Machine Learning Methods for Predicting the Field Compressive Strength of Concrete

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Abstract

This research analyzes the compressive strength behavior of field-placed concrete (herein termed field concrete) as a function of mixture constituents. Compressive strength prediction of field concrete is inherently different and more challenging than that of laboratory concrete and merits its own analysis. In this work, we employ both field- and laboratory-obtained data to train and test machine learning models of increasing complexity for compressive strength prediction. This training and testing scheme enables determination of the best-performing model specific to field concrete. In this work, the random forest machine learning model for predicting field compressive strength generated the best performance; the RMSE, MAE, and R2 values were 730 psi, 530 psi, and .51, respectively. The methodological reasons for varying model performance are also examined. Finally, the ability of machine learning models trained on laboratory concrete data to predict the compressive strength of field concrete mixtures is evaluated and compared to those models trained exclusively on field concrete data. The analysis shows that the hybridization of field and laboratory data for building predictive models is a promising method for reducing common over-prediction issues caused by laboratory concrete models that are used in isolation.

Construction and Building Materials
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Keywords: concrete; compressive strength; machine learning; prediction; statistical modeling

1. Introduction

The 28-day compressive strength of concrete is a critical design parameter for reinforced concrete structures [1]. Empirical prescriptive- and performance-based mixture design methodologies remain the conventional means to obtain concrete mixture design proportions that meet minimum 28-day compressive strength requirements. However, numerical approaches for predicting the 28-day compressive strength of concrete are emerging in the literature. Accurate numerical estimation of the 28-day compressive strength of concrete is desirable because more precise prediction (1) provides assurance of concrete quality, (2) reduces the number of concrete batches that are needed to be tested to meet strength targets, and (3) enables a reduction in factors of safety. Recent computational studies have demonstrated the ability of advanced statistical modeling techniques to numerically predict concrete
compressive strength for laboratory-mixed concrete, termed laboratory concrete herein [2]–[13]. However, prediction of the 28-day compressive strength of concrete placed in the field on an actual construction site, termed field concrete herein, remains a challenge for the concrete industry due to variable environmental conditions and other uncertainties

1.1 Prediction challenges for field concrete mixtures

Estimating the 28-day compressive strength of concrete is a multifaceted problem. Complex physical and chemical interactions occur between concrete constituents, which, in turn, affect compressive strength. Therefore, nonlinear mathematical models are advantageous for accurately capturing all phenomena. As an example, consider the following physically intuitive correlations: compressive strength decreases (nonlinearly) as the water-to-cement ratio (w/c) increases [14], [15], while increasing air content for improved workability and freeze-thaw resistance reduces compressive strength [16]. Other correlations have not been as intuitively deduced to-date. For example, it is well known that the proportion of coarse-to-fine aggregate affects compressive strength, but the relationship has not been precisely determined due to confounding factors, such as particle size distribution, aggregate angularity, and water demand. Coarse aggregate, for example, may vary in nominal size, grading, chemical composition, shape, surface texture, and absorptivity [17]; these properties can impact the strength of the interfacial bonds between the aggregate and mortar, which, in turn, affect the compressive strength of concrete. Furthermore, the addition of supplementary cementitious materials (SCMs), like fly ash, slag, and silica fume, also introduce new, complex, and nonlinear relationships to compressive strength because of complex factors, such as fineness, chemical variability, and pozzolanic reactivity [18], [19]. Additionally, the fineness and mineral composition of fly ash and slag can be highly variable, depending on the original industrial source and additional processing steps [20].

The conditions of the job site at which field concrete is mixed and placed are also highly variable and lead to high variability in field compressive strength compared to laboratory concrete. For instance, it is commonplace for the environmental conditions at construction sites to be loosely controlled. Here, temperature, humidity, and inclement weather can all affect concrete curing and the final compressive strength [21], [22]. Such variabilities do not exist in laboratory concrete mixing, which suggests that accurate prediction of the compressive strength of field concrete is a more challenging problem compared to compressive strength prediction of laboratory concrete.

1.2 Machine learning methods for compressive strength prediction

Because of the physical limitations described above, there is growing interest in predicting concrete compressive strength using machine learning (ML) models for both field and laboratory concrete mixtures [23], [24]. ML models predict compressive strength (i.e., the target variable) from the types and quantities of the mixture ingredients (i.e., the input variables). Using pairs of data of the form [input
variables, target variable], a model is trained from a collected dataset and learns the relationship between the target and input variables without constraint on prior intuitive understanding. The vast majority of this type of research has been performed on laboratory concrete, which, as discussed, suggests limitations on the actual usefulness of these models for predicting the compressive strength of field concrete, given the myriad of convoluting factors.

Prior research in ML methods for compressive strength prediction has been limited to testing ML methods using laboratory data to determine best-possible prediction models for concrete compressive strength. A particularly popular ML algorithm is artificial neural networks (ANNs). The first study of ANNs by Yeh et al. [25] employed ANNs on a dataset of over 1000 laboratory concrete mixture designs. Since then, other researchers have reported ANN studies with coefficients of determination (R²) of up to 0.999 [2], [8], [10], [26], [28]. However, a significant number of ANN studies employ less than 100 experimental data points, which may not sufficiently sample the predictor variable space. While ANNs are a flexible and powerful ML method, it suffers from the need to train a large number of parameters. For small datasets (as is common for field concrete), ANNs can quickly overfit the data, which leads to strong training set performance but poor generalization performance on new datasets. Other ML methods that appear in the literature include support vector machines (SVM) [25], [26] and decision tree-based models [13], [29]. Studies that employ these methods are less common than ANN studies, due to the historical alignment of compressive strength prediction and ML methods.

Some narrower-scope prediction studies that used ML have focused on modeling concrete mixtures that contain particular mixture ingredients, such as fly ash [28], blast-furnace slag [30], recycled aggregate [31], silica fume [32], and metakaolin [33]. This body of research generates models that are useful for predicting compressive strength when specific constituents are included. However, this approach narrowly tailors the model to the particular dataset and, thus, is less useful when either mixture ingredients or external conditions (possibly unmeasured) may change.

A recent study by Young et al. considered field concrete data and compared the predictive performance of four ML models for predicting both field and laboratory concrete [23]. This study found that variance can be significantly better explained in the laboratory concrete dataset, which is compatible with the idea that laboratory concrete has fewer uncontrolled variables. The study determined that the four ML methods investigated exhibited equivalent predictive performance for field concrete – a somewhat unintuitive result, given that the four methods employed do not share common assumptions about the underlying data. In this study, it is also of note that the laboratory and field datasets contained different mixture ingredients (i.e., input variables). For example, the laboratory concrete dataset included blast-furnace slag, while the field concrete dataset does not, making an apples-to-apples comparison difficult between models for both laboratory and field concrete.
1.3. Innovative contribution/knowledge gaps
Despite a large body of research in this area of study, the challenge of training a ML model for accurate prediction of concrete compressive strength remains relevant. More specifically, two significant gaps exist in the literature. First, prior studies are not well-grounded in best-practice methods of the ML community. The standard procedure in ML is to generate a pipeline of methods that increase in complexity [34]. The reason for this is two-fold: (1) while powerful, ML methods often search a large model space and may miss simple solutions recognized by the researcher and (2) the failure of simpler models is typically caused by a failure in model assumptions that reveals previously hidden details about the data interactions and non-linear behavior observed in the system. These failures can thus be used to inform the appropriate choice of ML tools for further development. Second, consensus on the best model architectures for predicting the compressive strength of field concrete has not yet been reached.

To this end, this study aims to address the aforementioned knowledge gaps and is particularly focused on approaches for accurate prediction of the compressive strength of field concrete. First, we employ the standard ML procedure of testing models of increasing complexity in order to determine the best-performing model for field concrete. This procedure enables us build on past research by discussing why certain ML methods are particularly well-suited for the concrete compressive strength prediction problem. The field concrete dataset in this study contains 1681 concrete mixtures and was collected by the Colorado Department of Transportation (CDOT). The laboratory concrete dataset in this study was obtained from the University of California, Irvine Machine Learning Repository, which contains data for more than 1000 mixtures [35].

Following the analysis of the field concrete dataset we evaluate the ability of ML models learned on laboratory concrete data to predict the compressive strength of field concrete mixtures. For this analysis, we perform the same ML procedure for the laboratory data and select the best-performing model. This model is then used to predict the compressive strength of field concrete mixtures and the relative model performance is analyzed. It is hypothesized that the laboratory ML model performance will be less-than satisfactory for predicting field compressive strength. Consequently, this work includes an analysis of laboratory models that are supplemented with varying percentages of field data in order to determine if such hybridized datasets can improve performance of laboratory concrete models.

2. Machine Learning (ML) Methods
As discussed in the introduction, this paper builds a pipeline of ML methods with increasing complexity, such that the underlying structure in the training data can be stepwise analyzed. First, in Section 2.1, we describe the ML methods used in the pipeline. We introduce linear methods (i.e., linear regression, polynomial regression), transformed linear methods (i.e., kernelized support vector regression, kernelized
Gaussian process regression), and non-linear methods (i.e., regression trees, boosted trees, random forest). In general, simple models are introduced first, and subsequent models increase in complexity. The simplest methods (e.g., linear regression) tend to require the most assumptions about the underlying data structure, and the most complex methods (e.g., boosted trees) require few assumptions about the underlying structure of the data. Second in Section 2.2, we analyze the utility of predictive models trained on laboratory concrete data for predicting field concrete strength. Third, in Section 2.3, we introduce the performance measures used to evaluate the effectiveness of each model: the coefficient of determination ($R^2$), root mean squared error (RMSE), and mean absolute error (MAE). Last, in Section 2.4, overfitting is discussed, which occurs when a model not only captures the desired qualities in the data, but also begins to exactly model the training data itself. An overfitted model is undesirable because it lowers the predictive performance on “unseen” testing data. In other words, overfitted models do not generalize well to real-world cases. In this analysis, we describe and utilize nested cross-validation as a means reduce overfitting. Reserved testing data is used for final determination of the best-performing model.

### 2.1 ML Methods

All models were created in the R Project for Statistical Computing [36]; in addition, Table 1 lists the ML methods employed in this study, as well as the specific package and function used for model training. For each ML method, we discuss parameter tuning and the intuitive meaning of the parameters.

<table>
<thead>
<tr>
<th>Model Type</th>
<th>R Package</th>
<th>R Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linear Methods</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Linear regression</td>
<td>stats</td>
<td>lm</td>
</tr>
<tr>
<td>Polynomial regression</td>
<td>stats</td>
<td>lm</td>
</tr>
<tr>
<td>Transformed Linear Methods</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kernelized support vector regression</td>
<td>kernlab</td>
<td>ksvm</td>
</tr>
<tr>
<td>Kernelized Gaussian process</td>
<td>kernlab</td>
<td>gausspr</td>
</tr>
<tr>
<td>Non-Linear Methods</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Regression trees</td>
<td>rpart</td>
<td>rpart</td>
</tr>
<tr>
<td>Random forest</td>
<td>randomForest</td>
<td>randomForest</td>
</tr>
<tr>
<td>Boosted trees</td>
<td>gbm</td>
<td>gbm</td>
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</tbody>
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#### 2.1.1 Linear Regression

The simplest model to apply and analyze is linear regression. In addition to providing useful understanding of the data, linear regression also serves as a good baseline from which other techniques can be evaluated. Linear regression is a model that describes the output (target) variable as a linear combination of the predictor variables [37]. This linear combination is a hyperplane in N-dimensional
space, where N is the number of coefficients in the model. The model solution is the hyperplane that minimizes the squared error between the observed output and the predicted output. Mathematically, the solution is described as:

\[
y = x^T \beta,
\]

Eq. 1

where \( x \) is the input vector, \( \beta \) is the N-dimensional vector of coefficients (parameters) for the linear model, and \( \hat{y} \) is the predicted output variable from the model. The underlying assumption in linear regression is that the relationship between the predictor variables and the output variable is linear. Moreover, the model assumes that predictor variables are independent from one another, and the resulting residuals, the difference between the predicted and observed output variables, are both homoscedastic (i.e., have constant variance) and normally distributed. When these assumptions are violated, it indicates that a linear model is not appropriate. When such violations occur, it is reasonable to use transformations on the input data to try to reduce or eliminate the violation in assumptions. Failure of such methods to improve the resulting model error and reduce violation of the assumptions means that the dataset requires more complex non-linear models.

### 2.1.2 Multivariate Polynomial Regression

Multivariate polynomial regression (called polynomial regression in this study) uses nth degree polynomials of the input variables to predict the output variable. Polynomial regression is a generalization of Eq. 1; however, each \( x \) term may be: (1) an original predictor variable (e.g., \( x_1 \)), (2) a pure higher-order term of one predictor variable (e.g., \( x_1^2 \)), or (3) an interaction term between two or more predictor variables (e.g., \( x_1 x_2^2 \)).

A generalized example of an expanded second-order polynomial solution with two predictor variables (for simplicity) is described by:

\[
\hat{y} = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{11} x_1^2 + \beta_{22} x_2^2 + \beta_{12} x_1 x_2
\]

Eq. 2

The transformation of the predictor variables allows for modeling of higher-order relationships and modeling interactions between the input variables; Eq. 2, for example, shows a parabolic relationship. When the original predictor variables are transformed, they are called “features.” This term, also commonly applied to all input variables of the models, denotes the fact that the inputs have been transformed from their original space. In this analysis, polynomials up to third-order are employed, where third order is chosen due to limits in computational power. Since polynomial regression is a form of linear regression, the same assumptions are required—more specifically, independence of the input features,
homoscedasticity of the residuals, and normality of the residuals. Note, however, these assumptions apply to the transformed features and not the original data space.

2.1.3 Kernalized Regression Methods

Kernalized regression methods utilize two mathematical concepts applied in tandem – a transformation of the predictor variables and the pairing of the new predictors with a regression method. These pairings can then be analyzed in order to determine which (if any) kernel and regression assumptions fit the data well.

Kernels are a set of transformations that can be used to map the original predictor variable space to a high-dimensional feature space [34]. Here, this mapping is more complex than the polynomial mappings in the previous section, and all mappings are the result of extensive previous research effort [38]. Each kernel has its own set of tuning parameters that must be optimized. This paper compares four kernel transformations, including: linear kernel, radial basis function (RBF) kernel, sigmoid kernel, and polynomial kernels (up to order 4). The model order of the polynomial kernels is only limited by the available computational power.

Kernel transformations have the form:

\[ k(x, x') = \langle \phi(x), \phi(x') \rangle \quad \text{Eq. 3} \]

where \( k \) is the kernel function, \( x \) and \( x' \) are N-dimensional input vectors (N is the number of predictor variables), and \( \phi \) is a mapping from m dimensions to an m-dimensional space. Note that \( \langle \phi(x), \phi(x') \rangle \) denotes the inner product between the two mappings and can be thought of as a measure of similarity between the two transformed vectors. The kernel tuning parameters are optimized in tandem with the optimization of a regression model. This optimization is discussed below. Table 2 provides the kernel transformation equations and kernel tuning parameters used in this study.

<table>
<thead>
<tr>
<th>Name</th>
<th>Kernel Transformation</th>
<th>Tuning Parameters</th>
</tr>
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<tbody>
<tr>
<td>Linear</td>
<td>( k(x, x') = \langle x, x' \rangle )</td>
<td>n/a</td>
</tr>
<tr>
<td>Radial Basis Function</td>
<td>( k(x, x') = \exp(-\gamma |x - x'|^2) )</td>
<td>( \gamma )</td>
</tr>
<tr>
<td>Polynomial</td>
<td>( k(x, x') = (\gamma (x, x') + r)^d )</td>
<td>( \gamma, r, \text{and } d )</td>
</tr>
</tbody>
</table>

The transformed variables (features) can be utilized with any regression method. The concept here is that parameters for both the kernel transformations and the regression methods are tuned simultaneously such that the cross-validated model error is minimized. When there are multiple tuning parameters, a grid search technique is employed in order to find near-optimal parameter values. In this paper, two kernelized regression methods are tested: support vector regression and Gaussian process regression.
Support vector regression (SVR) is a version of support vector machines (SVM) used for regression purposes (rather than classification) [39]. The regression model generated by SVR depends on only a subset of the dataset, and these data points are deemed support vectors. When an SVR model is trained, support vectors are the points from the dataset that produce error values (ε) larger than a prescribed threshold value. SVR model training generates values for $\beta_m$ (the coefficients for the transformed support vectors) and $\beta_0$ (the intercept). This occurs via minimization of Eq. 4 using gradient descent:

$$\min_H H(\beta_m, \beta_0) = \sum_{i=1}^{N} V(y_i - \hat{y}) + \frac{\lambda}{2} \sum \beta_m^2$$  \hspace{1cm} \text{Eq. 4}

Here, $V(\epsilon)$ is the prescribed error measure, $y$ is the observed target variable, and $\lambda$ is a regularization parameter that serves as a degree of importance given to large error values. When $\lambda$ increases, large errors are more greatly penalized in the model; this parameter can be tuned using cross-validation. In this study the SVR is paired with the aforementioned kernel transformations in order to examine the utility of transformations of the predictor variables.

The second regression method that is employed with the kernel-transformed data is Gaussian process regression (GP). GP can be thought of as the Bayesian interpretation of linear regression. Rather than assuming that the relationship between the predictor variables and the target variable has the prescribed linear functional form (e.g. $\hat{y} = x^T \beta$), GP simply assumes that the data can be represented as a sample from a multivariate Gaussian distribution and that the mean of this distribution is zero. This approach is “less parametric” in the sense that the model is more loosely defined. Using GP, the predictions of the target variable are made using the conditional probability, $p(y_*|y)$. In short: given the data, how likely is a certain prediction for $y_*$? Here, note the subtle difference between $\hat{y}$ and $y_*$. $\hat{y}$ represents a predicted target variable from a model, and $y_*$ represents a distribution of possible outputs from the model. In the case of GP, $\hat{y}$ is the expected value of $y_*$, $E[y_*] = E[y_*|y]$.

Given the assumed Gaussian distribution, the matrix of all predictor variables in the dataset ($X$), the output vector ($y$) and the new matrix of data inputs, the goal is to make a prediction on the new set of data points ($x_*$). The derived conditional distribution has the form,

$$y^*|y \sim N(K, K^{-1}y, K_* - K K^{-1} K_*^T),$$  \hspace{1cm} \text{Eq. 5}

where $K$, $K_*$, and $K_*$ are the covariance matrices resulting from $k(x, x')$, $k(x, x_*)$, and $k(x_*, x_*)$, respectively. The prediction, $\hat{y}$, is the expected value of this distribution, which can be reduced to the equation below.

$$\hat{y} = K K^{-1} y$$  \hspace{1cm} \text{Eq. 6}
Since GP employs only the assumption of a Gaussian distribution and the covariance matrices for model formulation, no tuning parameters are necessary for this regression method beyond those required for the choice of kernel. The performance of GP allows us to assess the veracity of the Gaussian distribution assumption for the data under multiple different transformations of the predictor variables. If none of the above regression methods can adequately model the output variable, models with no linearity assumptions (e.g. regression trees, artificial neural networks) are reasonable model options consider.

2.1.4 Regression Trees

The goal of a regression tree is to generate partitions in the predictor variables such that the target variable can be predicted based on the partitions among the input variables. Figure 1a provides a simple illustration of regression tree “nodes” (i.e., partition rules) and “leaves” (i.e., terminal nodes that lead to one output value). For instance, in the example provided in Figure 1a, there are two predictor variables ($x_1$ and $x_2$). The “root node” (the uppermost blue ellipse) is a rule that partitions the data along $x_1$. For this node, if $x_1$ is greater than 7.5, then the predicted output (in red) is 0.8. However, if the value of $x_1$ is less than 7.5, then one must proceed to the next node in the tree. This process continues until a predicted output variable is reached. For the same example regression tree, Figure 1b demonstrates that a regression tree partitions the predictor variables into rectangular spaces; the and the predicted output is the same value throughout each of these rectangular cells.
Figure 1: (a) Diagram of an example regression tree model with two predictor variables, $x_1$ and $x_2$. (b) This diagram shows the same decision tree using the two predictor variables as axes. It helps visualize the rectangularity of the target variable predictions when simple regression trees are employed. Within each rectangle the predicted target variable would be the same.

Training a regression tree is performed by selecting partitions in succession using a criterion of variance reduction in the target variable [40]. Since each successive partition is always chosen such that the variance of the target variable is reduced, regression trees are prone to overfitting the data. To prevent overfitting, a variety of regularization techniques can be employed. This study minimizes the cost complexity function, which places a penalty for each additional node that is selected for the model. As shown below, the cost complexity function $R_a(T)$ has two terms that influence its value:

$$R_a(T) = R(T) + \alpha \cdot f(T),$$  \hspace{1cm} Eq. 7

Where $R(T)$ is the training error, $f(T)$ is the number of leaves in the regression tree, and $\alpha$ is the regularization parameter that is determined via cross-validation [41]. In Section 2.3 the cross-validation procedure used in this study is thoroughly discussed.

Regression trees have the advantage that they do not assume linearity in the data, and, therefore, no complex data transformations are needed. Overall, this approach is simpler than linear methods, but it requires careful consideration so as not to overfit the data. Regression trees also implicitly select variables, which means that a trained regression tree will show variables that have more importance for predicting the target variable in earlier nodes in the tree. Lastly, regression trees are interpretable and can provide some insight on the dataset being analyzed.

A disadvantage of simple regression trees is that they suffer from model instability; in other words, small changes to the dataset might create a completely different set of partitions, and, consequently does not lead to the best-performing model. For this reason, more complicated tree-based methods are often considered that are more stable. Random forest and boosted trees are examples of more complex tree-based methods that aim to reduce this instability and are discussed in the subsequent sections.

2.1.5 Random Forest

Random forest is a method that builds an ensemble of regression trees in order to reduce the instability of individual trees. Random forest utilizes two strategies for improving the instability issue. First, it employs the concept of “bootstrap aggregation” (sampling with replacement) in order to generate many similar datasets that were sampled from the same original dataset. These datasets each lead to an individual tree within the ensemble. Second, it incorporates randomness during tree-learning in order to reduce the
correlation between each tree within the ensemble. For instance, when generating new nodes (for individual trees within the random forest), only a subset of the original predictor variables is selected as the set of candidate variables on which to partition the data. The variable value that minimizes variance in the output from these randomly selected predictors is the variable selected for that node. This process is repeated for all nodes in a regression tree and then for all regression trees in the random forest. For a random forest model, the tuning parameters are: the number of randomly selected predictors ($k$), the number of individual trees that are trained ($n$), and the tree depth ($d$) [42].

The advantage of the random forest method is that it significantly reduces the instability of simple regression trees. Furthermore, this method has been shown to minimize correlation between trees compared to other tree-ensemble methods (e.g. “bagging trees” that use only bootstrap aggregation and not random variable selection) [40]. One disadvantage of random forests is their reduction in interpretability compared to simple regression trees; random forests cannot be easily visualized and individual trees are often not good predictive models on their own. However, variable importance plots can reveal the relative importance of predictor variables.

### 2.1.6 Boosted Trees

Like random forest, boosted trees are an ensemble method for dealing with the instability and poor predictive performance of simple regression trees. Generally, the concept of “boosting” is an ensemble strategy that can be used to improve weak learning algorithms (e.g. regression trees) [43], [44]. Boosting can be applied to any weak learning algorithm but is commonly utilized for regression trees. The main concept of boosting is to build a model using the weak learning algorithm. Then another model is learned on the residuals from the first model. This step of model-building on the previous model’s residuals is repeated for a set number of iterations. Therefore, a boosted tree is simply a model where the weak learning algorithm used in each iteration is a regression tree.

Unlike random forest in which all trees are of the same importance, boosted trees are hierarchical, meaning that each tree layer is constructed recursively. The tuning parameters for boosted trees are: the number of trees, the interaction depth (maximum number of nodes per tree), the minimum number of observations per node (a stopping criteria used to prevent trees that have only one observation at each leaf), and the shrinkage rate (the rate at which the impact of each additional tree is reduced).

Boosted trees are similar to random forest in their advantages and disadvantages. Boosted trees tend to have high predictive performance on highly nonlinear datasets and can be successful on problems where there is unequal importance of predictor variables [34]. One disadvantage of boosted trees is that this method has low interpretability; it is difficult to gain much intuition of the patterns that the model has
learned or to determine why a boosted tree model is successful (or not) at predicting the target variable. This means that a strong ML pipeline must be used to train boosted trees to ensure that the approach has not overfit the data.

2.2 Testing of Laboratory and Hybrid Models for Field Concrete Strength Prediction

2.2.1 Laboratory Models

As was discussed in the introduction, many studies in the literature have developed ML models for predicting concrete compressive strength using laboratory concrete datasets. While these laboratory models report high predictive performance \([2], [4]–[8], [10], [26], [27]\), it has not yet been tested whether they are useful for predicting the compressive strength of field concrete. A significant novelty of this study is that laboratory models are tested to determine if they are, in fact, useful for predicting compressive strength when presented with other datasets – namely, field concrete data.

One issue preventing the direct testing of laboratory ML models from the literature is the use of concrete age as a predictor variable. In other studies, age is a convenient predictor variable because it can explain a high percentage of variance in compressive strength data. In other words, removing age as a predictor and using only the final compressive strength as the output causes the compressive strength problem to be significantly more difficult (i.e., model performance measures tend to be poorer). In this analysis, the desired model output is the final compressive strength (approximated by the 28-day strength) of a concrete mixture as a function of only the quantities of the mixture ingredients. Due to this difference between the prediction problem described herein and that of the literature, laboratory models for predicting the 28-day compressive strength using laboratory concrete data have been trained specifically for this study. Model utility is examined via the process described below and illustrated in Figure 2.

First, the aforementioned suite of ML models (i.e., linear regression, polynomial regression, kernel regression, tree-based models) is trained and tested using the laboratory data described in Section 3. The model with the best testing performance is selected. Then, the predictor variables from the field data are used as inputs and the performance measures and diagnostic plots for this new data shall be reported and analyzed.
2.2.2 Models Trained on Hybrid Data

It was hypothesized that the previously-described laboratory model will not satisfactorily predict the compressive strength of real concrete mixtures. Thus, an analysis of models trained on hybrid data (i.e., a dataset that is composed of both field laboratory data) is conducted to determine whether they can improve predictive performance compared to “pure” laboratory models.

Models trained on hybrid data are potentially valuable because there is an inherent tradeoff between the use of laboratory and field models for predicting real concrete compressive strength. On one hand, laboratory data is the cheapest and most accessible data to acquire. It is also the best method for exploring new and exotic concrete mixtures that are uncommon in industry. However, laboratory compressive strength data has the disadvantage that it does not reflect the full set environmental variables experienced by field concrete. Accordingly, it is expected that ML models trained on hybrid data may have the potential to improve the predictive performance of laboratory models.

In this novel hybrid approach, a percentage, $\alpha$, of the hybrid dataset is composed of the field data, and the rest is composed of the laboratory dataset. This procedure is used to determine if small amounts of field data can improve model performance. In order to determine the effect of variable amounts of field data, different $\alpha$ values are utilized (10%, 20%, 30%, 40%, and 50%). The model building process occurs, as follows, for each value of $\alpha$:

For each $\alpha$:  

**Figure 2.** Process for testing the predictive capability of laboratory models using field concrete data. The dotted outline indicates the laboratory model that is selected based on its performance measures.
1. Sort the field dataset in the order of lowest compressive strength to highest compressive strength and partition this sorted dataset into quintiles.

2. In order to ensure the field data portion of the hybrid data is well-sampled, randomly sample (in equal number) the appropriate number for points from the quintiles of the sorted field dataset. Randomly sample from the field dataset the appropriate number of points.

3. Use this hybrid data to train a cross-validated ML model. (The selection of ML model is determined by the best performing laboratory model.)

4. Use the remaining, unsampled field data to determine the average testing performance of the hybrid model. The performance measures described in the following section are reported.

5. Repeat steps 1-4 five times to find average performance measures for each $\alpha$.

2.3 Performance Measures

When training statistical, data-driven models, it is necessary to have a method to quantify the model performance so that hyperparameter tuning can be iterated to select the best possible model. There are several established metrics for determining predictive performance, each with advantages and disadvantages, which will be discussed below. Common quantitative performance measures common to regression modeling (rather than classification modeling) include the coefficient of determination ($R^2$), root mean square error (RMSE), and mean absolute error (MAE) [45], [46]. These metrics, coupled with model diagnostic plots and visualization of predicted versus observed output values, provide a comprehensive picture of a model’s performance.

$R^2$ is a measure of the proportion of the variance in the data that is explained by the model. Accordingly, $R^2$ is the ratio shown in Eq. 8, where $y_i$ is the observed value from the data, $\hat{y}_i$ is the predicted value from the model, and $\bar{y}$ is the average output from the data.

$$R^2 = \frac{\sum_i (y_i - \bar{y})^2}{\sum_i (\hat{y}_i - \bar{y})^2}$$

Eq. 8

The value of $R^2$ ranges from zero to one, with higher values indicating a better ability to explain the variance in the data with the model. However, $R^2$ is a measure of correlation, not accuracy, and should be used with other performance measures because it is dependent on the variance of the output variable.

The root mean square error (RMSE) indicates how concentrated the data is around the model fit. The RMSE is measured on the same scale as the output variable, and is always positive due to the squared residuals in its calculation. Using the RMSE accentuates the effect of outliers in the error metric. This means that if median error of the model (usually captured by the mean absolute error) is low, the RMSE
of the model can still be large due to the inability to model some outliers in the data. Given observed values, $y_i$, predicted values, $\hat{y}_i$, and $n$ observed values RMSE is calculated as:

$$RMSE = \sqrt{\frac{\sum_{i=1}^{n}(y_i-\hat{y}_i)^2}{n}}$$

Eq. 9

The mean absolute error (MAE) is a measure of prediction accuracy of a model that uses the absolute value of the errors rather than a squared value. The use of the absolute value reduces the influence of very large errors on the measure of performance. Thus, MAE is a measure of the median error of the model and is complimentary to the use of $R^2$ and RMSE.

$$MAE = \frac{\sum_{i=1}^{n}|y_i-\hat{y}_i|}{n}$$

Eq. 10

Like RMSE, MAE is measured on the same scale as the output variable, and a lower value indicates a better model fit. In addition, MAE values for a model are typically smaller than the RMSE value for the same model. In this paper we chose to use both RMSE and MAE in order to report both the median and mean error of each model.

2.4 Cross-validation

A critical issue to consider when training and comparing statistical and machine learning models is the prevention of overfitting. Overfitting is a problem for ML models that have a high capacity to learn non-linear relationships and are trained on datasets that do not contain a sufficiently large variance of the data (i.e., on datasets that are not rigorously sampled). When using iterative training methods such as grid search, a model is particularly prone to overfitting if the same data is used for the training and validation datasets. In this case, the resulting performance measures would indicate that the model has good predictive performance, but when these models are tested on new data, poor performance is observed.

To prevent overfitting, ML learning methods and pipelines can employ several strategies. The strategy employed herein, is called nested cross-validation (nested CV), which splits the data into “training”, “validation”, and “testing” datasets. In the “inner CV loop”, the performance measures are approximately optimized by fitting a model to each of several training datasets. Subsequently, the performance measures are directly optimized by selecting hyperparameters with each validation data set. In the “outer CV loop”, the testing error is estimated by averaging test set scores for several dataset splits. In order to prevent data leakage, it is critical that the trained models have never been exposed to the testing data.

When performing CV, the selection of the sizes of the training, validation, and testing sets is critical because this choice affects the bias/variance tradeoff for a given statistical model. To strike a balance between bias and variance error, this paper uses five folds (i.e., partitions) for both the inner and outer CV
loops which can generate a favorable bias/variance tradeoff according to the literature [34]. This choice results in 25 validation scores and 5 testing performance measure scores for each model.

3. Datasets

In this study, two datasets – field and laboratory concrete compressive strength data - are used. The field dataset is from the Colorado Department of Transportation (CDOT); it has 1681 mixture designs and corresponding compressive strength values. The mixture constituent variables in this dataset include masses of cement, fly ash, water, water-reducing admixtures (WRA), coarse aggregate, fine aggregate, and percent air entrainment. The laboratory dataset was obtained from the Machine Learning Repository at the University of California, Irvine [35]. This dataset contains over 1000 mixture designs and corresponding compressive strength values. However, it originally contained some mixtures that included blast-furnace slag (a mixture ingredient not included in the field dataset) as well as some mixtures in which the compressive strength was measured earlier than 28 days of curing. In order to reconcile these differences, only mixtures that do not include blast furnace slag and that measure compressive strength after 28 days are included in this analysis. This decision reduced the number of usable mixtures to 311. One last discrepancy is that the laboratory dataset does not report air entrainment values. It is not clear which of the following is true: a constant amount of air was entrained, no air was entrained, or variable amounts of air were entrained but not reported. Notably, this discrepancy does not prevent model training for either dataset. However, when the best laboratory predictive model is used to predict field compressive strength, the air entrainment predictor cannot be utilized.

Table 1 provides a statistical summary of the two datasets. The laboratory dataset has been converted to US customary units for ease of comparison. Note also that both datasets have used the Absolute Volume Method for proportioning concrete mixtures, which generates weights of ingredients on a cubic yard basis; this means that ingredient quantities are comparable between datasets.

Table 1: Statistical summary of laboratory- and field-acquired datasets

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Statistic</th>
<th>Cement</th>
<th>Fly Ash</th>
<th>Coarse Aggregate</th>
<th>Sand</th>
<th>Water</th>
<th>Air</th>
<th>WRA</th>
<th>Strength</th>
</tr>
</thead>
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<td></td>
<td>Units</td>
<td>lbs/yd³</td>
<td>lbs/yd³</td>
<td>lbs/yd³</td>
<td>lbs/yd³</td>
<td>lbs/yd³</td>
<td>Vol. %</td>
<td>Oz/yd³</td>
<td>psi</td>
</tr>
<tr>
<td>Lab</td>
<td>Mean</td>
<td>501</td>
<td>113</td>
<td>1678</td>
<td>1332</td>
<td>307</td>
<td>-</td>
<td>149</td>
<td>5357</td>
</tr>
<tr>
<td></td>
<td>Median</td>
<td>487</td>
<td>161</td>
<td>1689</td>
<td>1330</td>
<td>314</td>
<td>-</td>
<td>154</td>
<td>5362</td>
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<tr>
<td></td>
<td>Min</td>
<td>227</td>
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<td>1350</td>
<td>1001</td>
<td>236</td>
<td>-</td>
<td>0</td>
<td>1239</td>
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<tr>
<td></td>
<td>Max</td>
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<td>337</td>
<td>1896</td>
<td>1593</td>
<td>384</td>
<td>-</td>
<td>761</td>
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<tr>
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<td>Mean</td>
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<td>106</td>
<td>1697</td>
<td>1256</td>
<td>265</td>
<td>6.6</td>
<td>28</td>
<td>5938</td>
</tr>
</tbody>
</table>
4. Results and Discussion

In this analysis, we evaluate the predictive performance of the aforementioned ML models. The values for RMSE, MAE, and $R^2$ for all models are reported in Figure 3. Low values for RMSE and MAE, and high values for $R^2$ indicate better model performance, respectively. For simplicity of discussion, RMSE is used as the primary metric of performance. In addition, both the testing and validation performance is reported, which facilitates the discussion on overfitting in the models. These performance measures are plotted as boxplots to illustrate the range and variance of the error. The set of errors for each model is determined using a nested five-fold cross-validation, with five testing values and twenty-five training values for each model. Each model’s performance from a methodological standpoint is discussed in the sections to follow. The methodological and architectural reasons for each model’s performance are also examined.
(a) RMSE (psi) for different models:

- Linear
- Poly 2
- Poly 3
- SVR Linear
- SVR Poly 2
- SVR Poly 3
- SVR Poly 4
- GP Linear
- GP Poly 2
- GP Poly 3
- GP Poly 4
- GP RBF
- RT
- BT
- RF

(b) $R^2$ for different models:

- Linear
- Poly 2
- Poly 3
- SVR Linear
- SVR Poly 2
- SVR Poly 3
- SVR Poly 4
- GP Linear
- GP Poly 2
- GP Poly 3
- GP Poly 4
- GP RBF
- RT
- BT
- RF

(c) MAE (psi) for different models:

- Linear
- Poly 2
- Poly 3
- SVR Linear
- SVR Poly 2
- SVR Poly 3
- SVR Poly 4
- GP Linear
- GP Poly 2
- GP Poly 3
- GP Poly 4
- GP RBF
- RT
- BT
- RF
Figure 3. Boxplots of the three cross-validated performance measures – (a) RMSE, (b) $R^2$, and (c) MAE for all ML models. Both the training and testing performance measures are reported. The abbreviations are as follows: linear regression (Linear), polynomial regression (Poly), support vector regression (SVR), Gaussian process (GP), regression tree (RT), boosted tree (BT), random forest (RF). Kernels are referred to as follows: second-order polynomial (Poly 2), third-order polynomial (Poly 3), fourth-order polynomial (Poly 4), radial basis function (RBF). Note also that the testing RMSE and MAE scores for the polynomial regression models are so large that they do not fit on the page.

4.1 Linear Regression

Linear regression is the first model tested in this analysis. This model assumes that the predictors are independent and the residuals are homoscedastic and normally distributed. The performance of linear regression is used as a baseline for comparing model performance and for determining what other models may be more appropriate for the data. For the linear regression model, Bayesian information criterion (BIC) – a parsimonious model selection criterion – is employed to select important predictor variables. Of the seven mixture ingredients, BIC selects five of these as predictor variables (cement, fly ash, water, air, and WRA); this model has a mean testing RMSE, MAE, and $R^2$ of 803 psi, 582 psi, and 0.40, respectively. Of note is the relatively low value of $R^2$, which indicates that a linear model is only able to capture 40% of the variance in the data.

There are two possible reasons for the poor performance of this model. One reason is that there are strong predictor variables that were not measured in the dataset. Consequently, the model does not have all necessary information and is unable to perform well. A second possible reason is that the data does not fit the linear assumption of the model, that is, the assumption that the predictors are linearly to produce an output. These possibilities are further evaluated below in diagnostic plots.

Four diagnostic plots are shown in Figure 4. Figure 4a shows a plot of the residuals versus the predicted outputs; significant deviation of the smoothed red line indicates non-constant error variances and outliers. For this model, the smoothed average of the error variances indicates nearly constant error variance. The quantile-quantile (Q-Q) plot (Figure 4b) diagnoses the normality of the residuals. Normal residuals (in the statistical sense) lie along the dotted line; however, this figure indicates that there is some deviation from normality of the residuals among higher residual values. Figure 4c is a scale-location plot, which illustrates whether the homoscedasticity assumption is violated. For this plot, the residuals are standardized (to have a mean of zero and a variance of one) and the absolute value is taken. This plot shows that there is a slight increase in error variance with increasing compressive strength, which is indicative of minor heteroscedasticity. Lastly, Figure 4d shows the standardized residuals against their
leverage, which is helpful for indicating if particular points more strongly influence the regression. In this case, a few outlier points more highly influence the regression. However, the figure also plots contours of the Cook’s distance measure, which measures the effect of deleting a given observation. Cook’s distance is increased by both leverage and large residuals. Since no points have a Cook’s distance greater than 0.5, there is no great concern about large residuals also having too great of leverage over the fit.

One conclusion from the model diagnostics is that there are only minor assumption violations (non-normality of residuals and heteroscedasticity). Despite this result, the linear model retains poor predictive performance, which indicates that there are unmeasured variables needed for predicting compressive strength. Nevertheless, it is reasonable to investigate the use of other types of models to determine if improved performance can be achieved.

4.1.1 Polynomial Regression

Figure 4. Model diagnostic plots: (a) Residuals versus predicted plot to check for non-constant error variance for both positive and negative residuals, (b) Quantile-quantile plot to check normality of residuals, (c) Scale-location plot to inspect homoscedasticity, and (d) Residuals versus leverage plot to determine if any outliers severely impact the regression equation. The blue lines represent the smoothed average for each model diagnostic.
Polynomial regression introduces higher order terms and interaction terms between variables, which can sometimes improve model performance because they approximate unobserved phenomena. Here, the polynomial regression has potential because the linear regression analysis indicates a lack of the necessary predictors for improving model performance. In this analysis, polynomial regression is employed for second order and third order terms to determine if there is a physical basis for higher order variables or interaction terms.

One key aspect of polynomial regression is that the method acts like a feature selection method. In other words, a set of polynomial features is created, and then the features with the largest reduction in RMSE are kept for the final model. This is the method by which interaction terms are discovered. During the experiments in this paper, the following terms were discovered and included in the model: \((Water \times WRA) \times (Air)\) and \((Cement)^2 \times (Fly \ ash)\). The first feature is somewhat intuitive; it is expected that some interaction between water and WRA would be relevant. However, it is somewhat less intuitive that air content is also a part of this feature. The second feature is intuitive because it is expected that fly ash and cement would interactively have an impact on concrete compressive strength.

Promisingly, polynomials of order two and three decrease the training RMSE compared to the linear model by 2.0% and 2.8%, respectively. Given this trend, it’s likely that the RMSE of this model will decrease given unlimited computational power.

However, it is critical to also analyze the testing error. The testing error values for polynomial orders two and three are higher than the training error by 40.6% and 123.8% and are too large to reasonably fit on Figure 3. This result suggests that the polynomial regression models overfit the data as the polynomial order grows. Thus this model type is not suitable for compressive strength prediction in concrete.

4.2 Kernel Transformations and Regression

A different approach to discovering interactions and modeling unobserved phenomena is to use non-linear transformations of the data. Some of these are commonly known as kernel transformations. This section will survey techniques in using kernel transformations.

4.2.1 Support Vector Regression

Solving the regression problem using kernel transformations, support vector regression is a popular technique that has shown good results in the literature. In this paper, an array of kernels was tested in cross-validation. These kernels include the RBF kernel, and polynomial kernels (2, 3, and 4). One of the major goals of adaptive regression techniques like SVR is to discover any underlying structure in the data. Of the tested kernels, the RBF kernel has the greatest reduction in RMSE compared to linear
regression. Here, RBF SVR reduces the average RMSE by 2.9%. In contrast, the linear and polynomial kernels (orders 2, 3, and 4) reduce this error by -0.6%, -0.1%, 1.1%, and 0.2%, respectively. From this result, it is inferred that the RBF kernel generates the optimal hyperplane for linearly separable patterns among the tested kernels. The minimal improvement from polynomial kernels implies that the regression curve is not well-modeled by a polynomial.

The performance of SVR with RBF demonstrates that transformation of the predictor variables improves upon the linear regression baseline model. However, as will be demonstrated in section 3.3, further improvements in performance can be made with other models. One possible explanation for this behavior is that SVR can suffer from the curse of dimensionality in the sense that all terms in the transformed space are given equal weight, so the kernel cannot adapt itself to focus on the critical “subspaces” of the data [34]. Hastie et al. illustrates this concept via a prediction problem with four standard normal features (i.e., “real” features) with a polynomial decision boundary and six Gaussian random features (i.e., “noise” features) [34]. Although applying a polynomial kernel with SVR reduces the test error, the real features are drowned out by the noise features. In the example, kernelized SVR is unable to perform as well compared to when the real features are the only modeled features. We hypothesize that this behavior is also true in this case; the noise of irrelevant variables essentially overpowers the predictive capability of SVR to capture the true underlying behavior of field compressive strength.

4.2.2 Gaussian Process Regression

As is displayed in Figure 3 the GP training and testing performance show that the RBF kernel also generates the highest performance for GP for the kernels utilized in this study. Compared to the linear regression baseline, the GP with RBF-transformed data decreases the average testing RMSE by 3.6%. Utilizing the linear and polynomial (orders 2, 3, and 4) transformations, the reduction in RMSE is -0.1%, 1.0%, 2.5%, and 1.6% respectively. With these results, we can conclude that the same transformation (RBF) generates the hyperplane most suitable for use in both SVR and GP.

Moreover, this analysis shows that GP is preferred over SVR for this type of data due to its improved performance measures. We hypothesize that GP is a better-performing method (compared to SVR) due its further relaxation of the linearity assumption. Unlike GP, SVR retains the assumption that a transformation of the predictor space causes the data to be linearly separable. GP, on the other hand, makes predictions based on the maximum likelihood of an output given the data, normal parameter distributions, and penalty term that minimize the prediction error. The improved performance of GP over SVR indicates that model performance improves when no linearity assumption exists.

4.3 Tree-based Models
4.3.1 Simple Regression Trees

Unlike the aforementioned techniques, tree-based methods assume that the predictor variables may be partitioned repeatedly and that each final partition generates a different output value. For the simplest tree-based method (regression trees), the average testing RMSE indicates an increase of 6.9% compared to linear regression. We hypothesize that this result is due to the instability of regression trees. In other words, the constructed nodes for a tree may change significantly if the input training sample is slightly changed. Figure 3 illustrates the decreased performance of this model for all three metrics: RMSE, MAPE, and $R^2$.

Although the testing performance of the simple regression tree indicates it should not be used for prediction, the results of the model can be used to better understand the relative importance of certain variables for determining concrete compressive strength. In Figure 7, the nodes (e.g. Cement < 569 lbs.) and terminal node predictions (e.g. 4868 psi) are illustrated in the regression tree graph. Values of cement are the first and second nodes, as well as multiple nodes lower in the tree, which indicate the importance of cement quantity as a discriminating predictor variable for this tree. The next most important variable is the quantity of fly ash, which, like cement, has positive correlation with strength. All of the mixture ingredients appear in nodes in the tree, indicating that all are valuable for prediction.

Figure 5. This figure represents the best-performing simple regression tree graph for compressive strength prediction. Final predictions from each terminal node are shown in ellipses.
4.3.2 Boosted Trees

Boosted methods are used to reduce the instability of single trees. In this paper, the ensemble tree model reduced the average testing RMSE by 13.2% compared to the simple regression tree and by 6.9% compared to linear regression. For this dataset, boosted trees are the second best method for prediction based on the three performance measures. Notably, the average training RMSE for boosted trees (749 psi) is slightly lower than that of the random forest model (751 psi). However, the random forest model has the lower testing RMSE by 5.4%. Despite the nested cross-validation routine, it appears that the boosted tree model is slightly overfitted due to the higher value of testing RMSE compared to the training RMSE. Recall from section 2.1.6, that this method iteratively builds regression trees on the residuals from each consecutive tree. We hypothesize that the model has learned noise in the residuals rather than signal in the data, which has lead to lower testing performance.

4.3.3 Random Forest

Like boosted trees, the random forest model reduces the instability of simple regression trees by utilizing an ensemble of trees that utilize bootstrap aggregation and random variable selection. Consequently, the model decreases the average testing RMSE by 9.4% compared to the linear model. It also improves upon the testing RMSE of the simple regression tree by 20.0%. Furthermore, the average testing error is slightly lower than the average validation error (730 psi versus 739 psi) indicating that it is unlikely that the random forest model is overfitted. These testing and validation performance measures indicate that random forest is the best method for predicting compressive strength with this dataset. It has the lowest RMSE and MAE as well as the highest R^2 value (730 psi, 530 psi, and .51, respectively).

This result may be due to the ability of tree-based methods to learn inconsistent variable importance in the data. In other words, each tree, trained on a subset of the data might learn a slightly different set of variable importance weights. In aggregate, the random forest can then better predict the target variable. An example of inconsistent variable importance can be seen in Figure 5; for mixtures with cement quantities of less than 569 pounds, the next most important variable for determining strength is fly ash. In contrast, above 569 pounds, the next splitting criterion is an even higher quantity of cement. Not only do random forest models have the ability to learn inconsistent variable importance, they also reduce the instability of individual trees and reduce the potential for overfitting [47].

4.4 Prediction of Field Compressive Strength with Laboratory and Hybrid Models

4.4.1 Models Trained on Laboratory Data
As was discussed in Section 2, many studies in the literature have developed ML models for predicting concrete compressive strength using laboratory datasets. While these laboratory models report high predictive performance, it is relevant to consider whether they are useful for predicting field concrete strength.

Consequently, in this study, a suite of ML models (i.e., linear regression, polynomial regression, kernel regression, tree-based models) is trained and tested using the laboratory data described in Section 3. Among those tested, the highest-performing model for the laboratory dataset is the random forest model, in which the number of random variables selected at each node was 3, and the number of trees was 550 trees; this model achieves a testing $R^2$ value of 0.80.

Subsequent to the random forest model selection, the predictor variables from the field data have been used as inputs in the laboratory random forest model to determine how well the model can predict compressive strength of real concrete. The predicted output is plotted versus the observed field strength value in Figure 6. Points near the 1:1 line would indicate a high-performing model. This plot shows that despite its high performance using laboratory data, the laboratory model is not able to predict field strength to a high degree of accuracy; the RMSE for the field data is 1655 psi. Furthermore, this plot illustrates that, overall, the laboratory model tends to over-predict compressive strength. It is likely that this effect is due to the ideal curing conditions in the laboratory setting, which would tend to generate higher compressive strength values than if the same mixture was cured under highly variable environmental conditions.
4.4.2 Models Trained on Hybrid Data

As was described in Section 2.3.2, models employing hybrid training data are explored in order to determine if small amounts of field data can improve the performance of laboratory ML models for predicting compressive strength of field concrete. In this analysis, $\alpha$ values of 10%, 20%, 30%, 40%, and 50% replacement percentages are selected via the quintile sampling method discussed in section 2.3.2. The remaining, unused field data is to determine the average testing performance of each hybrid model.

As was hypothesized, the inclusion of small percentages of field data significantly reduces the RMSE, MAE and increases the $R^2$ (compared to a pure laboratory model). As is shown in Figure 7, the most significant model improvements occur with the addition of the initial 10% of field data, which reduces the RMSE by 43.0%. However, continued performance improvements occur with the additional supplementation of field data driving the models. Furthermore, Figure 8 illustrates via predicted vs. observed scatter plots how the addition of field data improves predictive performance. A model comprised of 100% field data, which was analyzed in Section 4.3, is the standard with which the hybrid models are compared in terms of the extent to which predictive performance could be improved. This analysis illustrates that ML modeling of hybrid training data is a promising area of research that improves upon the downsides of field models and laboratory models being used in isolation.

Future research in this area may explore different ML methods (i.e., models other than random forest) or other hybridization strategies for utilizing hybrid training data. In addition, it may be of interest to focus this modeling procedure on concretes with exotic mixture ingredients, which inherently have been rarely employed in industry, and thus, have few data points with which to model compressive strength.
Figure 7. Graphs illustrating the continued improvement in (a) RMSE, (b) $R^2$, and (c) MAE as additional field data is supplied to the model.
Figure 7. Scatter plots of predictive versus observed for ML models trained on hybrid data with the following percentages of field data: (a) 10%, (b) 20%, (c) 30%, (d) 40%, (e) 50%. Points lying near the one-to-one line indicate better model performance.

5. Conclusions

The goal of this work was to specifically analyze the compressive strength behavior of *field concrete* as a function of mixture ingredient quantities in order to determine how this prediction problem is inherently different and more challenging than prediction of the compressive strength of *laboratory concrete*. Furthermore, this work trained and tested a variety of ML models for predicting compressive strength of field concrete mixtures and determined which ML models are best suited for the data.
By analyzing the performance measures and a variety of diagnostic plots, the reasons for differing performance for field concrete ML models have been elucidated. For instance, from the linear regression model diagnostics, it was found that there are only very minor violations of linearity assumptions; this result indicated it is likely that important predictor variables are missing from the data. Further manipulation of the predictor space via polynomial regression and kernel transformation indicated that a transformed predictor space can improve predictive capability (via a 4% reduction in testing RMSE). Moreover, it was found that nonlinear models, specifically random forest, generated the best performance measures, which is attributed to its full rejection of linear assumptions and ability to learn inconsistent variable importance in the data.

It was also confirmed that, at the current time, the most accurate prediction of compressive strength of field concrete is achieved with models trained on field concrete data; however, ML models that employ hybrid training data show promise for significantly improving predictive performance of laboratory concrete models even when only small amounts of field concrete data are available. For instance, it was found that when only 10% of the training dataset was field concrete data points, the RMSE was reduced by 43%. Moving forward, this research could be extended to explore other ML models with the hybridized approach or applications when it is desirable to explore modeling of exotic concrete mixtures and ingredients.

Broadly, the results of this research support two main conclusions: (1) Prediction of field concrete strength requires the application of nonlinear ML models using field-specific data. In particular, advanced tree-based models, such as random forest, are high-performing even when field data is relatively less abundant than laboratory data. (2) Although there is value in testing and statistical model training for the strength prediction of laboratory concrete, these models should not be used for stand-alone prediction of field concrete strength because they do not capture the many convoluting factors of field concrete placement and curing. However, ML models that employ hybrid training data can significantly improve the predictive performance compared to laboratory concrete ML models due to the small amounts of field concrete data that is supplied.

References


