mmpf: Monte-Carlo Methods for Prediction Functions

by Zachary M. Jones

Abstract Machine learning methods can often learn high-dimensional functions which generalize well but are not human interpretable. The mmpf package marginalizes prediction functions using Monte-Carlo methods, allowing users to investigate the behavior of these learned functions, as on a lower dimensional subset of input features: partial dependence and variations thereof. This makes machine learning methods more useful in situations where accurate prediction is not the only goal, such as in the social sciences where linear models are commonly used because of their interpretability.

Many methods for estimating prediction functions produce estimated functions which are not directly human-interpretable because of their complexity: for example, they may include high-dimensional interactions and/or complex nonlinearities. While a learning method’s capacity to automatically learn interactions and nonlinearities is attractive when the goal is prediction, there are many cases where users want good predictions and the ability to understand how predictions depend on the features. mmpf implements general methods for interpreting prediction functions using Monte-Carlo methods. These methods allow any function which generates predictions to be be interpreted. mmpf is currently used in other packages for machine learning like edarf and mlr (Jones and Linder, 2016; Bischl et al., 2016).

Marginalizing Prediction Functions

The core function of mmpf, marginalPrediction, allows marginalization of a prediction function so that it depends on a subset of the features. Say the matrix of features $X$ is partitioned into two subsets, $X_u$ and $X_{−u}$, where the former is of primary interest. A prediction function $f$ (which in the regression case maps $X \rightarrow y$) where $y$ is a real-valued vector might not be additive or linear in the columns of $X_u$, making $f$ difficult to interpret directly. To obtain the marginal relationship between $X_u$ and $f$ we could marginalize the joint distribution so that we obtain a function $f_u$ which only depends on the relevant subset of the features:

$$f_u(x_u) = \int f(x_u, x_{−u}) P(X|X_{−u}) P(X_{−u}) dX_{−u}.$$

This, however, can distort the relationship between $X_u$ and $f$ because of the inclusion of dependence between $X_u$ and $X_{−u}$ (specifically $X_u|X_{−u}$), which is unrelated to $f$. An alternative is to instead integrate against the marginal distribution of $X_{−u}$ as in 2.1, as suggested by (Friedman, 2001):

$$\hat{f}_u(x_u) = \int f(x_u, x_{−u}) P(X|X_{−u}) P(X_{−u}) dX_{−u}.$$

To illustrate this point, suppose data are generated from an additive model, $f() = x_1 + x_2$ and $(x_1, x_2) \sim MVN(0, \Sigma)$ where the diagonal entries of $\Sigma$ are 1 and the off-diagonals are 0.5. That is, $(x_1, x_2)$ are correlated by construction. Now if we want to understand how $f$ depends on $x_1$ we could integrate against the true joint distribution, as in 2.1. However, this distorts the relationship between $x_1$ and $f$, because the conditional probability of $x_1$ given $x_2$ is higher at values of $(x_1, x_2)$ which are more extreme (due to their correlation). Since $x_2$ is related to $f$, this has the effect of distorting the relationship between $x_1$ and $f$, and, in this case, makes the relationship appear more extreme than it is, as can be seen in the left panel of Figure 1. This distortion of the relationship between $x_1$ and $f$ can be made more misleading if $x_2$ interacts with $x_1$ to produce $f$, or if $x_2$ has a nonlinear relationship with $f$, as can be seen in the right panel of Figure 1.

Integrating against the marginal distribution of $x_1$ recovers the true additive effect (left panel) and the average marginal effect ($x_1 + 0.5x_1x_2$, in the right panel) respectively.

Using mmpf

In practical settings we do not know $P(X)$ or $f$. We can use $\hat{f}$, estimated from $(X, y)$ as a plug-in estimator for $f$ and can estimate $P(X_{−u})$ from the training data, allowing us to estimate the partial dependence of $X_u$ on $\hat{f}$ Friedman (2001).
Figure 1: The marginal relationship between $x_1$ and $f$ as estimated by integrating against the marginal distribution of $x_2$ (the blue line) or the joint distribution of $(x_1, x_2)$ (the red line). The true relationship is shown by the dashed line. In the left panel $f$ is an additive function of $x_1$ and $x_2$ and in the right panel $x_1$ and $x_2$ interact via multiplication to produce $f$.

\[ f_u(x_u) = \sum_{i=1}^{N} \hat{f}(X_u, X_{-u}) \]

This the behavior of the prediction function at a vector or matrix of values for $X_u$, averaged over the empirical marginal distribution of $X_{-u}$.

The function marginalPrediction allows users to compute easily partial dependence and many variations thereof. The key arguments of marginalPrediction are the prediction function (predict.fun), the training data (data), the names of the columns of the training data which are of interest (vars), the number of points to use in the grid for $X_u$, and the number of points to sample from $X_{-u}$ $(n$, an integer vector of length 2). Additional arguments control how the grid is constructed (e.g., uniform sampling, user chosen values, non-uniform sampling), indicate the use of weights, and instruct how aggregation is done (e.g., deviations from partial dependence). Below is an example using the Iris data (Anderson, 1936):

```r
library(mmpf)
library(randomForest)

data(iris)
iris.features = iris[, -ncol(iris)] # exclude the species column
fit = randomForest(iris.features, iris$Species)

mp = marginalPrediction(data = iris.features, vars = "Petal.Width", 
n = c(10, nrow(iris)), model = fit, uniform = TRUE, 
predict.fun = function(object, newdata) predict(object, newdata, type = "prob"))

print(mp)
##
## | Petal.Width | setosa | versicolor | virginica |
## |-------------|--------|------------|-----------|
## | 0.1080000   | 0.6074333 | 0.2337333 | 0.1288133 |
## | 0.3066667   | 0.6074333 | 0.2337333 | 0.1288133 |
## | 0.6033333   | 0.6082667 | 0.2350533 | 0.1293200 |
## | 0.9000000   | 0.1707200 | 0.5997333 | 0.2295467 |
## | 1.1666667   | 0.1688133 | 0.5880800 | 0.2431067 |
## | 1.3333333   | 0.1640400 | 0.5619867 | 0.2431067 |
## | 1.4333333   | 0.1619867 | 0.5619867 | 0.2431067 |
## | 1.7000000   | 0.1640400 | 0.4242800 | 0.4116800 |
## | 1.9666667   | 0.1619867 | 0.2666667 | 0.6313467 |
## | 2.2333333   | 0.1619867 | 0.2047867 | 0.6332267 |
## | 2.5000000   | 0.1619867 | 0.2047867 | 0.6332267 |
```

In this case, $\hat{f}$ returns a probability of membership in each class for each value of the variable.
Figure 2: The expected value of $\hat{f}$ estimated by a random forest and then marginalized by Monte-Carlo integration to depend only on “Petal.Width.”

“Petal.Width.” This is computed based on the average prediction for each value of “Petal.Width” shown and all the observed values of the other variables in the training data. As can be readily observed, partial dependence can be easily visualized, as in Figure 2.

In fact, any function of the marginalized function $\hat{f}_u$ can be computed, including vector-valued functions. For example the expectation and variance of $\hat{f}_u$ can be simultaneously computed, the results of which are shown in Figures 3 and 4. Computing the variance of $\hat{f}_u$ can be used for detecting interactions between $X_u$ and $X_{\cdot u}$ (Goldstein et al., 2015). If the variance of $\hat{f}_u(X_u)$ is constant then this indicates that $X_{\cdot u}$ does not interact with $X_u$, since, if it did, this would make $\hat{f}$ more variable in regions of the joint distribution wherein there is interaction between $X_u$ and $X_{\cdot u}$.

```r
mp.int = marginalPrediction(data = iris.features, 
vars = c("Petal.Width", "Petal.Length"),
n = c(10, nrow(iris)), model = fit, uniform = TRUE,
predict.fun = function(object, newdata) predict(object, newdata, type = "prob"),
aggregate.fun = function(x) list("mean" = mean(x), "variance" = var(x)))
head(mp.int)
## 1: Petal.Width Petal.Length setosa.mean setosa.variance versicolor.mean 
## 2: 0.1 1.655556 0.9549867 0.0011619193 0.04448000 
## 3: 0.1 2.311111 0.9530933 0.0011317899 0.04637333 
## 4: 0.1 2.966667 0.4574667 0.0003524653 0.52816667 
## 5: 0.1 3.622222 0.4550400 0.0002619447 0.52472000 
## 6: 0.1 4.277778 0.4550400 0.0002576688 0.52472000 
## versicolor.variance virginica.mean virginica.variance 
## 2: 0.001141889 0.0005333333 0.00000239821 
## 3: 0.001141889 0.0005333333 0.00000239821 
## 4: 0.001154918 0.0143466667 0.00054076492 
## 5: 0.001016158 0.0143466667 0.00054076492 
## 6: 0.001556364 0.0202400000 0.00093196886 
```
Figure 3: The expected value of $\hat{f}$ estimated by a random forest and then marginalized by Monte-Carlo integration to depend only on “Petal.Width” and “Petal.Length.”

Figure 4: The variance of $\hat{f}$ estimated by a random forest and marginalized by Monte-Carlo integration to depend only on “Petal.Width” and “Petal.Length.” Non-constant variance indicates interaction between these variables and those marginalized out of $f$. 
Bibliography


Zachary M. Jones
Pennsylvania State University
University Park, Pennsylvania
United States
zmj@zmjones.com

The R Journal Vol. 10/1, July 2018