Spatially resolved characterization of the transcriptome and proteome promises to provide further clarity on cancer pathogenesis and etiology, which may inform future clinical practice through classifier development for clinical outcomes. However, batch effects may potentially obscure the ability of machine learning methods to derive complex associations within spatial omics data. Profiling thirty-five stage three colon cancer patients using the GeoMX Digital Spatial Profiler, we found that mixed-effects machine learning (MEML) methods may provide utility for overcoming significant batch effects to communicate key and complex disease associations from spatial information. These results point to further exploration and application of MEML methods within the algorithm development life cycle for clinical deployment.

Keywords: GeoMx Digital Spatial Profiler; Spatial Omics; Mixed Effects; Machine Learning; Boosting Methods; Tumor Metastasis; Colon Cancer

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1. Introduction

Practicing pathologists routinely order immunohistochemical stains to assess the spatial distribution of important biomarkers for disease prognostication. Recently, higher resolution spatially resolved technologies have emerged. Nature Methods declared “Spatially Resolved Transcriptomics” technologies as the method of the year in 2020. Although multiplexed immunohistochemistry and laser capture microdissection approaches once served as popular methods for assessing spatially resolved omics information, these approaches are largely intractable for high throughput analysis because quantitative analysis is hindered by significant tissue distortion from chemical restaining processes, insufficient multiplexing, and prohibitive costs. New spatial omics technologies can multiplex far more markers through targeted cleavage of oligonucleotide tags. For instance, the Nanostring GeoMX Digital Spatial Profiler (DSP) first utilizes multiple immunofluorescent (IF) antibody stains to highlight various cellular lineages, then uses UV-light to cleave attached oligos for quantification via the nCounter instrument or next-generation sequencing (NGS). A particular novelty of this technology is the ability to highlight regions of interest (ROI) through segmentation of the IF stains or selection of specific locations of interest. Informatics techniques developed for spatial data include estimation of: 1) which genes are spatially variable, 2) how spatial clustering patterns of expression may differ between cases and controls, 3) colocalization and coordination between cellular populations, accomplished using canonical protein markers or cellular deconvolution, and 4) integration with other imaging (e.g., morphology, architecture via hematoxylin and eosin stain, H&E) and single-cell multi-omics (e.g., RNASeq, ATACSeq; models of cell fate) which can further shed light on diagnosis and prognostication.

Prospective deployment of clinical decision aids can use low-cost tissue staining techniques (e.g., IHC) to profile protein markers deemed relevant through spatial omics analysis. However, existing approaches (e.g., differential expression and spatial autocorrelation) suffer from technical factors, including: reagent application, batch effects, and other forms of repeated measurements. These confounders have the potential to bias disease associations and preclude prospective classifier deployment. Compared to traditional penalized, generalized linear modeling approaches, algorithms that can use the complete multiplexed set of markers and their interactions (e.g., Random Forest models) may be best positioned to capture the disease pathogenesis yet are likely less resistant to batch effects. Thus, we need machine learning algorithms that can fully utilize spatial multiplexed information while being immune to batch effect corruption. In this study, we assess the potential for mixed-effects machine learning (MEML) methods to address this issue, capturing unintuitive spatial and multiplexed complexities, while respecting patient and batch level differences.

We assessed colorectal adenocarcinoma (CRC) cases, with or without nodal and/or distant metastasis using spatial omics technologies (Figure 1A-B). Prognosis / metastatic potential is well characterized based on presence of tumor invasion into distinct tissue layers (e.g., epithelium, lamina propria, submucosa, muscularis propria, pericolic fat, and serosa). Previous research has demonstrated the importance of specific somatic alterations (e.g., APC, MMR) and tumor-infiltrating lymphocytes (and their spatial coordination) as prognostic indicators. Here, we used the DSP to collect spatial expression of 39 immune markers across 840 ROIs and applied MEML to characterize factors relating to different macro-architectural contexts (e.g., markers related to tissue interface with tumor, inter; within tumor, intra; away from tumor, away) and metastasis. This study will serve as the backbone of an expanded investigation of spatial immune markers descriptive of nodal and distant metastasis for Colon cancer patients.
Figure 1: Overview of Experiment: Paired H&E and IF stains used to help select ROI (black squares) for: A) Non-metastatic patient, B) Metastatic patient; C) Tree boosting methods are combined with mixed effects modeling to adjust for D) patient/batch-level effects to yield E) significant disease associated interactions

2. Motivation for Comparison Study

First, we will provide a broad overview of common analytical techniques for spatial omics data, then discuss potential statistical oversights (e.g., modeling of statistical interactions and nonlinearities, batch effects/repeat measurements), to highlight the advantages of MEML methods.

2.1. Review of Prior Spatial Omics Analysis Methods

2.1.1. Assessment of spatially variable markers

Within a slide, there is spatial variation in marker abundance, such that regions of interest (ROI) that are closer together may exhibit similar gene/protein expression. Spatial autocorrelation is typically assessed using a Gaussian Process (GP), a mixed-effects model with a parameterized kernel. Included is an example of this model for multivariate normally distributed expression data:

\[
Y_i \sim MVNormal(\mu_i, \sigma^2) \\
\mu_i = \beta \vec{x}_i + b_{[i]} \\
\vec{b} \sim MVNormal(\vec{0}, \Sigma)
\]  

Where \( \vec{b} \) is a vector of random intercepts, here observed at locations \( \vec{s} \). The covariance matrix of the random effects, \( \Sigma \), is a kernel used to test spatial hypotheses; for instance, that the correlation in expression between two spatially dependent markers drops off with distance \( d \), \( \Sigma_{i,j} = a^2 \exp \left( \rho^2 d(\vec{s}_i, \vec{s}_j) \right) \). The target of inference would be \( \rho \), the degree to which the correlation in expression is retained as a function of the distance (\( d \); e.g. L2-norm) between the ROIs \( i \) and \( j \). Less parametric forms of this hypothesis can be tested through network autocorrelation and, more recently, through prediction models such as graph neural networks (GNN). GNNs can predict the expression patterns in a region and can thus be useful for increasing the resolution of the spatial expression patterns. As an example of the latter, the GNN would have the following generic form,
where the expression of neighboring (neighborhood \( N \)) cells/ROIs, \( j \), are used to predict the current ROI’s expression \(^{17,18} \): \( Y_j = \sigma(\sum_{j \in N} a_j Y_j) \).

2.1.2. Clustering of spatial patterns and differential expression

Other analyses expand the aforementioned GP models to estimate characteristic clustering patterns, enabling inter-slide comparison. Less parametric forms that take into account regional dependencies include graph neural networks, which can utilize aggregation/pooling mechanisms across nested observations to predict a single outcome while learning representative patterns.

Here, establishing differential expression for specific markers and applying penalized regression procedures across a set of markers are simpler analytic approaches that compare spatially variable markers across slides and differentiate disease status. The latter is formalized below, where \( Y_i \) represents whether the case underwent metastasis, predicted from markers \( \vec{x}_i \):

\[
Y_i \sim \text{Binomial}(1, p_{\text{METS}}) \\
\text{logit}(p_{\text{METS}}) = \beta \vec{x}_i
\]

(2)

In this simplified scenario, comparisons across spatial clustering patterns can be incorporated into this analysis by either stratifying the effects by assigned macro-architecture (e.g., intra, inter, away) or including an interaction term for the macro-architecture.

2.2. Motivation for Mixed Effects Machine Learning Approaches

The goal of this preliminary comparative study is to explore methods for classification of tissue macro-architecture/metastasis from spatially resolved expression patterns (Subsection 2.1.2) and to discuss opportunities for further classifier development. In this section, we discuss two deficiencies of existing methods.

Machine learning techniques search the ideal specification of interactions and nonlinear transformations that capture the relationship between the input and target of inference. For instance, classification and regression models (CART; e.g., Random Forest) may capture interactions between variables and nonlinear/discontinuous associations through conditional decision splits \(^{19} \). Interpretation methods (e.g. Shapley Additive Feature Explanations, SHAP) can derive etiological factors in the form of salient predictors and interactions \(^{20} \). As omics datasets are usually high-dimensional and multi-collinear, CART methods better spot relevant interactions in the data while managing the high-dimensionality.

However, these algorithms assume that observations are independent and identically distributed \((i.i.d.)\), which is invalid in the presence of repeated measurements, where a correlation or dependency structure exists between observations. In the case of spatial omics modalities, an ROI may exist within several levels of nested dependencies (e.g., multiple ROI per section, many sections per block and/or profiling batch). It is also possible that multiple blocks are extracted from different biopsies across space (sampling site) and time. These factors may potentially exacerbate the negative impact of technical effects on statistical inference. Even after documenting and controlling for batch effects, the degradation in performance resulting from the correlation between batch and outcome has not been thoroughly studied in this context \(^{21,22} \). Hierarchical modeling can leverage information across clusters of observations for efficient modeling of the fixed effects while respecting the dependency structure to avoid potentially biased inference. A random-intercept
model, modeling different baseline measurements per cluster, follows the form of a linear mixed-effects model, which is a special case of the GP model \(^{23}\):

\[
Y_i \sim MVNormal(\mu_i, \sigma^2)
\]

\[
\mu_i = \beta_0 x_i + \beta_1 x_i + b_{[i]}
\]

\[
\bar{b} \sim MVNormal(0, \sigma^2) \tag{3}
\]

Given that spatial omics data may require algorithms that leverage nonlinear, complex high-order interactions offered by machine learning while respecting the dependency structure of the data, practitioners of spatial omics analyses, specifically those developing classifiers, should consider machine learning models which utilize mixed-effects modeling. Several such methods have been developed in the previous decade, being usually formulated as \(^{16,24-26}\):

\[
Y_i \sim MVNormal(\mu_i, \sigma^2)
\]

\[
\mu_i = f_0(x_i) + b_{[i]}
\]

\[
\bar{b} \sim MVNormal(\bar{0}, \Sigma) \tag{4}
\]

This modeling approach learns to estimate a random-effect component, \(\bar{b}\), that allows the estimation of a fixed-effect component robust to clustering by replacing a linear fixed-effects component, \(\beta x_i\), with the fixed-effects machine learning model, \(f_0(x_i)\) (Figure 1C-D). Model fitting is typically accomplished by first removing the random effects from the outcome before fitting the fixed effects machine learning model, \(Y_i^* = Y_i - b_{[i]}\), and then estimating or drawing fixed effects components through frequentist or Bayesian procedures. This procedure is repeated until convergence of both the fixed and random effects components. Potential drawbacks of separating the fixed and random effect components include having to omit the random-effect component during inference, where in many cases out of cluster prediction yields similar results to the fixed effects counterpart, and that complexities in the random effects are not well captured (only random intercepts and slopes). Alternatively, clustering can be directly included in the decision splits of the model. For instance, for a given split, one can fit a generalized estimating equation model with a custom dependency structure to assess the ability to partition variable \(k\) with cutpoint \(a\): \(\logit(p_{ijk}) = \beta_0 + \beta_1 x_{ijk} \leq a + \epsilonij\), where \(\epsilonij \sim N(0, \Sigma)\) and the correspondent Wald statistic on \(\beta_1\) may inform the variable and value to split on \(^{27}\).

In sum, failing to consider the dependency structure in omics data risks introducing bias into the learning process of a machine learning model by learning properties of the batch rather than the inference target. As omics data becomes increasingly available at the single-cell and spatial resolutions, mixed-effect modeling approaches may be optimally positioned to capture potential dependency structures/repeat sampling in the data.

3. Materials and Methods

3.1. Data Acquisition and Preprocessing

Under IRB approval, pathology reports and slides were reviewed for colon adenocarcinoma cases at Dartmouth Hitchcock Medical Center (DHMC) from 2016 to 2019. A total of thirty-six biopsies were initially selected for spatial profiling, half of them exhibiting local invasion but no nodal or distant metastasis (no METS) and the other half of them having undergone nodal and/or distant metastasis (METS). The cohort was restricted to stage three assignments under the pTNM staging system, which balances the impacts of local invasion, nodal and distant metastasis for
prognostication. Cases were matched between the non-metastatic and metastatic groups based on tissue size (measured through connected component analysis of whole slide image), tumor grade, mismatch repair (MMR) dysregulation status, site of the tumor (e.g., left or right colon), age, and sex. Matching was achieved through conducting fisher’s exact tests and two-sample t-tests after iterative resampling. Tissue blocks were sectioned into five-micron sections. Sections were stained with fluorescent-labeled antibodies (highlighting tumor (PanCK), immune cells (CD45), and nuclei (SYTO13)) which are covalently linked to photocleavable oligonucleotide tags targeting a panel of Immune Cell Profiling and Tumor Immune Environment markers. These sections were visualized using immunofluorescent (IF) images provided by the digital spatial profiling (DSP) instrument. Subsequent sections were stained with hematoxylin and eosin (H&E) and scanned using the Leica Aperio-AT2 scanner at 20x. IF whole slide images (WSI) were stored in TIFF format (16-bit unsigned color channels; one channel per stain) and H&E images were stored in SVS format (8-bit color channels). IF and H&E images were viewed simultaneously by the pathologist using the ASAP annotation software, which allowed the pathologist to delineate immune populations within macroarchitectural regions of interest (intra or within tumor, inter or at interface with tumor, away from tumor) on the IF images using polygonal/spline annotations. For each slide, a semi-automated workflow placed eight regions of interest (ROI; square grids of maximal spatial dimensions allowable by the GeoMx DSP instrument) within each annotation region after removing ROI with low immune cell density as quantified by the presence of CD45 stain. The 24 ROIs per slide were mapped onto the IF slide, uploaded to the DSP instrument, and manually registered to the unmanipulated DSP IF images. ROI were placed onto a slide within the device corresponding to the semi-autonomous placements and manually adjusted to the nearest appropriate region by a practicing pathologist if any ROI were misplaced (e.g., incorrect region or low immune cell density). Within each ROI, immune cells were segmented by using color thresholding and connected component analysis to deduce which areas in the ROI had overlapping nucleus and immune stains while excluding the tumor stain. The oligo tags within the immune cell areas were cleaved using ultraviolet light, collected and counted with nCounter to quantify protein expression within the immune cell regions of all ROI. Four slides were assessed at a time due to the DSP’s batching mechanism. Additional cases were included/excluded either due to tissue lifting after cover-slipping procedures (leading to case removal) or to further balance subsequent batches, leading to the inclusion of thirty-five cases (840 ROIs).

A total of 28 ROIs were removed after quality control procedures, retaining 812 ROIs. Within each batch, we separately performed and compared expression after ERCC normalization, normalization based on nuclei area and count, and normalization to housekeepers and IgG isotype controls. Using the raw data, we compared log2-transformed counts of housekeeper markers to each other (S6, Histone H3, GAPDH) and separately compared counts of IgG isotope controls (Rb IgG, Ms IgG1, Ms IgG2a), also looking for significant batch effects. We used this information to sub-select markers that were consistent with other control markers and exhibited the lowest potential for batch effects (Housekeepers: GAPDH, Histone H3; IgG: Rb-IgG, Ms-IgG1), and normalized the expression of the other markers based on the geometric mean of these respective markers. A preliminary analysis demonstrated the lowest impact from technical factors when using IgG isotopes, so we normalized expression based on IgG isotope controls for the following experiments. ROIs were further labeled with positional coordinates, nuclei count and total area, MMR alteration status, age and sex, site of origin/metastasis, tumor grade, nodal and distant metastasis status, and
macro-architectural region (*intra, inter, away*). A total of 36 protein markers were selected for analysis, not including housekeeping genes. We emphasize that the collected data represents a preliminary analysis and does not represent the data for final classifier development. Nonetheless, the data collected may prove helpful for the comparison of different classification algorithms.

### 3.2. Experimental Design: Prediction Tasks and Modeling Approaches

We compared predictive performance of several mixed-effects machine learning (MEML), two fixed effects techniques (Random Forest, XGBoost) 28, and classical statistical methods. First, we treated the patient as a random intercept, whether the ROI could be localized to the tumor interface (*macro*), for both: a) out of cluster predictive accuracy (*macro-ooc*; tested on held out set of patients) and b) within batch accuracy (*macro-ws*; samples from the same batch were partitioned for testing). Then, using the sample collection batch as a random intercept, we sought to establish whether, based on the assigned macro-architectural region, colon cancer metastasis could be predicted (*mets*); first, predicting *mets* regardless of the assigned macro-architecture (*mets-overall*; here including whether ROI was intra- or inter- tumoral as additional predictors). We also stratified by macro-architecture, built classifiers to predict whether there was nodal or distant metastasis (*mets-intra, mets-inter, mets-away*) based on sampling the spatial markers localized to the site of origin. Our primary comparison was to compare out-of-cluster sampling performance with within-cluster sampling for the macro tasks. Since predicting out-of-cluster leads to the exclusion of random effects, we expect differences between fixed effects machine learning models and their mixed effects counterparts to be marginal. Thus, for the *mets* tasks, we focused on comparison of predictions within batch, as we wanted to shift focus from development of a classifier which can be used on external datasets to one where interrogation of important fixed effects predictors could potentially yield etiological factors worth follow up. Age and sex were included as model predictors. We provide brief overviews of selected modeling approaches and their implementations (R v3.6, Python v3.7):

1. **Gaussian Process Boosters (GPBoost):** tree boosting algorithms are additive models which fit weak learners to the residual between the outcome and sum of previous trees. Covariance parameters that account for random intercepts and arbitrary covariance functionals are updated at each boosting step via iterative minimization of the model likelihood and risk functional by gradient/newton descent. GPBoost models can generalize to classification tasks by regressing on a continuous latent outcome, dichotomized to yield binary outcomes. We model patients/batches as grouped random intercepts:

   \[
   Y_i \sim \text{Binomial}(1, p_i) \\
   \text{logit}(p_i) = f_0(x_i) + b_{[i]} \\
   f_0(x_i) = \sum_j^n f_j(x_i) \\
   b_{[i]} \sim N(0, \tau^2I) 
   \]

   (5)

2. **GPBoost with Nested Coordinates (GPBoost-Coords):** Similar to GPBoost, with a random intercept for patient/tissue section and grouped within-patient, a spatial Gaussian Process with covariance function of exponential form which captures spatial dependence between ROI:

   \[
   \text{logit}(p_i) = f_0(x_i) + b^{(1)}_{[i]} + b^{(2)}_{[i]} \\
   b^{(1)}_{[i]} \sim N(0, \tau^2I) \\
   b^{(2)}_{[cluster\ k]} \sim \text{MVNormal}(\bar{\theta}, \Sigma_{[cluster\ k]}) \\
   \Sigma_{[cluster\ k]}_{i,j} = a \times \exp(-\rho \times d(s_i, s_j)) 
   \]

   (6)
3. **Semi-Parametric Mixed Effects Bayesian Additive Regression Trees (SP-BART):** Bayesian additive regression trees are similar to boosting in that decision trees are being successively estimated, this time in the form of particle Gibbs sampling of decision trees, with priors set over tree parameters (splitting structure, predictor selection, tree depth, residual variance) as regularization and Bayesian backfitting techniques to draw new trees based on the residual. This semi-parametric approach uses Markov Chain Monte Carlo procedures to draw additive trees as the fixed effects model components (dbarts), with subsequent Hamiltonian Monte Carlo (HMC) draws via Stan, to update patient/batch-level random intercepts.

4. **Bayesian generalized linear mixed-effects model (BGLMM) with Horseshoe Prior:** Implementation of Bayesian logistic regression using the brms package, where model parameters/random intercepts are drawn using HMC, and horseshoe shrinkage prior with documented performance over Lasso priors to effectively remove irrelevant/collinear genes.

5. **Regularized BGLMM with Interactions (BGLMM-Int):** A hybrid approach between mixed-effects CART methods and generalized linear procedures utilizes SHAP to extract the top 10 salient interactions from the GPBoost model and adds proposed interactions (robust to clustering; Figure 1E) as terms in the BGLMM Horseshoe model, under the premise that these interactions help the GLMM recover any performance gap for their MEML counterparts.

6. **BGLMM with Gaussian Process (BGLMM-GP):** Similar to GPBoost-Cords, replacing the gradient boosting decision trees with a linear functional: \( \text{logit}(p_i) = \beta x_i + b^{(1)}_i + b^{(2)}_i \). To assess the predictive performance of each modeling approach, we performed ten-fold cross-validation and bootstrapped performance statistics (C-Statistic and Log-Loss) through iterative resampling of the held out validation sets of each of the cross-validation folds before averaging. We focused on sensible/coarse selection of hyperparameters: Salient predictors for the CART methods were extracted using SHAP to estimate the local importance of each predictor and summed to form global attributions. Interaction effects were extracted using SHAP’s TreeExplainer. After selecting predictors via Horseshoe Lasso variable selection and standardizing, posterior draws of unpenalized BGLMM model parameters were used to assess and visualize significant predictors.

4. **Results**

4.1. **Macro: Inter-Tumoral Prediction**

The ability to characterize ROI out-of-batch/sample (OOS) indicates the potential to assess new cases prospectively for macro-architectural characteristics. Here, we expected within-sample (WS) performance to exceed predictions made on held-out patients. As expected, the prediction of whether an immune cell ROI was embedded in the interface with tumor exhibited greater performance within-sample across all approaches. MEML methods, taken as a group, slightly outperformed fixed effects approaches for macro-architectural prediction (Table 1). Surprisingly, generalized linear mixed effects approaches outperformed both the fixed effects and MEML methods, with slightly greater performance assigned to the BGLMM model, which utilized interactions detected from the GPBoost model (Table 1). Adding spatial dependence to the model in the form of Gaussian Processes did not improve predictive performance for GPBoost and BGLMM.
Top predictors for tumor interface macro-architecture between the GPBoost model and BGLMM model were concordant (e.g., PanCk, CD127, CTLA4, CD44) and of similar directionality (Figure 2A,C). IgG isotope technical factor Ms-IgG2A was found to be in the top half of predictor for GPBoost, though effects were largely negligible and undetected by the BGLMM. Some of the interactions extracted from the GPBoost model were found to be some of the top predictors for the BGLMM-Int model (e.g., PanCk:CD68, PanCk:CD34) (Figure 2B–C; Table 2), indicating the relevance of MEML extracted interactions in capturing macro-architectural associations.

Table 1: Bootstrapped comparison of model performance, using C-Statistic/Area Under the Curve and Log-Loss

| Task      | C-Statistic ± SE | Fixed Effects | RF | XGBoost | MEML | Bayesian Generalized Linear Mixed Models |
|-----------|------------------|---------------|----|---------|------|------------------------------------------|
|           |                  |               |    |         |      | BGLMM | BGLMM-Int | BGLMM-GP |
| Macro     |                  |               |    |         |      |       |           |           |
| OOS       | 0.759±0.01       | 0.747±0.01    | 0.752±0.01 | 0.747±0.01 | 0.772±0.01 | 0.778±0.01 | 0.765±0.009 | 0.777±0.01 |
| WS        | 0.781±0.009      | 0.773±0.01    | 0.782±0.009 | 0.780±0.009 | 0.784±0.009 | 0.788±0.009 | 0.802±0.009 | 0.786±0.01 |
| METS      |                  |               |    |         |      |       |           |           |
| Overall   | 0.909±0.006      | 0.951±0.004   | 0.971±0.003 | n/a   | 0.897±0.006 | 0.862±0.008 | 0.896±0.006 | n/a         |
| Intra     | 0.834±0.014      | 0.848±0.014   | 0.89±0.011 | n/a   | 0.867±0.012 | 0.848±0.013 | 0.877±0.012 | n/a         |
| Inter     | 0.881±0.013      | 0.865±0.013   | 0.899±0.012 | n/a   | 0.874±0.012 | 0.836±0.014 | 0.861±0.012 | n/a         |
| Away      | 0.827±0.014      | 0.842±0.014   | 0.895±0.011 | n/a   | 0.887±0.011 | 0.885±0.012 | 0.885±0.011 | n/a         |

Figure 2: Macro Predictions: A) SHAP summary plot of top GPBoost terms; B) GPBoost SHAP partial dependency of CD68 interaction with PanCk; C) Posterior distributions of top predictors from unpenalized BGLMM-Int

4.2. METS: Nodal and Distant Metastasis Prediction

The greatest performance differences between fixed effects methods and MEML methods were noticed across all metastasis prediction tasks, where the GPBoost method obtained as little as a 2% boost in C-statistic to as much as a 9% performance gain versus RF and XGBoost. Within batch, it was easier to detect metastasis when all macro-architectural regions were utilized, while performance was roughly equivalent for each of the subregions (Table 1). Since performance of GPBoost exceeded that of all other approaches, the model may have learned significant interactions
between genes that were invariant to batch differences. Using the detected interactions to supplement the BGLMM model allowed the BGLMM interaction model to vastly exceed the performance of its non-interaction counterpart and obtain performance competitive with GPBoost (Table 1). Across all tasks, the BGLMM model mostly outperformed its CART counterpart BART.

GPBoost and BGLMM derived some concordant predictors (e.g. CD20, CD14), while others (e.g. FOXP3) were not found in the BGLMM model (Figure 3A,C). IgG isotope technical factor Ms-IgG1 was found to be in the top half of GPBoost predictors; effects were largely negligible and undetected by the BGLMM. The GPBoost model was also able to detect important interactions, (e.g. effect modification of markers CD20 and Fibronectin by age) (Figure 3B-C; Table 2).

Table 2: Top terms from BGLMM-Int models; GPBoost-extracted interactions are emphasized in bold

| OOS   | WS     | Overall     | Intra      | Inter      | Away      |
|-------|--------|-------------|------------|------------|-----------|
| PanCk | PanCk  | CD20:age    | Beta_2_microglobulin | CD68:CTLA4 | CD25      |
| CTLA4 | CD127  | CD14        | CD45       | CD11c      | CD44      |
| CD68/PanCk | CTLA4 | age         | CD8        | CTLA4      | FAP_alpha |
| CD34  | PanCk/CD68 | CD68   | FOXP3      | CD14       | Fibronectin|
| PanCk:CD27 | CD44 | CD11c       | Beta_2_microglobulin:FOXP3 | GZMB    | CD68      |
| CD68  | PanCk:CD34 | Fibronectin:age | age       | CD68       | age       |
| CD27  | PanCk:CD68 | KI_67        | CD8:CD45   | CD14:CD66b | FAP_alpha:CD163 |
| CD56  | age:sex | age         | age        | CD163      |           |
| CD34  | CD14/CTLA4 | PanCk | Beta_2_microglobulin | CD66b    | PanCk:age |
| PanCk:CD56 | Beta_2_microglobulin | GZMB:age     |           |           |           |

Figure 3: METS Predictions: A) SHAP summary plot of top GPBoost terms; B) GPBoost SHAP partial dependency of CD20 interaction with age; C) Posterior distributions of top predictors from unpenalized BGLMM-Int

5. Discussion

Spatial localization of gene and protein markers provides additional resolution on disease pathogenesis and etiology, which may soon lead to more informative and promising diagnostic and prognostic decision aids. While failing to account for repeated measurements in this setting may threaten to preclude classifier deployment, incorporating mixed-effects modeling into machine learning model development may derive potential additional etiological factors due to nonlinear interaction modeling while accounting for batch effects. In this study, we performed a preliminary
assessment and comparison of different generalized linear and CART models for deployment as classifiers for the assessment of macro-architectural characteristics and potential for undergoing nodal or distant metastasis. Results from our study indicate that classifier development for spatial omics technologies such as GeoMX DSP or Visium’s Spatial Transcriptomics may benefit by taking into account repeated measurements on the case and batch level. At minimum, mixed effects classification algorithms in spatial omics studies may detect complex gene-gene and gene-phenotype interactions that can be applied towards clinical findings (e.g., reportable risk estimates).

The added advantage of employing an MEML model is the ability to extract an interaction that the linear approach can utilize, which may outperform generalized linear approaches without interactions, often unfairly reported in algorithm comparison studies. In this study, using this hybrid approach of informing interactions through MEML methods yielded BGLMM models that could outperform fixed effect and MEML methods due to simpler decision boundaries (line instead of stepwise function) and less biased sampling of the posterior of the parameters.

We acknowledge a few study limitations. Namely, we performed very coarse hyperparameter searches and other notable MEML methods were omitted due to overlapping design. Additional algorithmic fine-tuning may demonstrate superior performance that is different from that reported in this work. The role of batch imbalance (association of batch with an outcome) has been well explored as it degrades the model performance in bulk expression datasets. However, its effects, particularly on intra-batch normalization and estimation of random effects, are unclear with the presented data and demand further exploration. We chose not to include batch effect correction using ComBat to limit the number of comparisons. Regarding the batch effects, other than modeling Gaussian Processes nested within random intercepts, we did not include any random slopes in any of the MEML and BGLMM models, nor did we consider multiple hierarchy levels simultaneously (random intercept for batch and patient/biopsy). METS predictions were made within-batch and are not an accurate reflection of deployment on new samples. Projective predictive feature selection can help select more relevant BGLMM predictors. We also noticed that there was no gain in performance from using the ROI spatial coordinates, which may have been an artifact of implementational difficulties, the low sample size of ROIs and/or selection/learning of improper kernels. Spatial autocorrelation is often studied in spatial omics analyses, and, in the future, we expect to include an in-depth assessment of spatial effects.

In the future, we aim to refocus these findings for a more clinical setting after further algorithmic fine-tuning and validate our research results using immunohistochemistry (IHC). For prospective deployment of these technologies, we envision IHC serving as a low-cost alternative and potentially less encumbered by batch effects to rapidly spatially assess markers that have demonstrated utility for identification of colon metastasis. Such data modalities require calibration of collection and analysis. For instance, this would require automation of macro-architectural segmentation (intra, inter, away), which can be accomplished through deep learning algorithms. We have yet to assess the potential for bias from semi-subjective/automated selection of ROI within these macro-automated regions. Application of chemical reagents demands further standardization to ensure meaningful deployment. Regarding further methods development, aggregation across macro-architectural regions may be explored using graph neural networks, which can pool spatial omics information across these regions to form a succinct vectorial representation of patient’s overall profile that can be integrated with H&E imaging modalities. Simulation studies may provide further validation of the utility of MEML methods for the assessment of spatial omics technologies.
6. Conclusion

In this work, we demonstrated the potential utility of MEML methods for the assessment of spatial protein markers collected using DSP on the Nanostring GeoMx. We illustrated that MEML methods obtain superior within-batch performance for Colon metastasis prediction versus fixed effects and generalized linear methods. Furthermore, MEML methods, when combined with generalized linear modeling, may lead to clearer communication of significant spatial disease associations in clinical research studies through the extraction of key complex interactions. Further standardization of data collection, normalization, and analysis are potential opportunities to consider spatial omics machine learning technologies for meaningful clinical deployment.

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8. References

1. Mars, V. Method of the Year: spatially resolved transcriptomics. Nat Methods 11, 9–14 (2014).
2. Zinn, A. et al. Integration of Digital Pathologic and Transcriptomic Analyses Connects Tumor-Infiltrating Lymphocytes Spatial Density With Clinical Response to BRAF Inhibitors. Front. Oncol. 10, (2020).
3. Jakubowska, A., Kodu, M., Koricinski, W., Karczag-Koruda, L. & Fassnacht, M. Tumor-infiltrating lymphocytes in primary tumors of colorectal cancer and their metastases. Exp Ther Med 8, 4904–4912 (2019).
4. P. D. et al. Advances in mixed cell deconvolution enable quantification of cell types in spatially-resolved gene expression data. (2020) doi:10.1101/2020.01.04.235168.
5. Sun, S., Zhu, J. & Zhou, X. Statistical analysis of spatial expression patterns for spatially resolved transcriptomic studies. Nature Methods 17, 193–200 (2020).
6. Camp, Z. & Niu, Q. Inferring spatial and signaling relationships between cells from single cell transcriptomic data. Nature Communications 11, 2084 (2020).
7. Zhang, M. et al. Spatially modulating sharing platforms: applications and analysis tools. Brief Bioinform. doi:10.1093/bib/bbaa145.
8. Dhillon, T. M. et al. COVID-19 tissue atlas reveals SARS-CoV-2 pathology and cellular targets. Nature 598, 107–113 (2021).
9. McNamara, K. L. et al. Spatial proteomic characterization of HER2-positive breast tumors through neoadjuvant therapy predicts response. Nat Cancer 2, 400–413 (2021).
10. Foster, D. S. et al. Integrated spatial multi-omics reveals fibrilloblast fate during tissue repair. Sci Adv. 2021.04.02.437928 (2021) doi:10.1126/sciadv.2021.04.02.437928.
11. Idee, G. E. et al. The Prognostic Implications of Tumor Infiltrating Lymphocytes in Colorectal Cancer: A Systematic Review and Meta-Analysis. Scientific Reports. 10, 3360 (2020).
12. Laassen, J. et al. Spatial and phenotypic immune profiling of metastatic colon cancer. JCI Insight 3, (2018).
13. Naruecho, I. P. et al. Automated Analysis of Lymphocyte Infiltration, Tumor Building, and Their Spatial Relationship Improves Prognostic Accuracy in Colorectal Cancer. Cancer Immunol Res 7, 609–620 (2019).
14. Naroea, I. P. et al. Spatial immune profiling of the colorectal tumor microenvironment predicts good outcome in stage II patients. npg Digital Medicine 3, 1–10 (2020).
15. Sigrist, F. Gaussian Process Boosting. arXiv:2004.02615 [cs, stat] (2021).
16. Song, Q. & Su, J. DTS: deconvoluting spatial transcriptomics data through graph-based artificial intelligence. Briefings in Bioinformatics (2021) doi:10.1093/bib/bbaa144.
17. Yuan, Y. & Bar-Joseph, Z. GCNs: graph convolutional networks for inferring gene interaction from spatial transcriptomics data. Genome Biology 21, 300 (2020).
18. Breiman, L. Random Forests. Machine Learning 45, 5–32 (2001).
19. Lundberg, S. M. et al. From local explanations to global understanding with explainable AI for trees. Nature Machine Intelligence 2, 56–67 (2020).
20. Parker, H. S. & Leib, J. T. The practical effect of batch on gene expression. Am J Pathol Gene Med Biol 1, Article 10 (2012).
21. Soneson, C., Gerstein, S. & Delorenzi, M. Batch Effect Confounding Leads to Strong Bias in Performance Estimates Obtained by Cross-Validation. PLOS ONE 9, e100315 (2014).
22. McShane, L. Statistical vetting: A brevior course with examples in R and Stan. CRC press, 2020.
23. Signoret, F. Latent Gaussian Model Boosting. arXiv:2105.08966 [cs, stat] (2021).
24. Hugan, A., Bellassance, F. & Larouque, D. Mixed-effects random forest for clustered data. Journal of Statistical Computation and Simulation 84, 1313–1328 (2014).
25. Chessa, T. & Guastella, C. XBosost: A Scalable Tree Boosting System. in Proceedings of the 22Nd ACM SIGKDD International Conference on Knowledge Discovery and Data Mining 785–794 (ACM, 2016).
26. Berger, E. et al. A Mixed-Effects Random Forest with Variable Selection for Stratified Categorical Outcomes. arXiv:1809.07982 (2018).
27. Kasper, A. & Bleich, J. btetMachine: Machine Learning with Bayesian Additive Regression Trees. arXiv:1312.2711 [stat] (2014).
28. Carpenter, B. et al. Stan: A Probabilistic Programming Language. vol. 76 (2017).
29. Breiman, L. Bagging Predictors. Machine Learning 24, 123–140 (1996).
30. Carvalho, F., Hoffman, M., Neal, R. & Zhang, D. Handling Sparsity via the Horseshoe. in Advances in Neural Information Processing Systems 25 (eds. Gao, Y. et al.) 1705–1713 (Curran Associates, Inc., 2012).
31. Carvalho, F., Polson, N. G. & Scott, J. G. Handling Sparsity via the Horseshoe. in Artificial Intelligence and Statistics 25 (eds. Larochelle, H. et al.) 161–169 (PMLR, 2012).
32. Pedregosa, F., et al. Scikit-learn: Machine Learning in Python. Journal of Machine Learning Research 12, 2825–2830 (2011).
33. Lundberg, S. M. & Lee, S.-I. A Unified Approach to Interpreting Model Predictions. in Advances in Neural Information Processing Systems 30 (eds. Guyon, I. et al.) 4765–4774 (Curran Associates, Inc., 2017).
34. Levy, J., Lebrun, H., Christensen, B., Tosicnic, T. D. & Bryan, J. Journey across epistromic’s third variables: an anesthesiologist’s guide to successfully navigating confounding, mediation, and effect modification. Regional Anesthesia & Pain Medicine (2021) In press--2021R019080.
35. Speer, J. L. et al. BMDM forest: A random forest method for modeling clustered and longitudinal binary outcomes. Chronomics and Intelligent Laboratory Systems 185, 122–134 (2019).
36. Pellagatti, M., Masi, C., Leva, F. & Pagantoni, A. M. Generalized mixed-effects random forest: A flexible approach to predict university student dropout. Statistical Analysis and Data Mining: The ASA Data Science Journal 14, 241–257 (2021).
37. Lin, S. & Luu, W. A New Multilevel CART Algorithm for Multilevel Data with Binary Outcomes. Multivariate Behavioral Research 54, 578–592 (2019).
40. Capitaine, L., Genuer, R. & Thiébaut, R. Random forests for high-dimensional longitudinal data. *Stat Methods Med Res* 30, 166–184 (2021).
41. Chen, C. et al. Removing Batch Effects in Analysis of Expression Microarray Data: An Evaluation of Six Batch Adjustment Methods. *PLoS ONE* 6, e17238 (2011).
42. Piironen, J., Paasiniemi, M. & Vehtari, A. Projective inference in high-dimensional problems: Prediction and feature selection. *Electronic Journal of Statistics* 14, 2155–2197 (2020).
43. Bartonicka, A., Wickham, S. R., Pat, N. & Conner, T. S. The value of Bayesian predictive projection for variable selection: an example of selecting lifestyle predictors of young adult well-being. *BMC Public Health* 21, 695 (2021).
44. Levy, J., Hankemeier, C., Barwick, C., Christensen, B. & Vaicikus, L. Topological Feature Extraction and Visualization of Whole Slide Images using Graph Neural Networks. *Proc. Symp Biocomput* 285-296 (2021) doi:10.1101/2020.08.01.231639.