular findings can be seen in the {region}?
41. What are the irregularities in the {region}?
42. What irregularities are present in the {region}?
43. What types of irregularities are visible in the {region}?
44. What type of irregularities is visible in the {region}?
45. What kind of irregularities are observed in the {region}?
46. What kinds of irregularities can be identified in the {region}?
47. What specific irregularities are detected in the {region}?
48. What specific types of irregularities are evident in the {region}?
49. What types of irregularities are evident upon examination of the {region}?
50. What types of irregularities can be seen in the {region}?

Presence

1. Can {abnormality} be identified in the {region}?
2. Can {abnormality} be observed in the {region}?
3. Can {abnormality} be detected in the {region}?
4. Can {abnormality} be seen in the {region}?
5. Can {abnormality} be founded in the {region}?
6. Can {abnormality} be recognized in the {region}?
7. Can we detect any signs of {abnormality} in the {region}?
8. Can we observe any signs of {abnormality} in the {region}?
9. Can we recognize any signs of {abnormality} in the {region}?
10. Can we see {abnormality} in the {region}?
11. Can we find {abnormality} in the {region}?
12. Can we detect {abnormality} in the {region}?
13. Can we observe {abnormality} in the {region}?
14. Can we identify {abnormality} in the {region}?
15. Is there any sign of {abnormality} in the {region}?

16. Is there any indication of {abnormality} in the {region}?
17. Is there any evidence of {abnormality} in the {region}?
18. Is there any suggestion of {abnormality} in the {region}?
19. Is there a clear sign of {abnormality} in the {region}?
20. Is there a clear indication of {abnormality} in the {region}?
21. Is there a clear evidence of {abnormality} in the {region}?
22. Is there a clear suggestion of {abnormality} in the {region}?
23. Is {abnormality} visibly present in the {region}?
24. Is {abnormality} clearly visible in the {region}?
25. Is there any visual evidence suggesting {abnormality} in the {region}?
26. Is there any indication of {abnormality} upon examination of the {region}?
27. Is there any indication of {abnormality} in the {region}?
28. Is there visual evidence of {abnormality} in the {region} on this scan?
29. Are there any visible indications of {abnormality} in this {region}?
30. Are there any visible cues indicating {abnormality} in the {region}?
31. Are there any visible indicators of {abnormality} in the {region}?
32. Are there any clear indications of {abnormality} in this {region}?
33. Are there any clear cues indicating {abnormality} in the {region}?
34. Are there any clear indicators of {abnormality} in the {region}?
35. Are there any indications of {abnormality} in this {region}?
36. Are there any cues indicating {abnormality} in the {region}?
37. Are there any indicators of {abnormality} in the {region}?
38. Are there any observable signs of {abnormality} in the {region}?
39. Are there any signs of {abnormality} in the {region}?
40. Are there any features of {abnormality} in the {region}?
41. Does the {region} show the presence of {abnormality}?
42. Does the {region} show the existence of {abnormality}?
43. Does the image suggest the presence of {abnormality} in the {region}?
44. Does the image suggest the existence of {abnormality} in the {region}?
45. Does the {region} exhibit any evidence of {abnormality}?
46. Does the {region} display any features suggestive of {abnormality}?
47. Does the {region} display any characteristics suggestive of {abnormality}?
48. Does the {region} exhibit any characteristics indicative of {abnormality}?
49. Does the {region} exhibit any features indicative of {abnormality}?
50. Does the visual features suggest the presence of {abnormality} in the {region}?

Location

1. Where is the {abnormality} located in the image?
2. Where can the {abnormality} be found within the image?
3. Where in the image is the {abnormality} located?
4. Where in the image is the {abnormality} localized?
5. Where in the image can the {abnormality} be found?
6. Where in the image does the {abnormality} appear?
7. Where in the image does the {abnormality} locate?
8. Where in the image does the {abnormality} locate?
9. Where specifically within the image is the {abnormality} located?
10. Where exactly within the image is the {abnormality} located?
11. Where exactly is the {abnormality} located in the image?
12. Where specifically is the {abnormality} located in the image?
13. Where exactly within the image is the {abnormality} localized?
14. Where specifically within the image is the {abnormality} localized?
15. Where within the image can the {abnormality} be precisely located?
16. Where exactly within the image does the {abnormality} present?
17. Where within the image does the {abnormality} specifically present?
18. Where in the image does the {abnormality} appear?
19. What is the location of the {abnormality} in the image?
20. What is the precise location of the {abnormality} in the image?
21. What is the specific location of the {abnormality} within the image?
22. What is the precise region of the {abnormality} in the image?
23. What is the specific region of the {abnormality} within the image?
24. What particular region within the image does the {abnormality} occupy?
25. What particular location within the image does the {abnormality} occupy?
26. What specific location within the image does the {abnormality} occupy?
27. What specific region within the image does the {abnormality} occupy?
28. What specific area of the image does the {abnormality} occupy?
29. What specific region of the image does the {abnormality} appear?
30. What specific spot within the image contains the {abnormality}?
31. What particular region of the image is affected by the {abnormality}?
32. What specific area within the image is impacted by the {abnormality}?
33. What specific region within the image is impacted by the {abnormality}?

34. What specific location within the image is impacted by the {abnormality}?
35. What particular region within the image is affected by the {abnormality}?
36. What particular area within the image is affected by the {abnormality}?
37. What particular location within the image is affected by the {abnormality}?
38. What specific region within the image does the {abnormality} affect?
39. What specific area within the image does the {abnormality} affect?
40. What specific location within the image does the {abnormality} affect?
41. What specific location within the image does the {abnormality} appear?
42. What specific region within the image does the {abnormality} appear?
43. What specific area within the image does the {abnormality} appear?
44. What particular spot within the image does the {abnormality} present?
45. What particular area within the image does the {abnormality} present?
46. What particular region within the image does the {abnormality} present?
47. What particular location within the image does the {abnormality} present?
48. What specific area within the image does the {abnormality} occur?
49. What specific location within the image does the {abnormality} occur?
50. What specific region within the image does the {abnormality} occur?

Size

1. What is the approximate size of the {abnormality} in the {region}?
2. What is the approximate scale of the {abnormality} in the {region}?
3. What is the approximate size range of the {abnormality} in the {region}?
4. What is the approximate magnitude of the {abnormality} in the {region}?
5. What is the approximate dimension of the {abnormality} in the {region}?
6. What is the approximate measurement of the {abnormality} in the {region}?
7. What is the estimated size of the {abnormality} in the {region}?
8. What is the estimated scale of the {abnormality} in the {region}?
9. What is the estimated size range of the {abnormality} in the {region}?
10. What is the estimated magnitude of the {abnormality} in the {region}?
11. What is the estimated dimension of the {abnormality} in the {region}?
12. What is the estimated measurement of the {abnormality} in the {region}?
13. What is the size assessment of the {abnormality} in the {region}?
14. What is the measurement of the {abnormality} in the {region}?
15. What is the scale of the {abnormality} in the {region}?
16. What is the size of the {abnormality} in the {region}?

17. What is the size range of the {abnormality} in the {region}?
18. What is the magnitude of the {abnormality} in the {region}?
19. What is the dimension of the {abnormality} in the {region}?
20. What is the overall size of the {abnormality} in the {region}?
21. What is the overall scale of the {abnormality} in the {region}?
22. What is the overall measurement of the {abnormality} in the {region}?
23. What is the overall size range of the {abnormality} in the {region}?
24. What is the overall magnitude of the {abnormality} in the {region}?
25. What is the overall dimension of the {abnormality} in the {region}?
26. What is the scale of the {abnormality} detected in the {region}?
27. What is the size of the {abnormality} detected in the {region}?
28. What is the size range of the {abnormality} detected in the {region}?
29. What is the measurement of the {abnormality} detected in the {region}?
30. What is the magnitude of the {abnormality} detected in the {region}?
31. What is the dimension of the {abnormality} detected in the {region}?
32. What is the scale of the {abnormality} appeared in the {region}?
33. What is the size of the {abnormality} appeared in the {region}?
34. What is the size range of the {abnormality} appeared in the {region}?
35. What is the magnitude of the {abnormality} appeared in the {region}?
36. What is the dimension of the {abnormality} appeared in the {region}?
37. What is the measurement of the {abnormality} appeared in the {region}?
38. How large is the affected {abnormality} area in the {region}?
39. How large is the observed {abnormality} area in the {region}?
40. How large is the {abnormality} observed in the {region}?
41. How large is the {abnormality} in the {region}?
42. How large does the {abnormality} appear in the {region}?
43. How large does the {abnormality} appear to be in the {region}?
44. How large is the {abnormality} area in the {region}?
45. How large is the affected {abnormality} area in the {region}?
46. How big is the affected {abnormality} area in the {region}?
47. How big is the observed {abnormality} area in the {region}?
48. How big is the {abnormality} observed in the {region}?
49. How big is the {abnormality} in the {region}?