ABSTRACT

Machine Learning has been applied to pathology images in research and clinical practice with promising outcomes. However, standard ML models often lack the rigorous evaluation required for clinical decisions. Machine learning techniques for natural images are ill-equipped to deal with pathology images that are significantly large and noisy, require expensive labeling, are hard to interpret, and are susceptible to spurious correlations. We propose a set of practical guidelines for ML evaluation in pathology that address the above concerns. The paper includes measures for setting up the evaluation framework, effectively dealing with variability in labels, and a recommended suite of tests to address issues related to domain shift, robustness, and confounding variables. We hope that the proposed framework will bridge the gap between ML researchers and domain experts, leading to wider adoption of ML techniques in pathology and improving patient outcomes.

1 INTRODUCTION

Pathology is often treated as a ground-truth for many serious diseases and a pathologist’s diagnosis is critical for a wide variety of tasks, including drug development and clinical diagnostics. The typical workflows of a pathologist are complicated, subjective and limited by what humans can evaluate. The use of machine learning in pathology has begun to transform the field with applications like clinical decision support tools (Campanella et al., 2019), improving prognostic utility over manual reads (Taylor-Weiner et al., 2021), automating slide scoring in clinical trials (Glass et al., 2021), novel biomarker discovery (Echle et al., 2021), and understanding tumor pathology (Jiang et al., 2020; Diao et al., 2021). These models require thorough validation and verification since they directly impact patient lives and their evaluation requires a careful look at the setting in which they will be deployed. Pathology images also differ from natural images in various ways - they are massive in resolution (billions of pixels) with each magnification-level containing markedly different information (Komura & Ishikawa, 2018), have different equivariances from natural images (Veeling et al., 2018) and contain stain variations which can confound models (Yagi, 2011; Madabhushi & Lee, 2016; Tellez et al., 2019). In addition to that, exhaustive labeling of these images is prohibitive as they require expert annotations and are time-consuming. Finally, building ML models for these problems requires understanding of biological causal structures to deal with spurious features (Castro et al., 2020). These differences necessitate model evaluation guidelines specific to pathology problems to prevent life threatening model failures.

In machine learning, dataset creation, evaluation setup, choice of metrics, and the data splits used for training, validation and testing have been areas of study in themselves, and the research community depends on agreed standards for evaluating novel methods and their utility in the real world. Previous work in medical imaging has highlighted problems in transferring these evaluation methodologies, making models brittle for deployment in real-world setting (Park & Han, 2018; England & Cheng, 2019; Varoquaux & Cheplygina, 2021; Reineke et al., 2021). In this work, we propose a guideline on ML model evaluation for pathology which go beyond standard ML metrics. We define how to setup
We divide model evaluation into three stages: evaluation setup, label collection, and evaluation metrics as shown in Figure 1. We provide pathology-specific recommendations for each of them in the following sections.

2.1 Evaluation Setup

2.1.1 Alignment of Evaluation Experiments with Model’s Intended Use

The applications of ML in pathology vary by their intended use, such as drug discovery, diagnostics, prognostics, decision-support, or triage. Therefore, their evaluation setup and metrics must account for the targeted use-case. For example, a clinical decision support tool is used by pathologists to look at predictions generated by the model to refine their diagnosis. The evaluation setup must measure the change in pathologist performance when assisted by the model instead of comparing model predictions against that of the pathologist.

Another example is the use of ML in building a triage tool for screening patients. In oncology, ML can be used to filter out obviously benign cases so that the pathologists prioritize malignant cases, thus improving patient care. In such scenarios, instead of using common metrics like accuracy or AUROC, one should use Precision@K%Recall due to the low tolerance for a false negative.

2.1.2 Enriched and Confounder-Aware Test Set Creation

Pathology samples come with biological and other metadata like patient-related (age, gender, ethnicity), image-related (scanner type, image format, magnification), specimen & stain related (tissue collection type, thickness of specimen, staining protocol), and clinical (disease stage, prescribed therapy, omics data). Low prevalence of certain metadata values coupled with small, imbalanced datasets in pathology often lead to overoptimistic results if the test set composition does not consider these factors.
not account for smaller substrata. This hidden stratification problem \citep{Oakden-Rayner2020} can result in the model not working well for patients belonging to a hidden substrata, which in turn deteriorates trust in such tools. Another stratum which needs more representation is samples close to the decision boundary where significant ambiguity in labels exist. This occurs since labels in pathology represent a biological process which is continuous. For example, along with benign and malignant there exist borderline cases which might be under-represented if real-world data distribution is replicated in test dataset.

A related problem is the presence of confounders in the test set which are spuriously predictive of the label. For example, in pancreatic cancer, \citep{Kather2020} show that KRAS mutation can be predicted from a pathology image. However, KRAS mutation is almost exclusively present in only one sub-type of pancreatic cancer (pancreatic ductal adenocarcinoma), thus becoming a strong biological confounder \citep{Waters2018}. Therefore, even a well performing mutation prediction model might be relying on pancreatic sub-type features instead of the KRAS mutation features. The fact that these confounders can also be partial, non-biological and specific to the dataset, further complicates matters \citep{Badgeley2019, Larrazabal2020, Zech2018}.

To ensure that the test-set is reliable and the model is predicting the variable of interest and not a confounder, we offer the following suggestions:

- Curate test sets such that they have enough representation of any sub-strata of data that we care about or that could be a confounder \citep{Seyyed-Kalantari2020}. This can be done through enrichment (oversampling) of rare substrata as mentioned in this FDA guide \citep{Food2017}. However practitioners should be aware of selection bias due to test-set enrichment \citep{Yu2020}.

- Incorporate stratified evaluation metrics as opposed to a single test-set wide global metric. For example, ensuring consistent performance across patient demographics or disease severity.

- Conduct a correlation check on all metadata values present in the dataset to see if any of them correlate highly with label values for the task of interest. For example, if all images positive for prostate cancer come from the same hospital, the model can learn to predict the hospital instead of the cancer, making it a potential confounder.

### 2.1.3 **EXTERNAL HOLD-OUT TEST SET**

Curated datasets are useful for research, but often suffer from the problem of a homogeneous test-set. The approach of collecting data from a single source and splitting it into train-val-test sets results in inflated performance since the models might not generalize to distributional shifts (like images from a new hospital) \citep{Tellez2019, Campanella2019}. Many popular pathology datasets \citep{Borkowski2019, Hosseiniv2019, Karimi2019} and studies \citep{Shao2021, Sudharshan2019} do not have externally held-out test sets from a new site which prevent an evaluation of the generalizability. We suggest using multiple held-out patient cohorts from different medical sites and demographics while constructing the test set \citep{Koh2021}. We also suggest checking for test data leakage in the form of patient samples being present across multiple splits. Finally, we urge the community to make such a testing setup the norm for model evaluation in pathology.

Figure illustrates the discordance among pathologists while grading NASH, an aggressive form of non-alcoholic fatty liver disease increasingly being seen as an epidemic in the US. In this case, NASH severity is found by performing a liver biopsy, viewing the extracted tissue through a microscope and observing signs of disease progression such as presence of fat, different kinds of inflammation etc.

What is most telling in this figure is the extremely low agreement among doctors while predicting disease progression even after using the most reliable way to do so, which is viewing the tissue extracted from the biopsy. The numbers on the left column denote the Kappa score for different morphological indicators of NASH (N denotes the number of patients), when the first study was done in 2005. The right column shows the agreement scores for the same from a 2020 study; there has been little or no change in these agreement scores among human experts even after 15 years.
(a) Figure illustrates the discordance among human experts while grading NASH from pathology images. NASH is an aggressive form of non-alcoholic fatty liver disease increasingly being seen as an epidemic in the US. Kappa score (agreement score) among pathologists for different morphological indicators of NASH are shown 15 years apart; there has been no improvement in pathologist consensus.

(b) Variations in pathology images across scanners and lab sites. The first row shows the same slide scanned with different scanners with visually perceptible differences. The second row shows slides collected from the same scanner but different lab sites, which differ visually due to staining protocols, choice of reagents, method of specimen preparation and so on. Models struggle to generalize to these variations.

Figure 2

2.2 EVALUATION LABELS

It is possible to get fast and cheap labels using crowdsourced tools for natural images, and the correctness of these labels is easily verifiable. In contrast, it is a lot harder to collect ground-truth labels for pathology. Pathology images have billions of pixels, millions of cells, and thousands of regions of interest, making label collection extremely resource-intensive. Annotators (typically certified pathologists) are in short supply and per unit cost of annotations is order of magnitude higher than natural images. Annotators also have their own biases (Aeffner et al., 2017). A more insidious problem is of the high ‘inter-annotator variability’ (Carrasco-Zevallos et al., 2021) as shown in Figure 2a and ‘intra-annotator variability’ (Kleiner et al., 2005; Davison et al., 2020): annotators disagree not only with each other but also with their past selves. These variabilities exist at different stages of label generation - they can originate from inherent ambiguity in the labeling system, different interpretations of the same labeling scheme by different experts, and the differences in the expertise of these annotators (Ozkan et al., 2016). The threshold for acceptable variability is subjective, and depends on factors such as the particular organ and disease.

We propose the following guidelines while collecting labels:

• Collect multiple annotations per image and measure the inter-annotator agreement. If possible, collect multiple annotations on the same image from the same annotator at adequately-spaced timepoints (called ‘washout period’) to measure intra-annotator variability. If these variabilities are high, the ML task might not be viable.

• Aggregate the labels to a single consensus if the intra and inter-annotator variabilities are low. Instead of comparing model performance with a single pathologist (Liu et al., 2017) or consensus, measure the agreement of the model with the consensus and judge it against inter-annotator consensus. This is more realistic and appreciates the difficulty of the task as judged by human’s inability to do it consistently (Raciti et al., 2020). Kappa (Linear, Quadratic) and Intraclass Correlation Coefficient (ICC) are popularly used to measure single and multi-rater agreement (Bulten et al., 2022; Koo & Li, 2016).

2.3 EVALUATION METRICS

2.3.1 REPRODUCIBILITY, REPEATABILITY, & ROBUSTNESS TESTS

Pathology images are heterogeneous with regard to staining, tissue thickness, and are prone to variations during tissue processing, cutting, staining, and digitization (Figure 2b). These variations cause minor visually perceptible changes to the image, but can drastically change the model prediction. Lack of robustness to these perturbations poses a major challenge to the wider adoption...
and use of ML in pathology, especially in a clinical setting [Schöning-Markiefka et al. 2021]. Accuracy based assessment of models, although necessary, is not sufficient for capturing model generalization. Following tests must be incorporated into the model evaluation framework to assess model’s robustness to these variations:

- **Reproducibility Tests:** Models must be tested under known sources of variations and evaluated for consistency and accuracy in performance. This includes testing by explicitly changing external variations such as staining & scanning, variations associated with hardware (non-determinism in GPU) and software packages. Models in pathology are usually trained on smaller patches of the image due to their large size [Shao et al. 2021; Taylor-Weiner et al. 2021]. Stochasticity introduced due to sampling the patches can lead to inconsistent predictions and robustness to this variability must also be reported.

- **Repeatability Tests:** Repeatability expresses the precision under the same operating conditions over a short interval of time [FDA 1995]. This test captures unknown & unaccountable variations that are not covered by reproducibility tests. Consistency in model predictions must be measured and reported by rescanning & rerunning inference on the same image multiple times, keeping all known sources of variations constant.

- **Robustness to Perturbations:** Creating test sets that capture all possible real-world variations of images is not practical. An alternative can be to evaluate robustness on synthetically generated perturbations inspired by real world variations to image samples, by measuring consistency of predictions across these perturbations [Hendrycks & Dietterich 2019; Schöning-Markiefka et al. 2021]. Models that are robust to synthetic variations have been shown to be robust to natural variations [Faryna et al. 2021].

### 2.3.2 Negative Control Tests

Cross domain differences in pathology images collected from different sources may confound the discovery of explanatory variables and lead to poor generalization. This phenomenon, called ‘batch effects’ [Goh et al. 2017], can lead to the model learning spurious features. Existing literature on batch effects show that models can predict patient age, sex, race, time of slide preparation, site, and scanner [Schmitt et al. 2021; Howard et al. 2020].

Although creating confounder-aware test sets can mitigate some of these issues, partial and hidden confounders may go unnoticed and affect model performance. Another approach to tackle this issue is to perform negative control tests which involve running experiments under conditions in which the model is not expected to produce correct predictions [Lipsitch et al. 2010]. We suggest the following strategies for designing negative control experiments for ML in pathology:

- **Train with regions of the image which have no discriminative information.** For example, a model trained to predict cancer using only background (non-tissue area) should perform poorly.

- **Train without regions which are essential for predicting the label.** For example, models for predicting cancer should perform poorly when trained and deployed on patches sampled from non-cancer regions.

Failure of a negative control test signals the presence of confounders and biases that create spurious correlations. These tests are not designed to detect all types of confounders, but they help detect the prominent ones and take appropriate actions.

### 3 Conclusion

Machine learning for pathology requires different evaluation standards due to differences in image characteristics and clinical use-case. Existing datasets and evaluation metrics prevalent within the ML community need to be rethought to fit pathology applications. We have observed that there exist two distinct communities, one of ML researchers focused on the development of novel methods and the other of computational pathology researchers focused on exploring and utilizing the power of ML in healthcare, and model evaluation differs between them materially. We argue that domain-specific knowledge from pathology concerning label reliability, test set enrichment, and biological confounders should permeate into ML research. Conversely, advancements in ML
such as synthetic data generation, robustness and sensitivity analysis, and out-of-domain testing deserve wider adoption in computational pathology. In this paper, we propose a set of guidelines for evaluating ML models in pathology which will help researchers create better datasets, metrics, and evaluation setup. We believe this will improve patient outcomes through ML applications in pathology and inform real-world motivated research.

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