Multi-label annotation of text reports from computed tomography of the chest, abdomen, and pelvis using deep learning

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Abstract

Background: There is progress to be made in building artificially intelligent systems to detect abnormalities that are not only accurate but can handle the true breadth of findings that radiologists encounter in body (chest, abdomen, and pelvis) computed tomography (CT). Currently, the major bottleneck for developing multi-disease classifiers is a lack of manually annotated data. The purpose of this work was to develop high throughput multi-label annotators for body CT reports that can be applied across a variety of abnormalities, organs, and disease states thereby mitigating the need for human annotation.

Methods: We used a dictionary approach to develop rule-based algorithms (RBA) for extraction of disease labels from radiology text reports. We targeted three organ systems (lungs/pleura, liver/gallbladder, kidneys/ureters) with four diseases per system based on their prevalence in our dataset. To expand the algorithms beyond pre-defined keywords, attention-guided recurrent neural networks (RNN) were trained using the RBA-extracted labels to classify reports as being positive for one or more diseases or normal for each organ system. Alternative effects on disease classification performance were evaluated using random initialization or pre-trained embedding as well as different sizes of training datasets. The RBA was tested on a subset of 2158 manually labeled reports and performance was reported as accuracy and F-score. The RNN was tested against a test set of 48,758 reports labeled by RBA and performance was reported as area under the receiver operating characteristic curve (AUC), with 95% CIs calculated using the DeLong method.

Results: Manual validation of the RBA confirmed 91–99% accuracy across the 15 different labels. Our models extracted disease labels from 261,229 radiology reports of 112,501 unique subjects. Pre-trained models outperformed random initialization across all diseases. As the training dataset size was reduced, performance was robust except for a few diseases with a relatively small number of cases. Pre-trained classification AUCs reached > 0.95 for all four disease outcomes and normality across all three organ systems.

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Background

Machine learning algorithms have demonstrated considerable potential as disease classifiers for medical images. However, the majority of algorithms are specialized for a single organ or disease making their utility narrow in scope. This limited scope is mainly attributed to a sparsity of training data, since curating datasets for image-based classifiers has traditionally relied on radiologist annotation of the disease or its sequelae. As an alternative to image-based labeling, automated extraction of disease labels from radiology report text has the potential to address this training data scarcity and to avoid human annotation efforts [1–4].

Rule-based algorithms (RBA) are a conventional method for mining report text that utilize simple logic based on pre-defined keywords or patterns. In a landmark study, Wang et al. [5] used a RBA to extract labels of 8 thorax diseases from 108,948 chest X-ray reports to effectively train an image-based disease classifier. Using a similar method for CT, Draelos et al. [6] demonstrated the broad applicability of RBA-obtained labels by mining the more complex reports associated with over 36,000 chest CT volumes to train a classifier for 83 chest abnormalities. However, a major limitation of RBAs is that their performance and scope is reliant on the completeness of dictionaries containing pre-defined keywords. Furthermore, the radiologist’s interpretation that accompanies a CT is usually composed in a free or semi-structured text form, rendering the extraction of disease labels using simple logical rules a nontrivial task [7].

To improve their utility, RBA-extracted labels can then be used to train neural networks that deviate from strict rules by learning salient semantic features, a form of natural language processing (NLP) [8, 9]. For example, Steinkamp et al. [10] trained a recurrent neural network (RNN) to classify disease in pathology reports written in unseen formats, suggesting the network had learned a generalizable encoding of the semantics. Building upon this NLP approach, Yuan et al. [11] combined a pre-trained word embedding model with a deep learning-based sentence encoder to classify pulmonary nodules in a diverse set of radiology reports from different universities. While promising, it is often difficult to determine which semantic or structural features of the reports that the model perceives as most salient. To improve the interpretability of NLP-based classifiers, an attention-guided RNN [12] can be used to project the attention vector onto report text [13], allowing the user to visualize the words that a model is giving the most weight to when classifying an abnormality. For example, Banerjee et al. [14] demonstrated that an attention-guided-RNN could be used to visualize synthesized information on pulmonary emboli from thoracic CT free-text radiology reports.

In this study, we propose a framework for automated, multi-disease label extraction of body (chest, abdomen, and pelvis) CT reports based on attention-guided RNNs trained on RBA extracted labels. For each organ system, a RNN was trained to classify the lungs/pleura, liver/gallbladder, kidneys/ureters as being positive for one or more of four different diseases or normal. Although there has been extensive work in radiology report labeling, to our knowledge, there are no related works that demonstrate the utility of an RBA to train deep learning-based NLP disease classifiers in such a breadth of organ systems, diseases, and body CT reports.

The main contributions of this study are threefold:

1. To develop a RBA that can meet the challenges of free-text narration in radiology CT reports.
2. To broaden the scope of our extracted labels by training attention-guided RNNs to perform multi-label disease classification of CT reports.
3. To determine alternative factors that influence disease classification performance including random vs. pre-trained embedding and different sizes of training dataset.

Materials and methods

Institutional Review Board (IRB) approval was obtained, and informed consent was waived for this retrospective study that was compliant with the Health Insurance Portability and Accountability Act. In this section, we first describe the dataset that was used. Then, we outline the development processes of our RBAs and the subsequent addition of attention-guided RNNs to enable multi-label classification of radiology reports. Figure 1 displays the overall workflow of this paper.
This proposed work was a considerable expansion of our two previous conference proceedings manuscripts [15, 16]. Our initial demonstration [15] was focused on the binary classification of organ systems as normal vs. abnormal rather than specific disease classification. While our more recent conference proceedings manuscript experimented on multi-disease annotation [16], model performance was evaluated by aggregating diseased classes into a single abnormal class. In the present study, we report disease-specific classification performance for each previous version and the final version used in this study. Compared to these previous implementations, we expanded the RBA dictionary by adding more terms, introducing wild-card entries to tackle mispellings or grammatical errors, and increased the total number of reports threefold.

**Dataset**
A total of 261,229 chest, abdomen, pelvis structured CT reports of 112,501 unique subjects between the years 2012 to 2017 were extracted from the health system of our institution with IRB approval and deidentified. A representative example of a radiology CT report is shown in Fig. 2, which contains protocol, indication, technique, findings, and impression sections. The distribution of CT protocols is shown in Fig. 3.

**Rule-based algorithms (RBA)**
A separate RBA was created for the lungs/pleura, liver/gallbladder, and kidneys/ureters. Each RBA was limited to the findings section of the CT reports to minimize the influence of biasing information referenced in other sections and to ensure that the automated annotation reflected image information in the current exam (e.g., indication for exam, patient history, technique factors, and comparison with priors). For example, the impression section could describe a diagnosis based on patient history that could not be made using solely image-based information. For the purpose of RNN training, reports were filtered by protocol name to ensure organ-relevant scans were used for each model. For example, only protocols that included the entire chest (CAP, C, CA, and CP) were used to train the lungs/pleura model. In order to select target disease and organ keywords for the RBA dictionary, we computed term frequency–inverse document frequency (TF-IDF) [17] on the findings sections of a random batch of 3500 radiology reports. Informed by the prevalence of organ and disease keywords, we intentionally selected the three organ systems and four abnormal findings for each system such that the labels varied widely in location, appearance, and disease manifestations. For lungs/pleura, the four findings selected were atelectasis, nodule/mass, emphysema,
and effusion. For liver/gallbladder; stone, lesion, dilation, and fatty liver. For kidneys/ureters; stone, lesion, atrophy, and cyst. A board-certified radiologist (G.D.R.) provided guidance to define the TF-IDF terms into several categories, specifically:

(a) single-organ descriptors specific to each organ, e.g., pleural effusion or steatosis,
(b) multi-organ descriptors applicable to numerous organs, e.g., nodule or stone,
(c) negation terms indicating absence of disease, e.g., no or without,
(d) qualifier terms describing confounding conditions, e.g., however, OR
(e) normal terms suggesting normal anatomy in the absence of other diseases and abnormalities, e.g., unremarkable.

Additional file 1: Appendix S1 displays the dictionary terms and their descriptor type for each organ system. Figure 4 displays an overview of the RBA’s flowchart and logic. Although a separate RBA was created for each organ system, the workflow was the same. After the dictionary was refined, report text was converted to lowercase and each sentence was tokenized. In summary, the RBA was deployed on each sentence, and the number of potential diseases was counted first using the logic for the multi-organ descriptor and then the single-organ descriptor. If no potential disease labels were detected, then the normal descriptor logic was finally applied to verify normality. This process was repeated for each disease outcome allowing a report to be positive for one or more diseases or normal. Note that in this study an organ system was defined as normal not only by excluding the four diseases studied but also in the absence of dozens of abnormalities and diseases states that were not otherwise
analyzed, as shown in Additional file 1: Appendix S1. If the RBA failed to categorize the report definitively as positive for disease or normal (e.g., there was no mention of the organ system), then the report was labeled as uncertain and was not included in this study.

Upon manual review, we observed that many reports were incorrectly labeled normal due to excessively long sentences, which were either complex sentences with multiple clauses or fused together due to grammatical errors (e.g., missing periods). Such sentences were impractical to analyze with simple logic, so each report sentence was subject to a length criterion threshold for the normal outcome, another feature which made this RBA noticeably different from previous implementations.

From the full set of 261,229 reports, the lungs/pleura RBA classified a total of 165,659 reports from 74,944 subjects, the liver/gallbladder RBA classified 96,532 reports from 50,086 subjects, and the kidneys/ureters RBA classified 87,334 reports from 46,527 subjects. Note that the full set of cases does not correspond to the sum of reports for each organ system due to overlap of disease labels, where a single subject could have multiple findings across multiple organ systems. Figure 5 displays the disease distribution by organ system. Reports were randomly divided by subject into subsets for training (70%), validation (15%), and testing (15%) the RNN model.

Since the RNN depends on labels generated by the RBA, we manually validated the quality of the RBA
labels. From the above test set, a test subset of 2158 (lungs/pleura = 771, liver/gallbladder = 652, kidneys/ureters = 749) CT reports were randomly selected, and 2875 labels (lungs/pleura = 1154, liver/gallbladder = 787, kidneys/ureters = 934) were manually obtained by a Master of Biomedical Science graduate with gross anatomy training (V.M.D.) who was supervised by a board-certified radiologist (G.D.R.). This reference set was used to compare performance of the final RBA against our previous versions.

Attention-guided RNNs and training
A separate RNN was trained for each organ system using the corresponding RBA-annotated reports. The neural networks (Fig. 1) used in this study consisted of an embeddings layer, Bidirectional Long-Short Term Memory (BiLSTM), attention mechanism, dense layer, and final classification layer [18, 19]. The BiLSTM layer was composed of 200 units and produces a sequential output. It was followed up by a 0.2 dropout layer to prevent overfitting. The attention mechanism began with a time-distributed dense layer, which received a sequential 3-dimensional input (batch size, maximum sequence length, 1), and computed the aggregation of each hidden state. Next, it was reshaped to 2-dimensional form (batch size, maximum sequence length) followed by softmax activation, which assigned weights to each hidden state to produce an attention vector. The dot product of the attention vector and sequential output of BiLSTM was the final output of the attention mechanism. It was then followed by dense and classification layers. Since outcomes for each disease were non-mutually exclusive, we used a weighted binary cross-entropy loss and modeled the outputs as independent Bernoulli distributions for each of the labels with sigmoid activation.

Pre-training, datasets, model implementation
In this study, we compared the multi-label classification performance of two embedding approaches: with embeddings pretrained on the PubMed+MIMIC-III [20] dataset, and without pretrained embeddings (randomly initialized embedding layer). Embeddings of 200 dimensions were used in both experiments. Afterwards, we analyzed the effect of training data size on classification performance by incrementally increasing the number of training cases from 20%, 40%, 60%, 80%, or 100% of the total dataset. To prepare the training data, a preprocessing step was applied to each report. All numbers and punctuation were removed from each “findings” section, and the text was then converted to lowercase and tokenized. The sequence of tokens was then zero padded to the length of 650 tokens per sample. The models were trained for 50 epochs using a batch size of 512. The models corresponding to the minimum of the validation loss were selected as final. In this study we used Adam optimizer and a constant learning rate of 0.0001. The models were implemented using Python TensorFlow framework. Training duration was approximately 30 min for each
model using 2 TITAN RTX GPUs. All models’ weights and code are publicly available at (https://gitlab.oit.duke.edu/railabs/LoGroup/multi-label-annotation-text-reports-body-CT).

**Model evaluation**

To compare our final RBA against previous versions, accuracy and F1 scores were reported using the manually obtained labels as the reference standard. Accuracy was used to assess the total correct labels for a given disease, and was calculated using true positive (TP), true negative (TN), false positive (FP), and false negative (FN) values:

\[
\text{Accuracy} = \frac{TP + TN}{TP + FP + TN + FN}
\]

The F1-score is defined as the harmonic mean of precision and recall (sensitivity) and was calculated as:

\[
\text{Precision} = \frac{TP}{TP + FP},
\]

\[
\text{Recall} = \frac{TP}{TP + FN},
\]

\[
F_1 = 2 \cdot \frac{\text{precision} \cdot \text{recall}}{\text{precision} + \text{recall}}.
\]

To compare the performance of randomly initialized versus pre-trained embeddings as well as different sizes of training data, receiver operating characteristic (ROC) area under the curve (AUC) and 95% confidence intervals (CI) were reported. The ROC curve is a plot of the false positive rate vs. the recall, and AUC is a summary metric used to report model performance. CIs were calculated using the DeLong method [21].

**Results**

Table 1 displays the labeling accuracy and F-score of previously reported RBAs and the final RBAs for the binary labels (present/absent) for each disease and organ system. Performance was calculated based on the manually annotated test set of 2,158 CT reports with 2,875 labels. The performance of the final RBAs were equal to or greater than both previously reported RBAs [15, 16] for all diseases, with accuracy ranging from 91 to 99% and F-score from 0.85 to 0.98.

Table 2 displays the ROC classification performance of the attention-guided RNN with and without pre-trained embedding when applied to the test set containing 48,758 reports (23,411 reports for lungs/pleura; 13,402 reports for liver/gallbladder and 11,954 reports for kidneys/ureters). Pre-trained embedding outperformed the models trained with randomly initialized embedding for all organ systems and diseases.

Figure 6 displays examples of the output vectors produced by the attention mechanism for each organ system. Figure 7 displays the classification performance of the attention-guided RNN with pre-trained embedding when different portions of training data were used. Figure 7a displays the number of reports used in the training

| Organ          | Label     | # Pos | Han et al. [15] | Faryna et al. [16] | Our Final RBAs |
|----------------|-----------|-------|-----------------|--------------------|---------------|
|                | Acc       | F-Score | Acc         | F-Score          | Acc              |
| Lungs/Pleura   | Atelectasis | 251    | 0.86          | 0.74              | 0.97           | 0.95           | 0.98           | 0.97           |
|                | Nodule     | 296    | 0.77          | 0.74              | 0.92           | 0.89           | 0.92           | 0.89           |
|                | Emphysema  | 193    | 0.82          | 0.45              | 0.98           | 0.96           | 0.99           | 0.98           |
|                | Effusion   | 205    | 0.82          | 0.53              | 0.84           | 0.58           | 0.98           | 0.97           |
|                | Normal     | 209    | 0.79          | 0.44              | 0.96           | 0.94           | 0.98           | 0.96           |
| Liver/Gallbladder | Stone    | 144    | 0.87          | 0.62              | 0.95           | 0.91           | 0.95           | 0.93           |
|                | Lesion     | 224    | 0.92          | 0.88              | 0.94           | 0.91           | 0.95           | 0.92           |
|                | Dilatation | 87     | 0.86          | 0.1               | 0.9            | 0.7            | 0.98           | 0.92           |
|                | Fatty      | 166    | 0.97          | 0.94              | 0.98           | 0.96           | 0.98           | 0.96           |
|                | Normal     | 166    | 0.94          | 0.9               | 0.95           | 0.9            | 0.96           | 0.93           |
| Kidneys/Ureters | Stone     | 174    | 0.91          | 0.82              | 0.93           | 0.85           | 0.93           | 0.85           |
|                | Atrophy    | 94     | 0.96          | 0.85              | 0.99           | 0.97           | 0.99           | 0.97           |
|                | Lesion     | 238    | 0.91          | 0.87              | 0.91           | 0.86           | 0.91           | 0.86           |
|                | Cyst       | 234    | 0.95          | 0.92              | 0.96           | 0.94           | 0.96           | 0.94           |
|                | Normal     | 194    | 0.94          | 0.89              | 0.96           | 0.92           | 0.96           | 0.92           |

Comparison of classification performance between previously reported RBAs and our final RBAs using the manually annotated test set. "# Pos" is the number of positive examples for that label in the test set, Acc = Accuracy. Bolded values represent an equivalent F1-Score or increase in performance.
dataset after randomly splitting in 20% increments for lungs/pleura, liver/gallbladder, kidneys/ureters. Figure 7b displays the classification performance after training with each increment. AUCs reached > 0.95 for all classes in each organ system when using the complete dataset in the pre-trained models. Although the performance tended to improve as more training samples were used, most labels showed a robust plateau such that performances were still within the confidence intervals for 100% of the data. The most notable drops in performance were classes with smaller sample size (e.g., stone and dilatation for liver/gallbladder and atrophy for kidneys/ureters).

### Discussion

Although deep learning-based disease classification algorithms have recently achieved exceptional accuracy, they often suffer from limited diversity of diseases and organ systems. This narrow scope is largely due to inadequate amounts of curated CT data where human-annotation efforts are required. As an alternative, the work described here sought to develop high-throughput, multi-disease label extractors for body CT reports that were broad in scope and could be easily adapted to new keywords and diseases. The utility of automated labeling has been demonstrated by efficiently annotating large radiology report datasets to develop image-based CT classifiers, even without specific knowledge regarding disease location [6, 22]. As the foundation of our NLP algorithm, we developed RBAs that utilized simple rules to extract precise labels from radiology reports with 91–99% accuracy for all four diseases or normal across all three organ systems.

However, the RBAs alone could not provide labels for our entire dataset because radiology reports often contain variability in writing, grammar, and even variation in descriptors for the same disease between radiologists [23]. To overcome this obstacle, we demonstrated that an attention-guided RNN can be trained using RBA-annotated reports to learn salient semantic features and generalize beyond simple rules or keywords to encompass more reports. Our final disease classification pipeline performed with an AUC of > 0.95 for all diseases and organ systems. Recent works investigating deep learning-based radiology report annotation have achieved similar performances, although the majority are limited to a specific disease or organ system [24]. Examples include classification of pulmonary emboli in thoracic CT reports with AUC from 0.93 to 0.99 [14], annotation of mammography reports with a keyword accuracy of 0.96 [25], and identification of femur fractures with an F1 score of 0.97 [26].

Interpreting radiology reports often requires knowing the underlying clinical context, because language seemingly associated with disease is often used to describe normal, clinically benign findings. These common findings can account for many false positive errors when compared to manual annotations that do take into account such clinical context. Compared to previously reported RBAs [15, 16], for example, our final lungs/pleura RBA extracted labels with higher accuracy except for lung nodule. That class had many false positives for

### Table 2 Classification performance of randomly initialized versus pre-trained embeddings

| Organ        | Label      | # Pos | Random Initialization (AUC) | Pre-trained (AUC) |
|--------------|------------|-------|----------------------------|-------------------|
| Lungs/Pleura | Atelectasis| 9329  | 0.9968 (0.9961–0.9974)      | 0.9973 (0.9967–0.9997) |
|              | Nodule     | 10,183| 0.9913 (0.9904–0.9922)      | 0.9935 (0.9928–0.9943) |
|              | Emphysema  | 3659  | 0.9972 (0.9963–0.9982)      | 0.9980 (0.9972–0.9987) |
|              | Effusion   | 5625  | 0.9975 (0.9970–0.9980)      | 0.9984 (0.9980–0.9989) |
|              | Normal     | 3110  | 0.9990 (0.9985–0.9995)      | 0.9990 (0.9982–0.9997) |
| Liver/Gallbladder | Stone     | 1981  | 0.7849 (0.7739–0.7059)      | 0.9761 (0.9721–0.9801) |
|              | Lesion     | 6463  | 0.9675 (0.9646–0.9700)      | 0.9946 (0.9936–0.9955) |
|              | Dilatation | 1497  | 0.8120 (0.8013–0.8228)      | 0.9926 (0.9906–0.9945) |
|              | Fatty      | 1795  | 0.9984 (0.9851–0.9917)      | 0.9991 (0.9986–0.9996) |
|              | Normal     | 3162  | 0.9745 (0.9716–0.9773)      | 0.9762 (0.9750–0.9774) |
| Kidneys/Ureters | Stone    | 2548  | 0.9562 (0.9514–0.9609)      | 0.9792 (0.9764–0.9819) |
|              | Atrophy    | 750   | 0.9523 (0.9436–0.9611)      | 0.9955 (0.9936–0.9973) |
|              | Lesion     | 4817  | 0.9757 (0.9731–0.9783)      | 0.9900 (0.9886–0.9915) |
|              | Cyst       | 4164  | 0.9862 (0.9843–0.9881)      | 0.9926 (0.9914–0.9939) |
|              | Normal     | 2048  | 0.9090 (0.8980–0.9028)      | 0.9980 (0.9980–0.9992) |

Classification performance of randomly initialized versus pre-trained embeddings for each disease. “# Pos” represents the number of positives for that label. Values are reported as area under the curve (AUC) with 95% confidence interval (CI). Bolded values represent an equivalent AUC or increase in performance.
“small, calcified lung nodules,” which are a common, benign finding. Similarly for liver/gallbladder, an increase in performance was seen for each class except for fatty liver disease, which had many false positives because a “small amount of fat adjacent to the falciform ligament” is also normal. Last, there was a related reason why our kidneys/ureters RBA labeling performance did not increase compared to previous work [16]. Sentences describing renal diseases often contained several abnormal labels that triggered false positives for a related class but not the key finding. For example, “calcified lesion is likely a cyst” was labeled as lesion rather than cyst, and “an inferior pole left renal lesion has some calcification” was labeled as stone instead of lesion. Such difficult examples demonstrate the need for more advanced interpretation.

Further inspired by the recent wide application of deep learning-based methods in different clinical NLP tasks [10, 27–33] and effective application of word embedding [34–36], we also experimented using a multi-label disease classifier with pre-trained embedding and randomly initialized embedding layers. As expected, the attention-guided RNNs with pretrained embedding outperformed the randomly initialized models in all classes across all organ systems. Additionally, we observed that performance improved steadily with increasing number of cases. The lower frequency classes seemed to be affected greatly compared to classes having high frequency, exemplified by atrophic kidneys where performance experienced a significant drop at around 500 cases (20% of total available cases) for training.

The body CT dataset used in this study was dominated by two types of exams: chest and abdomen-pelvis CTs. In many reports, one or more of the three organ systems were out of view and not mentioned at all by
the radiologist. For example, if a chest CT did not mention the kidneys, that would be labeled as uncertain by our RBA. However, in specific studies such as abdomen-pelvis CT, large organs such as the lung were often still described even if they were not completely visible e.g., “Limited view of the lung bases appear clear.” This short sentence would satisfy the logic of the RBA to label the report appropriately as normal for the lungs.

There were several limitations to this study. As a general limitation of RBA techniques, it was not possible to provide disease labels for all reports within our dataset. This was often because each sentence did not satisfy the pre-defined, strict rules. To mitigate this effect, future work should expand the dictionary through discovery of new and potentially uncommon language uses. Another limitation is that, unlike when radiologists annotate images manually, the labels derived from reports tend to describe all or much of an organ system (e.g., “bibasilar atelectasis”) and in some cases provide limited disease extent and location (e.g., “nodule measuring 1.8 × 2.1 cm on series 2 image 60”). Furthermore, our dataset suffered from notable class imbalance, including a low prevalence.

Fig. 7 Effect of different sizes of training data in the pretrained embedding models on classification performance. a Number of reports randomly split in 20%, 40%, 60%, 80% and 100% of total training dataset for each disease by organ system. b Performance of models on test-set trained with randomly split 20%, 40%, 60%, 80%, and 100% training data for each disease by organ system reported as AUC. Error bars represent 95% confidence intervals.
of normal cases as well as multi-fold differences between diseases, although this represented the natural prevalence within our study population. The dataset also came from a single health system, which comprises multiple hospitals but may share similarities in the reporting patterns for radiologists. Finally, this initial demonstration focused on building three separate classifiers rather than a single multi-organ model. Independent processing of diseases could have simplified the challenges imposed by multiple organ interaction, and in future work we will consider the feasibility of a single model, multi-organ approach.

Conclusions
The disease labeling pipeline described here offers numerous advantages. By using deep learning-based NLP, our algorithms were able to generalize beyond predefined rules and label a vast and heterogenous dataset as positive for one or more diseases or normal for three different organ systems. To the best of our knowledge, this was a first attempt in using RBA-extracted labels to train an attention-guided RNN to annotate a diverse set of diseases in a hospital-scale dataset of body CT reports. Ultimately, the work described here sought to facilitate future research in image-based disease classification algorithms by providing a general framework for labeling vast amounts of hospital-scale data in a manner that is both cost and time efficient.

Abbreviations
CT: Computed tomography; RBA: Rule-based algorithm; RNN: Recurrent neural network; ROC: Receiver operating characteristic; AUC: Area under the curve; CI: Confidence interval; NLP: Natural language processing; IRB: Institutional review board; TF-IDF: Term frequency–inverse document frequency; CAP: Chest, abdomen, and pelvis; C: Chest; AP: Abdomen-pelvis; A: Abdomen; P: Pelvis; CA: Chest-abdomen; CP: Chest-pelvis; BiLSTM: Bidirectional long-short term memory.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12911-022-01843-4.

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Author contributions
VMD drafted the manuscript, refined the dictionary, and manually labeled the radiology reports. FIT implemented the models and performed the analyses. KF and SH helped to develop the models. MAM contributed to the overall design, development, and writing of the manuscript. All authors have read and approved this manuscript.

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Availability of data and materials
The radiology reports used in the current study cannot be shared publicly because it is impractical to ensure the removal of all protected health information from the large amount of text data. Please contact the corresponding author for data requests. The code is publicly available at https://gitlab.oit.duke.edu/railabs/LoGro up/multi-label-annotation-text-reports-body-CT.

Declarations
Ethics approval and consent to participate
This study was approved by the IRB at Duke University under protocol # Pro00082329. Informed consent was waived by the IRB at Duke University for this retrospective study that was compliant with the Health Insurance Portability and Accountability Act. IRB approval included permission to access the raw data. All experiments were performed in accordance with relevant guidelines and regulations.

Competing for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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