To achieve minimum DNA input requirements for next-generation sequencing (NGS), pathologists visually estimate macrodissection and slide count decisions. Unfortunately, misestimation may cause tissue waste and increased laboratory costs. We developed an artificial intelligence (AI)-augmented smart pathology review system (SmartPath) to empower pathologists with quantitative metrics for accurately determining tissue extraction parameters. SmartPath uses two deep learning architectures, a U-Net based network for cell segmentation and a multi-field-of-view convolutional network for tumor area segmentation, to extract features from digitized H&E-stained formalin-fixed paraffin-embedded slides. From the segmented tumor area, SmartPath suggests a macrodissection area. To predict DNA yield per slide, the extracted features from within the macrodissection area are correlated with known DNA yields to fit a regularized linear model ($R = 0.85$). Then, a pathologist-defined target yield divided by the predicted DNA yield per slide gives the number of slides to scrape. Following model development, an internal validation trial was conducted within the Tempus Labs molecular sequencing laboratory. We evaluated our system on 501 clinical colorectal cancer slides, where half received SmartPath-augmented review and half traditional pathologist review. The SmartPath cohort had 25% more DNA yields within a desired target range of 100–2000 ng. The number of extraction attempts was statistically unchanged between cohorts. The SmartPath system recommended fewer slides to scrape for large tissue sections, saving tissue in these cases. Conversely, SmartPath recommended more slides to scrape for samples with scant tissue sections, especially those with degraded DNA, helping prevent costly re-extraction due to insufficient extraction yield. A statistical analysis was performed to measure the impact of covariates on the results, offering insights on how to improve future applications of SmartPath. With these improvements, AI-augmented histopathologic review has the potential to decrease tissue waste, sequencing time, and laboratory costs by optimizing DNA yields, especially for samples with scant tissue and/or degraded DNA.

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INTRODUCTION
Next-generation sequencing (NGS) has become an integral technique in the molecular diagnosis, prognosis, and treatment of cancer. To properly assess tumor tissue with NGS, solid samples must be dissected to meet minimum DNA input and tumor purity requirements. In standard practice, pathologists visually inspect hematoxylin and eosin (H&E)-stained, formalin-fixed, paraffin-embedded (FFPE) slides to determine how much tissue should be dissected and whether macrodissection is necessary to enrich for tumor cells. Besides meeting minimum input requirements, pathologists must also avoid recommending excessive dissection as tumor tissue is valuable and may be needed for further molecular tests. Tissue stewardship guidelines can help pathologists achieve this balance between sufficient and excessive dissection. However, following these suggestions using manual dissection techniques is difficult, and thus, there is an increasing need to optimize tissue extraction procedures as NGS becomes more relevant in clinical practice.

NGS pipelines have undergone tremendous advancements in the past decade, including the development of automated dissection systems for tissue extraction. Laser-capture microdissection was introduced about two decades ago, but has not been widely adopted in clinical laboratories because precise dissection of single tumor cells from FFPE slides is rarely necessary for clinical testing. Lower resolution mechanical macrodissection systems have also been developed as more clinically pragmatic alternatives. These systems can be combined with digital slide marking (digitally guided macrodissection), enabling integration with computer vision models for tumor enrichment. Several computer vision systems have been recently developed with the goal of estimating tumor-rich dissection areas from histopathology slides to meet tumor purity input requirements for molecular testing. However, no recommendation systems exist for estimating tissue quantity for minimum DNA input requirements, and thus even automated dissection systems rely on a pathologist to determine how many slides should be scraped. Unfortunately,
consequences of visual misestimation of tissue quantity include
sequencing failure, tissue waste, and increased laboratory costs
and turnaround times.

Here, we developed SmartPath: a computer vision-based
method to empower pathologists with quantitative metrics,
allowing them to accurately determine tissue input parameters
for desired DNA yields. Echoing design principles of artificial
intelligence (AI)-augmented pathology outlined by others25–29,
SmartPath functions as a pathologist-in-the-loop system rather
than a standalone predictor. Predictions are displayed in a
browser-based user interface (UI) viewed during pathology review.
The pathologist is free to recommend the predicted tissue input
parameters as presented, or to modify them based on their
expertise. We tested SmartPath in an internal trial to assess the
impact of AI-augmented pathology review in a real-world clinical
setting. We quantified immediate impacts of the AI-assistance on
tissue usage and the extracted DNA content, as well as on two
NGS workflow costs: the total number of extractions attempted
and the DNA extraction-to-sequencing time (T-seq). A thorough
statistical investigation of the impact of clinical covariates on
these metrics was conducted, revealing factors that influence NGS
success beyond tissue input parameters alone. Finally, recom-
mandations are made how to improve the system for future
applications.

MATERIALS AND METHODS
Model development
Before evaluating the SmartPath AI-augmented pathology review system
in an internal trial (see Methods: Internal Model Evaluation Trial), we first
developed the model through extensive validation experiments. The
underlying models can be grouped into two categories: feature generation
and DNA yield estimation. Feature generation aims to extract features from
a single H&E-stained histopathology whole-slide image (WSI). These
features are used to fit a DNA yield estimation model. At inference time,
the full pipeline from feature generation to DNA yield estimation is run to
produce predictions that augment pathologist tissue quantity selections to
achieve a total extracted DNA mass within a user-defined target range. This
modeling pipeline is summarized in Fig. 1A.

Feature generation pipeline. Feature extraction relies on pretrained tissue
and cell segmentation models. The tissue segmentation, based on a multi-
field-of-view network with a fully convolutional ResNet-18 backbone30,
produces segmentation maps of tumor-rich areas (AUC 0.947 for tumor
classification, Section S1.1). The cell segmentation model, based on the
U-Net architecture31, produces segmentations of cell nuclei throughout
the whole image (Section S1.2). These models are combined to assign
identities to tumor cells and lymphocytes (Section S1). Features are then
generated from these model outputs, consisting of four feature groups:
tumor shape, cell counts, cell nucleus shape, and cell nucleus texture,
totaling 3,461 features from each slide (Section S2).

Tissue and cell segmentation model comparison to pathologist
ground truth: Besides evaluating the classification accuracy of
the tissue segmentation model on pathologist annotations (see Section S1.1),
we also selected 334 slides (colorectal tumors metastasized to various
tissues) which were previously reviewed by pathologists and assigned
a visually estimated tumor percentage as part of our clinical NGS workflow.
We computed the correlation coefficient between model predicted tumor
percent and the visual estimate (Fig. 2A, R = 0.731). This modest
correlation is close to the inter-pathologist correlation for such visual
estimates of tumor percentage58. To evaluate the cell segmentation model,
we asked three pathologists to annotate all cells as tumor cell, lymphocyte,
or other within 300 fields of view sampled from 100 colorectal WSI5s (see
Section S1.2 for details). Correlation coefficients between predicted cell
counts and the consensus of the annotated cell counts were R = 0.834 for
all cells (Fig. 2B), R = 0.829 for tumor cells, and R = 0.728 for lymphocytes.

Macrodissection area masking during training and inference: In
our normal clinical workflow, approximately 30% of CRC samples are
macrodissected to ensure the extraction contains at least 20% tumor cells.
For this subset of slides, feature extraction was restricted to tissue within
the macrodissection area. Two strategies had to be developed for handling
these samples because the retrospective slides used for training already
had microdissection areas hand-drawn onto the slides by pathologists,
while the prospective slides processed during inference had not yet been
reviewed by a pathologist. To ensure that features from the training slides
were only from the hand-drawn region, we developed an ink detection
model which was post-processed to produce a macrodissection area mask
(Section S3).

During inference the macrodissection area was estimated from tissue
segmentation model predictions. The predicted tumor area was converted
to a binary mask and post-processed to produce a contour mimicking
hand-drawn macrodissection areas. Details on the implementation and
validation of this method are in Section S4.

DNA Yield Prediction
Training and Validation Sets for DNA Yield Prediction: The core
model underlying SmartPath is the prediction of DNA yield per slide using
linear regression on extracted imaging features. To acquire a training set, the
Tempus database was searched for slides scanned between January
2018 and January 2020 containing lung, breast, or colorectal cancer (CRC)
primary tumor tissue. Three different cancer types were used for training
because the average nuclear content per cell should be similar across
cancers, and therefore extracted cell-count features should follow the
same linear correlation with DNA yield regardless of cancer type. We
confirmed this assumption by estimating the yield per cell for each cancer
type (measured DNA yield per slide / predicted number of cells per slide)
and finding no statistical difference between the means. Using the three
cancer types also increased the training set size, which might help the
model better generalize to unseen data in the future.

Aspirates and cytology specimens were excluded from the training set,
as were slides with no recorded DNA mass or scraping, leaving a final
training set of 1605 slides. Approximately 28% were previously macro-
dissected, reflecting the rate at which samples are macrodissected in our
normal clinical workflow. Characteristics of the DNA yield prediction
training set are shown in Table S1. We also acquired a separate validation set of 332 retrospective samples
from the same database, restricted to only CRC tissues, which was used for
selecting a model with the best performing parameter combination (Table S2).
The validation set was enriched for macrodissected cases (57% were macrodissected) to ensure thorough evaluation of our macrodissec-
tion estimation algorithm. Characteristics of the validation set are shown in
Table S1.

Ground truth definition for DNA yield prediction model: The
ground truth for training the DNA yield prediction model was taken as the
extracted DNA yield from FFPE slides (see Section S5 for details on DNA
extraction procedure). Each slide in the training and validation sets was an
archival H&E slide representative of the unstained slides already extracted
and sequenced by our NGS laboratory. Although most underwent only one
DNA extraction attempt, some had multiple, in which case the imaged
slide may have been closer to the tissues used for the 2nd extraction
attempt. Therefore, the ground truth was defined as:

\[
\text{DNA yield} = \frac{\text{DNA1 yield}}{\text{Nslides, if only 1 extraction attempted}}
\]

\[
\text{DNA yield} = \frac{\text{DNA1 yield} + \text{DNA2 yield}}{\text{Nslides, for DNA1} + \text{Nslides, for DNA2}}, \text{if > 1 extractions attempted}
\]

Section S6 contains more details on the ground truth definition.

Parameter and feature exploration for final model selection: For
parameter exploration we used the full feature set (3641 features). Because
the number of features was larger than the number of samples in the training
set (1605), the linear model severely overfit and failed to
generalize without regularization. Optimal regularization parameters
were determined by parameter sweeping across L1 and L2 regularization
strengths. Each regularization was tested with natural log and Box-Cox
power transformations on features and ground truth. The parameter
combination with the best validation set performance (R = 0.818) was a log
transform and an L1 regularization with strength = 0.01 (see Table S2 for full
parameter exploration). Predictions of this optimal model are plotted against
our best training and validation ground truths in Fig. 2C, D.

To confirm that including all 3641 features was advantageous, we
performed a 200-fold cross validation using an 80/20 train/val split of
the training set using the optimal parameter combination found from
the exploration. We measured the mean coefficient magnitude across
folds for each parameter. The top 10 features accounted for 93.4% of the model coefficient magnitude and were from a combination of feature groups (cell counts, tumor shape, cell shape, texture), with the total cell count having the highest importance (Fig. S2). Further confirmation of the usefulness of keeping all features was done in a 200-fold cross validation experiment where cumulatively more features were included. Starting with just 1 feature, then progressively adding features by group, performance on the cross-validated training set and the withheld validation set increased as more features were included (Fig. S3). This confirmed that the inclusion of all features gave the best performing model.

We also explored the inclusion of categorical features, such as procedure type, tissue site, and institution. However, the inclusion of categorical features did not offer any significant boost in the performance of cross-validated models, and their model coefficients were consistently pushed towards 0 by regularization. Because the final model was to be run in a real-world scenario, where image artifacts may cause some features to have infinite or non-numeric values, additional steps were also taken to...
ensure that such situations were handled smoothly in the inference pipeline (Section S7).

**Target yields for number of slides prediction.** The goal of SmartPath is to recommend the number of scraped slides needed to achieve a DNA yield between 100–2000 ng. To convert the predicted DNA yield per slide into a recommendation of how many slides to scrape, we divided a target yield by the predicted yield per slide and rounded down to the nearest integer. This target yield is a tunable operating point of the algorithm. During the trial, the SmartPath system presented the number of slides needed to achieve a target yield of at least 100, 400, or 1000 ng. For details on how these target yields were selected, see Section S8.

We chose three target yields instead of one to give pathologists more flexibility. Because the relationship between number of slides scraped and DNA yield is linear, pathologists can also use the target yields to interpolate the recommended number of slides if they choose. This design choice emphasizes the principle of AI-augmented decision making, rather than AI automation.

**Internal model evaluation trial**

**Trial design.** To test the viability of our system in practice, we undertook an internal trial using clinical CRC samples to evaluate SmartPath compared to traditional (Trad) pathologist review. Trad and SmartPath workflows are summarized in Fig. 1B. Sample sizes of 250 SmartPath and 250 Trad samples were determined by power analysis at significance Level = 0.01 and power = 0.8 (Section S9). The internal trial was designed to be run in tandem with standard clinical workflow, mirroring every step until pathologist review. Before pathologist review, each FFPE block was cut into 20 sections and affixed to glass slides. One slide midway through the levels was stained for H&E and designated for pathologist review, while the others were designated for scraping. If the tumor was CRC primary and...
Internal trial performance evaluation metrics 

Table S4, the slides were flagged for trial enrollment and aggregated separately to avoid mixing with the rest of the clinical workflow. Enrolled samples were assigned in alternating order to Trad or SmartPath cohorts and assignments were recorded in a log, aiming to collect roughly equal numbers per day.

The Trad cohort H&E slides were a control group of samples which passed through our established pre-extraction workflow whereby a pathologist estimates tumor percentage by eye, marks a dissection area on the glass slide if needed, and recommends the number of slides to be scraped by an extraction technologist. All samples were still reviewed by a pathologist in a timely manner to not disturb the existing clinical sequencing workflow. Samples assigned to the Trad cohort were re-entered into the clinical workflow and path reviewed that same day.

The SmartPath cohort H&E slides were scanned on the Philips Ultra Fast Scanner (Philips, Eindhoven, The Netherlands) to produce a digitized WSI at \times 40 base magnification level (0.25 \mu m/\text{pixel}). Slide scanning automatically triggered feature generation and DNA yield prediction on the WSI (see S10). To minimize interference with existing clinical workflow, SmartPath-assisted review was conducted the morning after scanning, although in principle same day review is quite feasible because scanning and model deployment take only minutes to complete. After pathology review, the recommended number of slides were scraped, DNA was extracted (Section S5), and NGS was conducted using the Tempus XT platform\textsuperscript{8}. This process was repeated daily over the span of several months until 249 SmartPath and 252 Trad cohort samples were accumulated.

During SmartPath-assisted review, the pathologist viewed a custom-built UI (Fig. S5) displaying recommendations for macrodissection area and number of slides needed to achieve at least three possible target DNA yields of 100, 400, or 1000 ng. The pathologist had the option to accept or reject model recommendations. If they chose to accept, the desired target yield was selected in the UI. If they disagreed, they recorded in the trial log if the rejection was driven by clinical reasoning or model performance. In cases where the pathologist disagreed with the predicted macrodissection area, the pathologist drew their own microdissection area. A summary of the pathologist decisions made during the trial is shown in Table S4.

Internal trial data quality control. Of the 501 samples enrolled into the trial, 18 were rejected at pathology review due to insufficient tissue, 4 were erroneously enrolled either with incorrect cancer type or procedure type, 1 had an incorrect indication of number of slides scraped, and 2 were removed because their sequencing was delayed due to human error (Table S5). This left 476 samples for the overall analysis (233 Trad, 243 SmartPath). For analysis of T-seq, an additional 18 samples were dropped (7 Trad, 3.00% of population; 11 SmartPath, 4.53% of population) because they did not reach RNA sequencing due to failure downstream of extraction (either at library prep or hybridization steps, see Fig. 1B), and therefore did not have a defined sequencing time interval, leaving 458 samples (226 Trad, 232 SmartPath).

Statistical analysis of covariates

The FDA guidance for adjustment for clinical covariates in clinical trials (Docket number FDA-2019-D-0934) advises experimenters to identify the covariates expected to have an important influence on the primary outcome. The primary outcomes for the present work are the DNA yield and workflow costs. DNA yield may depend on sample age, as older samples may suffer from nucleotide degradation\textsuperscript{22}. It may also depend on the individuals involved in the extraction (i.e., the pathologist and technicians). The day on which the sample is extracted could have an impact on workflow turnaround time due to weekly lab scheduling cycles. We included these sample-level measures as covariates (Table 1) and also recorded several patient-level characteristics commonly reported in cancer studies (Table S5). Covariate imbalance was measured by the chi-squared test computed from contingency tables. Contingency tables were computed by cross-tabulating counts for each characteristic and chi-squared tests were performed in Python 3.7 using scipy.stats.contingency\textsuperscript{74}.

Analysis of covariance using generalized linear models. An analysis of covariance (ANCOVA) allows researchers to dissociate contributions of additional covariates from the treatment to the total variance. For the present application the treatment variable is the trial cohort (Trad or SmartPath) and the dependent variables are the following: DNA mass undershoot boolean (1 if \textless 100 ng, 0 otherwise), number of slides scraped (N slides), extraction count, and T-seq. Traditional ANCOVA is designed to run on normally distributed samples assuming linearity and homoscedasticity (constant variance across residuals). However, most of these metrics are not normally distributed, and thus appropriate distributions were chosen to model these dependent variables with generalized linear models (GLMs)\textsuperscript{35}. All ANOVA analyses were performed in R version 0.4.4.16. For details on GLM selection see Section S11.
Impact on extraction metrics

AI-assistance improved DNA yield within a target range of 100–2000 ng. The fraction of samples within the target range was significantly improved for the SmartPath cohort (Trad = 0.56+/− 0.064 vs SmartPath = 0.70+/− 0.058, P = 0.005, a 25% increase, Fig. 3A). This was primarily due to limiting over-extraction, as the fraction of samples with mass that overshoot the desired range was also significantly improved (Trad = 0.32+/− 0.06 vs SmartPath = 0.18+/− 0.049, P = 0.001, a 14% decrease, Fig. 3A). The fraction of samples that undershot the desired range was not improved overall.

Tissue characteristics, such as tissue area and DNA fragment quality, are known to impact tissue extraction14,37. We confirm that these effects exist in our data as well. In Fig. 3B, we subset the data into large and small tissue area groups (defined in Methods), revealing that reduction in overshoot was restricted to large tissues. When further subset by extraction quality (defined in Methods), the reduced overshoot effect was primarily seen in large tissues with high extraction quality (Fig. 3C, top). Therefore, AI-assistance helped pathologists preserve tissue use for samples that were already likely to succeed NGS. Subsetting also revealed a trend in reduction of the undershoot fraction for small tissues with low extraction quality (Fig. 3C, bottom). Although the difference was not significant (chi-squared P = 0.088), there were only 50 samples in this subset and the sample size may be underpowered to measure the effect. As discussed in subsequent sections, however, subsetting by small tissue area and low extraction quality showed significant improvements in other metrics.

AI-assistance fosters more efficient use of tissue slides. Across all CRC samples from the trial (n = 476), DNA yields <100 ng almost always resulted in multiple extraction attempts (Fig. S10). These results demonstrate the importance of better metrics for scraping parameters, as more slides should be scraped initially when lower DNA yields are expected in order to avoid repeating extraction. While NGS laboratories typically scrape 5–10 FFPE slides per extraction38, our AI model recommended a broader distribution of slides for scraping compared with the Trad cohort (Fig. 4A). Large tissues in the SmartPath cohort usually had only one or two slides scraped, thus conserving tissue in this subset. On the other hand, small tissues in the SmartPath cohort usually had >10 slides scraped (Fig. 4A, B). Therefore, while the mean number of slides was not significantly different between SmartPath and Trad cohorts across all tissue sizes (Fig. 4B), slides in the SmartPath cohort were used more efficiently. However, because the distribution of N slides was not normal, the median should also be considered. While the overall median number of slides scraped per sample in the SmartPath cohort was slightly higher than in the Trad cohort (7.7 ± 5.91 SmartPath vs 7.62 ± 3.0 Trad), the median in the SmartPath cohort was lower than in the Trad cohort (6 SmartPath vs 10 Trad).

Further subsetting the data by extraction quality also shows that SmartPath recommended fewer slides for large tissues regardless of extraction quality (Fig. 4C, low quality P = 0.07, intermediate quality P<0.01, and high quality P<0.01). An opposite trend was observed for small tissues with low and intermediate extraction quality, where more slides were recommended in the SmartPath cohort, although there was no significant difference (Fig. 4C bottom). For the subset of small samples with high extraction quality, the SmartPath and Trad cohort means were very similar. This could be desirable as high-quality samples are already likely to succeed NGS.

Impact of AI-assistance on NGS workflow costs

Number of extraction attempts is similar between cohorts. The number of extraction attempts is an important metric for workflow

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Table 1. Sample-level characteristics of evaluation trial dataset.

|                | Trad (N = 233) | SmartPath (N = 243) | Chi-sq. or t-test P-value |
|----------------|---------------|---------------------|-------------------------|
| **Sample age at extraction (days)** | 0.06 |
| Median          | 37.57         | 47.58               |
| Range           | 5.05–2443.01  | 5.58–3205.58        |
| **Procedure Type** | 0.19 |
| Biopsy (unspecified) | 68            | 93                  |
| Needle Biopsy   | 53            | 42                  |
| Resection       | 112           | 108                 |
| **Dissection**  | 0.73          |
| Macrodissected  | 119           | 130                 |
| Whole slide     | 114           | 115                 |
| **Pathologist** | < 1e-60       |
| A               | 100           | 1                   |
| B               | 47            | 205                 |
| C               | 36            | 0                   |
| D               | 24            | 4                   |
| E               | 23            | 1                   |
| F               | 3             | 34                  |
| **Extraction day of week** | 0.0008 |
| Monday          | 23            | 26                  |
| Tuesday         | 42            | 17                  |
| Wednesday       | 50            | 60                  |
| Thursday        | 36            | 37                  |
| Friday          | 30            | 58                  |
| Saturday        | 45            | 45                  |
| Sunday          | 7             | 2                   |

Sample characteristics with high cardinality (only showing N unique values)

|                | \(N\) (unique) |
|----------------|---------------|
| **Tissue site** | 38            |
| **Extraction tech** | 18            |

*Sample age is defined as the delta between time of first extraction attempt and time of sample collection. Because sample age is a continuous variable, a chi-squared test could not be performed. Instead, a t-test was performed on the log-transformed data.

Counts per category are shown for each characteristic grouped by Trad and Smart cohorts, except for characteristics with high cardinality which only show the number of unique categories. These counts define a contingency table for each covariate. A chi-squared test was run on each contingency table to obtain \(p\)-values assessing a significant difference between Trad and Smart cohorts. Sample-level data had no data missingness, and in some cases showed significant imbalance, as evidenced by the small chi-squared test \(p\)-values.

Univariate and multivariate GLMs: GLMs were fit only for the subset of the samples most in need of AI-assistance, namely small tissues with low extraction quality. Univariate GLMs were initially fit using the sample-level (Table 1) and patient-level characteristics (Table S5) as independent variables, but significant effects were not found for any of the patient-level characteristics. Multivariate models for each metric are built using only those variables with significant association in univariate tables (Tables S6–S9). For details on construction of univariate and multivariate models, see Section S13.
improvement because re-extractions are financially and temporally costly. The distributions of N extractions for SmartPath and Trad cohorts are shown in Fig. 5A. As these distributions were not normal, they are represented with Poisson distributions for calculation of statistics (see Methods). The distributions are dominated by cases with only one extraction, which is already the optimum. Overall, no significant difference in mean number of extractions per sample was observed between the SmartPath and Trad cohorts Fig. 5B. Grouping by extraction quality reveals that all high-quality samples were already performing at optimum for this metric, with only one extraction per sample, and intermediate-quality samples were performing near optimum. On the other hand, the SmartPath cohort had a decreased mean extraction count for low-quality samples, albeit not significant (Poisson rate ratio $P = 0.212$).

Further subsetting by tissue area reveals that the decrease in mean extraction count per sample approaches significance (Poisson rate ratio $P = 0.052$) for low-quality samples with small tissue area (Fig. 5C, bottom). This result suggests that AI-assistance may be useful in preventing re-extractions for low-quality samples with small tissue area, which are the samples most in need of improvement. However, there was a significant increase in mean extraction count for large intermediate-quality samples in the SmartPath cohort (Poisson rate ratio $P = 0.017$), caused by four samples which had >1 extraction count. The higher extraction count may be partly due to the age of these samples. SmartPath cohort samples were on average older than the Trad cohort (Fig. S12A), and for large samples in the SmartPath cohort, those with intermediate quality were also the oldest (Fig. S12B). Older samples correlate with higher extraction count (Fig. S12D), likely because they tend to be more degraded.

AI-assistance reduced DNA sequencing time for low quality samples with small tissue areas. Figure 6A shows the distribution of T-seq in the SmartPath and Trad cohorts. Similar to extraction counts, there was no significant difference in the mean T-seq between the two cohorts (Fig. 6B; Trad $3.74 \pm 1.67$ days, SmartPath $3.89 \pm 1.67$ days). We expect T-seq to follow a similar trend as extraction count because they are strongly correlated (Fig. S11). High-quality samples showed almost no difference in T-seq between cohorts, a reflection of the fact that extraction count is already optimal for high-quality samples. Intermediate quality samples showed a significant increase ($P = 0.018$) for the SmartPath cohort, likely due to the same samples that drove up extraction count for this group. However, when subset by tissue area the T-seq for small low-quality samples was almost 2 days shorter in the SmartPath cohort compared with Trad (Fig. 6C, bottom; Trad $6.90 \pm 2.77$ days, SmartPath $4.97 \pm 2.06$ days, $P = 0.025$).
Univariate analysis of covariates on full trial dataset

To determine if the effects observed were due entirely to the experimental condition (SmartPath vs. Trad) alone, we considered covariates of the study as detailed in Table 1. Despite an attempt to randomize samples by alternating assignment to SmartPath and Trad cohorts each day (see Methods), imbalances were detected. Moderate imbalance was detected for sample age (chi-squared $P = 0.06$), while strong imbalance was detected for pathologist (chi-squared $P < 1e-60$), extraction day-of-week (chi-squared $P = 0.0008$), and extraction tech (chi-squared $P = 0.0001$).

While dataset imbalance in covariates suggests other sources of variability besides the experimental condition, it alone does not prove that they impacted the dependent metrics. With univariate GLMs (see Methods), we quantified the correlation each covariate had with the following four trial metrics: DNA Mass undershoot boolean (True if <100 ng), number of slides scraped, extraction counts, and T-seq. Summary statistics of univariate GLMs for each covariate are presented in Tables S6–S9.

For three of the trial metrics, undershoot Boolean, number of slides scraped, and extraction count (Tables S6–S8), the extraction quality ($P < 2e-16$, $P = 5.96e-05$, and $P < 2e-16$, respectively) and tissue area ($P = 4.01e-06$, $P < 2e-16$, and $P = 2.78e-06$, respectively) were more predictive than any of the covariates. These same three metrics also were significantly correlated with procedure type. This is expected, as procedure type is strongly correlated with tissue area, where needle biopsies tend to have much smaller area than resections.

However, for T-seq (Table S9) the extraction day-of-week was the most predictive variable ($P < 2e-16$). This is a known effect, where sequencing times for samples extracted later in the week tend to be longer than samples extracted earlier in the week (Fig. S13B, see Discussion). Despite the randomized trial design,

Fig. 4  AI assistance offers more nuanced suggestions of number of slides to scrape for extraction. A Distribution of N slides scraped plotted for all samples (left), only large tissues (middle), and only small tissues (right). Without AI-assistance, pathologists tended to recommend either 5 or 10 slides for scraping (Trad, orange), but with AI-assistance the distribution was much broader (SmartPath, blue). SmartPath distribution is shifted towards fewer slides for large tissues and more slides for small tissues. B Box plots comparing numbers of slides scraped for SmartPath and Trad cohorts grouped by large and small tissues. P-values were computed from t-tests on log-transformed data assuming unequal variance. White dots = mean. Horizontal black line = median C Truncated violin plots comparing number of sides scraped between SmartPath and Trad cohorts, grouped by large (top) and small (bottom) tissues and by extraction quality. Back boxes = 25% and 75% percentiles. White dots – medians. Samples that did not have recorded numbers of slides scraped were dropped. P-values for Welch’s two-sided t-test assuming unequal variance are displayed above each group. Bimodal distributions for the Trad cohort (orange) correspond to 5 and 10 slides.
samples in the SmartPath cohort tended to be extracted later in the week than samples in the Trad cohort (Fig. S13A).

For all metrics except for N slides, the extraction tech group univariate GLMs showed significance ($P < 0.05$). Although this variable has high cardinality (25 categories), the Akaike Information Criterion (AIC) in these cases was lower than some of the other variables (Tables S6–S8), suggesting that these were not random correlations. Imbalance in the extraction tech group was also meant to be eliminated by trial design but persisted despite our efforts.

Only one of the metrics, N slides, was significantly predicted by sample age. None of the metrics had strong correlation with pathologist.

**Multivariate analysis of covariates for subset of samples with small tissue area and low extraction quality**

Here we investigate the impact these covariates had on the main effect of the experimental variable, the trial cohort. In Figs. 3–6 we identified that AI-assistance was most effective for samples with small tissue areas and low extraction quality. As this is the most interesting subset of samples, we restricted the following multivariate analysis to this subset ($N = 50$). For modeling each outcome metric, we chose only those covariates significantly associated ($P < 0.05$) with the outcome metrics in the univariate analysis (Tables S6–S9). We excluded procedure type, as it is already strongly correlated with tissue area and no surgical resections are present in this subset. We also excluded the extraction tech group as this variable has very high cardinality (25 categories) relative to the number of samples in this subset.

To measure the influence of these covariates on the main effect of AI-assistance, we compared a univariate GLM using trial cohort as the independent variable, to the multivariate GLMs for each outcome metric (see Methods). In the interest of brevity and focus, we limited the multivariate analysis to only 4 outcome metrics, dropping the target and overshoot fractions from the analysis. Summary statistics of these GLMs are shown in Table 2.

For the small tissue area and low extraction quality data subset, the trial cohort alone was significantly predictive for both extraction count and T-seq ($P = 0.049$ and 0.026, respectively). The other two metrics, undershoot and N slides, had univariate associations with trial cohort approaching significance ($P = 0.055$ and 0.075, respectively). In all outcome metrics, the inclusion of covariates in multivariate GLMs raised the trial cohort $p$-values, suggesting that the main effect can be partially explained by the covariates. However, the increase in trial cohort $p$-value was moderate.

Inclusion of covariates increased the AIC for multivariate GLMs of undershoot from 54.57 to 55.04 and N slides from 65.52 to 74.11, indicating that covariates carry little additional information about these metrics, adding complexity without proportionally improving the fit. For both NGS workflow metrics, though, the AIC was reduced (extraction count: from 120.21 to 118.91, T-seq: from 51.90 to 45.73), indicating that the covariates improved the model without adding unnecessary complexity.

The only significantly predictive covariate in the multivariate GLMs was extraction day-of-week, which strongly associated with T-seq ($P = 0.006$). The correlation of extraction day-of-week with T-seq is a known effect in our NGS laboratory (Fig. S13). Although the cohort imbalance in day-of-week was meant to be eliminated by alternating assignment of samples to SmartPath and Trad cohorts each day (see Methods), unfortunately the imbalance persisted (Table 1). Overall, the multivariate analysis shows that the covariates considered here have a measurable impact on the main effect of trial cohort. The adjusted effect of trial cohort is weaker upon inclusion of covariates; however it is difficult to ascertain if this fully explains the main effect as the main effect is itself underpowered ($N = 50$). Future trials of this tool should

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**Fig. 5 Number of extractions needed to reach DNA sequencing.**

A The numbers of extraction attempts needed to reach DNA sequencing are counted for SmartPath and Trad cohorts. Only 4 samples in the total dataset had extraction count=4, and only one had extraction count=5. B Mean number of extractions grouped by extraction quality. Error bars are 95% confidence intervals produced with 1000 bootstraps. High-quality samples have no error bars because they are already at optimum, with only one extraction per sample, regardless of cohort. To compute $p$-values, Poisson distributions were fit to thezeroed distributions (see Section S11) of SmartPath and Trad cohorts and a test was performed to assess if the ratio of the two Poisson rates is statistically different from 1. C Same as B but grouped by large and small tissues.
ensure that sufficiently high numbers of samples are available in the small tissue and low extraction quality regimes and should more rigorously control for covariate imbalances.

**DISCUSSION**

Here, we developed SmartPath, a computer vision tool to assist pathologists in determining NGS tissue input parameters, and tested it in a real-world clinical setting. Compared to the group that received traditional pathology review, AI-assistance produced significantly more DNA yields falling within a target range of 100–2000 ng. The AI-assisted model also improved tissue stewardship by recommending scraping of more slides for samples with small tissue areas, likely preventing some re-extractions, but fewer slides for samples with large tissue area.

While scraping more slides for small tissues may seem counter to tissue stewardship, underpredicting could result in an insufficient yield, leading to re-extraction which is a further waste of tissue. There is a trade-off between initial tissue use and prevention of re-extraction. Our model design intention was to bias towards preventing re-extraction, but evidently it was not biased enough as overall undershoot fraction was not reduced in the SmartPath cohort. In retrospect, target yields of the model could have been tuned even higher to improve the undershoot fraction at the expense of the overshoot fraction. This was explored in a simulation (Fig. S9) which suggests that, had we scraped 1.4x more slides (scaling the target yields from 100, 400, 1000 to 140ng, 560, 1400 ng) the undershoot fraction for the SmartPath cohort could have dropped below that of the Trad cohort, while still maintaining a reduction in overshoot fraction. During the trial, the reviewing pathologists also noted that higher target yields would have been advantageous, and 82% of their selected target yields were for the highest available yield, 1000 ng (Table S4).

For the two NGS workflow costs, the extraction count and T-seq, no significant difference was found between the full populations of SmartPath and Trad cohorts, but improvements were seen in the subset of small samples with degraded DNA quality. It is known that NGS fails more often for smaller, poor-fragmentation-quality samples, and pathologists will attempt to reject such samples for insufficient tissue. Notably, the SmartPath cohort had only 2 samples rejected for insufficient tissue, while the Trad cohort had 16, because pathologists were willing to observe SmartPath predictions before rejecting a sample (Table S4). Despite including samples in SmartPath review that likely would have been rejected in Trad review, the mean extraction count for small, low-quality samples was in fact reduced (albeit not statistically significant, Fig. 5C). Additionally, we observed a significant reduction in T-seq for small, low-quality samples (Fig. 6C). Furthermore, the subset of high-quality samples had only one extraction attempt regardless of SmartPath or Trad treatment (Fig. 5B), which is already optimal, so the extraction count could not be further improved for these samples. This subset also represented most of the samples in the entire trial (only ~13% were low quality). Therefore, lack of reduced NGS workflow costs in the full population is largely due to over-representation of high fragmentation quality samples in our cohort.

Colorectal cancer was chosen for the trial for internal workflow considerations, but we expect that the present algorithm can be generalized to other cancer types by replacing the underlying tissue and cell segmentation models with tissue-specific models already in existence. Other cancer types, such as non-small cell lung cancer and especially pancreatic cancer, have higher rates of
were split between normal lab operation and conducting this trial.

A pathologist who was trained to use the UI at the start of the trial.

introduced for the SmartPath cohort.

and therefore a bias towards sequencing failure could have been

samples for sequencing due to old age. By chance, the SmartPath

unsuitable for NGS; however, much older samples have been

success. Evidence suggests that samples >7 years old are


tively impact patients
day-of-week as a predictor (Table 2).

the AI-assistance effect was reduced after inclusion of extraction

19 epidemic, and thus effects due to limited personnel were likely

place in Aug-Dec 2020 during the lockdown period of the COVID-

bene

low-fragmentation-quality samples (Fig. S15) and therefore may

benefit even more from the AI-augmented pathology review system, although additional research must be conducted to determine if this is the case.

The present work does not evaluate the impact of SmartPath on genetic variant calling because its scope is restricted to evaluation of a workflow improvement tool. However, future applications could conceivably improve accuracy of variant calling by recommending tissue samples with optimized DNA yield and tumor enrichment. To evaluate such a system, a study could be designed where consecutive sections from the same samples are evenly split between SmartPath and control groups, and genetic variants between groups are compared.

Proper randomization is necessary to eliminate all biases in trials, but for many real-world trials like ours, this is not possible due to external constraints. Future tissue recommendation models could improve upon the current work by taking these sources of bias into account as variables in the model itself and/or explicitly eliminating these effects through trial design. One bias identified was the strong influence of extraction day-of-week on T-seq, likely due to weekly batch effects and staffing cycles which cause sequencing times to be longer for samples extracted later in the week. This resembles a well-documented weekly phenomenon in healthcare, termed the “weekend effect.” The trial also took place in Aug-Dec 2020 during the lockdown period of the COVID-19 epidemic, and thus effects due to limited personnel were likely exaggerated. This bias was meant to be eliminated by the trial design by enrolling similar numbers of samples into SmartPath and Trad cohort per day, but multivariate analysis showed that the AI-assistance effect was reduced after inclusion of extraction day-of-week as a predictor (Table 2).

Sample age has a known effect on FFPE sample extraction efficiency. For example, a model combining imaging-based and biochemical predictors could be trained to provide a prior on sample quality. Training data for an imaging-based quality predictor could come from a combination of established DNA quality metrics, including fragment analyzer data, qPCR assays to measure the amount of amplifiable DNA in a sample, the DNA Integrity Number, and Genomic Quality Number.

Furthermore, the trial only incorporated AI into the initial screening of the sample but did not incorporate pathologist feedback for updating predictions, primarily to avoid disturbing the existing clinical workflow of our NGS laboratory. We envision a future pathologist-in-the-loop application, where pathologists may edit macrodissection areas and receive updated predictions in real time. Workflow improvements could also be made to maximize efficiency. For example, a model combining imaging-based measures with clinical data could flag samples up front that may need AI-assistance, while passing samples with high likelihood to succeed in NGS. Future models may also be trained to predict not only DNA yield, but RNA yield and library-prep success.

The SmartPath system was designed to support an existing NGS pipeline, which uses traditional scraping-based macrodissection to extract tissue within pathologist-defined tumor regions. A caveat of scraping-based macrodissection is that it may not always enrich the tumor percentage due to heterogeneity of cell types within

| Table 2. Statistics for covariates fit to trial metrics in samples with small tissue area and low extraction quality (N = 50). |
|---|
| **Outcome Metric** | **Univariate GLM (trial cohort only)** | **Multivariate GLM (trial cohort + covariates)** |
| **AIC** | **p-value** | **AIC** | **Trial Cohort p-value** | **Covariate** | **Covariate p-value** |
| Undershoot (True if <100 ng) | 54.57 | 0.055 | 55.04 | 0.072 | Day-of-week | 0.22 |
| N slides scraped | 65.52 | 0.075 | 74.11 | 0.14 | Sample age | 0.84 |
| Extraction count | 120.21 | 0.049* | 118.91 | 0.057 | Day-of-week | 0.12 |
| T-seq | 51.90 | 0.026* | 45.73 | 0.062 | Day-of-week | 0.006* |

*Indicates p-value significance below 0.05.

**AIC** - Akaike Information Criterion. The AIC is a fitness parameter that trades off the complexity of a model with how well the model fits the data. It can be interpreted as a measure of model parsimony, where lower values indicate a more parsimonious model. It is a relative measure, and thus can only be compared between models for a given metric.

**Covariates** are chosen based on significant association in the univariate analysis, hence not every outcome metric is modeled with the same covariates.

**Because Pathologist is a categorical variable and thus has p-values for each category, only the most significant p-value is shown.
tumor areas. Future applications of SmartPath could integrate with laser-capture microdissection systems to guide single-cell-based microdissection.

By addressing these limitations, future applications of SmartPath may provide a viable alternative to manual estimation by accurately predicting tissue quantities needed for adequate DNA yield. SmartPath could be useful in circumstances where access to pathologists is scarce, or for laboratories processing large volumes of tissue. Coupled with a digital slide viewer, such a system can support fully remote pathology review of digitized WSIs, allowing NGS laboratories to widen their access to reviewing pathologists. Integration of SmartPath with automated microdissection systems could allow for tissue extraction workflows which are almost entirely automated, with a pathologist needed only to approve or modify input parameters, and potentially be economically and clinically beneficial for NGS laboratories.

DATA AVAILABILITY
Most data generated or analyzed during this study are included in this article and its supplementary information files. Raw data are not made available due to their proprietary nature, but are available from the corresponding author on reasonable request.

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1802
B.L. Olsinski et al.
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AUTHOR CONTRIBUTIONS
BLO, RDJ, AW, and BMM performed the study concepts and design. BLO, RJ, ABT, RPJ and MCS performed the development of methodology and writing, review, and revision of the paper. BLO and IH wrote and validated the algorithm code. MC, CW, and LS built the browser-based UI and coordinated deployment to production. RDJ and AW supervised the validation trial. BLO and BMM reviewed samples in both SmartPath and Trad arms during the validation trial. BLO, ABT, and RPJ provided acquisition and analysis of the validation trial results. BLO, ABT, and RPJ provided interpretation of data and statistical analysis. All authors read and approved the final paper.

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