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Original Research Article

Classification of cervical biopsy free-text diagnoses through linear-classifier based natural language processing



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ABSTRACT

Routine cervical cancer screening has significantly decreased the incidence and mortality of cervical cancer. As selection of proper screening modalities depends on well-validated clinical decision algorithms, retrospective review correlating cytology and HPV test results with cervical biopsy diagnosis is essential for validating and revising these algorithms to changing technologies, demographics, and optimal clinical practices. However, manual categorization of the free-text biopsy diagnosis into discrete categories is extremely laborious due to the overwhelming number of specimens, which may lead to significant error and bias. Advances in machine learning and natural language processing (NLP), particularly over the last decade, have led to significant accomplishments and impressive performance in computer-based classification tasks. In this work, we apply an efficient version of an NLP framework, FastText™, to an annotated cervical biopsy dataset to create a supervised classifier that can assign accurate biopsy categories to free-text biopsy interpretations with high concordance to manually annotated data (>99.6%). We present cases where the machine-learning classifier disagrees with previous annotations and examine these discrepant cases after referee review by an expert pathologist. We also show that the classifier is robust on an untrained external dataset, achieving a concordance of 97.7%. In conclusion, we demonstrate a useful application of NLP to a real-world pathology classification task and highlight the benefits and limitations of this approach.

Key messages

Linear classifiers using natural language processing algorithms successfully assigned accurate biopsy categories to free-text biopsy interpretations, with a concordance greater than 99.6% on validation set data and 97.7% on untrained external data.

Introduction

In the past 40 years, the incidence and mortality of cervical cancer decreased significantly in the United States due to successful cervical cancer screening programs. Cytology is the mainstay of cervical cancer screening programs around the world; the practice of cytology informs crucial subsequent clinical management and further testing through well-validated clinical decision algorithms.¹ Persistent infection with high-risk human papillomavirus (hrHPV) causes cervical cancer and precancerous lesions. Retrospective review of cervical cytology and HPV test results in correlation with follow-up cervical biopsy diagnoses may inform updates to clinical decision algorithms.^{2,3} Our group previously published several studies of this nature.^{4–9} As our dataset of cervical biopsy diagnoses increased in size, manual categorization of the free-text biopsy diagnosis into discrete categories proved increasingly challenging.

There is increasing interest in encoding pathology free-text information by extracting clinically relevant information from data-rich pathology reports.¹⁰ Some well-tested approaches have focused on using textprocessing algorithms to identify and amend report defects in surgical pathology, such as errors in voice recognition¹¹ and data tabulation.¹² These approaches encompass a wide variety of text processing strategies, including regular expressions, term detection and tokenizing, support vector machines, statistical approaches, and formal parsing. The comparison of these approaches is difficult due to lack of validation studies, highquality and annotated datasets with abundant textual diversity, and welldefined consensus approaches for performance evaluation.¹³

The field of natural language processing (NLP) consists of multiple models that have been used to increase classification accuracy in myriad applications ranging from quality improvement in factories, anomaly detection in national security, and photo recognition on social networks. These NLP models use algorithms to transform free-text data into encoded data, which is more reliable, less error-prone, and less costly to search than free text.¹⁴ Two primary approaches exist: rules-based approaches, which

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involve manual encoding of a set of rules designed to process a particular item using the grammar and heuristic rules appropriate for that dataset, and statistical-based approaches, which learn relationships present within existing data (training set) and apply it to previously unseen data (validation set). Rules-based approaches, which comprise word/phrase matching approaches such as regular expressions, SQL queries, and ontology matching,¹⁰ have wide-ranging applicability in several areas of pathology including automating tissue bank annotation,¹⁵ coding and retrieving surgical pathology reports,¹⁶ and extracting Cancer Registry data.¹² However, robustness to natural language variation is a challenge for any rules-based approach. Additionally, depending on the specific rules-based approach used, the ordering of rules can have significant and unforeseen effects on the output, due to the fundamentally iterative nature of most rules-based approaches.

Natural language models using statistical-based approaches, introduced through advances in machine-learning algorithms and statistical inference,¹⁷ were first introduced in the late 1980s, but with exponential increases in computational power and memory, are increasingly practical and useful for real-world datasets. These are further subdivided into 2 dominant approaches. Representation models learn relationships between elements in the input through a series of transformations such as component and cluster analysis before a classification/prediction step. In contrast, deep learningbased models utilize neural networks to extract relevant features from large training sets to feed into multi-layered (feed-forward) perceptron models, which are simplified models of biological neurons. For natural language processing, these 2 approaches translate into linear classifiers and deep classifiers, respectively. Linear classifiers assign labels to words by extracting relevant features from the set of input sentences, which are fed to a shallow classification algorithm (support vector machine; SVM) to obtain a categorical output. Deep classifiers process the input sentences with several layers of feature extraction, and backpropagate the features to deeper layers of the network by convolution, which may offer better classification performance. However, deep classifiers can have significant computational costs, often requiring expensive graphics processing units (GPUs) or other massively parallel computing devices, and often require larger, more diverse training data for acceptable classification performance.¹⁸

There is increasing focus within the field to develop more efficient linear classifiers with classification performance comparable to deep approaches while using much less computational power that can run even on inexpensive smartphones.¹⁹ A set of popular approaches uses distributed representations of word vectors that are indifferent to word ordering to create a statistical model of multiple word phrases, or "n-gram", frequency that can be used in downstream classification tasks. Specifically, unlike simple "bag of words" models, multiple-word n-grams enable local word ordering to be captured, as exemplified by the different meanings of the phrases "work to live" and "live to work". As originally described in the skipgram model and further refined by biased sampling against frequent words (Word2Vec,²⁰ Google) and capturing local word ordering (FastText,^{19,21} Facebook), these more efficient approaches have dramatically improved the scalability and usability of natural language processing in various portable applications, including internet search, image recognition, and content tagging. In particular, FastText, a library for efficient text classification and representation learning developed by Facebook (R) Research, builds on the improvements of linear classifiers while making significant improvements in classification accuracy through fast loss approximation (linearly decaying learning rate). This allows it to approximate the performance of conventional DNN classifiers (char-CNN, VDCNN) and comparable linear classifiers (Tagspace) with an order of magnitude of performance improvement.^{19,22}

In this project, we focus on implementing unsupervised, order-invariant word vectors to classify cervical biopsy free-text diagnoses into discrete pre-defined categories, and compare this approach with our previously implemented rules-based regular expression classifier. We also investigate discrepancies between categories generated by NLP and by the rulesbased classifier, with referee pathologists determining the "ground-truth" classification of these cases.

Methods

Dataset

The dataset is comprised of 59 751 free-text gynecologic biopsy diagnoses received from BioReference laboratories and interpreted at Houston Methodist Hospital from 2013 to 2018 (Fig. 1). These free-text diagnoses were labelled into discrete diagnosis categories using the rules-based classifier described below. An NLP classifier was created by training on a subset of the labelled data, then used to predict the label on the remaining data. Additionally, a dataset comprising 6672 free-text gynecologic biopsy diagnoses interpreted at Houston Methodist Hospital in 2020 was used for external validation (true holdout). This external dataset comprises interpretations from many different pathologists than the training and validation datasets, and was extracted from a different Laboratory Information System (LIS).

Rules-based classifier

Regular expressions were used to construct a rules-based classifier to generate preliminary discrete diagnosis labels. Briefly, the rules-based approach is a Java class file that accepts an input of diagnoses separated by newlines, lowercases all alphanumeric characters, applies a series of string-matching functions in an if/then loop to assign discrete categories (BiopsyClass), and converts the output into a separate text file.

The string-matching functions are sequenced to capture more significant diagnoses (e.g. invasive carcinoma, HSIL) before less significant diagnoses (e.g. LSIL, HPV effect) (Fig. 2). The classifier mapped the free-text biopsy diagnosis into a granular discrete class and a more general class using the BiopsyClass and MappedBiopsyClass vocabularies, respectively



Fig. 1. Overview and project workflow. 59 751 deidentified biopsy diagnoses from 5 years of cervical biopsy specimens at BioReference Laboratories, and the associated biopsy class and mapped biopsy class. Biopsy diagnoses are preprocessed to decrease uninformative variation in the text, and the FastText algorithm was run with varying proportions of the data as training and validation sets. The results were compared to rules-based regular expressions to assess accuracy, discrimination threshold, and any discrepancies between the two methods were examined.



Fig. 2. Overview of the rules-based classifier. The rules-based classifier does basic pre-processing of interpretation text followed by a string matching approach that eliminates diagnoses based on severity (carcinoma, followed by HSIL, followed by LSIL). An example is provided for a sample text interpretation (inset box).

(Table 1). The BiopsyClass vocabulary is a more granular classifier that distinguishes between Cervical Intraepithelial Neoplasia (CIN) 2, CIN3, adenocarcinoma in situ and carcinoma, and the MappedBiopsyClass vocabulary is a more general classification label.

NLP classifier

Pre-processing

The free-text diagnoses were pre-processed to reduce uninformative variation. Newlines were removed. Punctuation was preserved, but tokenized (a space was placed before and after each punctuation symbol). The text was transformed into lowercase. Multiple training and validation sets were constructed by randomly sampling without replacement a proportion r out of the 59 751 total diagnoses for the training set, where r ranged from 0.01 (1% of dataset used for training) to 0.95 (95%) with a step size of 0.01, for 5 run replicates each, for a total of n = 480 runs (see Bootstrapping below). For comparative performance evaluation with the rules-based approach and for discrepancy analysis, training was performed using a randomly sampled 80% of the total dataset.

Table 1

Vocabularies used for natural language processing categorization.

BiopsyClass	MappedBiopsyClass	Description			
Benign	Benign	Benign lesion or normal tissue			
CA:AD	CA	Carcinoma, adenocarcinoma			
CA:ADIS	HSIL+	Adenocarcinoma in situ			
CA:ENDOMETRIAL	CA:ENDOMETRIAL	Carcinoma, endometrial			
CA:NOS	CA	Carcinoma, not otherwise specified			
CA:SC	CA	Carcinoma, small cell			
CA:SQ	CA	Carcinoma, squamous cell			
Endometrium	Endometrium	Benign endometrium			
		High-grade squamous intraepithelial			
HSIL:CIN2	HSIL+	lesion (CIN2)			
		High-grade squamous intraepithelial			
HSIL:CIN3	HSIL+	lesion (CIN3)			
LSIL	LSIL	Low-grade squamous intraepithelial lesion			
		Vaginal squamous intraepithelial			
VaIN	VaIN	neoplasia			

Model training

For each training data set, we created an NLP classification model using FastText 0.2.0 on a Linux (Ubuntu 18.04) virtual machine. Settings used included epochs of 5–50, a learning rate of 0.1, and nGrams of 1–2 (bigrams). Accuracy probabilities and prediction labels on the validation set were generated using the "predict-prob" function. Selection of invariant constants (hyperparameters) was assisted by use of the "autotune" feature in FastText on a subset of hyperparameters (number of epochs, learning rate, and nGrams). The BiopsyClass was used as the training and prediction label for the algorithm. To reduce pathologist interpretative variance as a confounding factor (e.g. CIN2 vs CIN3 for HSIL), the more general MappedBiopsyClass was used to compare the performance of both the rules-based and NLP algorithms. Unless otherwise specified, training was performed on 80% (47 800 free-text diagnoses) of the randomly shuffled dataset using 20% (11 951 free-text diagnoses) as the validation set.

Model validation

Bootstrapping

To assess the stability of classifier performance based on choosing discrete training set entries, sampling with replacement (bootstrapping) was performed. The algorithm was run for 5 randomly generated training and validation steps for each proportion r, where r ranged from 0.01 to 0.95 with a step size of 0.01, for a total of n = 480 runs. Sensitivity of bootstrapping results to n-Gram size was assessed by varying n-Grams chose from 1 to 5, and re-running the bootstrapping process. The arithmetic mean of precision@1 was plotted for each selected proportion of the training set.

Hyperparameter optimization

FastText offers automatic hyperparameter optimization, allowing for semi-automated tuning of classifier hyperparameter such as learning rate, epoch sizes, and nGram sizes. The hyperparameter optimization algorithm sequentially runs training and validation classification and iteratively selects the parameters that produces the highest F1-score (combined precision and recall). The default parameters are nGrams = 1 and epoch = 5. Learning rate, epoch size (5–50), nGram length (from 1 to 5), array

J.W.-C. Hsu et al.

dimension size (1–100), and bucket size were selected as optimizable hyperparameters, and performed on a randomly sampled 80% training / 20% validation dataset. Optimized classification F1-score was achieved with an epoch size of 20, nGram length of 2, and default learning rate (0.1) with a dimension size of 100 and bucket size of 100 000.

K-fold cross validation

Cross validation using sequentially sampled portions of the data were used to assess classifier stability and check for spurious dependence on the sequence of the underlying data. The biopsy predictions were randomly shuffled, then sequentially partitioned into 5 folds, using 20% of the data as a hold-out set for each iteration. For each iteration, validation was performed on the remaining 80% of the dataset. Precision and recall @ 1 was evaluated with default (nGrams=1, epoch=5) and optimized (nGrams=2, epoch=20) FastText parameters, as described in the hyperparameter optimization section.

Analysis and discrepancies

Manual review of discrepant (NLP-predicted label differs from rulesbased predicted label) cases was performed by 2 pathologists (PAC, YG), who assigned the correct ("ground-truth") label for each discrepant case after mutual discussion. The results of the discrepancy review were designated as NLP (when only NLP was correct), Rules (when only the regular expression classifier was correct), or Neither (when neither NLP nor Rules were correct). Optimized NLP hyperparameters were used to train the NLP classifier on the training dataset with manually corrected "groundtruth" labels before external validation (see below).

External validation

As described in the discrepancy analysis above, free-text diagnoses were labelled into discrete diagnosis categories by rules-based classifier followed by manual review (to generate a "ground-truth" label). Optimized (nGrams = 2, epoch = 20) NLP classifier hyperparameters were used, and the NLP classifier trained previously was used on this dataset to predict classes with no hyperparameter tuning or modification of the training set. Any discrepancies between the "ground-truth" label and NLP predictions were analyzed.

Results

NLP classification concordance with regular expressions

Overall classification concordance with rules-based regular expressions was excellent at 99.3%; however, there was considerable heterogeneity within each BiopsyClass, ranging from 100% of diagnosis correctly classified to 57.9% for vulvar intraepithelial neoplasia (VIN) specifically (Fig. 3).

Parameter stability and effect of training parameters

K-fold cross validation

Precision and recall @1 with default (nGrams = 1, epoch = 5) and optimized (nGrams = 2, epoch = 20) FastText parameters remained stable between 0.987–0.989 and 0.994–0.995, respectively (Fig. 4). Run-to-run variation in class distribution remained minimal.

N-gram analysis

In addition to automatic hyperparameter optimization, the performance of the NLP classifier at various N-gram lengths was assessed using manual parameters (nGrams 1 to 5). A nGram setting of 2 noticeably increased precision and recall accuracy, especially for "rare" diagnoses such as carcinoma (CA) and vulvar intraepithelial neoplasia (VIN) with no increases in the training set (Fig. 5A) compared to nGram of 1. With only 1% of sampled training data, the algorithm was able to achieve more than 95% concordance with rules-based regular expressions (Fig. 5B).



Fig. 3. Overall and per-class classification accuracy. Training performed on 80% (47 800 free-text diagnoses) of the randomly shuffled cervical biopsy dataset using original default (nGrams = 1, epoch = 5) and optimized (nGrams = 2, epoch = 20) FastText parameters. Accuracy denotes percentage of NLP classifications concordant with rules-based regular expressions ("ground truth").

Discrepancy compilation and analysis

99 classification discrepancies in the validation set were identified. Of these, 39/99 (39%) were decided in favour of the NLP algorithm, 55/99 (56%) in favor of rules-based regular expressions, and 5/99 (5%) of results were incorrect by both NLP and rules (Fig. 6A, B). Examples of results from each category are given (Fig. 6C) and encompass problematic areas for both approaches such as descriptive "hedging-type" diagnoses as well as rarely encountered diagnoses such as stratified mucin-producing intraepithelial lesions (SMILe). These difficult classifications are reflected in the lower confidence scores (below 0.95) that the diagnoses were assigned to by the NLP algorithm.

External validation

The NLP classifier with optimized FastText parameters was used for external validation of 6672 free-text gynaecologic biopsy diagnoses from January 1, 2020 to December 31, 2020, achieving a concordance rate with ground-truth labels of 97.7% (6516/6672 correct MappedBiopsyClass). Examples of discrepant results are shown (Fig. 7) and shows some of the difficulties encountered including descriptive diagnoses as well as weighing of poorly represented diagnosis such as VIN in the training set.

Discussion

In this study, we demonstrate that a fast, efficient NLP technology package (FastText) can be successfully trained on a large collection of cervical



Fig. 4. 5-fold cross validation of cervical biopsy classification using FastText with original default (black) and optimized (red) parameters. Cross validation using sequentially sampled portions of the data were used to assess classifier stability. Precision and recall @1 with original default (nGrams = 1, epoch = 5) and optimized (nGrams = 2, epoch = 20) FastText parameters were assessed for each of 5 folds, encompassing 20% validation dataset for each fold.

J.W.-C. Hsu et al.



Fig. 5. Classification accuracy versus training set proportion for various nGram values. Training set proportion versus classification accuracy (concordance with rules-based regular expressions) for randomly sampled training set data. (A) 1%–95% of randomly sampled training set data for nGram of 1. 96.3% classification accuracy was obtained using 4% sampling of training set data (box). (B) 1–10% of randomly sampled training set data for nGram values from 1 to 5 (coloured lines). 95.3% classification accuracy was obtained using 1% sampling of training set data a nGram value of 2 (box).

biopsy free-text interpretation, and via supervised learning, classify biopsies with greater than 99% accuracy, which remains at up to 98% on external untrained data. We also demonstrate the sensitivity of the algorithm to training parameters such as n-gram length and training epochs, with an n-gram length of 2 with concomitantly longer training intervals to optimize classification accuracy. We also demonstrate the informative nature of discrepancy analysis (identifying interpretations where the NLP predicted class does not match the rules-based generated class), and how delegating these cases to a referee decision can identify weaknesses of both models and further refine classification accuracy while only minimally increasing manual workload.

FastText offers a number of optimizable parameters "out of the box" that significantly affects classification performance, including learning rate; number of training epochs; and the use of bi-grams, tri-grams, or

Α	Referee diagnosis					Number of			of	С	Pred	iction		Referee
	oonoordon	+ wi	th.							Interpretation	Rules	NLP	Confidence	Category
	CONCOLUAI	L WI	ui.				Gas	585		Referee concordant with Rules-based categories	jory			
	Neither						Ę	5		scanty atypical squamous cells favor Low grade squamous	VIN	1.011	0.50) /INI
	NLP						3	9		Intraepitnellal lesion (LSIL, valn I).	VIN	BENIGN	0.50	VIN
	Rules						5	5		Squamous cell carcinoma in situ, at leastRare atypical glandular cells, favor neoplastic	HSIL+	CA	0.82	HSIL+
	Total discr	ера	nc	ies	;		9	9		Low grade squamous intraepithelial lesion (CIN 1) with microfocal moderate dysplasia/CIN 2	HSIL+	LSIL	0.87	HSIL+
В										Invasive squamous cell carcinoma, well differentiated, arising in a background of high grade squamous intraepithelial lesion (HSIL, CIN 3). Tumor invades at least 1.5 mm deep.Deep / endocervical margin is positive for invasive carcinoma.Endocervical and ectocervical surgical marqins are positive for HSIL.	СА	HSIL+	0.96	CA
			Re	fer	ee	Cat	ea	orv		Referee concordant with NLP-based catego	rv			
				-				Î I		Features consistent with exogenous progestin effect	BENIGN	ENDOMETRIUM	0.75	ENDOMETRIUM
				METRI/	m					Minute fragment of squamous epithelium with mild atypia (see microscopic description)Fragments of superficial endometrial tissue (see microscopic description)	ENDOMETRIUM	BENIGN	0.30	BENIGN
		enign	-	A:ENDO	Idometr	SIL+	Ħ	z	otal	Scant transformation zone mucosa with squamous metaplasia, inflammation, and reactive changesNegative for dysplasiaBenign endometrial tissue, proliferative pattern	ENDOMETRIUM	BENIGN	1.00	BENIGN
NLP P	rediction	ä	ບີ	3	ш	Ξ	2	5	Ĕ	Low grade squamous intraepithelial lesion (LSIL, CIN 1) Endocervical	DENION	1.011	0.72	LOU
Benign		1	3	6	10	13	3	Т	36	Minute focus of High Grade Squamous Intraepithelial lesion Confirmed	DEINIGIN	LƏIL	0.73	LOIL
CA			0			4			5	on P16/ki67 block positivitybenign endocervical mucosa	BENIGN	HSIL+	0.96	HSIL+
CA:END		1	2	4		4			3	Referee discrepant with both categories				
Endome	trium	5	-	1		1		•	/	Stratified mucin-producing intraepithelial lesion (SMILe; see comment)	BENIGN	CA	0.34	HSIL+
HSIL+		8	1			4.4		2	1/	polypoid fragment of vulvar skin with focal features compatible with				
LSIL		16				11		3	30	human papillomavirus intection/ condyloma accuminatum. see note note: Immunostain for Ki-67 (a proliferation marker) has been				
VIN Total		31	12	7	10	30	3	6	99	reviewed. A few suprabasilar keratinocytes show positive nuclear staining with Ki-67. This staining pattern supports the above diagnosis.	BENIGN	LSIL	0.99	VIN

Fig. 6. Discrepancy analysis. All discrepant results in the validation dataset were manually reviewed by 2 pathologists (PAC, YG), and a referee diagnosis generated after mutual discussion. (A) Number and proportion of diagnoses concordant with the referee diagnosis for NLP, rules-based regular expression, or neither. (B) Confusion matrix of discrepancies between NLP-called categories with referee categories. Number of discrepant cases for each pair indicated in grid. (C) Example free-text interpretations of discrepancies, and corresponding MappedBiopsyClass classes for NLP, rules, and the referee. The classification confidence score given by FastText is also provided.

Α	Poforoo Catogor	B	Interpretation	Ground Truth	NLP	Confidence
		,	Ectocervix and transformation zone mucosa with changes consistent with HPV infection Minute fragment with HSIL/CIN2-3 P16/Ki-67 is confirmatory Immature squamous metaplasia	HSIL+	LSIL	0.96
	Beniç CA IDON HSIL LSII VIN	Tota	Adenocarcinoma and Small cell neuroendocrine carcinoma	CA:AD	CA:ENDOMETRIAL	0.97
NLP Prediction	CA:EN End		endocervical mucosa. Please correlate clinically and with endometrial stripe thickening as material may not be entirely		DENION	0.00
Benign	9 6 55	70	representative of the process	ENDUMETRIUM	BENIGN	0.82
CA		0	CIN-3/CIS with extensive endocervical gland involvement No			
CA:ENDOMETRIAL	1 3	4	invasive carcinoma identified No endocervical glandular		DENICH	0.00
Endometrium	4 1	5	dysplasia identified Dense chronic inflammation	HOILT	DEINIGIN	0.99
HSIL+	4 1 17 2	24	HSIL and AIS	HSIL+	VIN	0.97
LSIL	5 11	16	Rare strips of atypical endocervical cells worrisome for AIS	CA:ADIS	LSIL	1
VIN	36 1	37	Low-grade squamous intraepithelial in transformation zone			
Total	14 4 0 10 53 73 2 1	56	Benign ecto and endocervix	LSIL	BENIGN	0.92

Fig. 7. External validation. The external dataset comprises 6672 free-text gynecologic biopsy diagnoses interpreted at Houston Methodist Hospital in 2020 by several different pathologists, and extracted from a different LIS. Ground-truth labels were compiled after manual review, as performed in the discrepancy analysis. (A) Confusion matrix of discrepancies between NLP-called categories with referee categories. Number of discrepant cases for each pair indicated in grid. (B) Example free-text interpretations of discrepancies, and corresponding MappedBiopsyClass classes for NLP and referee categories.

other n-grams to generate the classifier. Compared to baseline parameters, the choice of bi-grams (nGrams = 2) with longer training epochs (epoch = 20) improved classification accuracy markedly; however, n-grams greater than 2 and longer epoch settings did not significantly improve results, but contributed to longer processing times. A possible explanation could be that more text diversity (larger sample sizes or multi-institutional data) would be needed for the substantially higher token count with higher nGram settings to be adequately trained, or that bi-grams already sufficiently capture training text with limited input diversity (such as pathology reports of only gynaecologic tissue).

Overall, NLP can achieve excellent concordance with a manually iterated rules-based approach to categorization, and in certain instances can achieve superior performance for diagnoses with variation not captured by regular expressions, such as different syntax or phrasing. Of note, the NLP approach is robust to training with a very limited dataset (1% of all data), which achieved greater than 95% concordance with rules-based approaches (Fig. 5B), and is chiefly limited by misclassification of rarer diagnoses. In other words, for a dataset of size 10 000, examining only 100 entries was sufficient to capture the majority of within-dataset variation. Conversely, increasing the proportion of training set sampled beyond 80% did not improve classification performance, as it is likely that the algorithm is overfitted with such a high proportion of training data (Fig. 5A); this may explain why increasing epochs further or similar hyperparameter changes did not measurably affect classification accuracy. Diagnoses with low frequency pose challenges for rules-based approaches because they may not have a rule coded for that scenario; the machine learning-based models are also challenged because the labelled training data may not have representative cases included or labelled correctly, and these diagnostic challenges are reflected by the relatively low confidence that the NLP algorithm assigns. These results could likely be improved by biased or class-imbalanced sampling of less common diagnoses/tokens, as demonstrated by similar applications of NLP in non-pathology fields.^{25,26} For rules-based approaches, which require continuous revision and updating to handle "edge cases" and changes in syntax, these challenges are comparable and also require a skilled pathologist/informatician to make the necessary updates.

Robustness of all classifiers, whether rules-based or machine-learning, to unknown incoming data is always a concern. As we show in external validation, the performance of the NLP classifier degrades somewhat but remains at 97.7% (Fig. 7). The challenges of handling unknown data are especially acute in token-based approaches such as FastText, where syntactical variations, misspellings, and changing clinical guidelines, such as emphasis on endocervical sampling, increase token diversity and adversely affect classification performance. These challenges are not limited to NLP, as rules-based classifiers also require continuous revision, maintenance, and ongoing validation to be robust to incoming data. Arguably, retraining/fine-tuning an NLP classifier has the potential to introduce less

systematic bias compared to manually adding new rules to a rules-based classifier, and could be performed as part of a semi-automated pipeline by pathologists without much experience in machine learning. Further pre-processing of incoming data could also be a valuable approach to reduce token variability for both approaches; minimum pre-processing was intentionally performed for this study to minimize subjectivity and to make a direct comparison between the rules-based classifier and NLP classifier informative.

In addition to labelling, confidence scores generated by the FastText algorithm are valuable for downstream interpretation tasks. These confidence scores are not available in the rules-based approach. Triaging diagnoses with lower confidence scores for human review, as well as incorporating these diagnoses into the model after retrospective review, can be an effective way to further improve model robustness. For example, rerunning the model with corrected discrepant annotations for SMILe (stratified mucin intraepithelial lesion) and vulvar skin with condyloma acuminatum correctly produced the classifications "HSIL" and "VIN" respectively in a subsequent validation run (Fig. 6C), and this process could be iterated to further improve classification as discrepant annotations are manually reviewed. In addition, NLP-based approaches offer the unparalleled advantage of fast retraining by simply re-processing the revised "ground-truth" classifiers generated by referee pathologists to further improve performance for real-world use, which supports an iterative training approach to handle unexpected alterations in incoming data.

Evaluating the suitability of natural language processing technologies for the practice of pathology is multi-factorial. Compared to manual approaches, NLP effectively eliminates inter-observer variability, and compared to manually curated rule-based systems, also reduces "design bias", a problem inherent to systems such as regular expressions. However, a successful implementation of this technology in a working pathology practice involves more than dataset selection, training, and validation; the context and method that NLP is integrated into daily workflow is also crucial to consider. The actual correctness of so-called "ground-truth" classifiers should also be considered; inter-observer variability among practicing pathologists can be significant for descriptive, complex, or rarely seen diagnoses^{23, 24}; and low prediction confidence from NLP-based models can effectively signal the presence of these so-called "problematic cases" by automatically flagging them for post-prediction manual review.

Conclusion

In this study, we demonstrated a successful implementation of NLPbased classifiers into study of a large-scale pathology dataset. These classifiers have significant advantages compared to rules-based classifiers such as classification performance, minimizing design bias in rule selection, and robustness to minor variations in free-text data. Integration of these classifiers into an interpretative workflow requires additional considerations such as LIS compatibility, flagging of problematic or "low confidence" bases, and refinement of the user experience during operation of the classifier, which remain the areas to be explored further.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi. org/10.1016/j.jpi.2022.100123.

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