

# Modeling Lake Trophic State: A Data Mining Approach

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## Abstract

Productivity of lentic ecosystems has been well studied and it is widely accepted that as nutrient inputs increase, productivity increases and lakes transition from low trophic state (e.g. oligotrophic) to higher trophic states (e.g. eutrophic). These broad trophic state classifications are good predictors of ecosystem health and ecosystem services/disservices (e.g. recreation, aesthetics, fisheries, and harmful algal blooms). While the relationship between nutrients and trophic state provides reliable predictions, it requires *in situ* water quality data in order to parameterize the model. This limits the application of these models to lakes with existing and, more importantly, available water quality data. To expand our ability to predict in lakes without water quality data, we take advantage of the availability of a large national lakes water quality database, land use/land cover data, lake morphometry data, other universally available data, and modern data mining approaches to build and assess models of lake trophic state that may be more universally applied. We use random forests and random forest variable selection to identify variables to be used for predicting trophic state and we compare the performance of two models of trophic state (as determined by chlorophyll *a* concentration). The first model estimates trophic state with *in situ* as well as universally available data and the second model uses universally available data only. For each of these models we used three separate trophic state categories, for a total of six models. Overall accuracy for the *in situ* and universal data models ranged from xx% to xx% and xx, xx, and xx described the most variation in trophic state. For the universal data only models, Overall accuracy ranged from xx% to xx% and xx, xx, and xx described the most variation in trophic state. Lastly, it is believed that the presence and abundance of cyanobacteria is strongly associated with trophic state. To test this we examine the association between estimates of cyanobacteria biovolume and the measured and predicted trophic state. Expanding these preliminary results to include cyanobacteria taxa indicates that cyanobacteria are significantly more likely to be found in highly eutrophic lakes. These results suggest that predictive models of lake trophic state may be improved with additional information on the landscape surrounding lakes and that those models provide additional information on the presence of potentially harmful cyanobacteria taxa.

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## 1. Introduction

Productivity in lentic systems is often categorized across a range of trophic states (e.g. the trophic continuum) from early successional (i.e. oligotrophic) to late successional lakes (i.e. hypereutrophic) (Carlson 1977). Lakes naturally occur across the range of trophic state and higher primary productivity is not necessarily a predictor of poor ecological condition. Lakes that are naturally oligotrophic occur in nutrient poor areas or have a more recent geologic history. These lakes are often found in higher elevations, have clear water, and are often favored for drinking water or direct contact recreation (e.g. swimming). Lakes with higher productivity (e.g. eutrophic lakes) have greater nutrient loads, tend to be less clear, have greater density of aquatic plants, and often support more diverse and abundant fish communities. Lakes will naturally shift to higher trophic states but this is a slow process. Given this fact, monitoring trophic state allows the identification of rapid shifts in trophic state or locating lakes with unusually high productivity (e.g. hypereutrophic). These cases are indicative of lakes under greater anthropogenic nutrient loads, also known as cultural eutrophication, and are more likely to be at risk of fish kills, fouling, and harmful algal blooms (Smith 1998, Smith et al. 1999, 2006). Given the association between trophic state and many ecosystem services and disservices, being able to model trophic state could allow for estimating trophic state in unmonitored lakes and provide a first cut at identifying lakes with the potential for harmful algal blooms and other problems associated with cultural eutrophication.

Cyanobacteria are an important taxonomic group associated with harmful algal blooms in lakes. Understanding the drivers of cyanobacteria presence has important implications for lake management and for the protection of human and ecosystem health. Chlorophyll *a* concentration, a measure of the biological productivity of a lake, is one such driver and is largely, although not exclusively, determined by nutrient inputs. As nutrient inputs increase, productivity increases and lakes transition from low trophic state (e.g. oligotrophic) to higher trophic states (e.g. hypereutrophic). These broad trophic state classifications are associated with ecosystem health and ecosystem services/disservices (e.g. recreation, aesthetics, fisheries, and harmful algal blooms). Thus, models of trophic state might be used to predict things like cyanobacteria.

We have three goals for this preliminary research. First, we build and assess multiple models of lake trophic state using a full suite of data including *in situ* water quality and universally available data

(e.g. landscape data). Second, we assess the accuracy of predicted trophic state in lakes with only the universally available data. Lastly, we explore associations between trophic state and cyanobacteria to explore.

## 2. Methods

### 2.1. Data and Study Area

We utilize four primary sources of data for this study, the National Lakes Assessment (NLA), the National Lake Cover Dataset (NLCD), modeled lake morphometry, and estimated cyanobacteria biovolumes (Homer et al. 2004, USEPA 2009, Xian et al. 2009, Hollister and Milstead 2010, Hollister et al. 2011, Beaulieu et al. 2013, Hollister 2014). All datasets are national in scale and provide a unique snapshot view of the condition of lakes in the United States’.

The NLA data were collected during the summer of 2007 and the final data were released in 2009. With consistent methods and metrics collected at 1056 locations across the conterminous United States (Figure 1), the NLA provides a unique opportunity to examine broad scale patterns in lake productivity. The NLA collected data on biophysical measures of lake water quality and habitat. For this analysis we primarily examined the water quality measurements from the NLA (USEPA 2009). Adding to the monitoring data collected via the NLA, we use the 2006 NLCD data to examine the possible landscape-level drivers of trophic status in lakes. The NLCD is a nationally collected land use land cover dataset that also provides estimates of impervious surface. We collected total land use land cover and total percent impervious surface within a 3 kilometer buffer surrounding the lake to examine larger landscape-level effect (Homer et al. 2004, Xian et al. 2009). We also used various measures of lake morphometry (i.e. depth, volume, fetch, etc.) as they are important in understanding lake productivity, yet many of these data are difficult to obtain for large numbers of lakes over broad regions. To add this information we modeled lake morphometry (Hollister and Milstead 2010, ???, Hollister et al. 2011, Hollister 2014). Lastly, to explore associations between trophic state and cyanobacteria, we used estimates of cyanobacterial biovolume calculated by Beaulieu *et al.* (2013). Cyanobacteria biovolumes are a truer measure of cyanobacteria dominance than abundance as there is great variability

in the size within and between species. We have consolidated the taxa level estimates from Beaulieu *et al.* (2013) and summed that information on a per-lake basis.

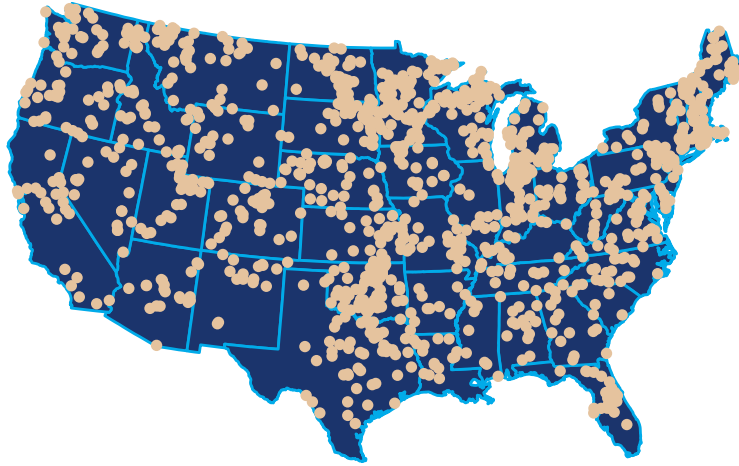


Figure 1: Map of the distribution of National Lakes Assessment Sampling locations

## 2.2. Predicting Trophic State with Random Forests

Random forest is a machine learning algorithm that aggregates numerous decision trees in order to obtain a consensus prediction of the response categories (Breiman 2001). Bootstrapped sample data is recursively partitioned according to a given random subset of predictor variables and completely grown without pruning. With each new tree, both the sample data and predictor variable subset is randomly selected.

While random forests are able to handle numerous correlated variables without a decrease in prediction accuracy, unusually large numbers of related variables can reduce accuracy and increase the chances of over-fitting the model. This is a problem often faced in gene selection and in that field, a variable selection method based on random forest has been successfully applied (Díaz-Uriarte and De Andres 2006). We use `varselRF` in R to initially examine the importance of the water quality and GIS derived variables and select a subset, the reduced model, to then pass to random forest (Díaz-Uriarte 2010).

Using R's `randomForest` package, we pass the reduced models selected with `varSelRF` and calculate confusion matrices, overall accuracy and kappa coefficient (Liaw and Wiener 2002). From the reduced model random forests we collect a consensus prediction and calculate a confusion matrix and summary stats.

### 2.3. Model Details

Using a combination of the `varSelRF` and `randomForest` we ran models for six combinations of variables and trophic state classifications. These combinations included different combinations of the Chlorophyll *a* trophic states (Table 1) along with all variables and the GIS only variables (i.e. no *in situ* information). The six model combinations were:

1. Chlorophyll *a* trophic state - 4 class = All variables (*in situ* water quality, lake morphometry, and landscape)
2. Chlorophyll *a* trophic state - 3 class = All variables (*in situ* water quality, lake morphometry, and landscape)
3. Chlorophyll *a* trophic state - 2 class = All variables (*in situ* water quality, lake morphometry, and landscape)
4. Chlorophyll *a* trophic state - 4 class = All variables (lake morphometry, and landscape)
5. Chlorophyll *a* trophic state - 3 class = All variables (lake morphometry, and landscape)
6. Chlorophyll *a* trophic state - 2 class = All variables (lake morphometry, and landscape)

Trophic State (4)	Trophic State (3)	Trophic State (2)	Cut-off
oligo	oligo	oligo/meso	$\leq 0.2$
meso	meso/eu	oligo/meso	$>2-7$
eu	meso/eu	eu/hyper	$>7-30$
hyper	hyper	eu/hyper	$>30$

Table 1: Chlorophyll a based trophic state cut-offs

### 3. Results

#### 3.1. Model 1: 4 Trophic States ~ All Variables

The selected variables that made up Model 1 were Potassium, Nitrogen:Phosphorus, Total Nitrogen, Total Phosphorus, Total Organic Carbon, Turbidity, Ecoregion, Organic Ions, Dissolved Organic Carbon, and Maximum Lake Depth (Table 2). Total accuracy for Model 1 is 0.667% and the Cohen’s Kappa is 0.546 (Table 3).

Variable	Percent
K	1.00
NPratio	1.00
NTL	1.00
PTL	1.00
TOC	1.00
TURB	1.00
WSA_ECO9	1.00
ORGION	0.29
DOC	0.18
DEPTHMAX	0.03

Table 2: Variable selection results for Model 1

Oligo	Meso	Eu	Hyper	class.error
135	58	4	1	0.32
42	235	76	9	0.35
2	70	217	47	0.35

Oligo	Meso	Eu	Hyper	class.error
0	3	68	175	0.29

Table 3: Random Forest confusion matrix for Model 1

Lastly, turbidity, total phosphorus, total nitrogen, and total organic carbon were the most important predictors of the 4 classes of trophic state (Figure 2).

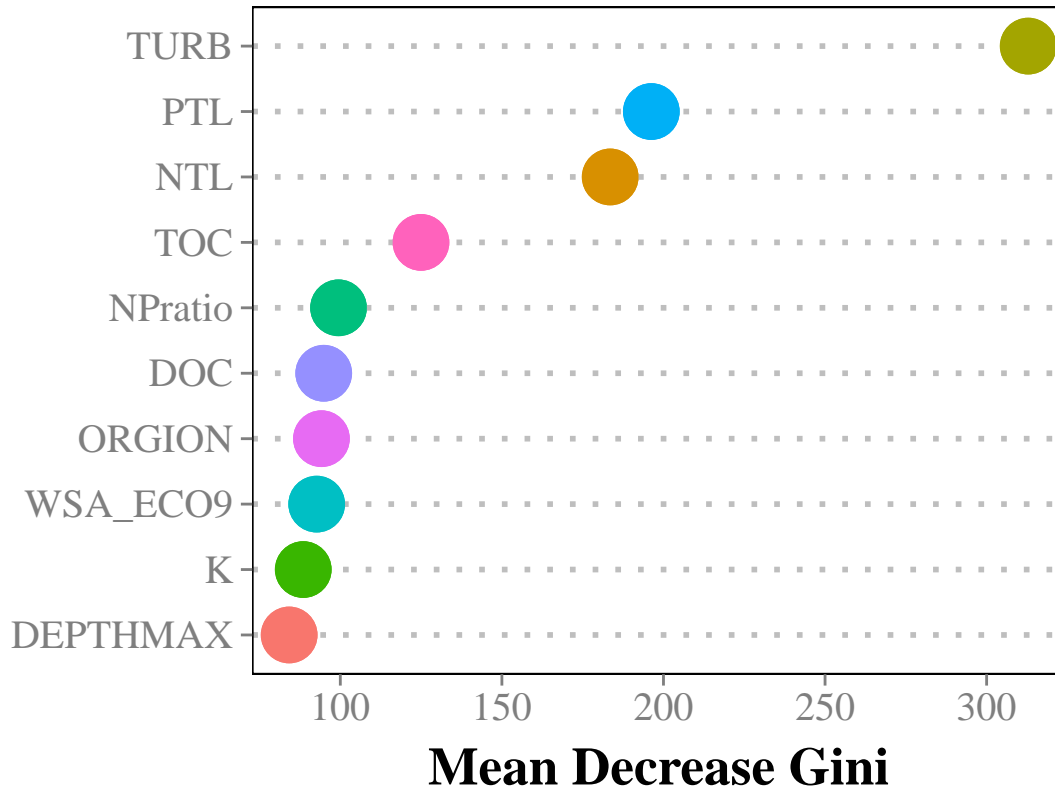


Figure 2: Importance plot for Model 1

### 3.2. Model 2: 3 Trophic States ~ All Variables

Total accuracy for Model 2 is 0.799% and the Cohen's Kappa is 0.618.

Variable	Percent
DOC	1.00

Variable	Percent
K	1.00
NTL	1.00
ORGION	1.00
PTL	1.00
TOC	1.00
TURB	1.00
WSA_ECO9	1.00
DEPTHMAX	0.98
NPratio	0.76
AlbersX	0.48
CropsPer_3000m	0.27
ELEV_PT	0.16
AlbersY	0.05
NH4	0.05
PH_FIELD	0.01
EvergreenPer_3000m	0.01

Table 4: Variable selection results for Model 2

Oligo	Meso/Eu	Hyper	class.error
121	75	0	0.38
40	609	40	0.12
0	72	173	0.29



Oligo	Meso/Eu	Hyper	class.error
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Table 5: Random Forest confusion matrix for Model 2

### 3.3. Model 3: 2 Trophic States ~ All Variables

Total accuracy for Model 3 is 0.87% and the Cohen's Kappa is 0.741.

Variable	Percent
K	1.00
NPratio	1.00
NTL	1.00
PTL	1.00
TOC	1.00
TURB	1.00
WSA_ECO9	1.00
ORGION	0.99
DEPTHMAX	0.96
DDs45	0.90
ELEV_PT	0.85
DOC	0.58
AlbersX	0.06
AlbersY	0.03
Na	0.03

Variable	Percent
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Table 6: Variable selection results for Model 3

Oligo/Meso	Eu/Hyper	class.error
489	71	0.13
77	505	0.13

Table 7: Random Forest confusion matrix for Model 3

#### 3.4. Model 4: 4 Trophic States ~ GIS Only Variables

Total accuracy for Model 4 is 0.482% and the Cohen's Kappa is 0.292.

Variable	Percent
AlbersX	1.00
CropsPer_3000m	1.00
EvergreenPer_3000m	1.00
MeanDepthCorrect	1.00
WSA_ECO9	1.00
AlbersY	0.35
ELEV_PT	0.02

Table 8: Variable selection results for Model 4

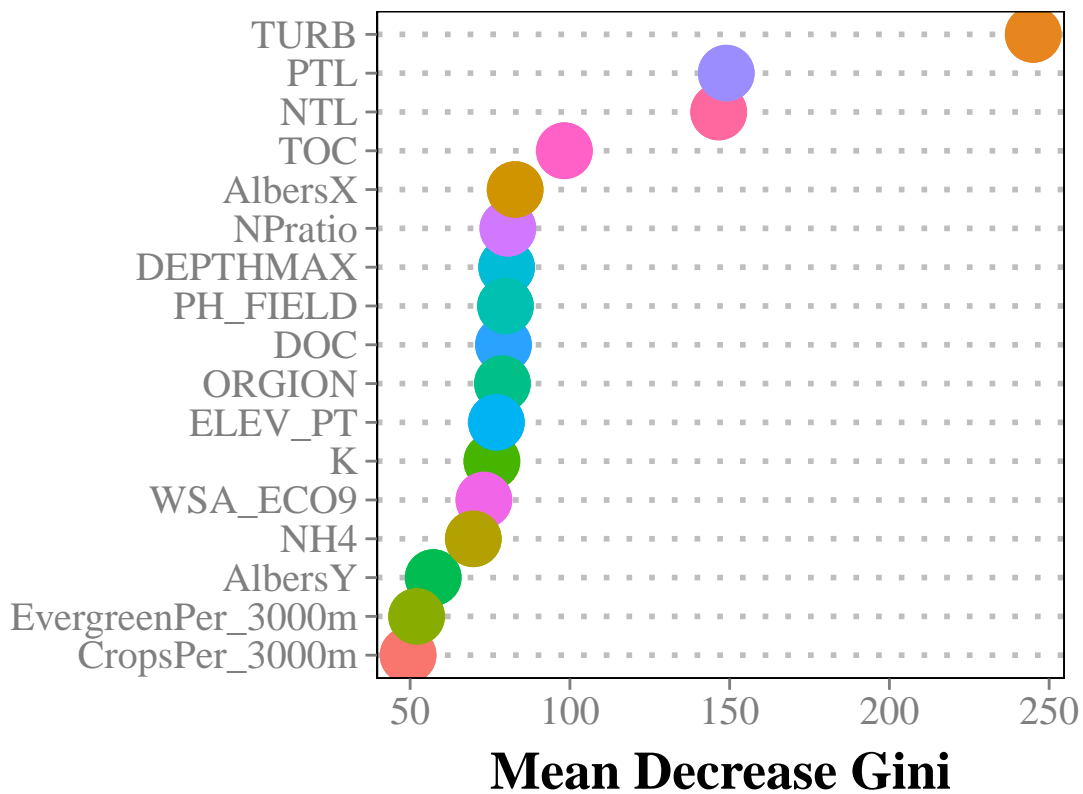


Figure 3: Importance plot for Model 2

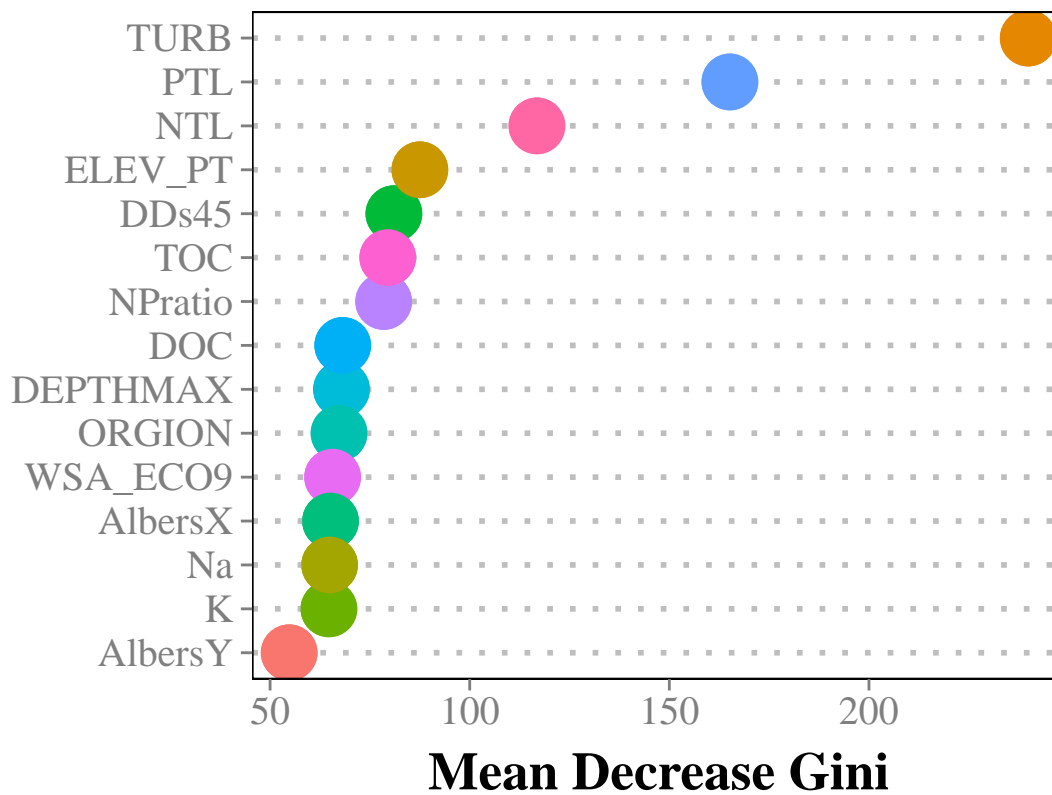


Figure 4: Importance plot for Model 3

Oligo	Meso	Eu	Hyper	class.error
95	73	27	2	0.52
48	201	80	32	0.44
20	114	124	77	0.63
2	36	79	129	0.48

Table 9: Random Forest confusion matrix for Model 4

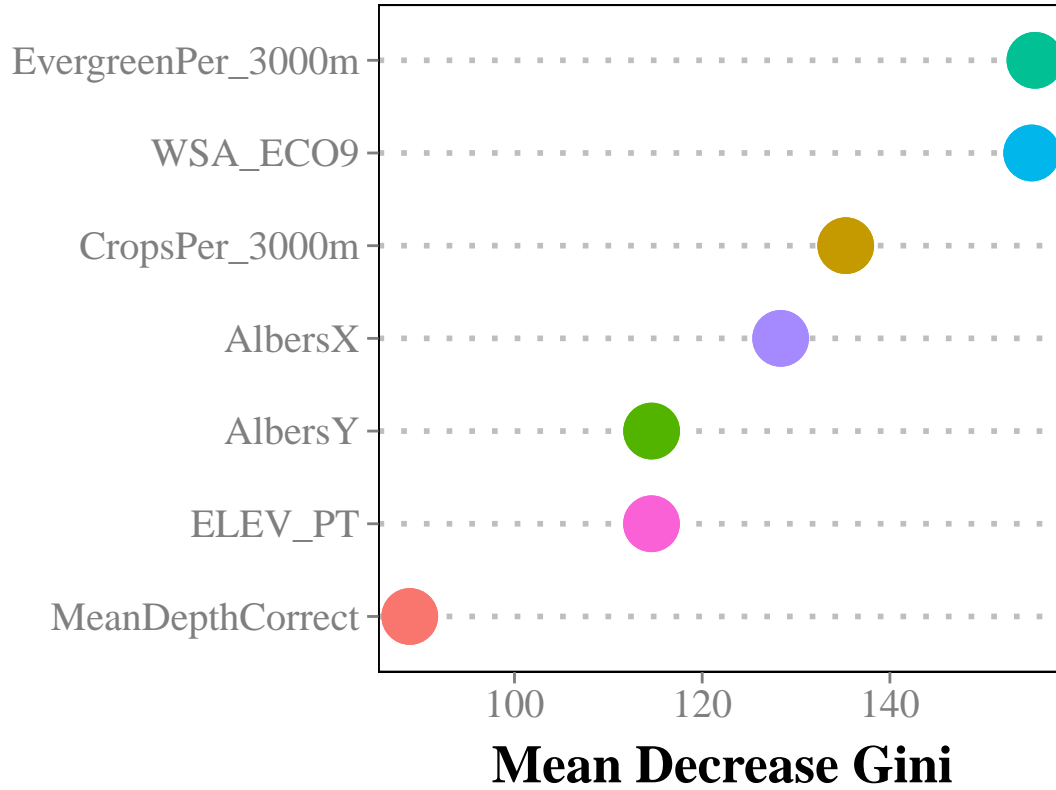


Figure 5: plot of chunk Importance\_Model4

:Importance plot for Model 4

### 3.5. Model 5: 3 Trophic States ~ GIS Only Variables

Total accuracy for Model 5 is 0.673% and the Cohen's Kappa is 0.343.

Variable	Percent
AlbersX	1.00
AlbersY	1.00
CropsPer_3000m	1.00
EvergreenPer_3000m	1.00
MaxDepthCorrect	1.00
MeanDepthCorrect	1.00
WSA_ECO9	1.00
ELEV_PT	0.97
DeciduousPer_3000m	0.94
ShrubPer_3000m	0.21
WoodyWetPer_3000m	0.11
DevOpenPer_3000m	0.10
VolumeCorrect	0.04

Table 10: Variable selection results for Model 5

Oligo	Meso/Eu	Hyper	class.error
79	116	1	0.6
48	582	66	0.16
0	141	105	0.57

Table 11: Random Forest confusion matrix for Model 5

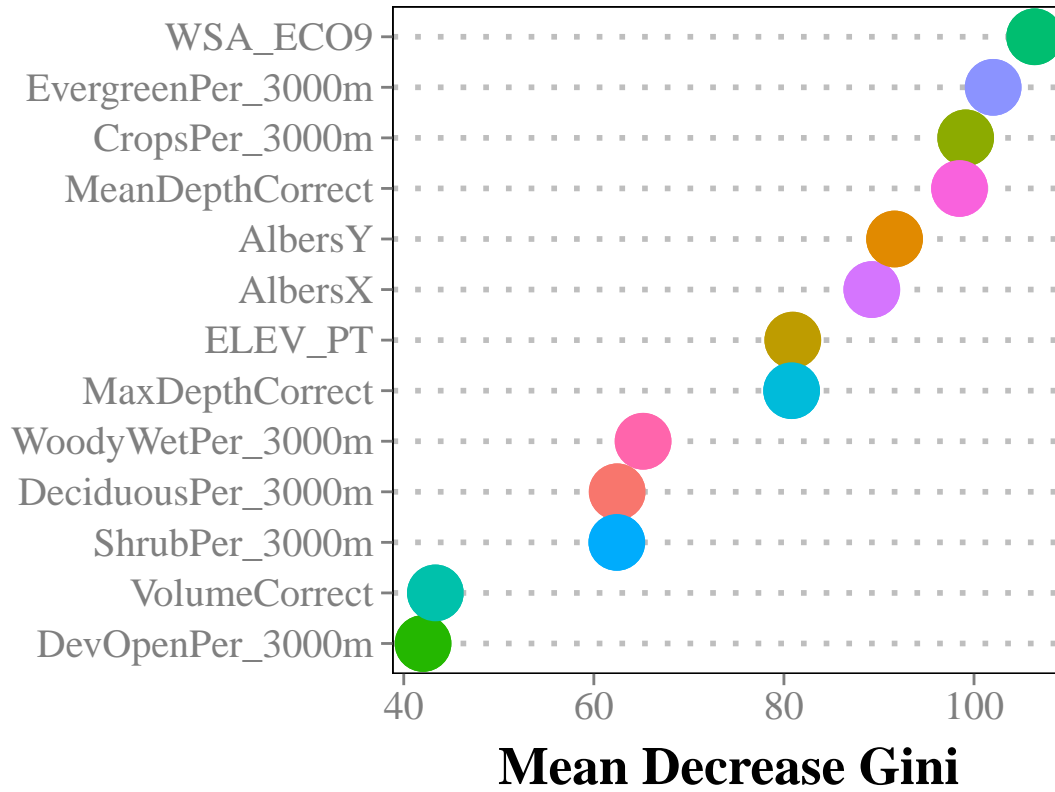


Figure 6: Importance plot for Model 5

### 3.6. Model 6: 2 Trophic States ~ GIS Only Variables

Total accuracy for Model 6 0.758% and the Cohen's Kappa is 0.517.

Variable	Percent
AlbersX	1.00
CropsPer_3000m	1.00
DDs45	1.00
ELEV_PT	1.00
EvergreenPer_3000m	1.00
MeanDepthCorrect	1.00
WSA_ECO9	1.00
AlbersY	0.98

Variable	Percent
MaxDepthCorrect	0.98
DeciduousPer_3000m	0.92
DevOpenPer_3000m	0.67
BASINAREA	0.31
PercentImperv_3000m	0.01

Table 12: Variable selection results for Model 6

Oligo/Meso	Eu/Hyper	class.error
428	129	0.23
146	435	0.25

Table 13: Random forest confusion matrix for Model 6

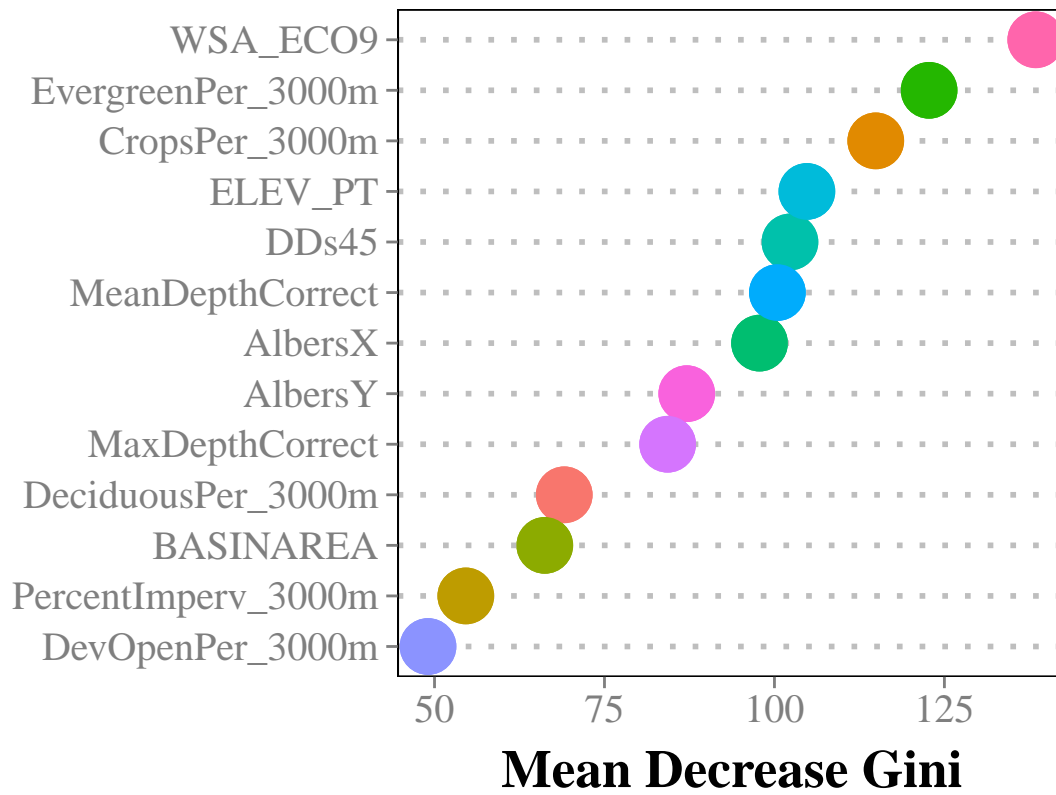
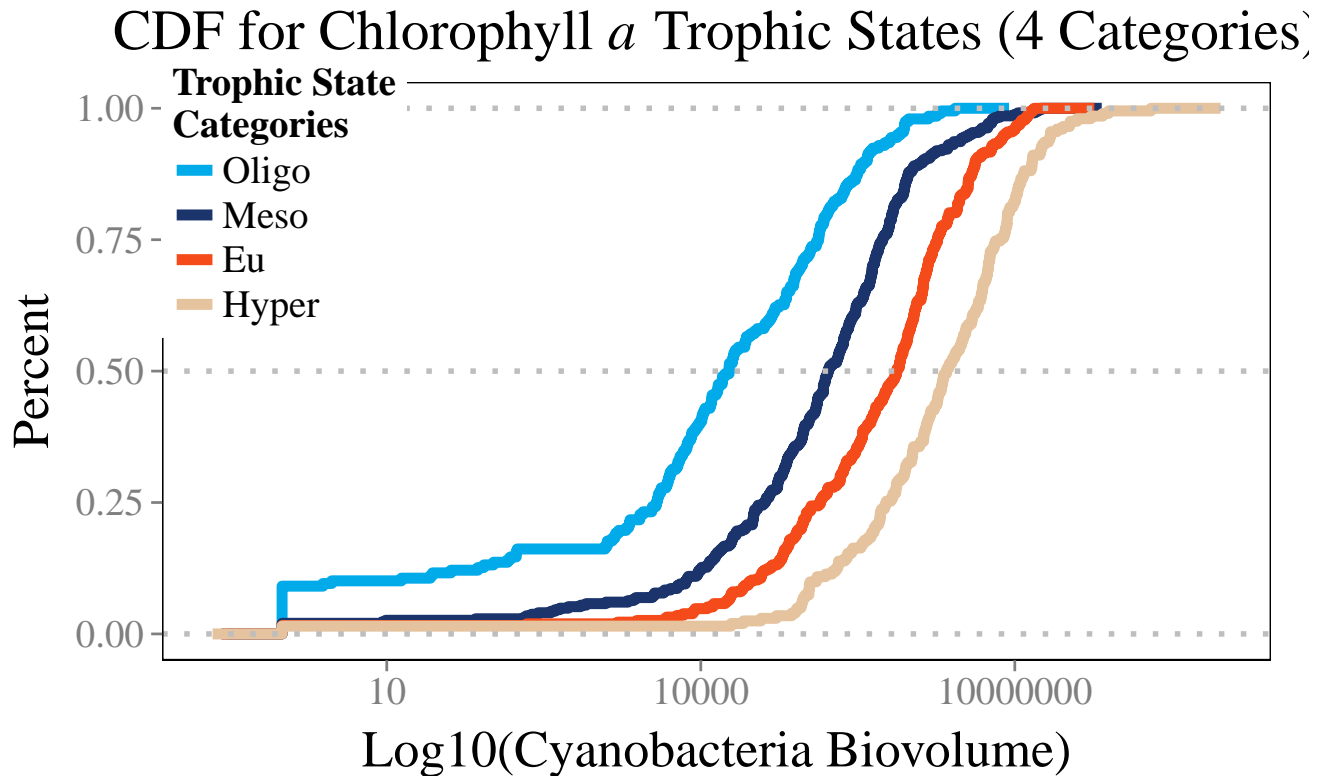
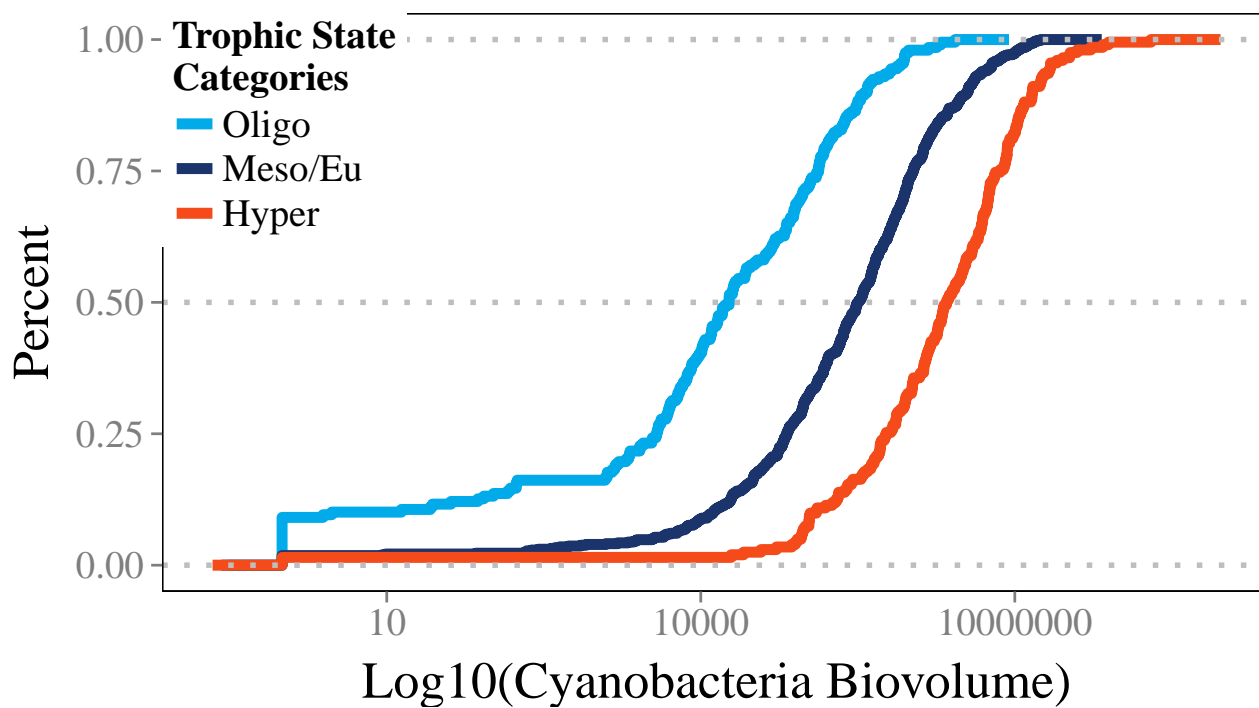


Figure 7: Importance plot for Model 6

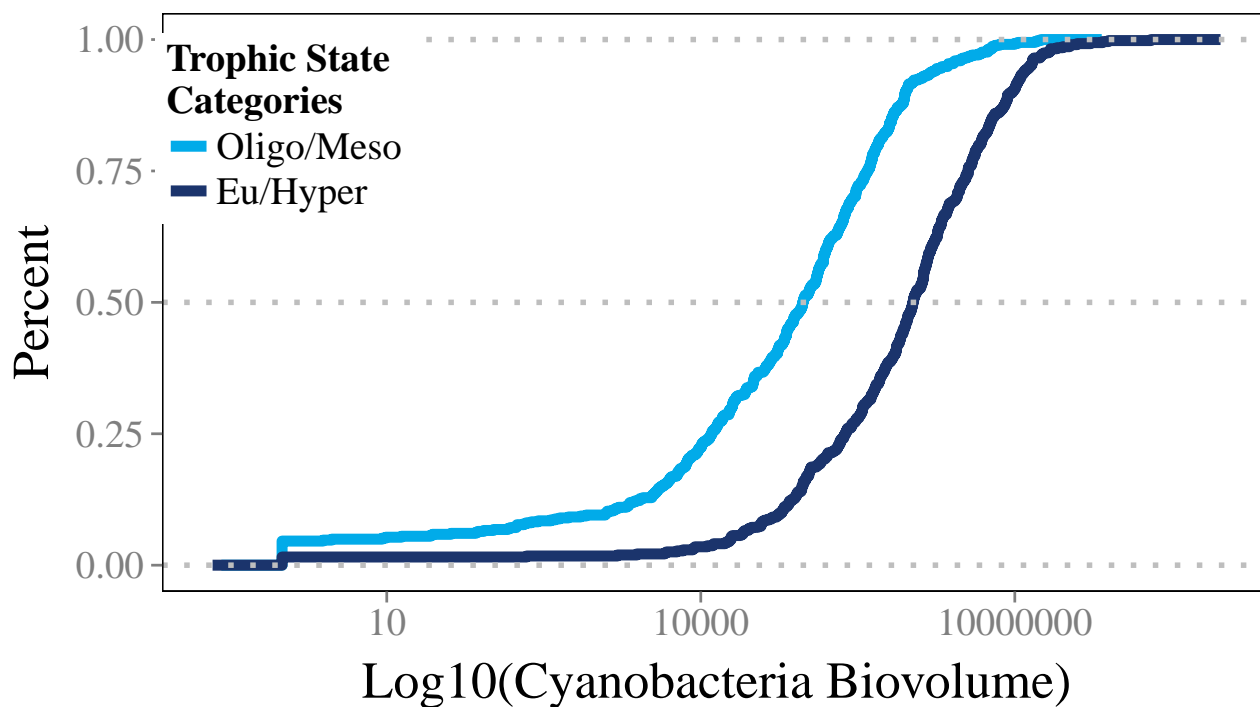




### CDF for Chlorophyll *a* Trophic States (3 Categories)

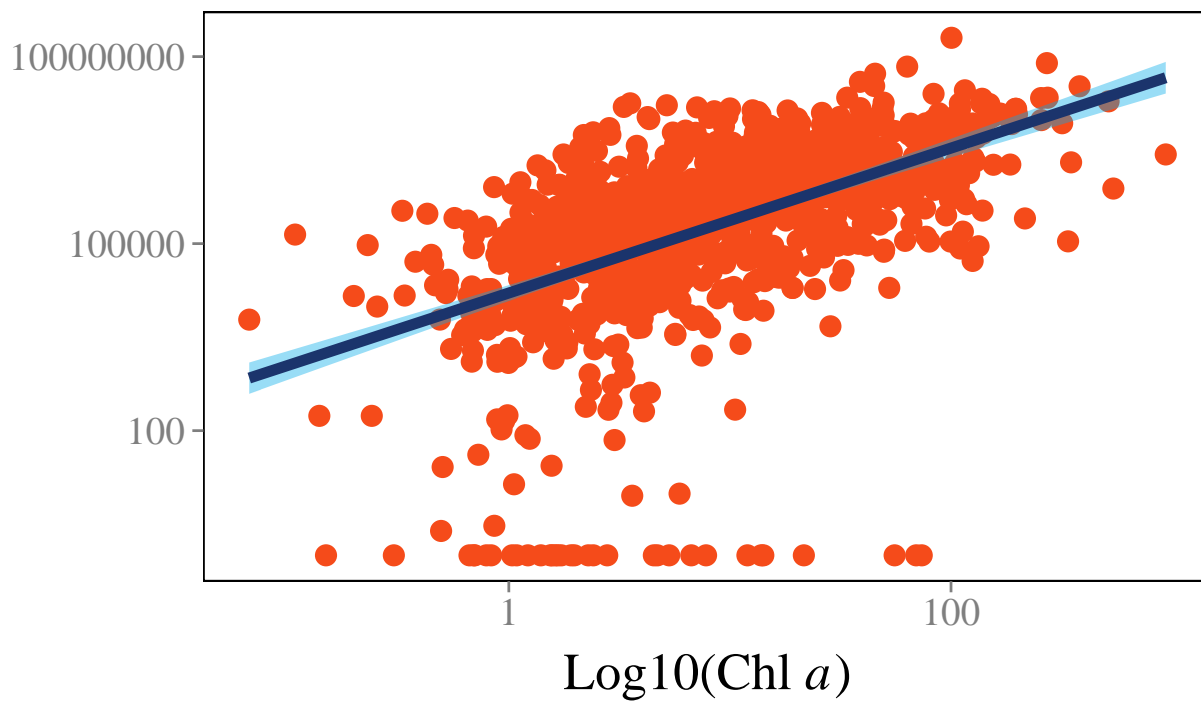


## CDF for Chlorophyll *a* Trophic States (2 Categories)



$\log_{10}(\text{Cyanobacteria Biovolume} + 1)$

## Chlorophyll *a* and Cyanobacteria Relationship



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