# **Regression in terms of Trees and Forests**

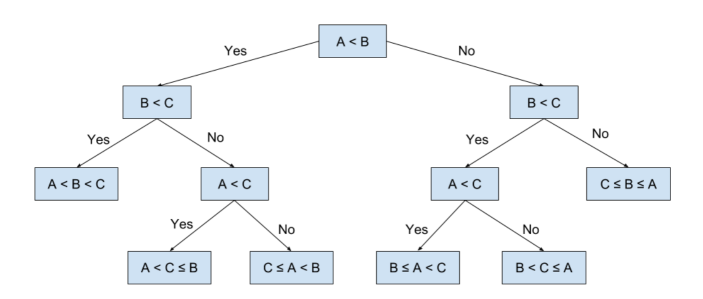
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Over the last few decades, the statistical learning/inference community has experienced a boom. With the advent of Al/ML, people have increasingly turned to sophisticated methods like Deep Learning and Reinforcement Learning. However, for a wide range of tasks of practical use, "classical" algorithms still fare sufficiently well. And what is more important -- the theory of these classical estimators is fairly well understood. We now know how they work, why they work, and how they can be modified effectively and deterministically for various use-cases. This makes their study not only relevant from a historical perspective, but also from a scientific perspective because they form that part of a data-scientists toolbox where he/she is aware of the underlying theory (as opposed to DL, which is still mostly a black-box), and consequently can wield them to considerable effect.

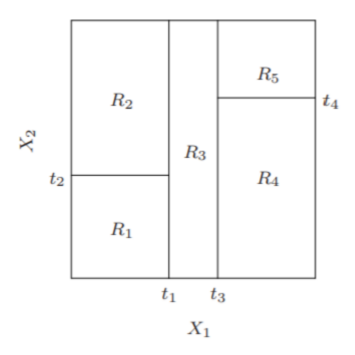
The most popular and well studied of these classical estimators/classifiers is perhaps Regression (and its many variants). In this article we delve deeper into a form of Regression that uses Decision Trees, an extremely powerful idea in Classical ML. Towards the end of this article we will also take a look at simple ensemble methods, like the Random Forest.

### **Decision Trees**

What are decision trees? Perhaps we are better off answering this with a picture:



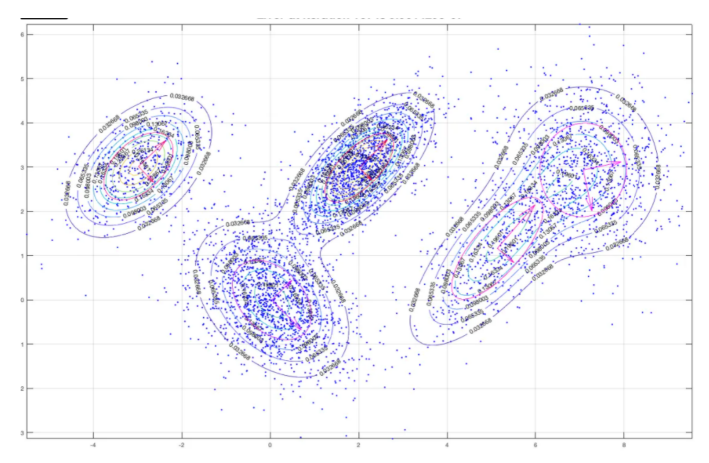
We think of A,B,C as attributes of a data-point. Each node represents a question (say A < B?, A+2B>9? etc.) which has TRUE/FALSE answer. The two edges out of the node represent these two answers. Each leaf will essentially represent a set of inequalities. This means that a leaf node corresponds to the region (in the domain) defined by these inequalities; like the following figure:



## Regression

The decision tree will split the dataset into many parts, by determining optimal attributes and corresponding splitting values. Given a new datapoint, we can now figure out to which leaf this point belongs and report the average of the decision values of all the datapoints belonging to this leaf.

Why are decision trees better than simple regression? Well consider the following data-points:



As we can see, there are different regions with different characteristics. If we used plain linear regression, it would attempt to treat these differing regions as the same, thereby causing a large error (on both test and training data), simply because it is not capable of capturing this feature of a dataset. Decision trees however

can divide the plane into regions and fit different structures on different parts, thereby giving a much better accuracy. This theoretical prediction is indeed also carried over in practice.

## **Constructing the Decision Tree**

The central idea lies in how we split a given node. We iterate over all the features and calculate the splitting value and impurity corresponding to that feature and **select the one providing the minimum impurity**, which is defined as the **weighted sum of the sample variances of the two child nodes**. This can be calculated by first sorting all the datapoints present in that node based on the corresponding feature values and then making a linear traversal.

Initially we would place all the points in the right child. At every step, we would move the current datapoint from the right node to the left node and make the required changes in the sample variances of the children nodes.

So how can we actually construct such a tree? The crux of the algorithm lies in coming up with the best splits at each node. The splits are identified with the sole purpose of minimizing the impurity at the nodes.

Now how to define this 'impurity' given a set of data points at a node?

For regression purposes variance of the data has proved to be a good measure of impurity. For our use case we find the split that minimizes the weighted sum of variance at each child node generated.

At each step we choose a leaf  $v_i$  and for each attribute  $d_i$  we calculate a constant  $c_i$  which is the split value that results in minimum impurity if the split is made on this attribute. This means that if a node is split at attribute  $d_i$  with the split value  $c_i$ , the node will be split into leaves one with values  $x_{d_i} \leq c_i$  and the other with  $x_{d_i} > c_i$ .

At this point we make a core observation: An optimal c for any attribute d can be chosen to be one of the  $(x_i)_d$ .

Why does this hold?

Suppose the values of  $(x_i)_k$  are sorted as  $z_1 \le z_2 \le z_3 \le \cdots \le z_N$  where N is the number of datapoints. For a split value c such that  $z_i \le c < z_{i+1}$  we have the following: all the values  $\le z_i$  belong to the left child and the rest to the right child. But this split remains same for any value of c in  $[z_i, z_{i+1})$ . So we can just take c to be  $z_i$  itself.

This makes life a bit simpler as for calculating the optimal split value for some attribute  $d_i$  we just need to iterate over all  $z_i$  and pick the value that results in the most variance reduction.

Now the only thing that remains is to choose an attribute to split on. This happens to be trivial as we just need to iterate over all attributes and choose the one that reduces the variance most. The optimal split for each attribute is calculated by following the steps explained above.

Okay, so far we have seen how we can grow our decision tree starting from a single root node by splitting the nodes at each iterative step. How long do we keep doing this?

Should we do this till we reach pure splits and cannot split further? Then we would get 100% accuracy on the training data but will most definitely end up just learning all the training data and our model will not generalize well to unseen data.

It looks like we need some stopping criterion to prevent our model from overfitting.

At this point let us introduce the coefficient of variance:

$$ext{CoefVar} = rac{ ext{Standard Deviation}}{ ext{Mean}} = rac{\sqrt{rac{1}{n}\sum_{j}(y_{j} - ar{y})^{2}}}{ar{y}}$$

which is a measure of the variability of data relative to the mean.

At every node we calculate this coefficient and if it falls below a certain threshold, i.e. the variability of data has reduced enough, the node is not split any more.

## **Code Walkthrough**

We begin with a single node, and iterate over all possible attributes and threshold values (Function *build* beginning from *line 140 below*) and pick the split leading to the minimum impurity. Function \_getimpurity (*line 65*) computes the impurity for a given attribute. Function \_get\_decision\_value\_for *feature* (*line 23*) computes the threshold splitting value correspoding to a feature. Once these are evaluated, \_divide *data* (*line 98*) is

# **Pruning**

Pruning is yet another technique to prevent overfitting of decision trees. It reduces the size of decision trees by removing parts of the tree that do not provide enough power (as opposed to the time/overfitting cost) to classify instances. Decision trees are the most susceptible out of all the machine learning algorithms to overfitting and effective pruning can reduce this possibility. As the name implies, pruning involves cutting back the tree. After a tree has been built (and in the absence of early stopping discussed below) it may be overfitted. The recusive algorithm will then repeatedly partition data into smaller and smaller subsets until those final subsets are homogeneous in terms of the outcome variable. In practice this often means that the final subsets (known as the leaves of the tree) each consist of only one or a few data points. The tree has learned the data exactly, but a new data point that differs very slightly might not be predicted well. This is why pruning is an important technique, in both theory and practice.

## **Code Walkthrough**

The code for pruning is implemented below in function prune\_decision\_tree (line 161 - 183 below)

- Consider an internal (non-leaf) node v, and let  $S_v$  be the sample variance of  $y_i$  of all datapoints contained in this node v. (line 170 172)
- Let  $S_v^\prime$  be the sum of the sample variances of  $y_i$  of all leaves which are descendants of v. (line 174)
- Clearly  $S_v^\prime < S_v$ , because splitting "improves" the sample variance.
- But, to penalize large structures, we look at  $P_v = S_v S_v' \lambda |T_v|$ , where  $|T_v|$  is the size of the subtree of v, and  $\lambda$  is a **pruning factor**. Larger  $\lambda$  results in more pruning. (*line 176*)
- If  $P_v < 0$ , we decide that this subtree must be pruned. Intuitively, we prune when pay more because of the size of the subtree than we gain because of variance reduction. (line 178 183)

```
import numpy as np
import sklearn.metrics as metrics
class TreeNode:
   def __init__(self):
        self.predicted value = None
        self.decision feature = None
        self.decision value = None
        self.left node = None
        self.right node = None
class DecisionTree:
   def init (self):
        self.root node = None
        self.min_coef_of_var = None
        self.reductions = None
   def fit(self, X, Y, min coef of var = 0, pruning factor = 0):
        self.min_coef_of_var = min_coef_of_var
        self.root node = self.build(X, Y)
        self.prune decision tree(X, Y, self.root node, pruning factor)
        self.reductions = np.zeros(len(X[0]))
        self.reductions = self.calculate reductions(X, Y, self.root node, self.r
eductions)
   def get decision value for feature(self, X, Y, decision feature):
        Z = np.empty((0, 2), float)
        for i in range(len(Y)):
            Z = np.append(Z, [[X[i, decision feature], Y[i]]], axis = 0)
        Z = np.sort(Z, axis = 0)
        left cardinality = 0
        right cardinality = 0
        left_sum = 0
        right sum = 0
        left sum of squares = 0
        right sum of squares = 0
        for i in range(len(Z)):
            right cardinality += 1
            right_sum += Z[i, 1]
            right sum of squares += Z[i, 1] ** 2
        current best_impurity = right_sum_of_squares
        current best decision value = Z[0, 0] - 1
        for i in range(len(Z)):
            left cardinality += 1
            left sum += Z[i, 1]
            left sum of squares += Z[i, 1] ** 2
            right_cardinality -= 1
            right_sum -= Z[i, 1]
            right sum of squares -= Z[i, 1] ** 2
```

```
impurity = left_sum_of_squares - left_sum * left_sum / left_cardinal
ity
             if (right cardinality != 0):
                 impurity += right_sum_of_squares - right_sum * right_sum / right
cardinality
             if (impurity < current best impurity):</pre>
                 current best impurity = impurity
                 current best decision value = Z[i, 0]
        return current best decision value
    def get impurity(self, X, Y, decision feature, decision value for feature):
        left cardinality = 0
        right cardinality = 0
        left sum = 0
        right sum = 0
        for i in range(len(Y)):
             if (X[i, decision_feature] <= decision_value_for_feature):</pre>
                 left cardinality += 1
                 left sum += Y[i]
             else:
                 right cardinality += 1
                 right sum += Y[i]
        if (left cardinality != 0):
             left mean = left_sum / left_cardinality
        else:
             left mean = 0
        if (right cardinality != 0):
             right mean = right sum / right cardinality
        else:
             right mean = 0
        impurity = 0
        for i in range(len(Y)):
             if (X[i, decision feature] <= decision value for feature):</pre>
                 impurity += (Y[i] - left mean) ** 2
             else:
                 impurity += (Y[i] - right mean) ** 2
        return impurity
    def divide_data(self, X, Y, decision_feature, decision_value):
        number of features = len(X[0])
        X_left = np.empty((0, number_of_features), float)
        X_right = np.empty((0, number_of_features), float)
        Y_{left} = np.empty(0, float)
        Y right = np.empty(0, float)
        for i in range(len(X)):
             if (X[i, decision_feature] <= decision_value):</pre>
                 X_{\text{left}} = \text{np.append}(X_{\text{left}}, [X[i]], axis = 0)
                 Y_{left} = np.append(Y_{left}, [Y[i]], axis = 0)
             else:
                 X \text{ right} = \text{np.append}(X \text{ right}, [X[i]], axis = 0)
                 Y_right = np.append(Y_right, [Y[i]], axis = 0)
        return X_left, Y_left, X_right, Y_right
```

```
def build(self, X, Y):
        if (len(X) == 0):
            return
        root node = TreeNode()
        node\ mean = np.mean(Y)
        root_node.predicted_value = node_mean
        node\ deviation = np.std(Y)
        # Do not split
        if (node deviation / node mean < self.min coef of var):</pre>
            root_node.decision value = 0
            root node.decision feature = 0
            return
        if (np.amin(Y) == np.amax(Y)):
            root node.decision value = 0
            root node.decision feature = 0
            return root node
        number of features = len(X[0])
        current best feature = -1
        current best decision value = -1
        current best impurity = -1
        for i in range(number of features):
            decision value for feature = self.get decision value for feature(X,
Y, i)
            impurity of decision value = self.get impurity(X, Y, i, decision val)
ue for feature)
            if (current best feature == -1 or impurity of decision value < curre</pre>
nt best impurity):
                current best feature = i
                current best decision value = decision value for feature
                current_best_impurity = impurity_of_decision_value
        root node.decision feature = current best feature
        root node.decision value = current best decision value
        X_left, Y_left, X_right, Y_right = self.divide_data(X, Y, root_node.deci
sion_feature, root_node.decision_value)
        if (len(Y left) == 0 or len(Y right) == 0):
            return root node
        root_node.left_node = self.build(X_left, Y_left)
        root_node.right_node = self.build(X_right, Y_right)
        return root_node
    def prune decision tree(self, X, Y, root node, pruning factor):
        if (root node == None):
          return 0
        X_left, Y_left, X_right, Y_right = self.divide_data(X, Y, root_node.deci
sion feature, root node.decision value)
        left tree size = self.prune decision tree(X left, Y left, root node.left
node, pruning factor)
        right tree_size = self.prune_decision_tree(X_right, Y_right, root_node.r
ight_node, pruning_factor)
```

```
tree_size = 1 + left_tree_size + right_tree_size
        impurity = 0
        for y in Y:
            impurity += (y - root node.predicted value) ** 2
        divided impurity = self.get impurity(X, Y, root node.decision feature, r
oot node.decision value)
        pruning value = impurity / len(Y) - divided impurity / len(Y) - pruning
factor * tree_size
        if (pruning value < 0):</pre>
            root node.left node = None
            root node.right node = None
            return 1
        else:
            return tree size
    def calculate reductions(self, X, Y, root node, reductions):
        if (root node == None):
            return reductions
        impurity = 0
        for y in Y:
            impurity += (y - root_node.predicted value) ** 2
        for i in range(len(X[0])):
            decision_value_for_feature = self.get_decision value for feature(X,
Y, i)
            impurity of decision value = self.get impurity(X, Y, i, decision val
ue_for_feature)
            reductions[i] += impurity - impurity of decision value
        X left, Y left, X right, Y right = self.divide data(X, Y, root node.deci
sion feature, root node.decision value)
        reductions = self.calculate reductions(X left, Y left, root node.left no
de, reductions)
        reductions = self.calculate reductions(X right, Y right, root node.right
node, reductions)
        return reductions
    def get reductions(self):
        return self.reductions
    def predict_single(self, X):
        current node = self.root node
        while (True):
            if (X[current node.decision feature] <= current node.decision value</pre>
):
                if (current node.left node != None):
                    current_node = current_node.left_node
                else:
                    return current_node.predicted_value
            else:
                if (current node.right node != None):
                    current_node = current_node.right_node
                else:
                    return current_node.predicted_value
```

```
def predict(self, X):
    Y = np.empty(0, float)
    for x in X:
        Y = np.append(Y, [self.predict_single(x)], axis = 0)
    return Y

decision_tree = DecisionTree()
```

We will use Boston housing data from sklearn.datasets for testing our algorithm.

### In [15]:

```
from sklearn.datasets import load_boston
data = load_boston(return_X_y=False)
X = data.data
y = data.target
from sklearn.model_selection import train_test_split
X_train, X_test, y_train, y_test = train_test_split(X, y, test_size = 0.2, rando
m_state = 1)
```

We fit our decision tree to the training data, and check our performance on the test data.

#### In [16]:

```
decision_tree.fit(X_train, y_train, min_coef_of_var = 0.025, pruning_factor = 0)
y_test_pred = decision_tree.predict(X_test)
metrics.r2_score(y_test, y_test_pred)
```

#### Out[16]:

#### 0.7114290465682265

We see that the score of our predictions after training the data on training set is (~71%).

#### In [17]:

```
decision_tree.get_reductions()
Out[17]:
```

```
array([13399.20701222, 5186.7937058, 6077.93066069, 3654.6684283
7, 15268.52069709, 20767.48739927, 10509.61121374, 9736.4801611
7, 5424.56393685, 6267.04097296, 9541.52648792, 3903.7219425
1, 21012.15264691])
```

#### In [18]:

```
decision_tree.fit(X_train, y_train, min_coef_of_var = 0.025, pruning_factor = 0.
4)
y_test_pred = decision_tree.predict(X_test)
metrics.r2_score(y_test, y_test_pred)
```

#### Out[18]:

#### 0.7428687746808185

The score is significantly improved (from  $\sim$ 71% to  $\sim$ 74%) if we use pruning (with pruning factor = 0.4). Next we check how scikit-learn fares on the same test:train split.

#### In [ ]:

```
from sklearn.tree import DecisionTreeRegressor
dt = DecisionTreeRegressor()
dt.fit(X_train, y_train)
```

#### Out[]:

#### In [ ]:

```
y_test_pred_dt = dt.predict(X_test)
metrics.r2_score(y_test, y_test_pred_dt)
```

#### Out[]:

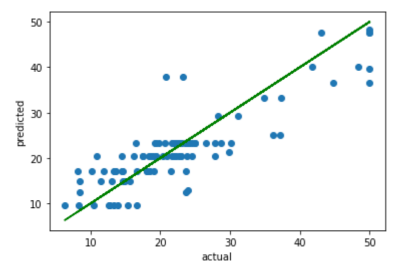
#### 0.8055224772577356

As we see, scikit-learn's implementation gives a score of (~80%). Their implementation is more advanced than our implementation which uses nothing but the absolute basics of decision trees.

Now, we investigate how our algorithm performed using a graphical approach.

## In [30]:

```
import matplotlib.pyplot as plt
plt.plot(y_test, y_test, color = "green")
plt.xlabel("actual")
plt.ylabel("predicted")
plt.scatter(y_test, y_test_pred)
plt.show()
```

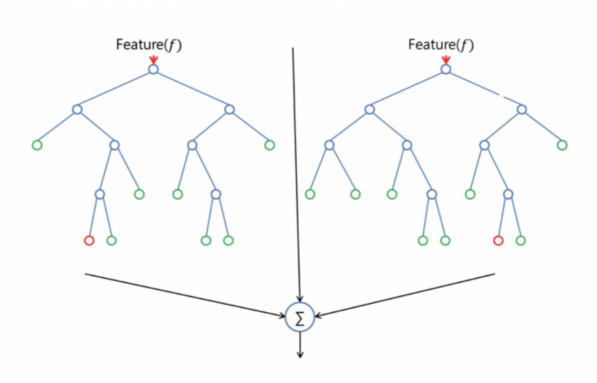


We plot the predicted value of price of house on the y-axis and actual value on the x-axis. Ideally all the points should lie on the y = x line. We observe that most of the points lie close to the y = x line. We also observe that there are some horizonal lines in the plot implying same predicted value for a range of actual values. This is because we implemented a basic decision tree. We will see later that these horizontal lines disappear when we use more advanced techniques like the random-forest.

## **Random Forests**

Now we take a look at simple ensemble methods using decision trees. The most famous of these is the classical Random Forest, using Bagging / Random Subspace projection methods.

But, what is a *random forest*? A random forest is simply a collection of trees trained on random subsets of the data (either resampled data points and/or subset of attributes). Here is a picture of a 2-tree random forest:



Another question is, how do we actually *use* this structure to generate predictions. In the simplest form, this prediction is simply done by averaging the answers from the trees. Note that we use averaging here because the problem we are solving is **regression** as opposed to **classification**.

Of course, this is not the only method that can be used. Other methods involve boosting trees, which train trees on the residual from other trees, but we do not delve into this matter here.

#### Construction

Now let us take a look at how we construct these forests. The code for the algorithm described here is included below.

Firstly, note that we take the algorithm for Decision Trees as a black-box (in our code, we use the Decision Trees above). Given a training dataset, we form a fixed (hyperparameter) number of trees (*lines 21-30 below*). For each tree, we choose a resampled (with replacement) subset of the datapoints (*line 23*), and a subset of the attributes (*line 24*). The size of these subsets are controlled by the hyperparameters. Then we train the tree on this data (*line 28*).

The first technique of using resampled data is very similar to the bootstrap method, and can also be used for constructing <u>confidence intervals</u> for the answers. It is also referred to as **bagging** *I* **bootstrap-aggregating** in the ensemble learning literature.

The second technique of choosing a random subset of the attributes is part of a generic technique for dealing with high-dimensional data, called **random subspace projection**. The name is so because choosing a particular set of attributes/coordinates is equivalent to projecting the high-dimensional datapoints to smaller subspaces.

**Note:** In our use-case we do not use random-subspace projection, because we have only a few attributes (~15), in which the projection method doesn't help much.

### **Prediction**

The prediction is implemented in lines *34-43*. It is a simple algorithm in our case: just average the answers from all the trees.

```
from numpy.random import choice, seed
class RandomForest:
    trees = []
    importance = None
    def init (self):
        self.trees = []
    def fit(self, X, y, num trees = 1, frac samples = None, frac attrs = None):
        seed(0)
        N, total attributes = X.shape
        self.importance = np.zeros(total attributes)
        if frac samples == None:
            frac samples = 1.0
        if frac attrs == None:
            frac attrs = 1.0
        for i in range(num trees):
            new tree = DecisionTree()
            sample = choice(range(N), size = int(N * frac samples))
            attrs = choice(range(total attributes), size = int(total attributes
* frac attrs))
            #print(attrs, end=", ")
            dataset = X[sample, :] #[:, attrs]
            ys = y[sample]
            new tree.fit(dataset, ys, min coef of var = 0.025, pruning factor =
0)
            self.importance += new tree.get reductions()
            self.trees.append((new_tree, attrs))
        self.importance /= num_trees
    def predict(self, X):
        N, _ = X.shape
        answer = np.zeros(N)
        for (t, attrs) in self.trees:
            answer += t.predict(X) # [:, attrs])
        return answer / len(self.trees)
```

Now, we train the random forest on our training data, using a 0.8 fraction of (resampled) data-points. However, we choose not to use to the projection method, for reasons described above.

```
In [27]:
```

```
rf = RandomForest()
rf.fit(X_train, y_train, num_trees = 30, frac_samples = 0.8, frac_attrs = 1.0)
```

And then we test it on our test-data.

#### In [28]:

```
y_test_rf = rf.predict(X_test)
y_test_rf
metrics.r2_score(y_test, y_test_rf)
```

#### Out[28]:

#### 0.8545010166573305

We see a really good improvement from the raw Decision Tree score (~72% to ~85%). Let us also compare it to the scikit-learn implementation (which is more advanced than ours):

#### In [ ]:

```
from sklearn.ensemble import RandomForestRegressor
rfsk = RandomForestRegressor(bootstrap=True)
rfsk.fit(X_train, y_train)
```

#### Out[]:

#### In [ ]:

```
y_test_rfsk = rfsk.predict(X_test)
metrics.r2_score(y_test, y_test_rfsk)
```

#### Out[]:

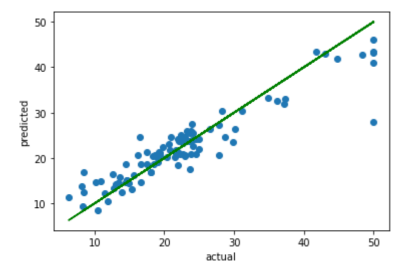
#### 0.9099455853887437

We see that scikit-learn manages an accuracy of ~91%. Our 85% is not that worse, given that our implementation is very bare-bones, and mostly proof-of-concept. This shows the power of the very idea of random forests (and in general, ensemble learning).

Finally, we plot our predictions as opposed to the ground truth (x = y).

#### In [31]:

```
plt.plot(y_test, y_test, color = "green")
plt.xlabel("actual")
plt.ylabel("predicted")
plt.scatter(y_test, y_test_rf)
plt.show()
```



We see that a large chunk of the predicted points is really close to the ground truths. Also, we observe no horizontal segments like in the case for Decision Trees. This is because of the averaging step in the Random Forests, and contributes immensely to the accuracy.

# Variable Importance

Now we take a look at the importance of variables. For this purpose, we compute the following quantity for each tree T in the ensemble:

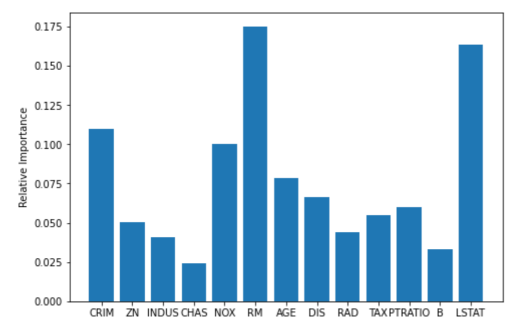
 $\mathrm{VarReduction}_i = \sum_{x \in T} \mathrm{Optimal\ Variance\ Reduction\ in\ node\ } x$  by attribute i and then denote

 $Importance_i = Average of VarReduction_i$ 

This quantity will be the importance of attribute i. We plot these quantities in a relative sense.

## In [34]:

```
importance = rf.importance
relative_importance = importance / np.sum(importance)
fig = plt.figure()
ax = fig.add_axes([0,0,1,1])
ax.bar(data.feature_names, relative_importance)
plt.ylabel('Relative Importance')
plt.show()
```



The following are the meanings of the attributes:

```
CRIM - per capita crime rate by town
ZN - proportion of residential land zoned for lots over 25,000 sq.ft.
INDUS - proportion of non-retail business acres per town.
CHAS - Charles River dummy variable (1 if tract bounds river; 0 otherwis e)
NOX - nitric oxides concentration (parts per 10 million)
RM - average number of rooms per dwelling
AGE - proportion of owner-occupied units built prior to 1940
DIS - weighted distances to five Boston employment centres
RAD - index of accessibility to radial highways
TAX - full-value property-tax rate per $10,000
PTRATIO - pupil-teacher ratio by town
B - 1000(Bk - 0.63)^2 where Bk is the proportion of blacks by town
LSTAT - % lower status of the population
MEDV - Median value of owner-occupied homes in $1000's
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So we can see that the most important attribute is RM, which is the average number of rooms. This makes sense, because the number of rooms is an important factor. The 2nd most important factor is LSTAT, which is the percentage of the population which is of lower status. This can be interpreted to mean that the price is lower in underdeveloped regions. And the 3rd most important factor is crime rate (CRIM), which too makes a lot of sense. On the other end, the CHAS variable is about the Charles River. It doesn't make too much of an impact on the price.