

RESEARCH ARTICLE

Information theory approaches to improve glioma diagnostic workflows in surgical neuropathology

Lokman Cevik1 | Marilyn Vazquez Landrove2 | Mehmet Tahir Aslan1 | Vasili Khammad3 | Francisco Jose Garagorry Guerra4 | Yolanda Cabello-Izquierdo1 | Wesley Wang1 | Jing Zhao5 | Aline Paixao Becker1 | Catherine Czeisler1 | Anne Costa Rendeiro6 | Lucas Luis Sousa Veras6 | Maicon Fernando Zanon7 | Rui Manuel Reis78 | Marcus de Medeiros Matsushita9 | Koray Ozduman10 | M. Necmettin Pamir10 | Ayca Ersen Danyeli11 | Thomas Pearce12 | Michelle Felicella13 | Jennifer Eschbacher14 | Naomi Arakaki15 | Horacio Martinetto15 | Anil Parwani1 | Diana L. Thomas1 | José Javier Otero1

1Department of Pathology, The Ohio State University Wexner Medical Center, Columbus, Ohio, USA
2Mathematical Biosciences Institute, The Ohio State University, Columbus, Ohio, USA
3Peoples’ Friendship University of Russia, Moscow, Russia
4Facultad de Medicina, UdeLaR, Cátedra de Anatomia Patológica, Hospital de Clínicas Manuel Quintela, Universidad de la República, Uruguay
5Department of Biomedical Informatics, The Ohio State University College of Medicine, Columbus, Ohio, USA
6Pathology Department, Barretos Cancer Hospital, Barretos, Brazil
7Molecular Oncology Research Center, Barretos Cancer Hospital, Barretos, Brazil
8Life and Health Sciences Research Institute (ICVS), School of Medicine, University of Minho, Braga, Portugal
9MultiPat Laboratório de Anatomia Patológica, Campinas, Brazil
10Department of Neurosurgery, Acibadem MAA University, Istanbul, Turkey
11Department of Pathology, Acibadem University School of Medicine, Istanbul, Turkey
12Division of Neuropathology, Department of Pathology, University of Pittsburgh Medical Center, Pittsburgh, Pennsylvania, USA
13Division of Neuropathology, Department of Pathology, University of Texas Medical Branch, Galveston, Texas, USA
14Department of Pathology, Barrow Neurological Institute, St. Joseph’s Hospital and Medical Center, Phoenix, Arizona, USA
15Departamento de Neuropatología y Biología Molecular, Instituto de Investigaciones Neurológicas Dr Raúl Carrea (FLENI), Buenos Aires, Argentina

Correspondence
José Javier Otero, Department of Pathology, The Ohio State University College of Medicine, 4166 Graves Hall, 333 W 10th Avenue Columbus, OH 43210, USA. Email: jose.otero@osumc.edu

Abstract
Aims: Resource-strained healthcare ecosystems often struggle with the adoption of the World Health Organization (WHO) recommendations for the classification of central nervous system (CNS) tumors. The generation of robust
INTRODUCTION

The integration of molecular subtyping in brain tumor diagnosis for patients in low-resource settings represents a huge inequity in clinical care and is the premier leadership challenge facing the neuropathology community. Without a doubt, the integration of molecular data with histology has improved diagnosis and prognostication and assisted with neuro-oncology and radiation oncology treatment planning [1]. However, the deployment of these technologies globally is fraught with caveats. Per recent reports, the United States has less than 450 formally trained neuropathologists (and fewer with molecular diagnostics experience), and the number of pathologists per capita shows a negative trajectory over time [2]. This trend is worse throughout the world. Globally, pathologist training is under-resourced, pathology practice is poorly compensated, dedicated subspecialty training is nonexistent, and poorly implemented laboratory procedures for basic histochemical and immunohistochemical assays dampen the prospects of molecular assay deployment. Although some of the technical costs of molecular assays have declined, low-resource healthcare ecosystems also suffer from a paucity of clinical bioinformaticists [3]. This lack of human resource investment in clinical bioinformaticists may even be a more dire threat to molecular pathology deployment than the distribution of sequencing technologies. Furthermore, as primary brain cancers are rare, brain tumor-specific immunohistochemical biomarker testing is not commonly available even in some academic pathology laboratories. These realities directly conflict with the diagnostic recommendations put forth in the WHO Classification of CNS Tumors. Solutions that attempt to overcome the lack of investment in reagents, physical resources, and human resources are needed. For these reasons, a strong interest within neuropathology has emerged to implement machine learning/artificial intelligence-based workflows in routine clinical practice.

Alternative solutions, including centralized diagnostic services, have been attempted. For instance, Crosier et al. reported on a centralized neuropathology review strategy for medulloblastoma in the United Kingdom which evaluated histology, RNaseq, and DNA methylation data of ~80 medulloblastoma patients per year [4]. They reported a modified risk stratification

---

Diana L. Thomas, Division of Neuropathology, Department of Pathology, The Ohio State University College of Medicine and Nationwide Children’s Hospital, 700 Children’s Dive, Columbus, OH 43205, USA. Email: diana.thomas@nationwidechildrens.org

Funding information
National Institute of Neurological Disorders and Stroke. Grant/Award Number: R03-NS116334
for 29% of patients and provided a critical feasibility analysis of this model. However, such a centralized review may not be feasible in all healthcare ecosystems. For instance, in the United States, DNA methylation profiling is not widely available or clinically validated in many laboratories. Expanding testing to all brain tumors is highly problematic. Chief among these challenges is the well-known problem that diagnostic tests lose their predictive values when disease incidence decreases in the sampling population. This phenomenon most commonly occurs when increasing the number of subjects being tested without regard to pretest probability thresholds. Furthermore, in some clinical paradigms, next-generation sequencing (NGS) has shown to be cost-effective only at specific pretest probability thresholds [5]. Triaging tests based on the incorporation of clinical, histologic, and immunohistochemical findings have been proposed as a cost-effective solution [6].

With these concerns in mind, we focused on generating simple, easily deployable tools for assessing the most critical investment strategies for low-resource laboratories. To achieve this, we utilized information theory to identify the most critical data required for diagnostics and posit that simple information theory calculations can aid in determining investment strategies for biomarker acquisition.

2 | MATERIALS AND METHODS

2.1 | Brain cancer simulated population model

Our brain cancer population simulation is composed of 51 diagnostic entities obtained from the WHO Classification of CNS Tumors [7]. Data on these features were extracted from the scientific literature to generate a simulation of patient populations with these entities. The clinical features, the histologic features, immunohistochemical markers, and molecular features were delineated in Supporting Information with all references (summary of each entity is available in Figures S1–S51). The importance of each feature in the diagnosis of this simulation model was performed by separating all the data into a training set (70%) and a test set (30%). The randomForest algorithm was called from R’s randomForest package using the default values, and the efficiency of the model was tested by evaluation of a confusion matrix by the confusionMatrix function call in R’s caret, which showed that this model had an accuracy of 0.95 (95% CI = [0.952, 0.9564], no information rate = 0.068, and \( p < 10^{-15} \)). We then split the data frame by diagnosis and for each diagnosis, passed the random forest classifier followed on the original data frame. For each feature of the data frame, we scrambled each diagnostic feature by performing a random sampling of the original data frame containing all diagnoses and saved the value of the accuracy generated by random forest classifier. These data are presented in Table S1.

2.2 | Real-world data sets for information theory validation

Nine different data sets from different countries were used for the calculations of Bayesian probabilities and mutual information. The numbers of available cases and details were delineated in Table 1.

2.3 | Age distribution and probability calculation

Age distribution of IDH1/2-mutant gliomas was presented as density plots. For the Bayesian probability plots, probabilities of 1p/19q codeletion in IDH1/2 mutant tumors given age and gender were calculated using the “naivebayes” package in R with a Laplace correction for smoothing.

2.4 | Information theory calculations

Information theory, initially developed by Claude E. Shannon at Bell Labs in 1948 [8], rests on the notion that information can be objectively quantified into units called bits (or nats, where 1 nat = log2(e)*bit). Mutual information is the exact opposite term of entropy, where entropy represents the uncertainty of the information. Maximum mutual information indicates all information quantity needed for the 1p/19q codeletion status (or diagnosis) in our example. For simplicity, the provided information with features is represented as a percentage of maximum mutual information (as in Figures 1, 2, and 5). Details were delineated in Supporting Information.

2.5 | Image acquisition

Hematoxylin–eosin (H&E)-stained sections and p53 and Olig2 immunohistochemistry-stained sections of IDH-mutant diffuse gliomas were selected for image segmentation. Details were delineated in Supporting Information.

2.6 | Segmentation methods (unsupervised and supervised) and evaluation of segmentation fidelity

Two types of segmentation methods were used in our analysis: unsupervised (Deconvolution combined with Otsu thresholding, K-means, and the Cut-Cluster-Classify) and supervised methods (Trainable Weka
Segmentation and U-Net). We made a ground truth using GIMP and R for the evaluation of segmentation fidelity and U-Net training. Details were delineated in Supporting Information.

2.7 Evaluation of segmentation fidelity

For our evaluation, we only have two regions of interest: cell nuclei (stained in purple for H&E images and brown for Olig2 and p53 images) and background. Since one of the challenges of segmenting our data set comes from the various levels of noise, we also developed an image noise assay to test how robust the methods are to noise. Percentage ratios of added noise were selected as 0, 5, 10, 15, 25, 50, 75, 90 to evaluate the whole pattern of noise resistance. We report accuracy and intersection over union (IoU). Details were delineated in Supporting Information.

2.8 Data processing

2.8.1 Condensation of data and probability calculation of Olig2 features

Eighty-nine features were extracted from the nuclei in each channel of the images using the masks after the segmentation with the deconvolution method. After feature extraction, we excluded incomplete nuclei in the images for a better representation. Extracted features are shape, moment, pixel intensity, and Haralick’s features. We had 213 features left after the elimination of repeated features in different channels. After principal component analysis (PCA) application, 8 principal components (PCs) in Olig2 features and 9 principal components (PCs) in H&E features were selected for the representation of 90% variation of features. We calculated the sample density of nuclei in a multidimensional space of PCs. Three hundred and fifty nuclei were sampled in each case for density calculation of features extracted from H&E- and Olig2-stained images. The mean, median, and standard deviation values of the distances to 10th (d10), 20th (d20), 40th (d40), and 60th (d60) closest nuclei were calculated in a distance map as “density features.” The histogram parameters, such as mean, standard deviation, kurtosis, and skewness, on selected 3 shape features (major axis, eccentricity, and area) were used as “histogram features.” The mutual information of these density and histogram features of H&E- and Olig2-stained slides were used for feature selection.

2.8.2 Automated p53 positivity workflow

An automated positivity workflow for p53 immunoreactivity was developed for objective quantification. The detailed workflow was explained in the Results section. Briefly, we did segmentation on 1 image from positive control, 1 image from negative control, and 4 images from the tumor on the slide. After feature extraction, we labeled nuclei in the positive and negative control and used these labels as ground truth to make an RF modeling for the prediction of nuclei in the tumor images. After prediction, the positivity percentage was calculated as a percentage of positive nuclei to the total. We compared the automated positivity workflow results of p53 with human observers’ evaluations. These human observers were practicing neuropathologists at different hospitals who utilized slightly distinct interpretation workflows for p53 immunohistochemistry. Each neuropathologist interpreted the p53 immunohistochemistry...
FIGURE 1  Different clustering patterns with clinical, histologic, immunohistochemical, and molecular information on the brain cancer population simulation and information gain in the glioma simulation model for clinical decision-making. Dimensionality reduction by principal component analysis is shown in A1–A4, and by t-stochastic neighbor embedding in B1–B4, with the features delineated on the top of each graph. Each color represents a unique diagnosis in the WHO classification scheme. (A1 and B1) Dimensionality reduction with clinical features alone demonstrates only a few visible clusters. (A2 and B2) Incorporating clinical history with histology generates the commencement of clear layering in PCA and discrete clusters with t-SNE. (A3 and B3) Inclusion of immunohistochemical data improves the capacity of discerning clusters. (A4 and B4) The additional molecular features does not significantly improve the clustering. (C) Individual information gains with all clinical, histologic, immunohistochemical, and molecular features in the glioma simulation model. Age, site, and Ki67 are the features that have the most amount of necessary information for the diagnosis. (D) Information gains with clinical, histologic, immunohistochemical, and molecular information. Histology provides most of the necessary information for the diagnosis. Combining the clinical data with histology and immunohistochemistry provides more than 95% of the necessary information, whereas adding molecular data provides minimal information gain. (E) IDH1/2-mutated tumors were subsetted, and the mutual information of the features on the X-axis was quantified as % information on the Y-axis. The conditional information function call was called on data the 1p19q status.
as indicative of “mutant” or “wild type” in 82 cases of p53 images. After completion, the same 74 cases (8 cases do not have ATRX images) were evaluated with ATRX images to detect any bias of ATRX information on p53 positivity results. For models utilized in Figure 6, the continuous scale %p53 positive cells obtained from our automated workflow was utilized.

3 | RESULTS

3.1 | Information theory enhances understanding of individual feature impact in diagnostic neuropathology

Our ultimate goal is to generate tools and workflows that improve pathologist decision-making throughout the world, with a particular emphasis on resource-poor settings. To this end, we generated a brain cancer population simulation, whereby we obtained specific features known to neuropathologists who are capable of distinguishing tumors and performed a thorough review of the scientific literature to obtain the data upon which to build this simulation (see Supporting Materials and Methods). We utilized three easily obtainable clinical features (age, neuroanatomical site, and gender), histologic features, and immunohistochemical stains (GFAP, ATRX, Ki67, IDH1 R132H, P53, CD34, EMA, Olig2, reticulin, and synaptophysin). We also utilized the results of basic molecular assays that are commonly reimbursed by private insurers, public aid, and Medicare, in the United States. All features for each entity simulated are delineated in the Supporting Materials and Methods. We first performed dimensionality reduction using principal component analysis (PCA) as well as a nonlinear algorithm, the tSNE, on our simulated brain tumor population (Figure 1A,B). We note that plotting tumors only on clinical features (age, site, and gender) with tSNE and PCA results in no discernable cluster generation (Figure 1A1,B1).

![Figure 2](image)

**FIGURE 2** A global IDH-mutant (Grades 2 and 3) data set was used for the application of information theory as real-world data and the importance of selected features for 1p/19q codeletion status. (A) IDH-mutant astrocytoma and oligodendroglioma cases have a different distribution of age. Oligodendroglioma cases are more common in advanced age. (B) Probability of being an oligodendroglioma increases with age. (C) There is not a significant difference in the gender for the probability of 1p/19q codeletion. (D) Correlation plot of features as a dummy variable. p53 and ATRX mutation have a strong correlation and negative correlation with 1p/19q codeletion. Gender does not show any correlation with other features including 1p9q codeletion. (E) Dimensionality reduction of the features with a PCA shows similar results to the correlation matrix, which is similar variation between the p53 and ATRX mutation. (F) p53 and ATRX mutation information provide significant information for the diagnosis. Adding ATRX to p53 mutation provides additional information even though there is marked redundancy between these features. Permutation analysis was used for significance analysis. *p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001; ****p ≤ 0.0001.
from H&E-stained sections, one is able to visibly appreciate the generation of specific clusters of entities (Figure 1A2,B2). The addition of immunohistochemistry further increased the clusters relative to clinical features plus immunohistochemistry. However, the addition of molecular features did not necessarily increase the clustering upon visual inspection (compared Figure 1A4,B4 with Figure 1A3,B3). These data are in-line with well-known concepts in the neuropathology community that emphasize the integration of clinical context with histology. We further conclude that this simulation provides a useful in silico framework to test workflow concepts by Monte Carlo-like simulations prior to real-life deployment.

Then we generated a thought experiment where we imagined a scenario where the biopsy/resection procedure represented a conduit of data transmission from the patient to the pathologist. This scenario is analogous to the flow of information in information processing and amenable to information theory calculations (see Section 2). We evaluated all the features delineated in our brain cancer simulation model and quantified the mutual information (Figure 1C). We note that age, neuroanatomical site, and Ki67 proliferation index were the features that carried the most amount of information in the glioma simulation model. A combination of individual features showed that histology data provided nearly 80% of the information necessary for diagnosis (Figure 1D). Surprisingly, basic clinical information (age, site, and gender) provided just over half of all information necessary for a diagnosis, whereas molecular data alone provided less than 20% of the information needed. Combining clinical data with histology data and immunohistochemistry increased the information to over 95%, with only minimal information gained by the molecular data. To identify the most critically important features for each diagnosis in our simulation, we generated a randomForest based workflow that identified the most critical diagnostic features in this simulation (Table S1). After evaluating the information theory calculations and the results of our randomForest analysis, we note a significant asymmetry in the type of information contained in patient data, with the neuroanatomical site, age, and Ki67 labeling index being the most important features in diagnosis. This asymmetry produces a significant tendency to generate bias in machine learning-based diagnostic algorithms. In order to illustrate how such simulations can be used for strategic planning, we subsetted the simulated brain cancer population for IDH-mutated tumors and performed information theory calculations to determine the utility of age, gender, site, IDH status, 1p/19q status, and immunohistochemical detection of Olig2, GFAP, CD34, synaptophysin, p53, and ATRX (Figure 1E). We noted that p53 and ATRX status provided a significant quantity of information as to 1p/19q status. We further conclude that basic information theory calculations permit the rapid acquisition of which features are most impactful for diagnostic neuropathology.

3.2 | Age, p53, and ATRX status have significant contribution in the diagnosis of IDH-mutant tumors

Then we focused on IDH-mutant tumors to validate these information theory-focused workflows. To achieve this, we took advantage of publicly available data sets from the Cancer Genome Atlas and the Chinese Glioma Genome Atlas as well as data from hospitals in Turkey, the United States, Argentina, and Brazil to provide a global perspective (see map in Figure S52). The data sets collected included patient age (continuous numeric), gender (binary), p53 (binary), ATRX (binary), and 1p/19q codeletion status (binary), summing to 1390 total IDH-mutant gliomas. We noticed that IDH-mutant gliomas showed a different distribution of patient age. Generating a probability distribution of diagnosis as a function of age, we noticed an increase in the probability of 1p/19q codeletion with advanced age (Figure 2A,B). Confirming other studies, we found no association of gender with 1p/19q codeletion status (Figure 2C).

We then evaluated the relationship of these biomarkers to 1p/19q codeletion status by two linear methodologies: correlation matrix and principal component analysis (PCA). To achieve this, we generated dummy variables of the features. Figure 2D shows that ATRX and p53 mutations show a high correlation with each other and, as expected, are highly negatively correlated with 1p/19q codeletion status. Then we interrogated the data variance by PCA. As shown in Figure 2E, PC1 and PC2 account for over 70% of the variance in the data set. By plotting the eigenvectors along PC1 and PC2, we can appreciate the significant redundancy between ATRX and p53 status in the PC1/2 graph (note the near superimposition of the ATRX and p53 eigenvectors in Figure 2E). Also, the eigenvector for gender shows a nearly orthogonal orientation relative to 1p/19q codeletion eigenvector, indicating no association. These evaluations raise the possibility that data for P53 and ATRX show significant redundancy, and if so, assessing both biomarkers would be unnecessary in diagnostic workflows. To test this hypothesis, we performed information theory calculations of these parameters (Figure 2F). We found a statistically significant increase in information when both ATRX and p53 were evaluated together, a finding in-line with current cIMPACT-NOW clinical practice [9]. This means new information gain is present when combining both p53 and ATRX data points. We conclude that information theory analysis is capable of determining if biomarkers showing significant correlation and covariance add significant information when used in combination. It also provides significant additional insight into linear computational methods.
3.3 Auto-adjustable image analysis workflows for surgical neuropathology

We next sought to evaluate the information gain extractable from tissue morphology. A well-appreciated morphologic distinction between IDH-mutant gliomas is the propensity of the former to show significantly greater nuclear pleomorphism. We first evaluated Olig2 expression by immunohistochemistry, a biomarker diffusely expressed in infiltrating gliomas [10–12], and which was a commonly utilized biomarker among our global collaborative group. Olig2's nuclear localization also facilitates image segmentation. We then evaluated a variety of supervised and unsupervised segmentation algorithms, which included (1) matrix deconvolution with OTSU thresholding [13, 14], (2) WEKA segmentation [15], (3) k-means segmentation, (4) Cut-Cluster-Classify segmentation [16], and (5) a convolutional neural network with a UNET architecture [17]. We also performed a separate analysis utilizing a UNET algorithm trained with experimental noise (see Figure S57 for Image Noise Assay workflow and comparison of segmentation methods).

Our data were derived from the OSU patient archives. We note that matrix deconvolution with OTSU thresholding showed high accuracy up until 25% noise for both H&E staining and Olig2 immunohistochemistry (Figure S57B,C, blue line), whereas UNET trained with significant noise was able to maintain high accuracy with high levels of noise. However, when evaluating IoU, both methods showed significant drop-off with more than 25% noise (Figure S57B,C). We conclude that for simple segmentation processes where the high signal to noise exist, matrix deconvolution with OTSU thresholding is the simplest approach.

We then generated image masks of all Olig2-stained nuclei and overlaid the image masks onto the red, green, and blue image matrices to extract features of cell nuclei. These features included shape, pixel intensity, and pixel texture representing over 200 features per nucleus. To reduce the dimensions, we performed PCA which provided 90% of the variance in 8 principal components for Olig2 staining. Evaluation of morphologic features in some cases showed a tendency for oligodendroglioma cases to show higher kurtosis in histograms describing nuclear shape features (Figure 3B). However, the absolute values of these parameters differed on a case-by-case basis, and we were not able to use these raw measurements. We also felt that the pleomorphism among cells likely also included marked variance in DAB-staining intensity and texture (e.g., compare astrocytoma staining pattern for Olig2 in Figure 3A). We, therefore, evaluated the principal components in multidimensional space (Figure 3C shows the workflow). An example of our analysis is shown in Figure 3D for a sample case of IDH-mutant astrocytoma and oligodendroglioma. Note that the IDH-mutant astrocytoma shows a wide scatter of coordinates, thus showing a lower data density. In contrast, oligodendroglioma is more heavily clustered in one area. This dense packing of the oligodendroglioma nuclei represents a reflection of the nuclear monomorphism in oligodendroglioma relative to astrocytoma. We calculated the mean Euclidean distance of each cell to its 10th, 20th, 40th, and 60th neighbor in the multidimensional space (Figure 3E). Data density calculations from Euclidean Distance capture the nuclear pleomorphism differences between IDH-mutant gliomas. We further posit that this methodology, which is based on within-sample differences, represents a superior metric for evaluating morphology as it would be more resistant to interlaboratory methodologies.

We next focused our analysis on features of p53 immunohistochemistry, an important biomarker for glioma classification. Intense p53 nuclear immunoreactivity is utilized as a surrogate for TP53 mutation but suffers significant limitations. In addition to some inactivating TP53 mutations resulting in complete negative staining [18, 19], Takami, et al. demonstrated that in addition to immunoreactivity intensity, percent of immunoreactivity was also a critical feature [20]. However, other studies have set a cutoff of positivity for p53 positivity by IHC as 10% regardless of intensity in other tumor paradigms [21]. Generating a universal intensity and texture cutoff for p53 immunohistochemistry is challenging due to variance in laboratory practices. In our institution, we routinely perform our glioma-related
p53 immunohistochemistry with a serous carcinoma positive control and a benign lymph node as a negative control, mounted on the same slide. We utilized these internal controls as our metric for the designation of p53 positive or negative for each case (see the workflow in Figure 4A). The mean accuracies of random forests
for each case are 0.991 (95% CI: 0.985–0.996). Once the model is trained, we passed it on to the actual data from the patient to generate the prediction. For benchmarking, we compared the performance of our model to two human neuropathologists (see Table 2 for concordance data). We note that Lin’s concordance for interobserver neuropathology recordings showed overlap in the 95% CI of neuropathology observer 1: automated workflow and neuropathology and observer 2: automated workflow. These data indicate that the automated workflow shows similar a concordance level to neuropathologists as neuropathologists concord with each other. We next tested the extent to which mean pixel intensity extracted from the nuclei were distinct between the astrocytoma and oligodendroglioma cases in the red, green, and blue channels (Figure 4B). Mean pixel intensity distribution showed a clear distinction between the tumor nuclei and control nuclei for the modeling. Figure 4C demonstrates the distribution of p53 positive cells, as determined by our random forest classifier for each case. The variance between the p53 quantification is quite large in the astrocytoma cases, whereas in the oligodendroglioma, the majority of the cases showed low %p53 detection. Based on these data, we then generated a probability function using the naïve Bayes function in R (Figure 4D). As %p53 positivity increases, the probability of 1p/19q codeletion in our data set decreases in an exponential manner.

**Figure 5A** demonstrates information theory calculations for this p53 continuous-scale data set. Automated p53 quantification using this workflow represents an objective alternative to subjective p53 interpretation. Also, the addition of ATRX information to p53 positivity percentage provides a significant information increase for diagnosis. We also used a mutual information approach to determine the significant features in the histogram and density features of H&E- and Olig2-stained images (Figure 5B,C). There are 12 significant Olig2 features (Figure 5C), whereas 3 significant H&E features were noted (Figure 5B).

### 3.4 | Informatics-based clinical practice guidelines for the evaluation of IDH-mutant tumors

We next trained a random forest classifier that used the significant Olig2 features of data density, p53 labeling (continuous scale quantification), and a combined ATRX/age feature on 82 cases. The ATRX/age feature was obtained from probability estimates of a random forest classifier trained on 1107 cases of the global data set after splitting the train (70%) and test (30%) sets, where 1p/19q codel status was modeled as a function of ATRX and patient age. This ATRX/age feature was calculated...
We note that in our OSU archival data set, missing (3 cases) or equivocal (2 cases, 1 on 82 cases, 1 on validation data set) ATRX status existed, and so we imputed the ATRX status based on a random forest classifier trained on the global data set, where ATRX status was modeled as a function of age and p53 status. For our 82 cases, we used an 80% threshold to convert the p53 continuous scale to a binary result (positive or negative) for this prediction. Then we tested the random forest classifier on the testing set (~30%, representing 25 cases). Our results showed an accuracy of 0.92%, \( p = 0.000022 \). We then implemented this algorithm on a series of 11 diagnostically challenging validation cases (Figure 6A and summarized in Table S2), most of which represented tumor recurrences. Detailed clinical histories of these patients are delineated in the

**TABLE 2** Comparison of p53 scoring systems

| Comparison                                      | CCC   | 95% CI   |
|------------------------------------------------|-------|----------|
| Observers–Workflow comparison                  |       |          |
| Observer 1 vs. Workflow                        | 0.60  | 0.46–0.72|
| Observer 2 vs. Workflow                        | 0.56  | 0.44–0.67|
| Observer 1 vs. Observer 2 vs. Workflow          | 0.63  |          |
| Observer–Observer with ATRX comparison          |       |          |
| Observer 1 vs. Observer 1 with ATRX            | 0.66  | 0.51–0.77|
| Observer 2 vs. Observer 2 with ATRX            | 0.93  | 0.90–0.95|
| Observer 1–Observer 2 comparison               | 0.78  | 0.70–0.85|
| Observer 1 vs. Observer 2                       | 0.73  | 0.61–0.82|

**FIGURE 5** Information theory of features (age, gender, p53 continuous scale, and ATRX) on the OSU data set and significance of H&E and OLIG2 features for diagnosis of IDH-mutant Gliomas. (A) Comparison of information gained on 82 cases. p53 positivity percentage from automated workflow and binary ATRX mutation results from the reports show the highest significant information gain for the diagnosis. Adding ATRX information to the p53 positivity percentage provides significant information although there is a big redundancy between them. (B and C) 10 Olig2 features provide significant information for the diagnosis although there is only 1 H&E feature that provides significant information. These 10 significant Olig2 features were used for modeling in later applications.

\[ \text{Information (percent of total)} \]

\[ n = 82 \text{ patients} \]

\( \text{H&E features} \)

\[ n = 67 \text{ patients} \]

\( \text{OLIG2 features} \)

\[ n = 82 \text{ patients} \]
FISH false positives. 1p/19q FISH false positives are the codeletion assay so as to avoid the potential of 1p/19q triggering a chromosomal microarray or NGS-based 1p/19q grade morphology, a model prediction of 0.5 should rate low-grade astrocytoma without further workup. If the low-grade 1p/19q codeletion, the case can be easily diagnosed as an mutant tumor, if our model prediction is less than 0.5 for (Figure 6B). We note that our algorithm is quite robust of expensive nonreimbursable modern molecular tests guideline to facilitate decisions related to the ordering. Based on these results, we propose a clinical practice formulation framework upon which to determine the features that offer the most information to reach an integrated neuropathologic diagnosis. We tested this hypothesis first with a highly comprehensive simulation of brain cancer patients and validated this approach using real-world data. We developed, validated, and deployed a simple image analysis workflow that could be adopted in resource-poor settings using open-source software as well as translated to a variety of diagnostic applications and proposed a clinical practice guideline recommendation in the evaluation of IDH-mutant tumors. We note instances in which there is a discordance between 1p19q FISH and 1p19q analysis by chromosomal microarray or next-generation sequencing as false-positive cases with regard to the diagnosis of oligodendroglioma.

 Supporting Information. Our model incorrectly classified four astrocytoma cases (Case 2, Case 3–2, Case 4, and Case 10). All the incorrectly classified cases showed either high-grade architectural features (necrosis and microvascular hyperplasia) or represented recurrent cases. Based on these results, we propose a clinical practice guideline to facilitate decisions related to the ordering of expensive nonreimbursable modern molecular tests (Figure 6B). We note that our algorithm is quite robust for low-grade cases. In the setting of a low-grade IDH-mutant tumor, if our model prediction is less than 0.5 for 1p/19q codeletion, the case can be easily diagnosed as an astrocytoma without further workup. If the low-grade glioma shows a prediction of >0.5, 1p/19q codeletion by FISH is a sufficient diagnostic test to diagnose accurately IDH-mutant astrocytoma and oligodendroglioma. For IDH-mutant tumors that are recurrent or show high-grade morphology, a model prediction of >0.5 should trigger a chromosomal microarray or NGS-based 1p/19q codeletion assay so as to avoid the potential of 1p/19q FISH false positives. 1p/19q FISH false positives are the

4 | DISCUSSION

In this study, we implemented, for the first time, information theory approaches as a conceptual and practical framework upon which to determine the features that offer the most information to reach an integrated neuropathologic diagnosis. We tested this hypothesis first with a highly comprehensive simulation of brain cancer patients and validated this approach using real-world data. We developed, validated, and deployed a simple image analysis workflow that could be adopted in resource-poor settings using open-source software as well as translated to a variety of diagnostic applications and proposed a clinical practice guideline recommendation in the evaluation of IDH-mutant tumors. We note instances in which there is a discordance between 1p19q FISH and 1p19q analysis by chromosomal microarray or next-generation sequencing as false-positive cases with regard to the diagnosis of oligodendroglioma.
that few emerging concepts in modern pathology have experienced as much hype as machine learning and AI (for instance, see Ref. [22] and related its editorial Ref. [23]). We propose that our information theory approach represents a simple way for resource-challenged healthcare ecosystems to make informed decisions on reagent and testing platform investments. We further propose that in silico simulation, such as performed by our glioma population simulation, provides an inexpensive framework amenable to Monte Carlo-like simulation and permits the testing of how specific datapoints may affect the differential diagnosis. We posit that generating such Monte Carlo-like simulations can provide multiple insights to other surgical pathology paradigms. The workflow presented in the Supporting Information can easily be substituted by other tumor types and/or other types of information by creating an \( m \times n \) matrix in Excel with each row corresponding to a specific tumor entity, each column a feature/biomarker, and each cell a proportion of cases showing positivity for the designated feature/biomarker. Such a table would then be imported into Rstudio or Python to generate the simulation using a sampling function as delineated in the methods.

4.1 Implications of information theory on diagnostic neuropathology

The glioma diagnosis simulation and our information theory analysis raise some very interesting points regarding the future development and deployment of clinical diagnostic aids for neuropathology. At the forefront of our findings is the identification that simple information provides the majority of information. Specifically, basic clinical history, radiographic documentation of tumor site, and histologic findings carried the most information. This is rather intuitive, but these findings are of utmost importance for low-resource healthcare ecosystems. These findings are in-line with recommendations for low-to-middle-income countries (LMIC) that have emphasized local healthcare ecosystems to embrace a culture of quality in pathology laboratories, in particular, by adopting appropriate standard operating procedures, quality control, and quality assurance practices [24]. Underscoring this point, a comprehensive study by Fleming et al. generated a recommended capabilities list for Pathology Laboratories in LMICs. Chief among these recommendations included high-quality histologic/cytologic services as well as training for laboratory staff and physician-pathologists [25, 26]. Our findings from the brain cancer population simulation underscore the key importance of quality histology in diagnostic neuropathology. Building upon these recommendations, we would further advocate that evaluation of clinical, histologic, and biomarker features by information theory could provide a quantitative framework from which to make data-driven decisions on diagnostic services.

Our work raises significant concerns with ML-/AI-based clinical diagnostic aids in neuropathology. First, most brain cancer data set structures include high-resolution genomic data on very few samples. Pooling data sets from various centers results in missing data elements that are often beyond the capabilities of imputation for clinical outcomes research [27]. In addition, as demonstrated in our glioma simulation, the quantity of information present is highly asymmetrical, and many brain tumor entities are exceedingly rare. This results in ML-/AI-based algorithms, nearly all of which utilize some version of information theory, to focus on age, neuroanatomical site, and proliferative index for diagnosis. As a result, predictive algorithms may generate adequate differential diagnoses, but these would be of minimal utility in routine practice. Our approach to overcoming this problem is to identify specific, granular diagnostic conundrums as candidate branch points in clinical decision-making, where ML/AI can make an impact. This permits the focused evaluation of biomarker information with several advantages over traditional analyses. PCA and correlation plots indicated such a high correlation and covariance between p53 and ATRX status that it raised the possibility that these two biomarkers were fully redundant, legitimately questioning employing both antibodies. In contrast, using an information theory approach, we found that the incorporation of both biomarkers simultaneously resulted in a statistically significant information gain. This synergy in information gain was masked using traditional linear analyses. Similar evaluations could be employed to evaluate additional frequently paired biomarkers.

4.2 Simple image analysis workflows deployable throughout the world

The advent of CNN networks and image recognition has revolutionized the modern world, and it will continue to provide significant impacts to diagnostic pathology. However, its deployment in the practice of pathology globally has several limitations. First and foremost, although pathologists worldwide have access to brightfield microscopes equipped with cameras, few laboratories contain whole slide imaging (WSI)-capable scanners. In addition to capturing all of the tissue examined, WSI has the added benefit of imposing a standard operating procedure (SOP) in image acquisition, whereas traditional photomicroscopy is nearly impossible to be replicated from one session to another. In our study, we utilized tiles of image patches taken from WSI images to mimic pathologists capturing images on their desktop microscopes. A second challenge is that the lack of adherence to SOPs for histopathology across the globe results in highly variable interlaboratory fixation and staining protocols [28]. Although several attempts to correct this have been published (e.g., Ref. [29]), the nonlinearity in
histologic color maps has posed a significant challenge when controlling for fixation and staining conditions across labs [30] and may be unsurmountable in labs with high batch variance in staining. For this reason, we focused on incorporating self-normalizing workflows when dealing with color intensity and color texture features. One simple approach is to use internal known positive controls, such as p53 immunohistochemistry in ovarian serous carcinoma [31] with a known negative control (in our case, a benign lymph node), and train within each case an algorithm based on pixel intensity and texture features. This internal slide control provides an unbiased methodology. A second approach we took was using the concept of data density within multidimensional space for the evaluation of nuclear pleomorphism in glioma. These analyses can be performed across histopathology labs as they would only require image acquisition using the same objective magnification and numerical aperture.

4.3 | Machine learning-workflow improvement to WHO and cIMPACT-NOW clinical guidelines

Our test-case scenario dealt with the need to confirm 1p/19q codeletion status in IDH-mutant gliomas. In many healthcare ecosystems, even in the United States, no reimbursement for chromosomal microarray to confirm 1p/19q codeletion exists. In contrast, 1p/19q codeletion testing by FISH, although suffering in positive predictive value relative to more advanced techniques, reimburses routinely. This leads to a situation in which pathologists must make a decision to perform unreimbursed reflex testing in a serial fashion to confirm 1p/19q codeletion, further delaying treatment planning by neuro-oncology/radiation oncology. The cIMPACT guidelines, which are anticipated to be incorporated into the 2021 WHO classification, represent a significant improvement to prior recommendations [9,32]. Specific to our test case at hand is that current cIMPACT guidelines dictate that documentation of TP53 mutation and concurrent ATRX mutation within an IDH-mutant tumor excludes the diagnosis of oligodendroglioma. A specific problem arises when pathologists identify an IDH-mutant tumor, and either ATRX immunohistochemistry is unavailable, or ATRX immunohistochemistry is equivocal (as was shown in Case 4 of Figure 6A) or negative due to copy number loss involving the ATRX locus in a tumor with 1p/19q codeletion. Furthermore, the designation of TP53 mutation subjectively by immunohistochemistry is fraught with caveats [33]. We developed a predictive model as a clinical diagnostic aid that incorporates Olig2 features, % of cells with strong p53 immunoreactivity, and the probability based on age and ATRX status. The goal of this model is to provide guidance in the event that ATRX immunohistochemistry is either unavailable or equivocal. Our validation set (Figure 6) included 5 false-positive 1p/19q FISH results, only two of which could have been prevented by adherence to cIMPACT-NOW guidelines, whereas our improved guidelines handle all cases in the validation set and would prevent all false-positive 1p/19q in the assays we analyzed.

ACKNOWLEDGMENTS

DT and JJO acknowledge the National Institute of Neurological Disorders and Stroke (NINDS) R03-NS116334. JJO and CMC acknowledge support from the Ohio State University-Translational Data Analytics Institute (OSU-TDAI) seed grant.

CONFLICT OF INTEREST

The authors have no conflicts of interest to declare that are relevant to the content of this article.

AUTHOR CONTRIBUTIONS

Clinical context and clinical informatics: Diana L. Thomas, José Javier Otero, Lokman Cevik, Mehmet Tahir Aslan, Wesley Wang; Data analysis and data visualization: Catherine Czeisler, José Javier Otero, Lokman Cevik, Mehmet Tahir Aslan, Wesley Wang; Data acquisition: Francisco Jose Garagorry Guerra, Yolanda Cabello-Izquierdo, Wesley Wang, Jing Zhao, Aline Paixao Becker, Catherine Czeisler, Anne Costa Rendeiro, Lucas Luis Sousa Véras, Maicon Fernando Zanon, Rui Manuel Reis, Marcus de Medeiros Matsushita, Koray Ozduman, M. Necmettin Pamir, Ayca Ersen Danyeli, Thomas Pearce, Michelle Felicella, Jennifer Eschbacher, Naomi Arakaki, Horacio Martinetto, Aníl Parwani; Writing—original draft preparation: Diana L. Thomas, José Javier Otero, Lokman Cevik, Marilyn Vazquez Landrove, Wesley Wang; Writing—review and editing: all authors; Mathematical and statistical modeling: Marilyn Vazquez Landrove, Jing Zhao.

ETHICAL APPROVAL

This study involved an analysis of retrospectively collected individual patient information and digital images in patients with IDH-mutant gliomas. This study was approved by the ethics board of the Ohio State University and all the local ethics boards.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are openly available in “figshare” at http://doi.org/10.6084/m9.figshare.16493964.

ORCID

Wesley Wang https://orcid.org/0000-0001-7904-9015
Aline Paixao Becker https://orcid.org/0000-0002-8695-810X
Rui Manuel Reis https://orcid.org/0000-0002-9639-7940
Ayca Ersen Danyeli https://orcid.org/0000-0001-8015-9916
José Javier Otero https://orcid.org/0000-0003-0680-4520
REFERENCES

1. Horbinski C, Ligon KL, Brastianos P, Huse JT, Venere M, Chang S, et al. The medical necessity of advanced molecular testing in the diagnosis and treatment of brain tumor patients. Neuro Oncol. 2019;21(12):1498–508.

2. Robboy SJ, Gross D, Park YJ, Kittrie E, Crawford JM, Johnson RL, et al. Reevaluation of the US pathologist workforce size. JAMA Netw Open. 2020;3(7):e2010648.

3. Chasapi A, Promponas VJ, Ouzounis CA. The bioinformatics wealth of nations. Bioinformatics. 2020;36(9):2963–5.

4. Crosier S, Hicks D, Schwalbe EC, Williams D, Leigh Nicholson S, Smith A, et al. Advanced molecular pathology for rare tumours: a national feasibility study and model for centralised medulloblastoma diagnostics. Neuropathol Appl Neurobiol. 2021;47(6):736–47.

5. Torchia MT, Austin DC, Kunkel ST, Dwyer KW, Moschetti WE. Next-generation sequencing vs culture-based methods for diagnosing periprosthetic joint infection after total knee arthroplasty: a cost-effectiveness analysis. J Arthroplasty. 2019;34(7):1333–41.

6. Walsh K, Kheng YC, Oniscu A, Harrison DJ, Wallace WA. Could molecular pathology testing in lung cancer be more cost-effective? J Clin Pathol. 2016;69(10):938–41. https://doi.org/10.1136/jclinpath-2016-203811

7. Louis DN, Olgaki H, Wiestler OD, Cavenee WK, World Health Organization, International Agency for Research on Cancer. WHO classification of tumours of the central nervous system. Lyon: International Agency for Research on Cancer; 2016.

8. Shannon CE. A mathematical theory of communication. Bell Syst Tech J. 1948;27(3):379–423.

9. Brach Allade K, Colman H, Holland EC, Louis DN, Jenkins RB, et al. cIMPACT-NOW update 3: recommended diagnostic criteria for “Diffuse astrocytic glioma, IDH-wildtype, with molecular features of glioblastoma, WHO grade IV”. Acta Neuropathol. 2018;136(5):805–10.

10. Otero JJ, Rowitch D, Vandenberg S. OLIG2 is differentially expressed in pediatric astrocytic and in ependymal neoplasms. J Neurooncol. 2011;104(2):423–38.

11. Ligon KL, Alberta JA, Kho AT, Weiss J, Kwaan MR, Nutt CL, et al. Advanced molecular pathology for rare tumours: a national feasibility study and model for centralised medulloblastoma diagnostics. Neuropathol Appl Neurobiol. 2021;47(6):736–47.

12. Ligon KL, Alberta JA, Kho AT, Weiss J, Kwaan MR, Nutt CL, et al. The medical necessity of advanced molecular testing in the diagnosis and treatment of brain tumor patients. Neuro Oncol. 2019;21(12):1498–508.

13. Chasapi A, Promponas VJ, Ouzounis CA. The bioinformatics wealth of nations. Bioinformatics. 2020;36(9):2963–5.

14. Crosier S, Hicks D, Schwalbe EC, Williams D, Leigh Nicholson S, Smith A, et al. Advanced molecular pathology for rare tumours: a national feasibility study and model for centralised medulloblastoma diagnostics. Neuropathol Appl Neurobiol. 2021;47(6):736–47.

15. Arganda-Carreras I, Kaynig V, Rueden C, Eliceiri KW, Cardona A, et al. Trainable Weka Segmentation: a machine learning tool for microscopy pixel classification. Bioinformatics. 2017;33(15):2424–6.

16. Vazquez Landrove MY. Consistency of density based clustering can serve as a surrogate marker for TP53 mutations in ovarian carcinoma: an immunohistochemical and nucleotide sequencing analysis. Mod Pathol. 2011;24(9):1248–53.

17. Takami H, Yoshida A, Fukushima S, Arita H, Matsushita Y, Nakamura T, et al. Revisiting TP53 mutations and immunohistochemistry—a comparative study in 157 diffuse gliomas. Brain Pathol. 2015;25(3):256–65. https://doi.org/10.1111/bpa.12173

18. Coates AS, Millar EKA, O’Toole SA, Molloy TJ, Viale G, Goldhirsh A, et al. Prognostic interaction between expression of p53 and estrogen receptor in patients with node-negative breast cancer: results from IBCSG Trials VIII and IX. Breast Cancer Res. 2012;14(6):R143.

19. Nagendran M, Chen Y, Lovejoy CA, Gordon AC, Komorowski M, Harvey H, et al. Artificial intelligence versus clinicians: systematic review of design, reporting standards, and claims of deep learning studies. BMJ. 2020;368:m689.

20. Rampton V. Artificial intelligence versus clinicians. BMJ. 2020;369:m1326.

21. Sayed S, Lukande R, Fleming KA. Providing pathology support in low-income countries. J Glob Oncol. 2015;1(1):3–6.

22. Fleming KA, Naidoo M, Wilson M, Flanagan J, Horton S, Kuti M, et al. High-quality diagnosis: an essential pathology package. In: Jamison DT, Gelband H, Horton S, Jha P, Laxminarayan R, Mock CN, Nugent R (eds). Disease control priorities: improving health and reducing poverty. Washington, DC: The International Bank for Reconstruction and Development/The World Bank; 2017.

23. Fleming KA, Naidoo M, Wilson M, Flanagan J, Horton S, Kuti M, et al. An essential pathology package for low- and middle-income countries. Am J Clin Pathol. 2017;147(1):15–32. https://doi.org/10.1093/ajcp/aqw143

24. Jakobsen JC, Gluud C, Weterslev J, Winkel P. When and how should multiple imputation be used for handling missing data in randomised clinical trials – a practical guide with flowcharts. BMC Med Res Methodol. 2017;17(1):162.

25. Gray A, Wright A, Jackson P, Hale M, Treanor D. Quantification of histochemical stains using whole slide imaging: development of a method and demonstration of its usefulness in laboratory quality control. J Clin Pathol. 2015;68(3):192–9.

26. Bautista PA, Yagi Y. Staining correction in digital pathology by utilizing a dye amount table. J Digit Imaging. 2015;28(3):283–94.

27. Jakobsen JC, Gluud C, Wetterslev J, Winkel P. When and how should multiple imputation be used for handling missing data in randomised clinical trials – a practical guide with flowcharts. BMC Med Res Methodol. 2017;17(1):162.

28. Fauzi MFA, Gokozan HN, Elder B, Puduvalli VK, Pierson CR, Fauzi MFA, et al. Optimized p53 immunohistochemistry is an accurate predictor of TP53 mutation in ovarian carcinoma. J Pathol Clin Res. 2021;23(8):1231–51.

29. Köbel M, Piskorz AM, Lee S, Liu S, LePage C, Marass F, et al. Optimized p53 immunohistochemistry is an accurate predictor of TP53 mutation in ovarian carcinoma. J Pathol Clin Res. 2021;23(8):1231–51.

30. Kather JN, Weis C, Marx A, Schuster AK, Schad LR, Zöllner FG, et al. New colors for histology: optimized bivariate color maps increase perceptual contrast in histological images. PLoS One. 2015;10(12):e0154572.

31. Köbel M, Piskorz AM, Lee S, Liu S, LePage C, Marass F, et al. Optimized p53 immunohistochemistry is an accurate predictor of TP53 mutation in ovarian carcinoma. J Pathol Clin Res. 2021;23(8):1231–51.

32. Kather JN, Weis C-A, Marx A, Schuster AK, Schad LR, Zöllner FG, et al. New colors for histology: optimized bivariate color maps increase perceptual contrast in histological images. PLoS One. 2015;10(12):e0154572.

33. Koibel M, Piskorz AM, Lee S, Liu S, LePage C, Marass F, et al. Optimized p53 immunohistochemistry is an accurate predictor of TP53 mutation in ovarian carcinoma. J Pathol Clin Res. 2021;23(8):1231–51.

SUPPORTING INFORMATION

Additional supporting information may be found in the online version of the article at the publisher’s website.

Fig S1-S58

FIGURE S1 Anaplastic astrocytoma, WHO grade 2, IDH-mutated

FIGURE S2 Anaplastic astrocytoma, WHO grade 2, IDH–wild type

FIGURE S3 Anaplastic ependymoma, posterior fossa A, WHO grade 3
FIGURE S43 Paraganglioma
FIGURE S44 Pilocytic astrocytoma
FIGURE S45 Pilomyxoid astrocytoma
FIGURE S46 Pleomorphic xanthoastrocytoma, WHO grade 2
FIGURE S47 Rosette-forming glioneuronal tumour, WHO grade 1
FIGURE S48 Schwannoma, WHO grade 1
FIGURE S49 Solitary Fibrous tumour/hemangiopericytoma grade 1
FIGURE S50 Subependymoma
FIGURE S51 Subependymal Giant cell astrocytoma, WHO grade 1

FIGURES S1–S51 Summary of each entity in simulation model. (A) Age distributions, (B) Neuroanatomical distributions, (C) Ki67 distributions, (D) Distribution of histological features, (E) Distribution of immunohistochemical features, (F) Distribution of molecular pathology features. Data represented in panels D, E, and F are violin plots of the distribution of the histological, immunohistochemical, and molecular biomarkers. Each biomarker is listed as a binary readout of 0 = negative, 1 = positive. The violin shape provides an indication of the relative proportion showing positive or negative results for this biomarker. In cases where the literature does not support a distribution, (for instance, necrosis in astrocytoma, a notch is placed at the level of the 0)

FIGURE S52 Distribution of cases in the global IDH mutant (Grade 2 and 3) dataset. World map shows the distribution of the cases in the global dataset. TCGA study (purple circles) has cases from different areas of the world such as Italy, Germany, Australia, but mostly the United States. This overall distribution of global dataset shows a good variety of ethnicity and genetic background to represent a real-world data

FIGURE S53 Deconvolution with Otsu's thresholding and Watershed. The images were splitted into R, G and B channels. Then R (red), G (green) and B (blue) channels were converted to H (hematoxylin), D (DAB) and X channels. D channel was selected for brown color in OLIG2 staining. After application of median filtering, Otsu's thresholding was applied to the D channel. The holes in the nuclei were filled out and then the nuclei on the border were eliminated for a better representation of nuclei. As a last step, watershed function was applied for overlapping or contacting nuclei.

FIGURE S54 K-means clustering segmentation. The images were converted to a data frame using R, G and B pixel intensity values. All pixels were clustered into 3 clusters using k-means clustering method. Each pixel was labeled as cluster 1, cluster 2, or cluster 3. The cluster that contain specifically the nuclei was chosen as foreground and the other 2 clusters were accepted as background.

FIGURE S55 Cut-Cluster-Classify (CCC) segmentation method. (A) All patches in the image are viewed as high dimensional points and a threshold is applied to the sample density (Cut) to get clusters. After clustering (Cluster), the patches below the threshold were classified.
based on the clusters (Classify). (B) Then all pixels were voted for clusters based on cluster result on overlapping patches. The cluster that contains nuclei was accepted as foreground and the other one was accepted as background. 5x5 patches were used for the segmentation.

**FIGURE S56** Relationships of R channel pixel intensity values with G and B pixel intensity values for color of noise. Mean pixel intensity values of R channel are correlated to mean pixel intensity values of G and B channels. To better represent these spectrums, we used support vector machine (SVM) modeling to predict G and B values from R. The colors of noise are purple for H&E staining and brown for OLIG2 staining.

**FIGURE S57** Image noise assay workflow and comparison of supervised and unsupervised image segmentation methods for H&E and OLIG2 immunostaining. (A) An image noise assay was created for the fidelity evaluation of image segmentation methods to determine the best one for our purpose. We extracted the purple color for H&E staining and brown color for OLIG2 staining. A support vector model (SVM) was made to generate a noise similar to colors of the image. After generation of noise for a full image, the pixels were randomly sampled with certain percentages to implement the noise to the images. This workflow provided us to compare the performance of segmentation methods with different amount of noise. (B and C) We plotted segmentation performance by accuracy and intersection-over-union (IoU). Additionally, we added the U-Net trained with random noise generation as “U-Net with noise” to the comparison. In U-Net with noise, we trained a new model every time when we added noise to the image. (B) In H&E staining, U-Net models show highest accuracies independent from the noise percentage. On the other hand, U-Net method shows low IoU, when U-Net with noise has a better IoU values. Overall deconvolution method has the better performance for accuracy and IoU. (C) In OLIG2 staining, similar to the H&E, deconvolution and U-Net with noise methods have the highest performance regarding accuracy and IoU. Despite deconvolution method has a sharp decrease after 25% noise, U-Net with noise maintained high IoU even after 25% noise.

**FIGURE S58** (A) Gini importance of RF model for the probability of 1p19q codeletion using ATRX and age. (B) Gini importance of final RF model used for the 1p19q codeletion in validation data set. (C) Simulation of the model with different p53 percentages showing 1p19q codeletion probability in different scenarios.

**Table S1** Impact of scrambling features on accuracy of random forest model

**Supplementary Material**

**Table S2** Validation set data

**Supplementary Material**

---

**How to cite this article:** Cevik L, Landrove MV, Aslan MT, Khammad V, Garagorry Guerra FJ, Cabello-Izquierdo Y, et al. Information theory approaches to improve glioma diagnostic workflows in surgical neuropathology. Brain Pathol. 2022;32:e13050. doi:10.1111/bpa.13050