# Field Chlorophyll Analysis of Old and Young Leaves

Kenia E. Segura Aba

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This tutorial uses R version 4.0.0, RRPP version 0.5.2, tidyverse version 1.3.0, and kableExtra version 1.1.0.

1. load necessary packages

```
library(RRPP)
library(tidyverse)
library(kableExtra)
```

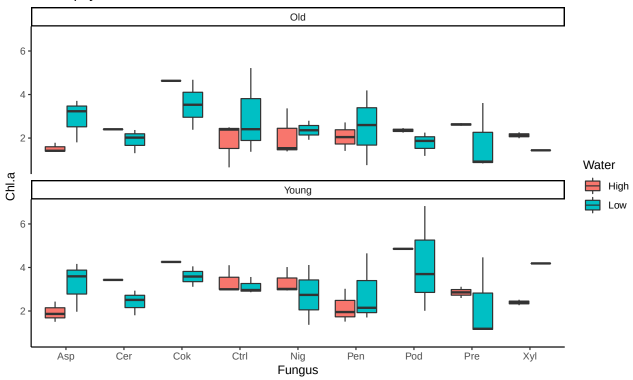
2. Upload data

```
path <- "~/Box/Summer 2018 TX Endo Field Samples and Analysis/Statistics/Old + Young Chlorophyll/"
chldata <- read.csv(paste(path, "chlorophyll_data.csv", sep = ""),
    header = T, row.names = 1)
# create a total chlorophyll column (a + b)
chldata$Total <- chldata$Chl.a + chldata$Chl.b</pre>
```

3. Calculate total chlorophyll content, summary statistics, and visualize data.

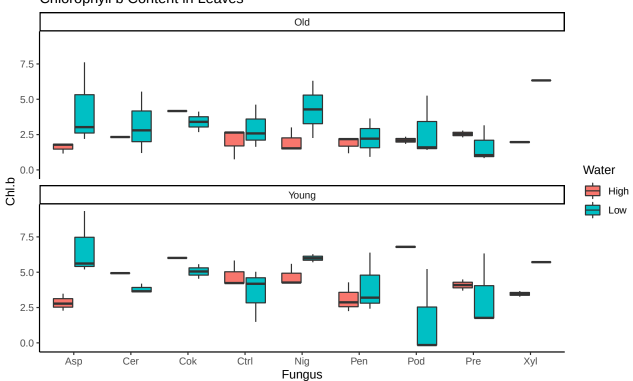
```
# plot chl a
chla <- chldata %>% ggplot(aes(x = Fungus, y = Chl.a, fill = Water)) +
    geom_boxplot() + theme_classic() + facet_wrap(~Age, dir = "v")
print(chla + labs(title = "Chlorophyll a Content in Leaves"))
```

### Chlorophyll a Content in Leaves



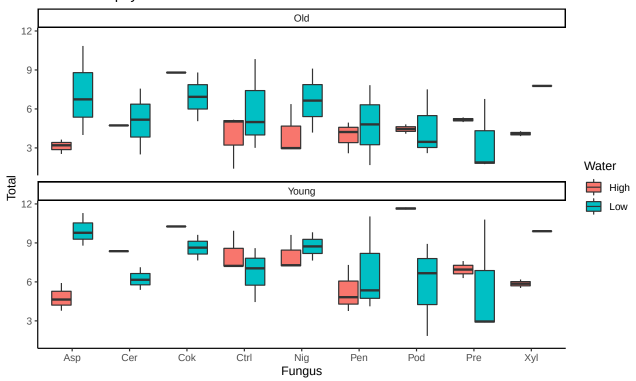
```
# plot chl b
chlb <- chldata %>% ggplot(aes(x = Fungus, y = Chl.b, fill = Water)) +
    geom_boxplot() + theme_classic() + facet_wrap(~Age, dir = "v")
print(chlb + labs(title = "Chlorophyll b Content in Leaves"))
```

#### Chlorophyll b Content in Leaves



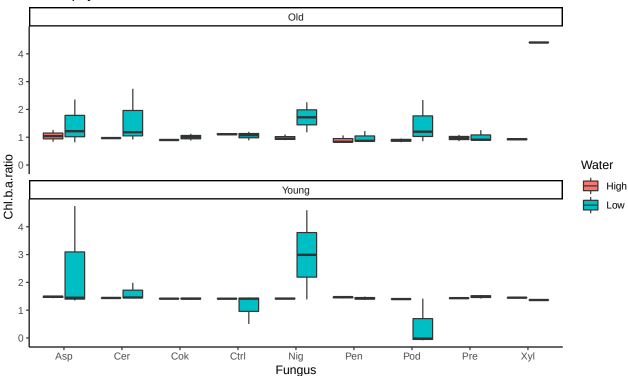
```
# plot total chlorophyll
tot <- chldata %>% ggplot(aes(x = Fungus, y = Total, fill = Water)) +
    geom_boxplot() + theme_classic() + facet_wrap(~Age, dir = "v")
print(tot + labs(title = "Total Chlorophyll Content in Leaves"))
```

#### Total Chlorophyll Content in Leaves



```
# plot chl b/a ratio
chlb.a <- chldata %>% ggplot(aes(x = Fungus, y = Chl.b.a.ratio,
    fill = Water)) + geom_boxplot() + theme_classic() + facet_wrap(~Age,
    dir = "v")
print(chlb.a + labs(title = "Chlorophyll b/a Ratio"))
```

#### Chlorophyll b/a Ratio



-		Chl a						
Water	Fungus	Min	Q1	Median	Q3	Max	Mean	SD
	Asp	1.3785	1.438375	1.64465	1.845525	2.4325	1.730400	0.3969141
	Cer	2.4034	2.659700	2.91600	3.172300	3.4286	2.916000	0.7249259
	Cok	4.2542	4.348175	4.44215	4.536125	4.6301	4.442150	0.2658014
	Ctrl	0.6543	2.416075	2.74235	2.996475	4.1046	2.605350	1.1327896
High	Nig	1.3885	1.886175	2.97785	3.276550	4.0201	2.710450	1.0410501
	Pen	1.4103	1.623000	1.99855	2.549825	3.0217	2.110083	0.6443822
	Pod	2.2508	2.407100	3.64395	4.842250	4.8829	3.605400	1.4465113
	Pre	2.5646	2.589350	2.64125	2.792350	3.1147	2.740450	0.2546096
	Xyl	1.9925	2.196050	2.26565	2.330000	2.5178	2.260400	0.2145433
	Asp	1.8032	2.282550	3.41230	3.680150	4.1644	3.077933	0.9724927
Low	Cer	1.3056	1.858450	2.19620	2.475375	2.9348	2.157850	0.5733045
	Cok	2.3816	2.930825	3.58330	4.209275	4.6790	3.556800	1.0136576
	Ctrl	1.3706	2.521075	2.91315	3.413525	5.2171	3.064117	1.2833496
	Nig	1.3661	1.783475	2.35900	3.125225	4.1147	2.549700	1.1977419
	Pen	0.7559	1.814325	2.37305	3.791275	4.6496	2.673950	1.4901283
	Pod	1.1815	1.904725	2.13075	3.335300	6.8223	2.971917	2.0599131
	Pre	0.8341	0.970475	1.15580	3.003875	4.4658	2.023350	1.5892062
	Xyl	1.4374	2.124750	2.81210	3.499450	4.1868	2.812100	1.9441194

		Chl b						
Water	Fungus	Min	Q1	Median	Q3	Max	Mean	SD
	Asp	1.1578	1.802800	2.06590	2.651275	3.4807	2.221967	0.8188023
	Cer	2.3279	2.979350	3.63080	4.282250	4.9337	3.630800	1.8425789
	Cok	4.1675	4.628375	5.08925	5.550125	6.0110	5.089250	1.3035514
	Ctrl	0.7493	2.652775	3.44605	4.228750	5.8297	3.391517	1.7540252
High	Nig	1.4517	1.897100	3.63340	4.264575	5.5941	3.350983	1.6580124
	Pen	1.1784	2.192475	2.23820	2.712925	4.2843	2.497917	1.0301252
	Pod	1.8431	2.224775	4.54235	6.762600	6.8523	4.445025	2.7190193
	Pre	2.3051	2.662850	3.23910	3.894375	4.4892	3.318125	0.9708800
	Xyl	1.9172	1.994675	2.64285	3.366050	3.6686	2.717875	0.8814486
	Asp	2.1927	3.570600	5.40600	7.116375	9.3309	5.496817	2.6947445
Low	Cer	1.1950	2.996500	3.61605	4.056800	5.5423	3.493850	1.4477185
	Cok	2.6757	3.764025	4.33190	4.794500	5.5670	4.226625	1.1983699
	Ctrl	1.4843	1.872325	3.38255	4.510275	5.0331	3.256267	1.5551574
	Nig	2.2579	4.842175	5.99005	6.284650	6.3091	5.136775	1.9392895
	Pen	0.9236	2.263875	2.80870	3.530525	6.3876	3.130333	1.8490598
	Pod	-0.1698	0.237575	1.50615	4.320025	5.2567	2.195317	2.4760652
	Pre	0.8355	1.220800	1.75545	2.810150	6.3290	2.479950	2.0534522
	Xyl	5.7141	5.869250	6.02440	6.179550	6.3347	6.024400	0.4388305

		Chl b/a						
Water	Fungus	Min	Q1	Median	Q3	Max	Mean	SD
High	Asp	0.8399000	1.0947000	1.345747	1.473605	1.513249	1.261981	0.2721681
	Cer	0.9686000	1.0861968	1.203794	1.321390	1.438987	1.203794	0.3326139
	Cok	0.9001000	1.0283148	1.156530	1.284744	1.412959	1.156530	0.3626463
	Ctrl	1.0730000	1.1162250	1.277402	1.411244	1.420294	1.261076	0.1690156
	Nig	0.8948000	0.9843500	1.245312	1.409754	1.446971	1.199054	0.2498186
	Pen	0.8204000	0.8932000	1.242076	1.456214	1.484446	1.182250	0.3142204
	Pod	0.8189000	0.9220250	1.175364	1.396577	1.403323	1.143238	0.3004410
	Pre	0.8585000	1.0282250	1.253842	1.427493	1.441321	1.201876	0.2815388
	Xyl	0.8925000	0.9447750	1.201136	1.444321	1.457067	1.187960	0.3023483
Low	Asp	0.8165000	1.2490922	1.397398	2.129357	4.745607	1.988317	1.4430095
	Cer	0.9153000	1.2432041	1.441539	1.852052	2.742500	1.617810	0.6550962
	Cok	0.8820000	1.0631250	1.248568	1.394481	1.457018	1.209038	0.2600366
	Ctrl	0.5008402	0.9324500	1.133150	1.357552	1.460632	1.087613	0.3580767
	Nig	1.1744000	1.3332090	1.821573	2.841354	4.594418	2.352991	1.5660008
	Pen	0.8517000	0.9573000	1.297849	1.408012	1.489756	1.204278	0.2805299
	Pod	-0.0844468	0.1964067	1.026300	1.360309	2.335800	0.949229	0.9190565
	Pre	0.8746000	0.9963500	1.335999	1.474218	1.545791	1.249302	0.2936034
	Xyl	1.3647763	2.1253072	2.885838	3.646369	4.406900	2.885838	2.1511063

		Total Chlorophyll							
Water	Fungus	Min	Q1	Median	Q3	Max	Mean	SD	
	Asp	2.5363	3.311200	3.71055	4.426775	5.9132	3.952367	1.184572	
	Cer	4.7313	5.639050	6.54680	7.454550	8.3623	6.546800	2.567505	
	Cok	8.7976	9.164500	9.53140	9.898300	10.2652	9.531400	1.037750	
	Ctrl	1.4036	5.068850	6.18840	7.225225	9.9343	5.996867	2.870463	
High	Nig	2.9146	3.833050	6.78635	7.260450	9.6142	6.061433	2.640505	
	Pen	2.5887	3.875550	4.52275	4.916675	7.3060	4.608000	1.573892	
	Pod	4.0939	4.631875	8.18630	11.604850	11.7352	8.050425	4.165380	
	Pre	4.9900	5.257525	5.82020	6.621250	7.6039	6.058575	1.167894	
	Xyl	3.9097	4.190725	4.90850	5.696050	6.1864	4.978275	1.063039	
	Asp	3.9959	7.250750	9.28470	10.581175	11.2971	8.574750	2.771967	
	Cer	2.5006	5.226200	5.77320	6.885400	7.5632	5.651700	1.806590	
Low	Cok	5.0573	7.002500	8.22835	9.009275	9.6197	7.783425	1.988842	
	Ctrl	3.0058	4.583650	6.01750	8.208475	9.8370	6.320383	2.620370	
	Nig	4.1805	6.777075	8.37355	9.282950	9.8183	7.686475	2.506609	
	Pen	1.6795	4.293075	5.07940	7.209275	11.0372	5.804283	3.241293	
	Pod	1.8412	2.814625	5.06620	7.297150	8.9249	5.167233	2.912251	
	Pre	1.7535	2.128350	2.91125	5.814025	10.7948	4.503300	3.583994	
	Xyl	7.7721	8.304300	8.83650	9.368700	9.9009	8.836500	1.505289	

4. Run Linear models in RRPP for each fungal treatment and estimate model coefficients. "d" is the amount of change in a variable for the coefficient indicated.

```
high <- chldata %>% filter(Water == "High") %>% droplevels.data.frame()
low <- chldata %>% filter(Water == "Low") %>% droplevels.data.frame()

# chl b/a ratio linear models and coefficient tests
highLM <- lm.rrpp(Chl.b.a.ratio ~ Block * Fungus * Age, data = high,</pre>
```

```
SS.type = "III", print.progress = F)
## Warning: Because variables in the linear model are redundant,
## the linear model design has been truncated (via QR decomposition).
## Original X columns: 36
## Final X columns (rank): 32
## Check coefficients or degrees of freedom in ANOVA to see changes.
summary(highLM)
##
## Linear Model fit with lm.rrpp
## Number of observations: 40
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
## Number of permutations: 1000
## Full Model Analysis of Variance
##
                        Df Residual Df
                                             SS Residual SS
                                                                   Rsq
## Block * Fungus * Age 31
                                     8 2.392622 0.01395011 0.9942033 44.26136
                        Z (from F)
                                         Pr(>F)
## Block * Fungus * Age 7.315093 0.0005714286
##
## Redundancy Analysis (PCA on fitted values and residuals)
##
                  Trace Proportion Rank
## Fitted
            0.06134928 0.9942034
## Residuals 0.00035770 0.0057968
## Total
            0.06170697 1.0000000
##
## Eigenvalues
##
##
                    PC1
             0.06134928
## Fitted
## Residuals 0.00035770
## Total
            0.06170697
coef(highLM, test = T)
## Linear Model fit with lm.rrpp
## Number of observations: 40
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
## Number of permutations: 1000
## Statistics (distances) of coefficients with 95 percent confidence intervals,
## effect sizes, and probabilities of exceeding observed values based on
```

```
## 1000 random permutations using RRPP
##
##
                                  d.obs UCL (95%)
                                                           Zd Pr(>d)
## (Intercept)
                            0.62593333  0.78962415  -1.7018084  0.952
## Block
                            0.21035000 0.20532837
                                                   1.7769127
                                                               0.032
## FungusCer
                            0.13231667 0.16036021
                                                   0.9805991 0.174
## FungusCok
                            0.06381667 0.09639785
                                                   0.8228150 0.198
## FungusCtrl
                            0.44870000 0.40717963
                                                   2.2324016 0.015
## FungusNig
                            0.14976667 0.20793797
                                                   0.7808900
                                                              0.230
## FungusPen
                            0.03556667 0.15704248 -0.5937618 0.657
## FungusPod
                            0.46796667 0.34985738
                                                   2.9073139 0.001
## FungusPre
                            0.11941667 0.16757535
                                                   0.9150558 0.196
## FungusXyl
                            0.19686667 0.22600022
                                                   1.3330707 0.104
## AgeYoung
                            0.76904060 0.71435667
                                                   2.4423778 0.011
## Block:FungusCtrl
                                                   2.3367838 0.009
                            0.19355000 0.17343434
## Block:FungusNig
                            0.10820000 0.13335851
                                                   0.8374439
                                                               0.210
## Block:FungusPen
                            0.08740000 0.10245408
                                                   1.1892266 0.135
## Block:FungusPod
                            0.34785000 0.27436479
                                                   3.0336971 0.001
## Block:FungusPre
                            0.09720000 0.11787687
                                                   1.0388508 0.154
## Block:FungusXyl
                            0.14065000 0.16774755
                                                   1.1616270 0.142
## Block:AgeYoung
                            0.16917263 0.13960447
                                                   2.6008443 0.005
## FungusCer:AgeYoung
                                                   1.4486806 0.100
                            0.12948086 0.14525467
## FungusCok:AgeYoung
                            0.08700861 0.11893465 1.0380282 0.157
## FungusCtrl:AgeYoung
                            0.41916536 0.27876256
                                                   3.5208619 0.002
## FungusNig:AgeYoung
                            0.09549141 0.18202537 0.3279167 0.329
## FungusPen:AgeYoung
                            0.02459043 0.27774227 -1.2738062 0.905
## FungusPod:AgeYoung
                            0.47760590 0.25875163
                                                   4.7649109
                                                              0.001
## FungusPre:AgeYoung
                             0.10072544 0.20149123
                                                   0.3180400 0.311
## FungusXyl:AgeYoung
                            0.16876311 0.26671137
                                                   0.7328366 0.215
## Block:FungusCtrl:AgeYoung 0.14707756 0.08482203
                                                   4.1435533 0.001
## Block:FungusNig:AgeYoung 0.05145228 0.08667462
                                                   0.6014386
                                                              0.252
## Block:FungusPen:AgeYoung 0.07179740 0.08862127
                                                    1.3539332
                                                              0.112
## Block:FungusPod:AgeYoung 0.31566678 0.14336820
                                                    5.8587068 0.001
## Block:FungusPre:AgeYoung
                            0.06524132 0.08701065
                                                   1.0789188 0.146
## Block:FungusXyl:AgeYoung 0.11646746 0.14606068
                                                   1.2626616 0.130
lowLM <- lm.rrpp(Chl.b.a.ratio ~ Block * Fungus * Age, data = low,</pre>
    SS.type = "III", print.progress = F)
##
## Warning: Because variables in the linear model are redundant,
## the linear model design has been truncated (via QR decomposition).
## Original X columns: 36
## Final X columns (rank): 34
## Check coefficients or degrees of freedom in ANOVA to see changes.
summary(lowLM)
##
## Linear Model fit with lm.rrpp
##
## Number of observations: 46
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
```

```
## Number of permutations: 1000
##
## Full Model Analysis of Variance
##
##
                        Df Residual Df
                                             SS Residual SS
                                                                  Rsq
                                                   4.291964 0.8996645 3.260568
## Block * Fungus * Age 33
                                    12 38.48417
                        Z (from F)
                                       Pr(>F)
## Block * Fungus * Age 1.697163 0.04057143
##
##
## Redundancy Analysis (PCA on fitted values and residuals)
##
##
                 Trace Proportion Rank
## Fitted
             0.8552037
                        0.8996645
## Residuals 0.0953770
                        0.1003355
                                     1
## Total
             0.9505807
                       1.0000000
                                     1
##
## Eigenvalues
##
##
                   PC1
## Fitted
             0.8552037
## Residuals 0.0953770
## Total
            0.9505807
coef(lowLM, test = T)
## Linear Model fit with lm.rrpp
## Number of observations: 46
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
## Number of permutations: 1000
## Statistics (distances) of coefficients with 95 percent confidence intervals,
## effect sizes, and probabilities of exceeding observed values based on
## 1000 random permutations using RRPP
##
##
                                   d.obs UCL (95%)
                                                            Zd Pr(>d)
                             2.604166667 2.380548 2.19819037
## (Intercept)
                                                                0.033
## Block
                             0.570500000 0.911889 0.67613518
## FungusCer
                             0.569933333 3.572471 -1.21800447
                                                                0.869
## FungusCok
                             1.963666667 2.973625 0.84748777
## FungusCtrl
                             1.861066667 2.960517 0.61208920
                                                                0.275
## FungusNig
                             0.194133333 2.542255 -1.05724107
                                                                0.877
## FungusPen
                             1.976066667 2.375479 1.30245979
                                                                0.119
## FungusPod
                             0.004266667
                                         4.284208 -2.20896620
                                                                1.000
## FungusPre
                                          2.385919 1.37534386
                             1.971166667
                                                                0.115
## FungusXyl
                             2.373233333 4.428246 -1.11996396
                                                                0.863
## AgeYoung
                             3.487937297 2.290608 3.53650583
                                                                0.003
## Block:FungusCer
                             0.210000000 1.701526 -1.30906211
                                                                0.906
## Block:FungusCok
                             0.812000000
                                          1.863189 0.07669004
                                                                0.406
## Block:FungusCtrl
                             0.724250000 1.615380 -0.12155632
                                                                0.525
## Block:FungusNig
                             0.029200000 1.169686 -1.23012054 0.951
```

```
## Block:FungusPen
                            0.746900000 1.387197 0.37657861
                            0.001950000 2.094904 -2.37351080
## Block:FungusPod
                                                               1.000
                            0.760600000 1.330731 0.47355806
## Block:FungusPre
                                                               0.298
## Block:AgeYoung
                            2.269119123 1.510589 3.55997849
                                                               0.001
## FungusCer:AgeYoung
                            1.380191764 3.848950 -0.17714962
## FungusCok:AgeYoung
                            4.137690011 3.997899 1.94003885
                                                               0.045
## FungusCtrl:AgeYoung
                            4.781024939 3.642115 2.88065883
                                                               0.005
## FungusNig:AgeYoung
                            0.471646461 3.513066 -0.85062696
                                                               0.768
## FungusPen:AgeYoung
                            4.241875805 3.355649 2.81425797
                                                               0.010
## FungusPod:AgeYoung
                            2.760631578 4.764139 0.33808800
                                                               0.364
## FungusPre:AgeYoung
                            4.264315316 3.303189 2.93256167
                                                               0.009
## FungusXyl:AgeYoung
                            1.823305527 5.201613 -1.57956277
                                                               0.947
## Block:FungusCer:AgeYoung 1.210536461 1.478774 1.27216653
                                                               0.120
## Block:FungusCok:AgeYoung
                            2.427236353 2.385815
                                                  1.99211764
                                                               0.046
## Block:FungusCtrl:AgeYoung 2.878650458 1.481022 5.00004366
                                                               0.001
## Block:FungusNig:AgeYoung 0.123682936
                                         1.594242 -1.01668608
                                                               0.859
## Block:FungusPen:AgeYoung 2.422709974
                                         1.459703
                                                  4.11219575
                                                               0.002
## Block:FungusPod:AgeYoung 2.419037276
                                         1.447938
                                                   4.03337324
                                                               0.002
                            2.421205917
## Block:FungusPre:AgeYoung
                                         1.458626 4.03168898
                                                               0.001
  5. Run ANOVA in RRPP for each linear model.
# chl b/a ratio anovas
highANOVA <- anova(highLM, effect.type = "F", error = c("Residuals",
    "Block:Fungus", "Block:Fungus:Age", "Residuals", "Block:Fungus:Age",
    "Block:Fungus:Age", "Residuals"))
summary(highANOVA)
## Analysis of Variance, using Residual Randomization
## Permutation procedure: Randomization of null model residuals
## Number of permutations: 1000
## Estimation method: Ordinary Least Squares
## Sums of Squares and Cross-products: Type III
## Effect sizes (Z) based on F distributions
##
##
                           SS
                                    MS
                                            Rsq
                                                               Z Pr(>F)
## Block
                    1 0.08849 0.088494 0.036772 50.7490
                                                                  0.003 **
                    8 0.07465 0.009331 0.031019 0.8201 -0.43603 0.656
## Fungus
## Age
                    1 0.12673 0.126734 0.052661 29.0071
                                                                  0.037 *
                    6 0.06827 0.011378 0.028368 6.5250 1.82704
## Block:Fungus
                                                                  0.022 *
## Block:Age
                    1 0.02862 0.028619 0.011892 6.5505
                                                                  0.046 *
## Fungus:Age
                    8 0.03616 0.004520 0.015025 1.0345 -0.42431 0.651
## Block:Fungus:Age 6 0.02621 0.004369 0.010893 2.5055 1.19833 0.106
                    8 0.01395 0.001744 0.005797
## Residuals
## Total
                   39 2.40657
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Call: lm.rrpp(f1 = Chl.b.a.ratio ~ Block * Fungus * Age, SS.type = "III",
      data = high, print.progress = F)
lowANOVA <- anova(lowLM, effect.type = "F", error = c("Residuals",</pre>
    "Block:Fungus", "Block:Fungus:Age", "Residuals", "Block:Fungus:Age",
    "Block:Fungus:Age", "Residuals"))
```

summary(lowANOVA)

```
##
## Analysis of Variance, using Residual Randomization
## Permutation procedure: Randomization of null model residuals
## Number of permutations: 1000
## Estimation method: Ordinary Least Squares
## Sums of Squares and Cross-products: Type III
## Effect sizes (Z) based on F distributions
##
##
                    Df
                            SS
                                   MS
                                                              Z Pr(>F)
                                           Rsq
## Block
                        0.651 0.6509 0.015217 1.8200
                                                                 0.183
                     1
## Fungus
                     8 10.522 1.3153 0.245979 3.9387
                                                        0.91855
                                                                 0.188
                        2.607 2.6069 0.060944 2.1006
## Age
                                                       1.15093
                                                                 0.087
## Block:Fungus
                        2.338 0.3339 0.054645 0.9336 -0.00297
                                                                 0.501
## Block:Age
                        5.149 5.1489 0.120369 4.1489
                                                                 0.040 *
                     1
## Fungus:Age
                     8
                        9.881 1.2351 0.230997 0.9953 -0.19437
                                                                 0.587
## Block:Fungus:Age
                    7
                        8.687 1.2410 0.203084 3.4698 1.84012 0.023 *
## Residuals
                    12 4.292 0.3577 0.100336
                    45 42.776
## Total
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Call: lm.rrpp(f1 = Chl.b.a.ratio ~ Block * Fungus * Age, SS.type = "III",
##
       data = low, print.progress = F)
  6. Test pairwise differences between least squares means. Similar to tukeyHSD function in the r stats
    package. The pairwise function will generate tables with confidence intervals and p-values for the
    pairwise statistic, Euclidean distance between least-squares means.
# chl b/a ratio pairwise
highpw <- pairwise(highLM, groups = high$Fungus)
summary(highpw, confidence = 0.95, stat.table = T)
## Pairwise comparisons
##
## Groups: Asp Cer Cok Ctrl Nig Pen Pod Pre Xyl
##
## RRPP: 1000 permutations
## LS means:
## Vectors hidden (use show.vectors = TRUE to view)
##
## Pairwise distances between means, plus statistics
##
                      d UCL (95%)
                                                Z Pr > d
## Asp:Cer 0.058187450 0.10362515 -0.0346201754
## Asp:Cok 0.105451323 0.14966563 0.0190512654
## Asp:Ctrl 0.000905116 0.03796745 -1.2585487742
                                                   0.968
```

0.529

0.519

0.506

0.479

## Asp:Nig 0.062926762 0.09451527 0.0231018162 ## Asp:Pen 0.079731152 0.10889033 -0.0410287190

## Asp:Pod 0.118743037 0.15222646 -0.0085941455

## Cer:Cok 0.047263873 0.10049838 -0.0224210471

0.060104737 0.09521596 -0.0286808290

0.074021132 0.10748402 -0.0087205741

## Cer:Ctrl 0.057282334 0.10313445 -0.0114821100 0.502

## Asp:Pre

## Asp:Xyl

```
## Cer:Nig 0.004739313 0.05099240 -1.0911271960 0.879
## Cer:Pen 0.021543703 0.06519094 -0.3260920691 0.566
## Cer:Pod 0.060555587 0.10432917 0.0136784160 0.472
## Cer:Pre 0.001917287 0.05199894 -1.2895651944 0.947
## Cer:Xyl 0.015833683 0.06386772 -0.5432613083 0.642
## Cok:Ctrl 0.104546207 0.14853223 0.0408905511 0.481
## Cok:Nig 0.042524561 0.08436076 -0.0262240862 0.471
## Cok:Pen 0.025720171 0.06928486 -0.1973884123 0.479
## Cok:Pod 0.013291714 0.06068414 -0.6399579370 0.674
## Cok:Pre 0.045346586 0.08764837 -0.0090404191
## Cok:Xyl 0.031430191 0.07696863 -0.1177374124
## Ctrl:Nig 0.062021646 0.09150534 0.0559327036
                                                 0.482
## Ctrl:Pen 0.078826036 0.10975971 -0.0076700792 0.492
## Ctrl:Pod 0.117837921 0.15166548 0.0198723805 0.491
## Ctrl:Pre 0.059199621 0.09263527 0.0006638177
                                                 0.502
## Ctrl:Xyl 0.073116016 0.10843133 0.0196979463
                                                 0.482
## Nig:Pen 0.016804390 0.04702013 -0.2927374501 0.553
## Nig:Pod 0.055816275 0.08712838 -0.0311029320 0.514
## Nig:Pre 0.002822025 0.04009023 -1.1086653292 0.896
## Nig:Xyl 0.011094370 0.04675439 -0.5807182213 0.670
## Pen:Pod 0.039011885 0.07138210 0.0136824892 0.491
## Pen:Pre 0.019626415 0.05276055 -0.2556348307 0.544
## Pen:Xyl 0.005710020 0.04093107 -0.8670715107 0.794
## Pod:Pre 0.058638300 0.09537530 0.0146895642 0.500
## Pod:Xyl 0.044721905 0.08076102 -0.0175396702 0.504
## Pre:Xyl 0.013916396 0.04935847 -0.4613067814 0.597
highpw2 <- pairwise(highLM, groups = high$Age)
summary(highpw2, confidence = 0.95, stat.table = T)
##
## Pairwise comparisons
## Groups: Old Young
##
## RRPP: 1000 permutations
##
## LS means:
## Vectors hidden (use show.vectors = TRUE to view)
## Pairwise distances between means, plus statistics
                    d UCL (95%)
                                         Z Pr > d
## Old:Young 0.4570754 0.4738713 0.01935818 0.493
lowpw <- pairwise(lowLM, groups = low$Fungus)</pre>
summary(lowpw, confidence = 0.95, stat.table = T)
##
## Pairwise comparisons
## Groups: Asp Cer Cok Ctrl Nig Pen Pod Pre Xyl
##
## RRPP: 1000 permutations
## LS means:
```

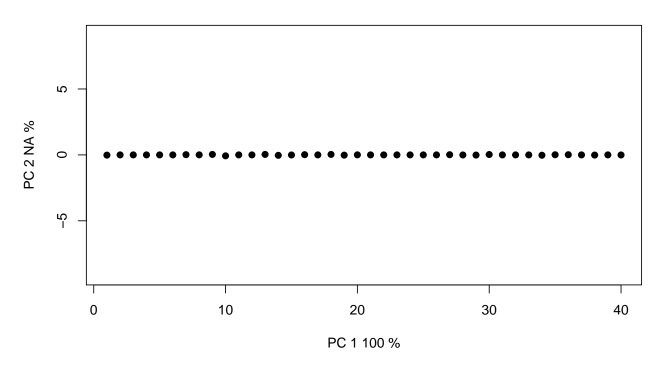
```
## Vectors hidden (use show.vectors = TRUE to view)
##
## Pairwise distances between means, plus statistics
                     d UCL (95%)
##
                                            Z Pr > d
## Asp:Cer 0.370507246 0.8824098 -0.146156889
## Asp:Cok 0.779278707 1.3245751 -0.021183465
## Asp:Ctrl 0.900704655 1.4303293 -0.006761711
## Asp:Nig 0.364673628 0.9346442 -0.174334354
## Asp:Pen 0.784038739 1.2714713 -0.002437905
                                               0.492
## Asp:Pod 1.039088154 1.5285736 0.032213714 0.502
## Asp:Pre 0.739014926 1.2570003 -0.006926196
## Asp:Xyl
           0.897521008 1.6447366 -0.020139560
                                               0.496
## Cer:Cok 0.408771461 1.0035870 -0.162227989
                                               0.505
## Cer:Ctrl 0.530197409 1.0699132 -0.053384542
## Cer:Nig 0.735180874 1.3234989 -0.011188988
                                               0.489
## Cer:Pen 0.413531493 0.9215101 -0.110791419
## Cer:Pod 0.668580908 1.1774047 0.012461459
                                               0.483
## Cer:Pre 0.368507680 0.8670062 -0.159698713
## Cer:Xyl 1.268028254 2.0177650 -0.008447710
                                               0.495
## Cok:Ctrl 0.121425948 0.6870629 -0.815420164
## Cok:Nig 1.143952335 1.7519492 -0.013823141
## Cok:Pen 0.004760032 0.6707432 -1.266044505
## Cok:Pod 0.259809447 0.8187707 -0.325649671
                                               0.557
## Cok:Pre 0.040263781 0.7021350 -1.087281063
                                               0.893
## Cok:Xyl 1.676799715 2.4871440 -0.020665013
## Ctrl:Nig 1.265378283 1.7925585 -0.008130811
## Ctrl:Pen 0.116665916 0.6867539 -0.724696498
                                               0.713
## Ctrl:Pod 0.138383499 0.6365635 -0.663442545
                                               0.688
## Ctrl:Pre 0.161689729 0.6769211 -0.555791625
                                               0.637
## Ctrl:Xyl 1.798225663 2.5619190 -0.016507705
## Nig:Pen 1.148712367 1.7013890 -0.002288398
## Nig:Pod 1.403761782 1.9756911 0.026647003
                                               0.487
## Nig:Pre
           1.103688554 1.6786208 -0.005586874
## Nig:Xyl 0.532847380 1.3596688 -0.157290367
                                               0.498
## Pen:Pod 0.255049415 0.7325595 -0.277786545
## Pen:Pre 0.045023813 0.6088501 -1.064300159
                                               0.879
## Pen:Xyl 1.681559747 2.4587113 -0.012092696
## Pod:Pre 0.300073228 0.8107839 -0.160157623
                                               0.494
## Pod:Xyl 1.936609162 2.7075396 0.009614820
                                               0.490
## Pre:Xyl 1.636535934 2.3869078 -0.015338941 0.491
lowpw2 <- pairwise(lowLM, groups = low$Age)</pre>
summary(lowpw2, confidence = 0.95, stat.table = T)
##
## Pairwise comparisons
## Groups: Old Young
## RRPP: 1000 permutations
## LS means:
## Vectors hidden (use show.vectors = TRUE to view)
## Pairwise distances between means, plus statistics
```

```
## d UCL (95%) Z Pr > d
## Old:Young 0.1482125 0.4289912 -0.2611132 0.542
```

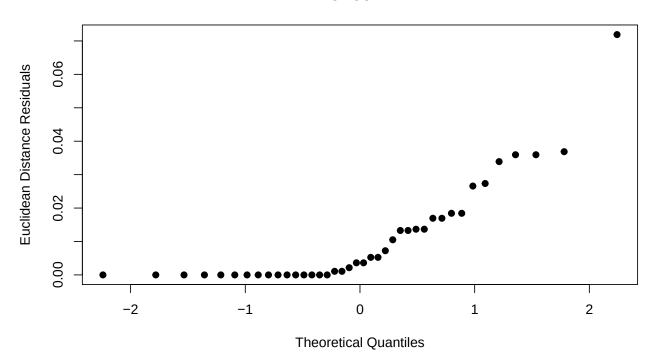
7. Examine RRPP plots to check for assumptions.

```
## chl b/a ratio residuals vs fitted values (homoscedasticity
## check)
hdiagnostics <- plot(highLM, type = "diagnostics")</pre>
```

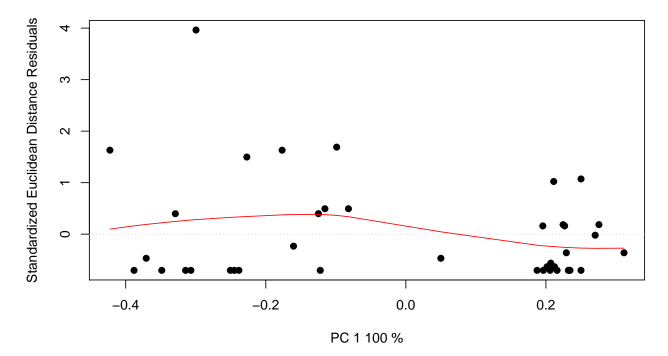
# **PCA** Residuals



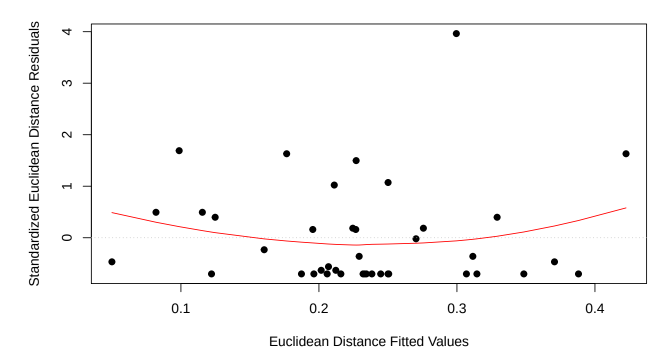
# Q-Q plot

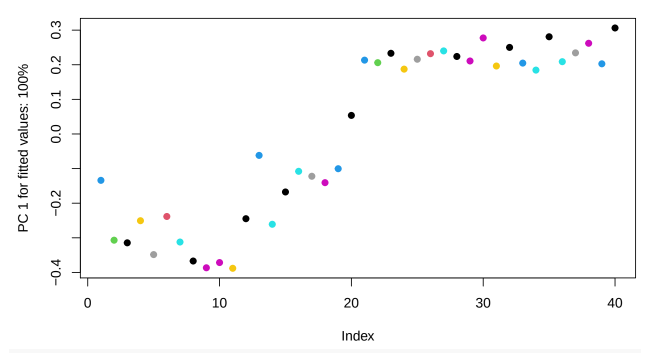


# Residuals vs. PC 1 fitted



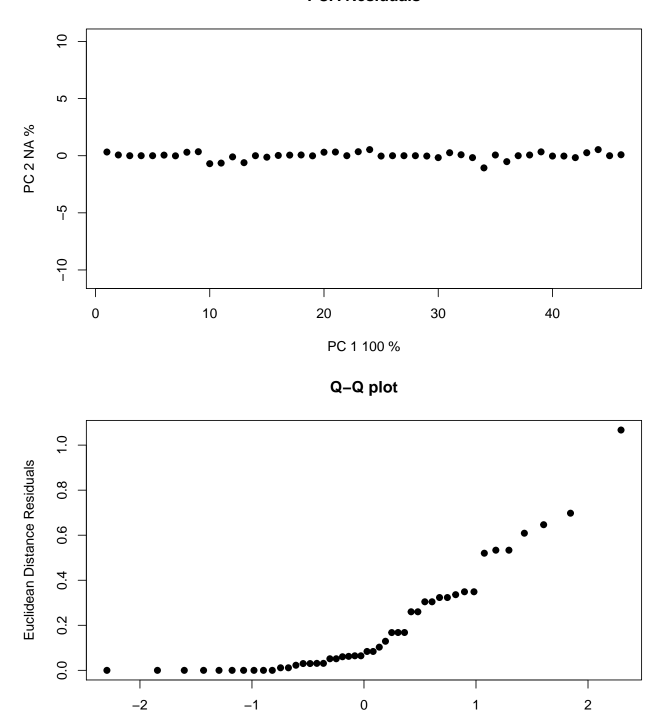
### Residuals vs. Fitted





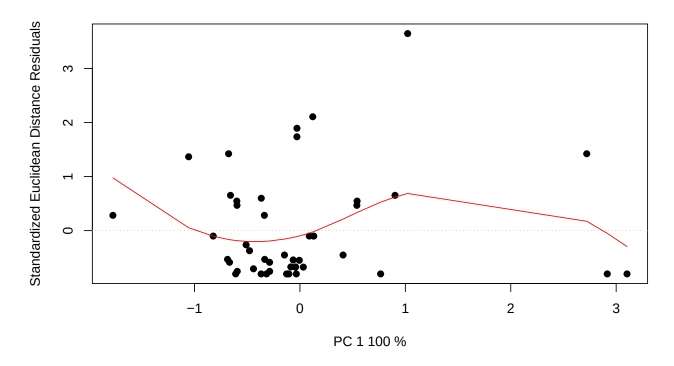
# residuals vs fitted values (homoscedasticity check)
ldiagnostics <- plot(lowLM, type = "diagnostics")</pre>

## **PCA Residuals**

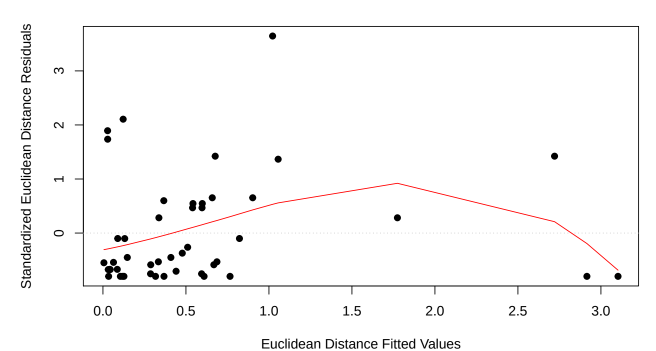


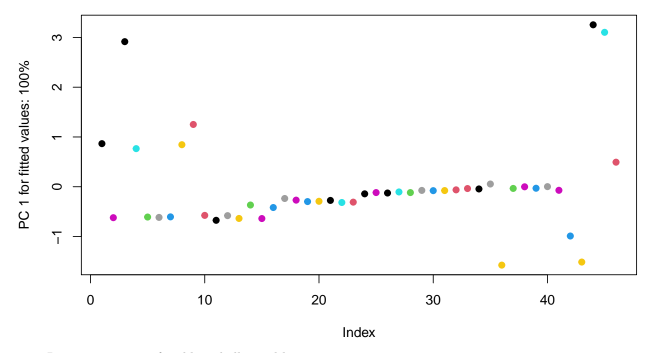
Theoretical Quantiles

### Residuals vs. PC 1 fitted



### Residuals vs. Fitted





8. Repeat steps 4 - 7 for chlorophyll a and b

```
## chl a linear models and coefficient tests
highLM <- lm.rrpp(Chl.a ~ Block * Fungus * Age, data = high,</pre>
    SS.type = "III", print.progress = F)
##
## Warning: Because variables in the linear model are redundant,
## the linear model design has been truncated (via QR decomposition).
## Original X columns: 36
## Final X columns (rank): 32
## Check coefficients or degrees of freedom in ANOVA to see changes.
summary(highLM)
##
## Linear Model fit with lm.rrpp
##
## Number of observations: 40
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
## Number of permutations: 1000
##
## Full Model Analysis of Variance
##
##
                        Df Residual Df
                                              SS Residual SS
                                                                    Rsq
                                      8 34.62868
                                                    4.896895 0.8761082 1.824918
## Block * Fungus * Age 31
                        Z (from F)
                                       Pr(>F)
## Block * Fungus * Age 0.9115409 0.1735714
##
##
## Redundancy Analysis (PCA on fitted values and residuals)
##
```

```
Trace Proportion Rank
## Fitted
             0.8879149 0.8761082
## Residuals 0.1255614 0.1238918
## Total
             1.0134763 1.0000000
                                     1
## Eigenvalues
##
##
                   PC1
## Fitted
             0.8879149
## Residuals 0.1255614
## Total
            1.0134763
coef(highLM, test = T)
##
## Linear Model fit with lm.rrpp
## Number of observations: 40
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
## Number of permutations: 1000
## Statistics (distances) of coefficients with 95 percent confidence intervals,
## effect sizes, and probabilities of exceeding observed values based on
## 1000 random permutations using RRPP
##
##
                                 d.obs UCL (95%)
                                                          Zd Pr(>d)
## (Intercept)
                             1.4887000 3.0853911 -0.91657749 0.849
## Block
                             0.0189000 0.7339735 -1.36102805
## FungusCer
                             0.8958000 1.5052538 0.45423624
## FungusCok
                             3.1225000 3.7312653 0.50712316
## FungusCtrl
                             0.4613333 1.9695736 -0.55591572
                                                             0.648
## FungusNig
                             2.5825333 2.5229932 1.84283976
## FungusPen
                             1.2428667 2.3952412 0.24622508
                                                             0.364
## FungusPod
                            1.1789000 2.3172255 0.29964699
                                                             0.316
## FungusPre
                             1.2563500 1.8762403
                                                  0.82941234
                                                              0.195
## FungusXyl
                             1.0466000 2.2303053 0.21596324 0.338
## AgeYoung
                             1.3735000 1.7854776 1.21858669
                                                             0.121
## Block:FungusCtrl
                             0.0708500 0.8329686 -1.00025230
## Block:FungusNig
                             1.0067000 0.9537625 2.06332649
                                                             0.037
## Block:FungusPen
                             0.3558500 0.9118538 -0.03534143
                                                             0.443
## Block:FungusPod
                             0.2273000 1.3362326 -0.77682599
## Block:FungusPre
                             0.0790500 0.9938551 -1.13348777
                                                              0.890
## Block:FungusXyl
                             0.2903000 1.4039738 -0.62786431
                                                              0.674
## Block:AgeYoung
                             0.4828500 0.8200662 0.63112918 0.231
## FungusCer:AgeYoung
                             0.1345500 1.9780155 -1.16347793
## FungusCok:AgeYoung
                             1.2665500 1.9560037 0.67878754
                                                             0.239
## FungusCtrl:AgeYoung
                             1.1549333 3.1216804 -0.25203936
                                                              0.546
## FungusNig:AgeYoung
                             2.1915667 2.9151877 1.07410148
                                                             0.158
## FungusPen:AgeYoung
                             0.8735667 2.3490271 -0.16738373
                             0.7334000 4.7852567 -1.13050431
## FungusPod:AgeYoung
                                                              0.862
## FungusPre:AgeYoung
                             1.7795000 2.4210114 1.10464050
                                                              0.147
## FungusXyl:AgeYoung
                             1.8918000 3.1764876 0.64418769
                                                              0.230
## Block:FungusCtrl:AgeYoung 0.0221500 1.0969676 -1.30517232
```

```
## Block:FungusNig:AgeYoung 1.5067000 1.1356602 3.00484209 0.013
## Block:FungusPen:AgeYoung 0.2853000 1.0946470 -0.49818084 0.625
## Block:FungusPod:AgeYoung 0.7454500 1.7882282 0.02809061 0.410
## Block:FungusPre:AgeYoung 0.8015500 1.0987716 1.02549553 0.167
## Block:FungusXyl:AgeYoung 1.0046500 1.8675746 0.44887360 0.288
lowLM <- lm.rrpp(Chl.a ~ Block * Fungus * Age, data = low, SS.type = "III",
   print.progress = F)
##
## Warning: Because variables in the linear model are redundant,
## the linear model design has been truncated (via QR decomposition).
## Original X columns: 36
## Final X columns (rank): 34
## Check coefficients or degrees of freedom in ANOVA to see changes.
summary(lowLM)
##
## Linear Model fit with lm.rrpp
##
## Number of observations: 46
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
## Number of permutations: 1000
## Full Model Analysis of Variance
##
                                             SS Residual SS
##
                        Df Residual Df
                                                                  Rsq
                                    12 67.19567
                                                   13.12627 0.8365793 1.861519
## Block * Fungus * Age 33
                        Z (from F)
                                      Pr(>F)
## Block * Fungus * Age 1.115489 0.1235714
##
## Redundancy Analysis (PCA on fitted values and residuals)
##
##
                 Trace Proportion Rank
## Fitted
             1.4932372
                       0.8365793
## Residuals 0.2916948
                        0.1634207
                                     1
## Total
             1.7849320
                       1.0000000
                                     1
##
## Eigenvalues
##
##
                   PC1
## Fitted
             1.4932372
## Residuals 0.2916948
## Total
             1.7849320
coef(lowLM, test = T)
##
## Linear Model fit with lm.rrpp
## Number of observations: 46
## Number of dependent variables: 1
```

```
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
## Number of permutations: 1000
##
## Statistics (distances) of coefficients with 95 percent confidence intervals,
## effect sizes, and probabilities of exceeding observed values based on
## 1000 random permutations using RRPP
##
##
                                d.obs UCL (95%)
                                                        Zd Pr(>d)
## (Intercept)
                            4.3430667 6.609287 -0.62588375 0.749
## Block
                            0.7142000 1.810957 -0.73988334
                                                           0.400
## FungusCer
                            2.7943333 4.862370 0.23929076
## FungusCok
                            2.6333333 4.084301 0.70598992 0.216
                          2.5017333 3.048351
                                               1.27451234 0.120
## FungusCtrl
## FungusNig
                          1.1112667 3.089896 -0.14606177 0.471
                          1.6043667
## FungusPen
                                      3.427425 0.20001063
                                                           0.366
## FungusPod
                          1.5069667 6.422746 -1.72609263 0.956
## FungusPre
                          0.2210000 2.806793 -1.03369918 0.862
## FungusXyl
                            2.1914667 3.982275 -0.34410244 0.639
## AgeYoung
                           1.0963333 2.885914 -0.05362694 0.424
## Block:FungusCer
                          0.8895000 1.872724 0.12187111 0.424
## Block:FungusCok
                           1.5832000 2.518236 0.74871140 0.213
## Block:FungusCtrl
                            1.2090500 1.446024 1.41494296 0.098
## Block:FungusNig
                            0.2778000 1.349741 -0.59721995 0.673
## Block:FungusPen
                           1.0023500 1.785917 0.40209781 0.309
## Block:FungusPod
                            0.1797000 2.580560 -1.94442949 0.978
## Block:FungusPre
                            0.6740500 1.693491 -0.14023799 0.498
## Block:AgeYoung
                            0.3849000 1.425723 -0.42277444 0.577
## FungusCer:AgeYoung
                            0.9018000 4.007211 -0.59667920 0.667
## FungusCok:AgeYoung
                            3.0812333 5.174928 0.64969711 0.233
## FungusCtrl:AgeYoung
                            4.2113000 4.220907
                                               1.92897216 0.052
## FungusNig:AgeYoung
                            1.1608667 4.044206 -0.39651985 0.576
## FungusPen:AgeYoung
                            1.2637000 4.089974 -0.31428285
                                                           0.542
## FungusPod:AgeYoung
                            2.8811000 5.491102 0.26335630 0.365
## FungusPre:AgeYoung
                            0.1191667
                                      3.962217 -1.21226643 0.950
## FungusXyl:AgeYoung
                            2.0379667 5.011630 -0.57240819 0.702
## Block:FungusCer:AgeYoung 0.3556500 1.906550 -0.71204832 0.721
## Block:FungusCok:AgeYoung 1.7435000 2.968737 0.58480712 0.249
## Block:FungusCtrl:AgeYoung 2.0082000 1.968025 2.03272731 0.043
## Block:FungusNig:AgeYoung 0.5530000 1.870426 -0.34142341 0.567
## Block:FungusPen:AgeYoung 0.6282000 1.904982 -0.22795535
## Block:FungusPod:AgeYoung
                            2.4821000
                                      2.013197 2.80958917
                                                           0.013
## Block:FungusPre:AgeYoung 0.1321000 1.821151 -1.10340450 0.894
# ratio anovas
highANOVA <- anova(highLM, effect.type = "F", error = c("Residuals",
    "Block:Fungus", "Block:Fungus:Age", "Residuals", "Block:Fungus:Age",
    "Block:Fungus:Age", "Residuals"))
summary(highANOVA)
##
## Analysis of Variance, using Residual Randomization
## Permutation procedure: Randomization of null model residuals
## Number of permutations: 1000
## Estimation method: Ordinary Least Squares
```

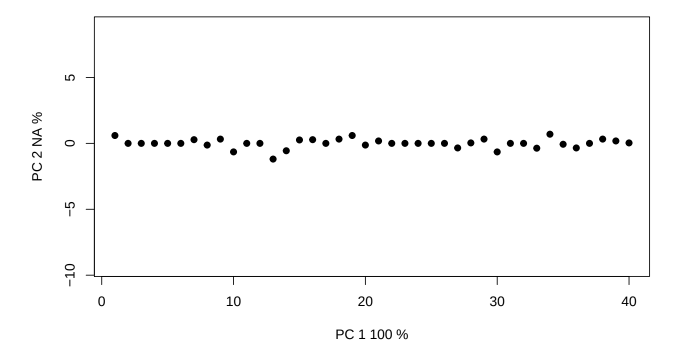
```
## Sums of Squares and Cross-products: Type III
## Effect sizes (Z) based on F distributions
##
##
                    Df
                           SS
                                   MS
                                                             Z Pr(>F)
                                           Rsq
## Block
                       0.001 0.00071 0.000018 0.0012
                                                                 0.832
## Fungus
                     8 6.835 0.85442 0.172936 3.6871
                                                       0.87411
                                                                0.202
                     1 0.404 0.40425 0.010228 1.3942
## Age
                     6 1.390 0.23174 0.035178 0.3786 -0.97017
## Block:Fungus
                                                                0.849
                       0.233 0.23314 0.005899 0.8041
## Block:Age
                     1
                                                                 0.326
                     8 2.416 0.30205 0.061134 1.0417 -0.22250
## Fungus:Age
                                                                0.555
## Block:Fungus:Age 6 1.740 0.28994 0.044013 0.4737 -0.82589
                                                                0.804
                     8 4.897 0.61211 0.123892
## Residuals
## Total
                    39 39.526
##
## Call: lm.rrpp(f1 = Chl.a ~ Block * Fungus * Age, SS.type = "III", data = high,
       print.progress = F)
lowANOVA <- anova(lowLM, effect.type = "F", error = c("Residuals",</pre>
    "Block:Fungus", "Block:Fungus:Age", "Residuals", "Block:Fungus:Age",
    "Block:Fungus:Age", "Residuals"))
summary(lowANOVA)
## Analysis of Variance, using Residual Randomization
## Permutation procedure: Randomization of null model residuals
## Number of permutations: 1000
## Estimation method: Ordinary Least Squares
## Sums of Squares and Cross-products: Type III
## Effect sizes (Z) based on F distributions
##
##
                    Df
                           SS
                                                               Z Pr(>F)
                                   MS
                                                    F
                                           Rsq
## Block
                     1 1.020 1.02016 0.012701 0.9326
                                                                  0.277
                     8 14.156 1.76956 0.176246 1.5751
                                                                 0.360
## Fungus
                                                       0.281274
                     1 0.258 0.25756 0.003207 0.2084
                                                       0.099191
## Age
## Block:Fungus
                     7 7.864 1.12346 0.097909 1.0271
                                                       0.174149 0.430
## Block:Age
                     1 0.148 0.14815 0.001844 0.1199
                                                                  0.625
## Fungus:Age
                     8 8.488 1.06096 0.105671 0.8586 -0.280167 0.612
## Block:Fungus:Age 7 8.649 1.23563 0.107684 1.1296 0.259904 0.404
## Residuals
                    12 13.126 1.09386 0.163421
## Total
                    45 80.322
## Call: lm.rrpp(f1 = Chl.a ~ Block * Fungus * Age, SS.type = "III", data = low,
       print.progress = F)
##
# pairwise
highpw <- pairwise(highLM, groups = high$Fungus)</pre>
summary(highpw, confidence = 0.95, stat.table = T)
##
## Pairwise comparisons
##
## Groups: Asp Cer Cok Ctrl Nig Pen Pod Pre Xyl
## RRPP: 1000 permutations
##
```

```
## LS means:
## Vectors hidden (use show.vectors = TRUE to view)
## Pairwise distances between means, plus statistics
                   d UCL (95%)
                                           Z Pr > d
## Asp:Cer 1.1856000 1.6802680 6.023151e-02 0.499
## Asp:Cok 2.7117500 3.2327485 -9.985210e-03 0.520
## Asp:Ctrl 0.8749500 1.2504739 -2.410592e-05 0.494
## Asp:Nig 0.9800500 1.3563340 -1.935596e-02 0.510
## Asp:Pen 0.3796833 0.7976021 -3.101756e-02 0.490
## Asp:Pod 1.8750000 2.2877578 7.267837e-04 0.503
## Asp:Pre 1.0100500 1.4345776 -4.036789e-03 0.500
## Asp:Xyl 0.5300000 0.9277738 3.401526e-02 0.483
## Cer:Cok 1.5261500 2.2614225 -5.739252e-02 0.521
## Cer:Ctrl 0.3106500 0.8291124 -2.545953e-01 0.540
## Cer:Nig 0.2055500 0.7366934 -4.863965e-01
                                             0.618
## Cer:Pen 0.8059167 1.3489886 4.596325e-02 0.493
## Cer:Pod 0.6894000 1.3019102 -7.582346e-02 0.518
## Cer:Pre 0.1755500 0.7570581 -6.165458e-01 0.669
## Cer:Xyl 0.6556000 1.2029159 -1.354628e-02 0.498
## Cok:Ctrl 1.8368000 2.3409031 -9.725432e-03 0.514
## Cok:Nig 1.7317000 2.2866762 3.812979e-03 0.507
## Cok:Pen 2.3320667 2.8531326 -1.451847e-02 0.513
## Cok:Pod 0.8367500 1.4065309 -2.298182e-02 0.512
## Cok:Pre 1.7017000 2.2537116 -5.963106e-03 0.505
## Cok:Xyl 2.1817500 2.7507572 -4.619412e-02 0.529
## Ctrl:Nig 0.1051000 0.5242413 -6.376140e-01 0.678
## Ctrl:Pen 0.4952667 0.8879894 -1.774451e-02 0.491
## Ctrl:Pod 1.0000500 1.4165075 7.627263e-04 0.494
## Ctrl:Pre 0.1351000 0.5655203 -5.899096e-01 0.648
## Ctrl:Xyl 0.3449500 0.7608720 -1.439114e-01 0.525
## Nig:Pen 0.6003667 0.9964252 -2.660414e-02 0.504
## Nig:Pod 0.8949500 1.3304591 1.686732e-02 0.506
## Nig:Pre 0.0300000 0.5371340 -1.199612e+00 0.932
## Nig:Xyl 0.4500500 0.9089682 -1.046968e-01 0.531
## Pen:Pod 1.4953167 1.9276026 -5.681110e-03 0.511
## Pen:Pre 0.6303667 1.0685515 -1.365553e-02 0.518
## Pen:Xyl 0.1503167 0.5969929 -5.105108e-01 0.622
## Pod:Pre 0.8649500 1.3545564 4.009262e-03 0.505
## Pod:Xyl 1.3450000 1.8303491 -4.279087e-02 0.528
## Pre:Xyl 0.4800500 0.9642244 -8.350287e-02 0.523
highpw2 <- pairwise(highLM, groups = high$Age)
summary(highpw2, confidence = 0.95, stat.table = T)
##
## Pairwise comparisons
## Groups: Old Young
## RRPP: 1000 permutations
##
## Vectors hidden (use show.vectors = TRUE to view)
##
```

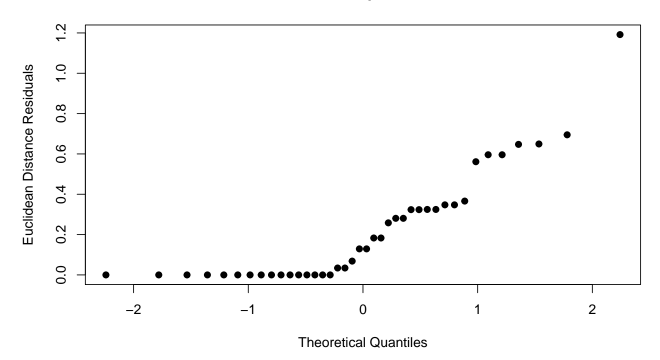
```
## Pairwise distances between means, plus statistics
##
                   d UCL (95%)
                                       7. Pr > d
## Old:Young 0.82123 1.025669 0.08700829
lowpw <- pairwise(lowLM, groups = low$Fungus)</pre>
summary(lowpw, confidence = 0.95, stat.table = T)
##
## Pairwise comparisons
## Groups: Asp Cer Cok Ctrl Nig Pen Pod Pre Xyl
## RRPP: 1000 permutations
##
## LS means:
## Vectors hidden (use show.vectors = TRUE to view)
## Pairwise distances between means, plus statistics
                    d UCL (95%)
                                             Z Pr > d
## Asp:Cer 0.92008333 1.5720712 0.0678586475
## Asp:Cok 0.47886667 1.2146186 -0.2178031399
## Asp:Ctrl 0.01381667 0.7543234 -1.2733739955
## Asp:Nig 0.52823333 1.2430587 -0.1043910973
## Asp:Pen 0.40398333 1.0515362 -0.1464333981
## Asp:Pod 0.10601667 0.8029328 -0.9219082734
## Asp:Pre 1.05458333 1.7086850 0.0187078538
## Asp:Xyl 0.26583333 1.2811022 -0.5864877562 0.671
## Cer:Cok 1.39895000 2.1300418 0.0096921857
## Cer:Ctrl 0.90626667 1.5517110 0.0479615991
                                               0.478
## Cer:Nig 0.39185000 1.1441704 -0.2766390792
## Cer:Pen 0.51610000 1.1546081 -0.0884521570
## Cer:Pod 0.81406667 1.4771202 0.0079406657
## Cer:Pre 0.13450000 0.8577604 -0.8048058541
                                               0.759
## Cer:Xyl 0.65425000 1.6480842 -0.1674366668
## Cok:Ctrl 0.49268333 1.2162097 -0.1920536284
                                               0.524
## Cok:Nig 1.00710000 1.7998018 -0.0286045174
## Cok:Pen 0.88285000 1.5932777 -0.0411620009
                                               0.518
## Cok:Pod 0.58488333 1.2904957 -0.1232107053
                                               0.520
## Cok:Pre 1.53345000 2.2981649 -0.0393285501
## Cok:Xyl 0.74470000 1.7063965 -0.1148949707
## Ctrl:Nig 0.51441667 1.2596794 -0.1404352867
                                               0.510
## Ctrl:Pen 0.39016667 1.0154965 -0.1829154233
                                               0.511
## Ctrl:Pod 0.09220000 0.8067514 -0.9225830360
## Ctrl:Pre 1.04076667 1.6999822 -0.0006409169
## Ctrl:Xyl 0.25201667 1.2827291 -0.6371056985
## Nig:Pen 0.12425000 0.9550303 -0.8730704753
                                               0.793
## Nig:Pod 0.42221667 1.1696425 -0.2618664491
## Nig:Pre 0.52635000 1.3379634 -0.1756051775
                                               0.524
## Nig:Xyl
           0.26240000 1.3637741 -0.6710819396
## Pen:Pod 0.29796667 0.9564494 -0.3786666724
                                               0.566
## Pen:Pre 0.65060000 1.3175833 -0.0759634282
## Pen:Xyl 0.13815000 1.2159924 -0.8514362884
                                               0.803
## Pod:Pre 0.94856667 1.6427931 -0.0482247913
                                               0.520
## Pod:Xyl 0.15981667 1.2255936 -0.8653035282
## Pre:Xyl 0.78875000 1.7594061 -0.1613061192 0.511
```

```
lowpw2 <- pairwise(lowLM, groups = low$Age)</pre>
summary(lowpw2, confidence = 0.95, stat.table = T)
##
## Pairwise comparisons
##
## Groups: Old Young
##
## RRPP: 1000 permutations
##
## LS means:
## Vectors hidden (use show.vectors = TRUE to view)
## Pairwise distances between means, plus statistics
##
                     d UCL (95%)
                                           Z Pr > d
## Old:Young 0.7019609 1.012106 0.06918075 0.465
# residuals vs fitted values (homoscedasticity check)
hdiagnostics <- plot(highLM, type = "diagnostics")</pre>
```

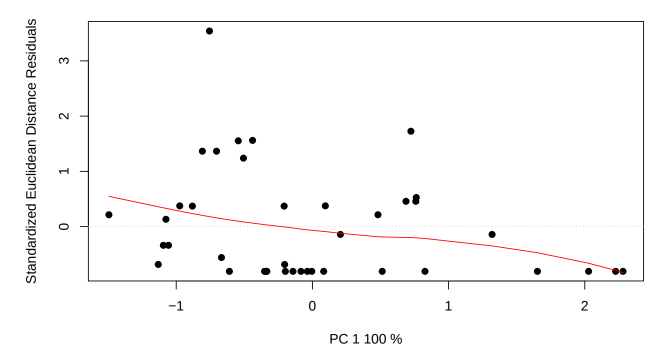
#### **PCA Residuals**



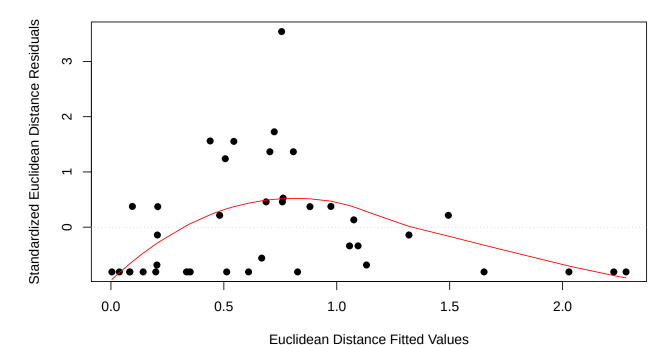
# Q-Q plot

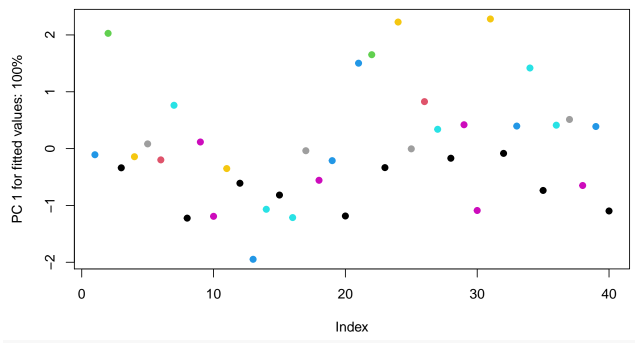


## Residuals vs. PC 1 fitted



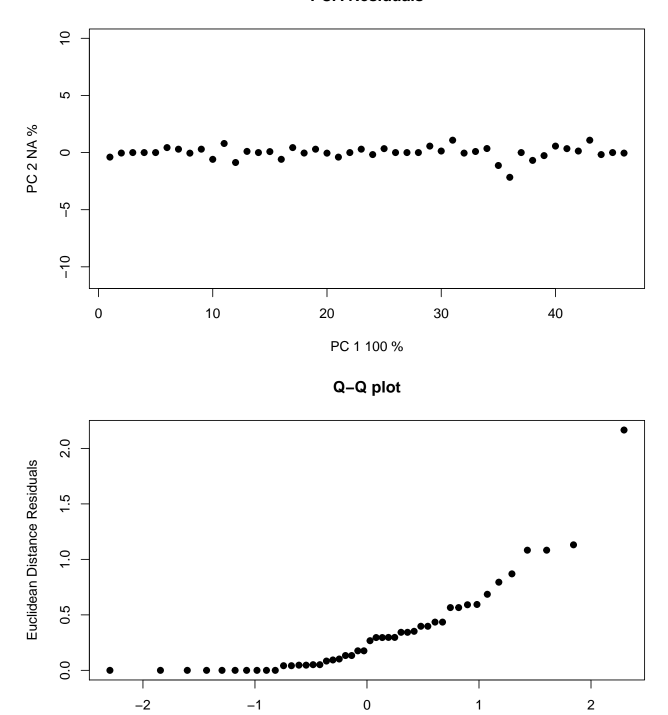
### Residuals vs. Fitted





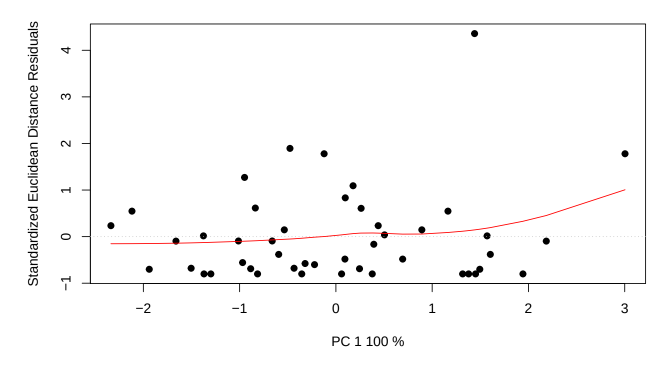
# residuals vs fitted values (homoscedasticity check)
ldiagnostics <- plot(lowLM, type = "diagnostics")</pre>

## **PCA Residuals**

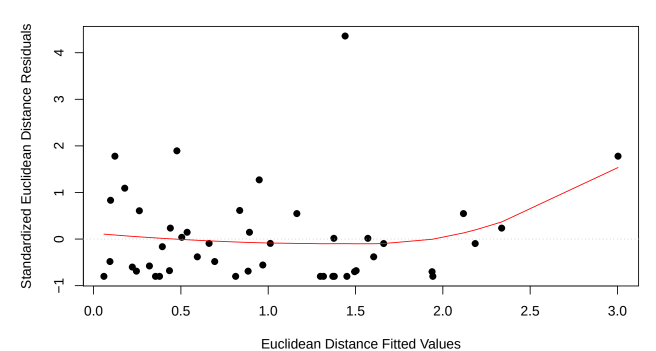


Theoretical Quantiles

### Residuals vs. PC 1 fitted



## Residuals vs. Fitted



```
4
PC 1 for fitted values: 100%
     က
     \alpha
     0
      7
     7
           0
                            10
                                                               30
                                              20
                                                                                 40
                                                   Index
## chl b linear models and coefficient tests
highLM <- lm.rrpp(Chl.b ~ Block * Fungus * Age, data = high,
    SS.type = "III", print.progress = F)
##
## Warning: Because variables in the linear model are redundant,
## the linear model design has been truncated (via QR decomposition).
## Original X columns: 36
## Final X columns (rank): 32
\ensuremath{\mbox{\#\#}} Check coefficients or degrees of freedom in ANOVA to see changes.
summary(highLM)
##
## Linear Model fit with lm.rrpp
## Number of observations: 40
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
## Number of permutations: 1000
## Full Model Analysis of Variance
##
##
                          Df Residual Df
                                                SS Residual SS
                                        8 87.50118
                                                       6.465898 0.9311898 3.492315
## Block * Fungus * Age 31
                          Z (from F)
                                          Pr(>F)
                            2.069305 0.02857143
##
  Block * Fungus * Age
##
##
## Redundancy Analysis (PCA on fitted values and residuals)
##
##
                  Trace Proportion Rank
              2.2436201 0.9311898
## Fitted
```

```
## Residuals 0.1657923 0.0688103
## Total
            2.4094123 1.0000000
                                     1
##
## Eigenvalues
##
                   PC1
##
## Fitted
            2.2436201
## Residuals 0.1657923
## Total
            2.4094123
coef(highLM, test = T)
## Linear Model fit with lm.rrpp
##
## Number of observations: 40
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
## Number of permutations: 1000
## Statistics (distances) of coefficients with 95 percent confidence intervals,
## effect sizes, and probabilities of exceeding observed values based on
## 1000 random permutations using RRPP
##
##
                                 d.obs UCL (95%)
                                                          Zd Pr(>d)
## (Intercept)
                             0.9718000 2.8583704 -1.03660693 0.852
## Block
                            0.3138000 0.6898212 0.18482158 0.372
## FungusCer
                            1.0423000 1.7033771 0.51606112 0.292
## FungusCok
                            2.8819000 3.5618688 0.56062421
                           1.0827667 2.5505618 -0.05217995 0.470
## FungusCtrl
## FungusNig
                            2.5081667 2.5226141 1.82220700
## FungusPen
                            0.9413333 2.3718940 -0.07297045 0.467
## FungusPod
                            1.8891000 2.6749685 0.92632436
                                                             0.198
## FungusPre
                            1.0948000 1.9614740 0.48818761
                                                             0.285
## FungusXyl
                            1.1520000 2.5970673 0.14132966
                                                             0.359
                            3.0766333 3.2160129 1.51609523
## AgeYoung
                                                             0.067
## Block:FungusCtrl
                            0.3293500 1.0469873 -0.31667387
                                                              0.535
## Block:FungusNig
                            1.0558000 1.0394326 1.93972982 0.046
## Block:FungusPen
                            0.3389500 1.0211341 -0.26347545
                                                             0.520
## Block:FungusPod
                            0.8227000 1.4470045 0.47080656
                                                             0.286
## Block:FungusPre
                            0.0753000 1.2622658 -1.26974634 0.936
## Block:FungusXyl
                            0.4171000 1.6212900 -0.49908304 0.613
## Block:AgeYoung
                            0.9157500 1.0320887 1.68338472 0.070
## FungusCer:AgeYoung
                            0.4449167 2.6884392 -1.00424383
                                                             0.816
## FungusCok:AgeYoung
                            0.3173833 1.9478067 -0.79718639 0.760
## FungusCtrl:AgeYoung
                             1.2420667 3.7434431 -0.45354092
## FungusNig:AgeYoung
                             1.8611000 3.8347514 0.13132629
                                                             0.412
## FungusPen:AgeYoung
                             0.4409667 2.6500513 -0.83491769
                                                             0.767
## FungusPod:AgeYoung
                             0.6755667 6.3330293 -1.59386532 0.944
## FungusPre:AgeYoung
                             1.8436833 2.8767622 0.79924471
## FungusXyl:AgeYoung
                             2.3386333 3.5882666 0.78755615
                                                             0.197
## Block:FungusCtrl:AgeYoung 0.1244500 1.2142553 -1.07014865
                                                              0.862
## Block:FungusNig:AgeYoung 1.6630000 1.3029613 2.83789016
                                                             0.018
## Block:FungusPen:AgeYoung 0.2330000 1.2264443 -0.75102567 0.732
```

```
## Block:FungusPod:AgeYoung 1.5442500 2.0239529 1.11457912 0.146
## Block:FungusPre:AgeYoung 1.0738000 1.2646536 1.42148600 0.104
## Block:FungusXyl:AgeYoung 1.4224500 2.1002427 0.86848907 0.189
lowLM <- lm.rrpp(Chl.b ~ Block * Fungus * Age, data = low, SS.type = "III",
    print.progress = F)
##
## Warning: Because variables in the linear model are redundant,
## the linear model design has been truncated (via QR decomposition).
## Original X columns: 36
## Final X columns (rank): 34
## Check coefficients or degrees of freedom in ANOVA to see changes.
summary(lowLM)
## Linear Model fit with lm.rrpp
##
## Number of observations: 46
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
## Number of permutations: 1000
## Full Model Analysis of Variance
##
##
                        Df Residual Df
                                             SS Residual SS
                                                                   Rsq
## Block * Fungus * Age 33
                                    12 183.9456
                                                   24.99044 0.8803919 2.676596
##
                        Z (from F)
                                       Pr(>F)
## Block * Fungus * Age 2.057026 0.02457143
##
## Redundancy Analysis (PCA on fitted values and residuals)
##
##
                Trace Proportion Rank
## Fitted
             4.087680 0.8803919
## Residuals 0.555343 0.1196081
                                    1
## Total
            4.643023 0.9999999
                                    1
##
## Eigenvalues
##
##
                  PC1
## Fitted
             4.087680
## Residuals 0.555343
## Total
            4.643023
coef(lowLM, test = T)
##
## Linear Model fit with lm.rrpp
##
## Number of observations: 46
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
```

```
## Number of permutations: 1000
##
## Statistics (distances) of coefficients with 95 percent confidence intervals,
## effect sizes, and probabilities of exceeding observed values based on
  1000 random permutations using RRPP
##
                                 d.obs UCL (95%)
##
                                                          Zd Pr(>d)
## (Intercept)
                             9.7034333 8.969915
                                                 2.28549354 0.020
## Block
                             2.7120500 2.096513 2.93023576
                                                             0.007
## FungusCer
                             3.7832333 4.811233 1.28983593
                                                             0.116
## FungusCok
                             4.1255333 5.680391 0.99550908 0.175
## FungusCtrl
                             3.7724667 5.120217
                                                 1.01606240 0.157
## FungusNig
                             1.3687333 5.163887 -0.41968895 0.574
## FungusPen
                             4.7273333 4.621536 1.99972450 0.044
## FungusPod
                             3.1066000 6.108060 0.20132282 0.384
## FungusPre
                             5.9124000 4.367254
                                                 2.98749742
                                                             0.011
## FungusXyl
                             0.6566833 3.774410 -0.80998918 0.744
## AgeYoung
                             6.7049333 4.886029 3.05596826 0.011
## Block:FungusCer
                             1.3418000 2.254783 0.60001739 0.254
## Block:FungusCok
                             1.2609500 3.573562 -0.22171967
                                                             0.519
## Block:FungusCtrl
                             1.2197000 3.018750 -0.18580118 0.514
                             0.6864500 2.404786 -0.35675722 0.540
## Block:FungusNig
## Block:FungusPen
                             1.3535000 3.028536 -0.03837009 0.465
## Block:FungusPod
                             0.7918000 3.711903 -1.13033119
                                                             0.859
## Block:FungusPre
                             1.6565000 3.030321 0.31874466 0.363
## Block:AgeYoung
                             4.5699500 2.540569 4.57087308 0.001
## FungusCer:AgeYoung
                             5.2040333 7.518396 0.96732553 0.174
## FungusCok:AgeYoung
                             7.7240333 8.049198 1.75260860 0.060
## FungusCtrl:AgeYoung
                             7.8890333 7.342928 2.13789398 0.034
## FungusNig:AgeYoung
                             3.7873833 6.916607 0.47405782 0.273
## FungusPen:AgeYoung
                             9.7007000 6.193270 3.58918820 0.003
## FungusPod:AgeYoung
                             7.1257000 9.183868 1.10856440 0.147
## FungusPre:AgeYoung
                            10.7553667 6.425508 4.09847422 0.001
## FungusXyl:AgeYoung
                             1.5143833 5.829198 -0.59593503 0.676
## Block:FungusCer:AgeYoung
                             3.5053500
                                       2.980275
                                                 2.44294656
                                                             0.024
## Block:FungusCok:AgeYoung
                             4.1488500 4.603948 1.55253934 0.086
## Block:FungusCtrl:AgeYoung 4.8520000 3.052348 3.86774216 0.002
## Block:FungusNig:AgeYoung
                             2.2579000
                                       3.200773 1.05591140 0.151
## Block:FungusPen:AgeYoung
                             5.1965000
                                        2.924345 4.47088715
                                                             0.001
## Block:FungusPod:AgeYoung
                             5.3413500 2.851511 4.56365093 0.001
## Block:FungusPre:AgeYoung
                             5.7951500 2.868602 5.09678018 0.001
# ratio anovas
highANOVA <- anova(highLM, effect.type = "F", error = c("Residuals",
    "Block:Fungus", "Block:Fungus:Age", "Residuals", "Block:Fungus:Age",
    "Block:Fungus:Age", "Residuals"))
summary(highANOVA)
## Analysis of Variance, using Residual Randomization
## Permutation procedure: Randomization of null model residuals
## Number of permutations: 1000
## Estimation method: Ordinary Least Squares
## Sums of Squares and Cross-products: Type III
## Effect sizes (Z) based on F distributions
```

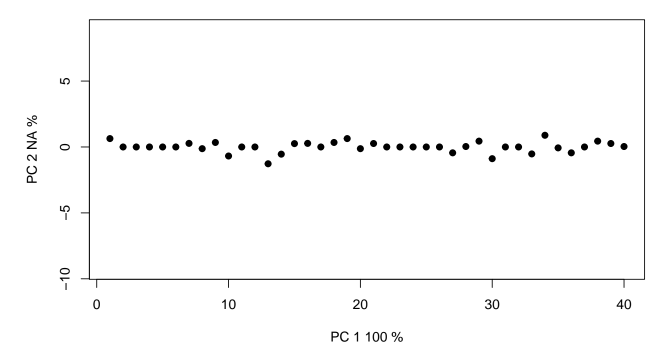
```
##
                                                            Z Pr(>F)
##
                   Df
                          SS
                                  MS
                                                   F
                                          Rsq
                                                                0.585
## Block
                    1 0.197 0.19694 0.002096 0.2437
                    8 5.524 0.69044 0.058781 2.7711
                                                               0.188
## Fungus
                                                      0.87223
## Age
                    1 2.028 2.02836 0.021586 5.0392 1.24671
                                                                0.049 *
                    6 1.495 0.24916 0.015909 0.3083 -1.29905
## Block:Fungus
                    1 0.839 0.83860 0.008924 2.0834
## Block:Age
## Fungus:Age
                    8 2.190 0.27375 0.023306 0.6801 -0.82424
                                                               0.801
## Block:Fungus:Age 6 2.415 0.40251 0.025701 0.4980 -0.78151 0.787
## Residuals
                    8 6.466 0.80824 0.068810
## Total
                   39 93.967
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Call: lm.rrpp(f1 = Chl.b ~ Block * Fungus * Age, SS.type = "III", data = high,
       print.progress = F)
lowANOVA <- anova(lowLM, effect.type = "F", error = c("Residuals",</pre>
    "Block:Fungus", "Block:Fungus:Age", "Residuals", "Block:Fungus:Age",
    "Block:Fungus:Age", "Residuals"))
summary(lowANOVA)
## Analysis of Variance, using Residual Randomization
## Permutation procedure: Randomization of null model residuals
## Number of permutations: 1000
## Estimation method: Ordinary Least Squares
## Sums of Squares and Cross-products: Type III
## Effect sizes (Z) based on F distributions
##
##
                   Df
                           SS
                                                              Z Pr(>F)
                                   MS
                                            Rsq
## Block
                       14.710 14.7104 0.070406 7.0637
                                                                 0.040 *
                      11.605 1.4506 0.055542 2.7036 1.62255
## Fungus
                                                                0.064 .
                        9.633 9.6335 0.046107 2.5619 1.11040
## Age
                    1
                                                                0.070 .
## Block:Fungus
                    7
                        3.756  0.5365  0.017976  0.2576  -1.63483  0.942
## Block:Age
                    1 20.884 20.8844 0.099956 5.5539
                                                                 0.017 *
## Fungus:Age
                    8 19.625 2.4531 0.093927 0.6524 -0.73125 0.784
## Block:Fungus:Age 7 26.322 3.7604 0.125983 1.8057 0.92115 0.167
## Residuals
                   12 24.990 2.0825 0.119608
## Total
                   45 208.936
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Call: lm.rrpp(f1 = Chl.b ~ Block * Fungus * Age, SS.type = "III", data = low,
##
       print.progress = F)
# pairwise
highpw <- pairwise(highLM, groups = high$Fungus)
summary(highpw, confidence = 0.95, stat.table = T)
##
## Pairwise comparisons
##
## Groups: Asp Cer Cok Ctrl Nig Pen Pod Pre Xyl
```

##

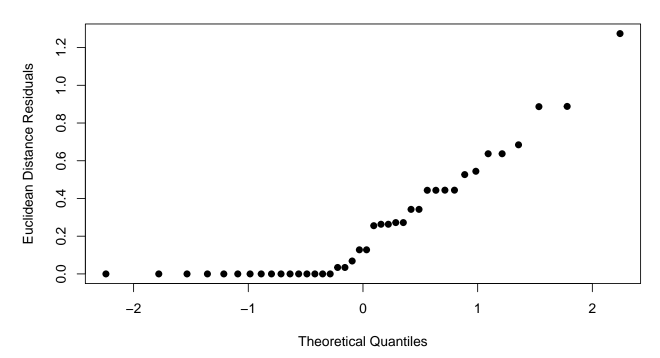
```
## RRPP: 1000 permutations
##
## LS means:
## Vectors hidden (use show.vectors = TRUE to view)
## Pairwise distances between means, plus statistics
                    d UCL (95%)
                                            Z Pr > d
## Asp:Cer 1.40883333 1.9945336 0.0637606595 0.492
## Asp:Cok 2.86728333 3.4661925 -0.0047963567
                                               0.500
## Asp:Ctrl 1.16955000 1.6112341 0.0083118362 0.501
## Asp:Nig 1.12901667 1.5695399 -0.0202340908
## Asp:Pen 0.27595000 0.7537367 -0.1784436293 0.502
## Asp:Pod 2.22305833 2.7169381 -0.0030759276 0.501
## Asp:Pre 1.09615833 1.5899385 0.0004934936 0.493
## Asp:Xyl 0.49590833 0.9758634 0.0048394708 0.481
## Cer:Cok 1.45845000 2.2753518 -0.0573008139
## Cer:Ctrl 0.23928333 0.8477858 -0.4895285835
                                              0.614
## Cer:Nig 0.27981667 0.8912138 -0.3951570849
## Cer:Pen 1.13288333 1.7543083 0.0489657952 0.505
## Cer:Pod 0.81422500 1.5207386 -0.0788821420
## Cer:Pre 0.31267500 0.9615748 -0.3437307668
## Cer:Xyl 0.91292500 1.5369945 0.0151171726
## Cok:Ctrl 1.69773333 2.2978103 -0.0105851449
                                               0.514
## Cok:Nig 1.73826667 2.3818621 0.0092136674 0.506
## Cok:Pen 2.59133333 3.2151287 -0.0190620434 0.517
## Cok:Pod 0.64422500 1.3009886 -0.0612888345 0.505
## Cok:Pre 1.77112500 2.4287079 -0.0046510510
                                              0.510
## Cok:Xyl 2.37137500 3.0356656 -0.0435889705
                                              0.517
## Ctrl:Nig 0.04053333 0.5294233 -1.0916038475
## Ctrl:Pen 0.89360000 1.3612861 -0.0121300105
## Ctrl:Pod 1.05350833 1.5421654 -0.0106100482
                                               0.518
## Ctrl:Pre 0.07339167 0.6205608 -0.9594614150
                                              0.814
## Ctrl:Xyl 0.67364167 1.1572582 -0.0523155704
## Nig:Pen 0.85306667 1.3181976 -0.0397347478
                                              0.519
## Nig:Pod 1.09404167 1.5924091 0.0141541023
## Nig:Pre 0.03285833 0.6157437 -1.2293129879
## Nig:Xyl 0.63310833 1.1638414 -0.0819411492 0.532
## Pen:Pod 1.94710833 2.4647373 -0.0216389701
                                              0.502
## Pen:Pre 0.82020833 1.3278977 -0.0181917671
## Pen:Xyl 0.21995833 0.7232532 -0.3870178122 0.572
## Pod:Pre 1.12690000 1.6920741 -0.0031561014 0.491
## Pod:Xyl 1.72715000 2.2883888 -0.0491816153 0.539
## Pre:Xyl 0.60025000 1.1499453 -0.0721971246 0.521
highpw2 <- pairwise(highLM, groups = high$Age)
summary(highpw2, confidence = 0.95, stat.table = T)
##
## Pairwise comparisons
##
## Groups: Old Young
## RRPP: 1000 permutations
##
## LS means:
```

```
## Vectors hidden (use show.vectors = TRUE to view)
##
## Pairwise distances between means, plus statistics
##
                   d UCL (95%)
                                       Z Pr > d
## Old:Young 2.190885 2.435993 0.08979591 0.452
lowpw <- pairwise(lowLM, groups = low$Fungus)</pre>
summary(lowpw, confidence = 0.95, stat.table = T)
##
## Pairwise comparisons
##
## Groups: Asp Cer Cok Ctrl Nig Pen Pod Pre Xyl
##
## RRPP: 1000 permutations
##
## LS means:
## Vectors hidden (use show.vectors = TRUE to view)
## Pairwise distances between means, plus statistics
                   d UCL (95%)
## Asp:Cer 2.0029667 2.942593 5.551760e-02 0.469
## Asp:Cok 1.2701917 2.369217 3.901236e-03 0.490
## Asp:Ctrl 2.2405500 3.266076 1.441241e-02 0.502
## Asp:Nig 0.3600417 1.512806 -5.837500e-01 0.646
## Asp:Pen 2.3664833 3.342638 2.283370e-02 0.486
## Asp:Pod 3.3015000 4.270812 6.647641e-02 0.481
## Asp:Pre 3.0168667 3.990030 1.870943e-02 0.487
## Asp:Xyl 0.5275833 2.076821 -4.892711e-01 0.623
## Cer:Cok 0.7327750 1.861699 -1.587278e-01 0.504
## Cer:Ctrl 0.2375833 1.328923 -7.667723e-01 0.742
## Cer:Nig 1.6429250 2.771172 1.270720e-02 0.504
## Cer:Pen 0.3635167 1.370296 -4.996732e-01 0.622
## Cer:Pod 1.2985333 2.343280 -3.979415e-03 0.491
## Cer:Pre 1.0139000 2.040683 -8.423441e-02 0.524
## Cer:Xyl 2.5305500 4.010692 -5.448366e-05
## Cok:Ctrl 0.9703583 2.064697 -9.973555e-02
                                             0.514
## Cok:Nig 0.9101500 2.057411 -1.106110e-01
                                             0.505
## Cok:Pen 1.0962917 2.213880 -5.475120e-02 0.509
## Cok:Pod 2.0313083 3.172132 2.763028e-02 0.477
## Cok:Pre 1.7466750 2.919395 -1.541431e-02 0.503
## Cok:Xyl 1.7977750 3.477338 -2.837168e-02 0.486
## Ctrl:Nig 1.8805083 3.010984 -1.697763e-02 0.506
## Ctrl:Pen 0.1259333 1.232834 -1.013516e+00 0.842
## Ctrl:Pod 1.0609500 1.987635 2.012947e-02 0.481
## Ctrl:Pre 0.7763167 1.758984 -1.125456e-01 0.516
## Ctrl:Xyl 2.7681333 4.353823 -2.837150e-02 0.500
## Nig:Pen 2.0064417 3.131585 -1.016671e-02 0.503
## Nig:Pod 2.9414583 4.058314 2.983836e-02
## Nig:Pre 2.6568250 3.885838 -1.353280e-02 0.491
## Nig:Xyl 0.8876250 2.485887 -2.303612e-01
## Pen:Pod 0.9350167 1.883259 3.707117e-03 0.473
## Pen:Pre 0.6503833 1.658783 -1.596798e-01 0.512
## Pen:Xyl 2.8940667 4.414800 -2.433315e-02 0.498
## Pod:Pre 0.2846333 1.233337 -5.811612e-01 0.652
```

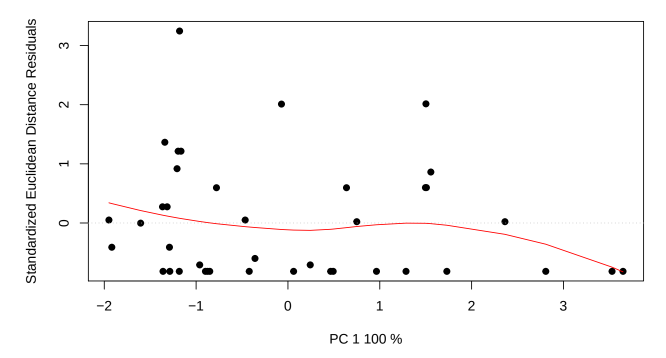
```
## Pod:Xyl 3.8290833 5.362483 6.367113e-03 0.475
## Pre:Xyl 3.5444500 5.145378 -2.662861e-02 0.503
lowpw2 <- pairwise(lowLM, groups = low$Age)</pre>
summary(lowpw2, confidence = 0.95, stat.table = T)
##
## Pairwise comparisons
##
## Groups: Old Young
## RRPP: 1000 permutations
##
## LS means:
## Vectors hidden (use show.vectors = TRUE to view)
## Pairwise distances between means, plus statistics
##
                    d UCL (95%)
                                          Z Pr > d
## Old:Young 1.035035 1.547157 0.008272272 0.502
# residuals vs fitted values (homoscedasticity check)
hdiagnostics <- plot(highLM, type = "diagnostics")</pre>
```



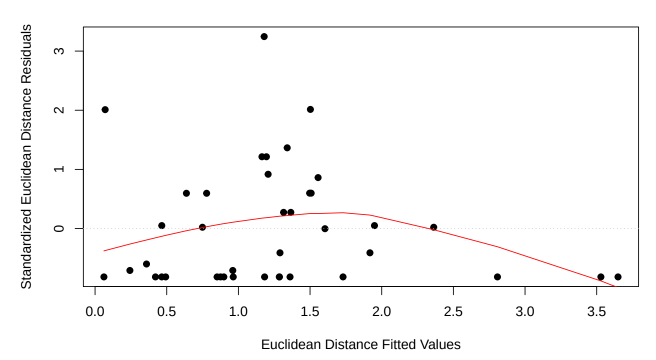
# Q-Q plot

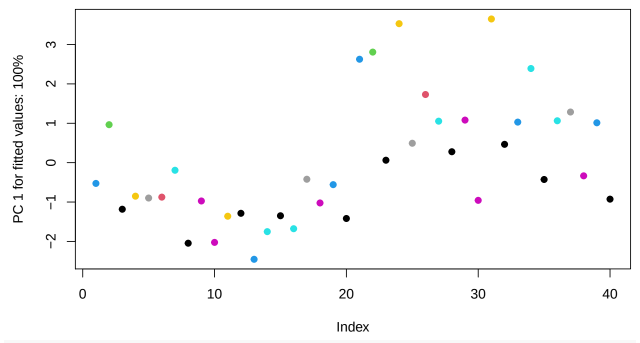


# Residuals vs. PC 1 fitted

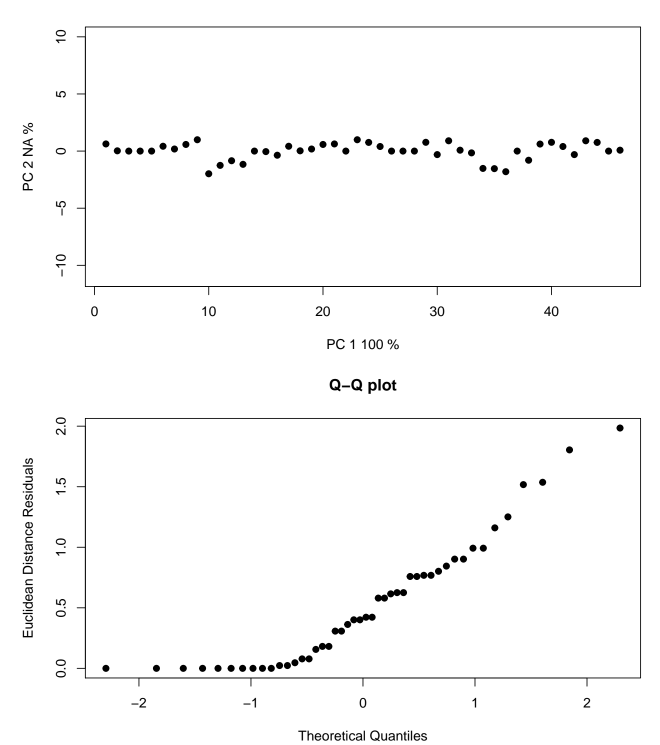


### Residuals vs. Fitted

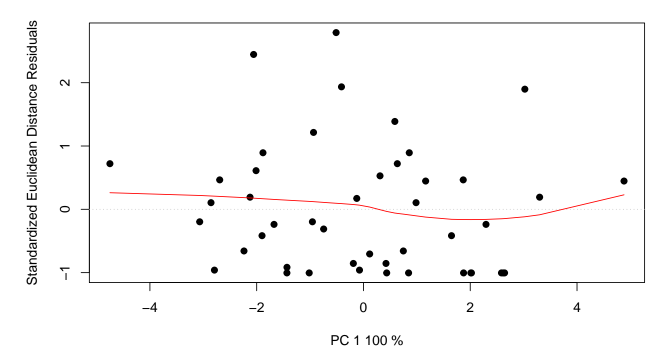




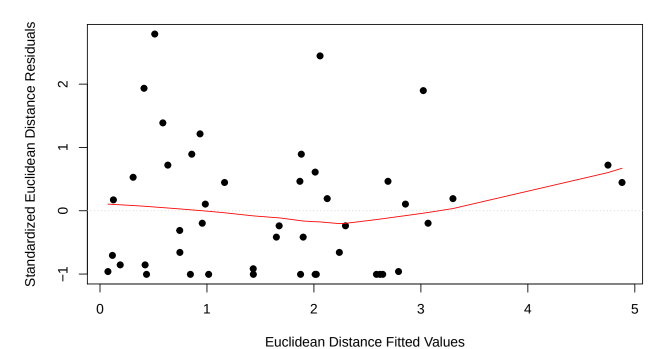
# residuals vs fitted values (homoscedasticity check)
ldiagnostics <- plot(lowLM, type = "diagnostics")</pre>



## Residuals vs. PC 1 fitted



## Residuals vs. Fitted



```
ဖ
PC 1 for fitted values: 100%
     \sim
     0
     7
           0
                           10
                                             20
                                                              30
                                                                               40
                                                 Index
## Total linear models and coefficient tests
highLM <- lm.rrpp(Total ~ Block * Fungus * Age, data = high,
    SS.type = "III", print.progress = F)
##
## Warning: Because variables in the linear model are redundant,
## the linear model design has been truncated (via QR decomposition).
## Original X columns: 36
## Final X columns (rank): 32
## Check coefficients or degrees of freedom in ANOVA to see changes.
summary(highLM)
##
## Linear Model fit with lm.rrpp
## Number of observations: 40
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
## Number of permutations: 1000
## Full Model Analysis of Variance
##
##
                         Df Residual Df
                                               SS Residual SS
                                                                     Rsq
## Block * Fungus * Age 31
                                       8 221.9666
                                                      22.53375 0.9078376 2.542041
                         Z (from F)
##
                                         Pr(>F)
## Block * Fungus * Age
                            1.50047 0.06657143
##
##
## Redundancy Analysis (PCA on fitted values and residuals)
##
##
                Trace Proportion Rank
## Fitted
             5.691452 0.9078375
```

```
## Residuals 0.577788 0.0921624
## Total
            6.269241 1.0000000
                                    1
##
## Eigenvalues
##
                  PC1
##
## Fitted
             5.691452
## Residuals 0.577788
## Total
             6.269241
coef(highLM, test = T)
## Linear Model fit with lm.rrpp
##
## Number of observations: 40
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
## Number of permutations: 1000
## Statistics (distances) of coefficients with 95 percent confidence intervals,
## effect sizes, and probabilities of exceeding observed values based on
## 1000 random permutations using RRPP
##
##
                                 d.obs UCL (95%)
                                                         Zd Pr(>d)
## (Intercept)
                             2.4605000 5.920830 -0.9732179 0.850
## Block
                             0.3327000 1.325547 -0.5496670
                                                             0.651
## FungusCer
                             1.9381000 3.191372 0.4908956
                                                             0.298
## FungusCok
                             6.0044000
                                        7.281553 0.5379335
                                                             0.279
                                       4.482527 -0.2560890
## FungusCtrl
                             1.5441000
                                                             0.531
## FungusNig
                             5.0907000
                                        5.022310
                                                 1.8473197
## FungusPen
                             2.1842000
                                       4.764053 0.1040508
                                                             0.404
## FungusPod
                             3.0680000
                                        4.986713 0.6383150
                                                             0.250
## FungusPre
                             2.3511500
                                        3.714938 0.7178551
                                                             0.238
## FungusXyl
                             2.1986000
                                       4.881486
                                                  0.1887340
                                                             0.340
                                        4.853749
                                                             0.076
## AgeYoung
                             4.4501333
                                                 1.5389731
## Block:FungusCtrl
                             0.4002000
                                        1.850631 -0.6059016
                                                             0.657
## Block:FungusNig
                             2.0625000
                                       1.979667 2.0092496
                                                             0.039
## Block:FungusPen
                             0.6948000
                                       1.937021 -0.1501771
                                                             0.486
## Block:FungusPod
                             1.0500000
                                        2.722623 -0.1071264
                                                             0.473
## Block:FungusPre
                             0.1543500
                                        2.258982 -1.2056489
                                                             0.912
## Block:FungusXyl
                             0.7074000
                                       2.973520 -0.5584964
                                                             0.632
## Block:AgeYoung
                             1.3986000 1.766544 1.3331748
                                                             0.114
## FungusCer:AgeYoung
                             0.5794667
                                        4.609162 -1.0592039
                                                             0.835
## FungusCok:AgeYoung
                             1.5839333 3.218044 0.2225087
                                                             0.368
## FungusCtrl:AgeYoung
                             2.3970000
                                       6.852218 -0.3590399
                                                             0.582
## FungusNig:AgeYoung
                             4.0526667
                                        6.681245 0.5568047
                                                             0.285
## FungusPen:AgeYoung
                             1.3145333
                                       4.967385 -0.5153642
                                                             0.617
## FungusPod:AgeYoung
                             1.4089667 11.155901 -1.3660827
                                                             0.902
## FungusPre:AgeYoung
                             3.6231833
                                       5.155174 0.9859370
                                                             0.158
## FungusXyl:AgeYoung
                             4.2304333
                                        6.711787 0.7340295
                                                             0.207
## Block:FungusCtrl:AgeYoung 0.1023000
                                        2.285465 -1.2404952
                                                             0.943
## Block:FungusNig:AgeYoung 3.1697000
                                        2.424762 2.9364990
                                                             0.014
## Block:FungusPen:AgeYoung 0.5183000
                                        2.332142 -0.6313884
                                                             0.674
```

```
## Block:FungusPod:AgeYoung 2.2897000 3.747401 0.6164059 0.238
## Block:FungusPre:AgeYoung 1.8753500 2.349132 1.2477925 0.126
## Block:FungusXyl:AgeYoung 2.4271000 3.890555 0.6786493 0.227
lowLM <- lm.rrpp(Total ~ Block * Fungus * Age, data = low, SS.type = "III",
    print.progress = F)
##
## Warning: Because variables in the linear model are redundant,
## the linear model design has been truncated (via QR decomposition).
## Original X columns: 36
## Final X columns (rank): 34
## Check coefficients or degrees of freedom in ANOVA to see changes.
summary(lowLM)
## Linear Model fit with lm.rrpp
##
## Number of observations: 46
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
## Number of permutations: 1000
## Full Model Analysis of Variance
##
##
                        Df Residual Df
                                             SS Residual SS
                                                                  Rsq
## Block * Fungus * Age 33
                                    12 314.3371
                                                   57.59527 0.8451458 1.984614
##
                        Z (from F)
                                       Pr(>F)
## Block * Fungus * Age 1.464465 0.08057143
##
## Redundancy Analysis (PCA on fitted values and residuals)
##
##
                Trace Proportion Rank
## Fitted
             6.985269 0.8451459
## Residuals 1.279895 0.1548542
                                    1
## Total
            8.265164 1.0000000
                                    1
##
## Eigenvalues
##
##
                  PC1
## Fitted
            6.985269
## Residuals 1.279895
## Total
            8.265164
coef(lowLM, test = T)
##
## Linear Model fit with lm.rrpp
##
## Number of observations: 46
## Number of dependent variables: 1
## Data space dimensions: 1
## Sums of Squares and Cross-products: Type III
```

```
## Number of permutations: 1000
##
## Statistics (distances) of coefficients with 95 percent confidence intervals,
## effect sizes, and probabilities of exceeding observed values based on
## 1000 random permutations using RRPP
##
                                d.obs UCL (95%)
##
                                                         Zd Pr(>d)
## (Intercept)
                            14.046500 14.147632 1.59282901 0.058
## Block
                             3.426250 3.290589 1.79950611
                                                            0.041
## FungusCer
                             6.577567 7.392581 1.47825353
                                                           0.087
## FungusCok
                             1.492200 7.273627 -0.61949886 0.663
## FungusCtrl
                             1.270733 6.319047 -0.70267111
                                                            0.714
## FungusNig
                             2.480000 5.762513 0.12206610 0.368
                             3.122967 6.178017 0.32631733 0.326
## FungusPen
## FungusPod
                             4.613567 6.470651 1.02121148 0.170
## FungusPre
                             5.691400 5.707320
                                                1.95829327
                                                            0.052
## FungusXyl
                             2.848150 4.217050 0.85427183 0.179
## AgeYoung
                             5.608600 5.811837 1.81566252 0.059
## Block:FungusCer
                             2.231300 2.607066 1.40979606 0.097
## Block:FungusCok
                             0.322250 4.824937 -1.15350193 0.902
## Block:FungusCtrl
                             0.010650 3.765990 -1.46488265 0.999
## Block:FungusNig
                             0.964250 2.661146 -0.14185727 0.477
                             0.351150 4.056759 -1.42661488 0.939
## Block:FungusPen
                             0.971500 2.845590 -0.16662065 0.485
## Block:FungusPod
## Block:FungusPre
                             0.982450 3.986420 -0.83398856 0.769
## Block:AgeYoung
                             4.185050 3.101306 3.02482067 0.011
## FungusCer:AgeYoung
                             6.105833 8.416641 1.03123275 0.172
## FungusCok:AgeYoung
                             4.642800 10.040343 0.21199777
                                                           0.354
## FungusCtrl:AgeYoung
                             3.677733 9.053234 0.01209651 0.435
## FungusNig:AgeYoung
                             4.948250 7.966459 0.66847968 0.234
## FungusPen:AgeYoung
                             8.437000
                                      7.970256
                                                 2.17918532 0.036
## FungusPod:AgeYoung
                             4.244600 8.901833 0.27889328 0.329
## FungusPre:AgeYoung
                            10.636200 7.897802 3.14051586 0.009
## FungusXyl:AgeYoung
                             3.552350 4.956430
                                                0.95239210 0.186
## Block:FungusCer:AgeYoung
                             3.861000
                                       3.629037
                                                 2.13877657
## Block:FungusCok:AgeYoung
                             2.405350 5.683113 0.05617713 0.407
## Block:FungusCtrl:AgeYoung 2.843800 3.739075 1.18005737 0.137
## Block:FungusNig:AgeYoung
                             2.810900 3.597535 1.23553528 0.125
## Block:FungusPen:AgeYoung
                             4.568300
                                       3.599418
                                                 2.77233699 0.019
## Block:FungusPod:AgeYoung
                             2.859250 3.692270 1.21155396 0.133
## Block:FungusPre:AgeYoung
                             5.663050 3.460660 3.91974843 0.001
# ratio anovas
highANOVA <- anova(highLM, effect.type = "F", error = c("Residuals",
    "Block:Fungus", "Block:Fungus:Age", "Residuals", "Block:Fungus:Age",
    "Block:Fungus:Age", "Residuals"))
summary(highANOVA)
## Analysis of Variance, using Residual Randomization
## Permutation procedure: Randomization of null model residuals
## Number of permutations: 1000
## Estimation method: Ordinary Least Squares
## Sums of Squares and Cross-products: Type III
## Effect sizes (Z) based on F distributions
```

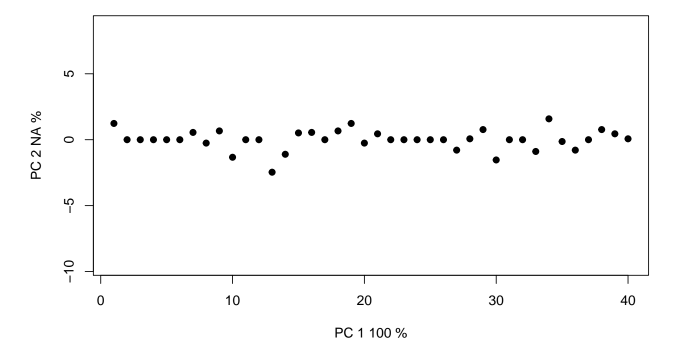
```
##
                                                            Z Pr(>F)
##
                   Df
                           SS
                                  MS
                                          Rsq
                                                   F
                        0.221 0.2214 0.000905 0.0786
## Block
                                                               0.693
## Fungus
                    8 24.327 3.0408 0.099495 3.3018 0.92054
                                                               0.182
## Age
                    1
                        4.244 4.2436 0.017356 3.1404
                                                                0.087
## Block:Fungus
                      5.526 0.9210 0.022600 0.3270 -1.18349
                                                               0.889
                    6
## Block:Age
                       1.956 1.9561 0.008000 1.4475
                    1
## Fungus:Age
                    8
                        8.799 1.0999 0.035989 0.8140 -0.56109
                                                               0.709
## Block:Fungus:Age 6
                       8.108 1.3513 0.033161 0.4797 -0.82034 0.795
## Residuals
                    8 22.534 2.8167 0.092162
## Total
                   39 244.500
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Call: lm.rrpp(f1 = Total ~ Block * Fungus * Age, SS.type = "III", data = high,
       print.progress = F)
lowANOVA <- anova(lowLM, effect.type = "F", error = c("Residuals",</pre>
    "Block:Fungus", "Block:Fungus:Age", "Residuals", "Block:Fungus:Age",
    "Block:Fungus:Age", "Residuals"))
summary(lowANOVA)
## Analysis of Variance, using Residual Randomization
## Permutation procedure: Randomization of null model residuals
## Number of permutations: 1000
## Estimation method: Ordinary Least Squares
## Sums of Squares and Cross-products: Type III
## Effect sizes (Z) based on F distributions
##
##
                   Df
                          SS
                                                            Z Pr(>F)
                                  MS
                                                   F
                                          Rsq
## Block
                       23.48 23.4784 0.063125 4.8917
                                                               0.059
                      15.97 1.9959 0.042930 1.7917 0.82125
## Fungus
                                                               0.174
                        6.74 6.7407 0.018123 2.4616 1.03071
## Age
## Block:Fungus
                    7
                        7.80 1.1140 0.020966 0.2321 -1.57930
                                                               0.949
## Block:Age
                    1 17.51 17.5146 0.047091 6.3962
                                                               0.007 **
## Fungus:Age
                    8 15.32 1.9152 0.041195 0.6994 -0.75229
                                                               0.789
## Block:Fungus:Age 7 19.17 2.7383 0.051536 0.5705 -0.64515
                                                               0.749
## Residuals
                   12 57.60 4.7996 0.154854
## Total
                   45 371.93
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Call: lm.rrpp(f1 = Total ~ Block * Fungus * Age, SS.type = "III", data = low,
##
       print.progress = F)
# pairwise
highpw <- pairwise(highLM, groups = high$Fungus)
summary(highpw, confidence = 0.95, stat.table = T)
##
## Pairwise comparisons
##
## Groups: Asp Cer Cok Ctrl Nig Pen Pod Pre Xyl
```

##

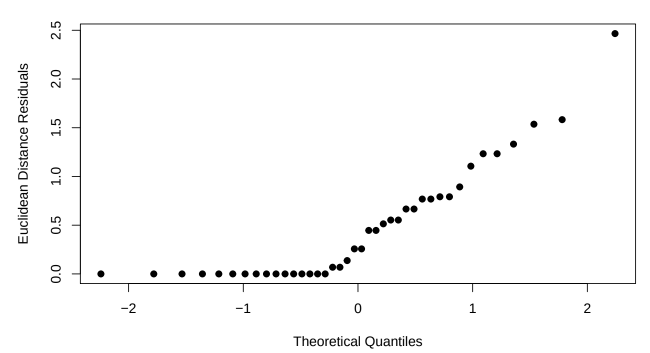
```
## RRPP: 1000 permutations
##
## LS means:
## Vectors hidden (use show.vectors = TRUE to view)
## Pairwise distances between means, plus statistics
                     d UCL (95%)
                                             Z Pr > d
## Asp:Cer 2.594433333 3.6721392 0.0623334604 0.488
## Asp:Cok 5.579033333 6.6767439 -0.0072307614 0.511
## Asp:Ctrl 2.044500000 2.8709959 0.0044865681
## Asp:Nig 2.109066667 2.9224115 -0.0199041407
## Asp:Pen 0.655633333 1.5546638 -0.0795169852
                                                0.495
## Asp:Pod 4.098058333 4.9833487 -0.0013203580
                                                0.505
## Asp:Pre 2.106208333 3.0177705 -0.0016048283
                                               0.507
## Asp:Xyl 1.025908333 1.9013216 0.0227288228
                                                0.481
## Cer:Cok 2.984600000 4.5300312 -0.0575604669
                                                0.523
## Cer:Ctrl 0.549933333 1.6776245 -0.3630493074
                                                0.572
## Cer:Nig 0.485366667 1.6374291 -0.4361145401
## Cer:Pen 1.938800000 3.0863939 0.0497056859
                                                0.502
## Cer:Pod 1.503625000 2.8260292 -0.0773612229
                                                0.523
## Cer:Pre 0.488225000 1.6991273 -0.4574227423
                                                0.612
## Cer:Xyl 1.568525000 2.7271735 0.0050097166
## Cok:Ctrl 3.534533333 4.6121712 -0.0102291978
                                                0.508
## Cok:Nig 3.469966667 4.6631701 0.0067391694
                                                0.512
## Cok:Pen 4.923400000 6.0655360 -0.0170282585
                                               0.511
## Cok:Pod 1.480975000 2.7002623 -0.0335369928
                                                0.505
## Cok:Pre 3.472825000 4.6739885 -0.0052763132
                                                0.508
## Cok:Xyl 4.553125000 5.7733336 -0.0449782246
                                                0.520
## Ctrl:Nig 0.064566667 0.9993244 -1.0898908451
                                                0.895
## Ctrl:Pen 1.388866667 2.2510710 -0.0113472493
                                                0.494
## Ctrl:Pod 2.053558333 2.9398952 -0.0053672177
                                                0.501
## Ctrl:Pre 0.061708333 1.0887264 -1.1653588504
                                                0.908
## Ctrl:Xyl 1.018591667 1.9314309 -0.0715412761
                                                0.521
## Nig:Pen 1.453433333 2.3123537 -0.0332901209
                                                0.513
## Nig:Pod 1.988991667 2.9118283 0.0155649776
## Nig:Pre 0.002858333 1.1281646 -1.3947397396
                                               0.996
## Nig:Xyl 1.083158333 2.0667876 -0.0894864502 0.534
## Pen:Pod 3.442425000 4.3985404 -0.0143118744
                                                0.504
## Pen:Pre 1.450575000 2.4117002 -0.0155429096
## Pen:Xyl 0.370275000 1.3041777 -0.4362776834
                                               0.602
## Pod:Pre 1.991850000 3.0452676 0.0002494841
## Pod:Xyl 3.072150000 4.1144333 -0.0463846136 0.532
## Pre:Xyl 1.080300000 2.1212680 -0.0767809966 0.519
highpw2 <- pairwise(highLM, groups = high$Age)
summary(highpw2, confidence = 0.95, stat.table = T)
## Pairwise comparisons
##
## Groups: Old Young
## RRPP: 1000 permutations
##
## LS means:
```

```
## Vectors hidden (use show.vectors = TRUE to view)
##
## Pairwise distances between means, plus statistics
##
                   d UCL (95%)
                                        Z Pr > d
## Old:Young 3.012115 3.462799 0.08881142 0.451
lowpw <- pairwise(lowLM, groups = low$Fungus)</pre>
summary(lowpw, confidence = 0.95, stat.table = T)
##
## Pairwise comparisons
##
## Groups: Asp Cer Cok Ctrl Nig Pen Pod Pre Xyl
##
## RRPP: 1000 permutations
##
## LS means:
## Vectors hidden (use show.vectors = TRUE to view)
## Pairwise distances between means, plus statistics
                   d UCL (95%)
## Asp:Cer 2.9230500 4.105414 0.085177960 0.463
## Asp:Cok 0.7913250 2.126375 -0.172584235
                                             0.514
## Asp:Ctrl 2.2543667 3.485428 0.020099936 0.503
## Asp:Nig 0.8882750 2.303592 -0.149355519
## Asp:Pen 2.7704667
                      3.991072 0.041620060
                                             0.478
## Asp:Pod 3.4075167 4.603394 0.085877001 0.457
## Asp:Pre 4.0714500 5.285035 0.026563125
## Asp:Xyl 0.2617500 2.253667 -0.956286762
                                             0.824
## Cer:Cok 2.1317250 3.426954 0.020790746
## Cer:Ctrl 0.6686833 1.855307 -0.223783754
                                             0.508
## Cer:Nig 2.0347750 3.312538 0.021003452
## Cer:Pen 0.1525833 1.489768 -0.971200177
                                             0.842
## Cer:Pod 0.4844667
                     1.732166 -0.461654419
                                             0.620
## Cer:Pre 1.1484000 2.418925 -0.127671539
                                            0.530
## Cer:Xyl 3.1848000 4.906868 -0.012847885
                                             0.496
## Cok:Ctrl 1.4630417
                      2.853609 -0.077576556
                                             0.532
## Cok:Nig 0.0969500 1.775473 -1.153234415
                                             0.922
## Cok:Pen 1.9791417 3.252532 -0.025517105
                                            0.524
## Cok:Pod 2.6161917 3.983672 0.019661318
                                            0.490
## Cok:Pre 3.2801250 4.638539 -0.031645138
                                             0.509
## Cok:Xyl 1.0530750 2.959145 -0.276003399
                                            0.547
## Ctrl:Nig 1.3660917 2.855899 -0.072110461
## Ctrl:Pen 0.5161000 1.806311 -0.402903536
                                            0.581
## Ctrl:Pod 1.1531500 2.306431 0.016278891
                                             0.474
## Ctrl:Pre 1.8170833 3.055947 -0.001732709 0.487
## Ctrl:Xyl 2.5161167 4.380340 -0.065556435
## Nig:Pen 1.8821917
                      3.211277 -0.022984604
                                            0.506
## Nig:Pod 2.5192417
                      3.864774 0.030450520
                                             0.499
## Nig:Pre 3.1831750 4.548475 -0.023441309
                                             0.500
## Nig:Xyl 1.1500250 3.067854 -0.260194969
                                             0.550
                      1.804311 -0.219321591
## Pen:Pod 0.6370500
                                             0.507
## Pen:Pre 1.3009833 2.571577 -0.040024804
                                             0.496
## Pen:Xyl 3.0322167 4.801769 -0.043255224
                                            0.523
## Pod:Pre 0.6639333 1.905160 -0.309884608 0.551
```

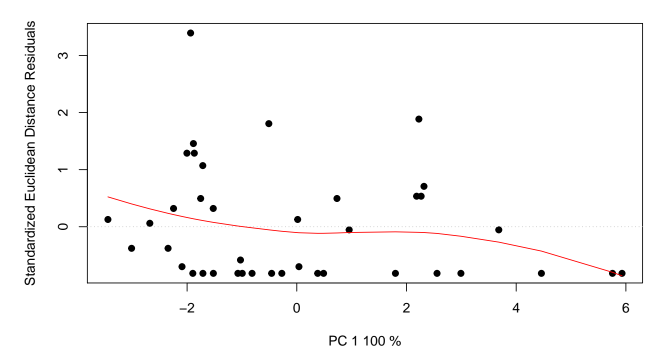
```
## Pod:Xyl 3.6692667 5.457112 -0.009533509 0.504
## Pre:Xyl 4.3332000 6.145788 -0.050869515 0.543
lowpw2 <- pairwise(lowLM, groups = low$Age)</pre>
summary(lowpw2, confidence = 0.95, stat.table = T)
##
## Pairwise comparisons
##
## Groups: Old Young
## RRPP: 1000 permutations
##
## LS means:
## Vectors hidden (use show.vectors = TRUE to view)
## Pairwise distances between means, plus statistics
##
                    d UCL (95%)
                                         Z Pr > d
## Old:Young 1.736996 2.317068 0.04409585 0.489
# residuals vs fitted values (homoscedasticity check)
hdiagnostics <- plot(highLM, type = "diagnostics")</pre>
```



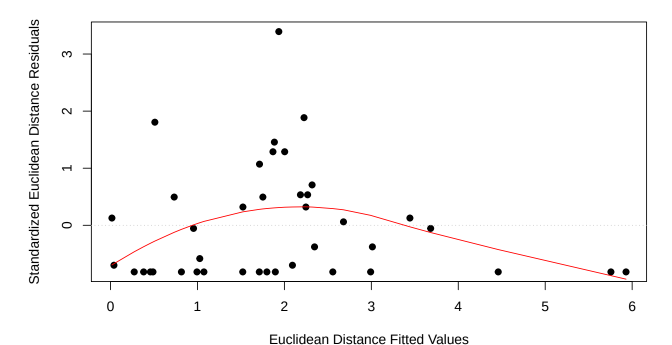


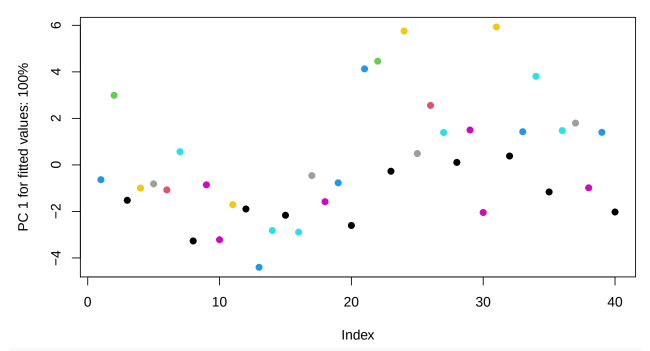


# Residuals vs. PC 1 fitted

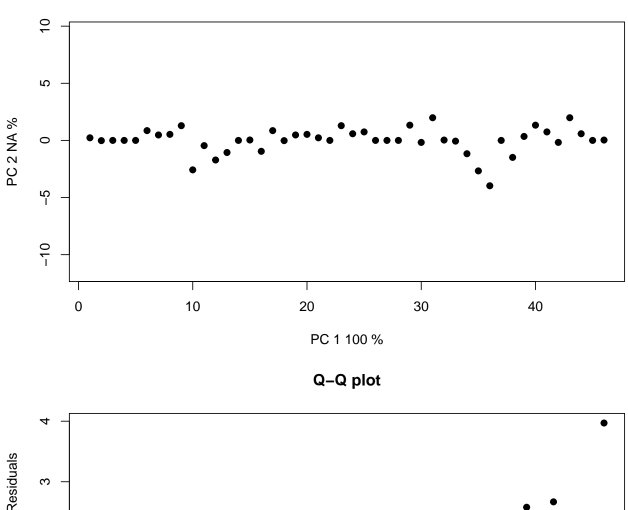


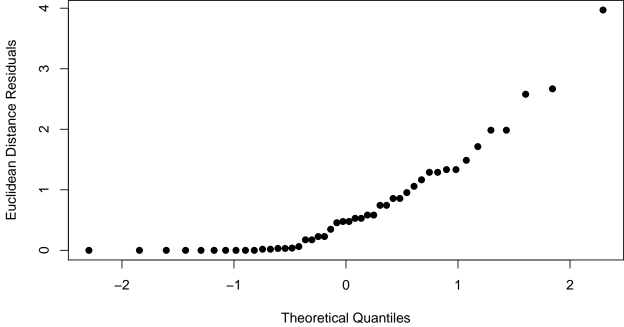
### Residuals vs. Fitted



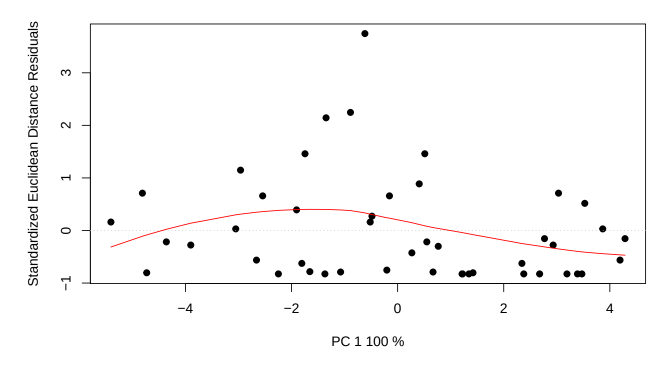


# residuals vs fitted values (homoscedasticity check)
ldiagnostics <- plot(lowLM, type = "diagnostics")</pre>

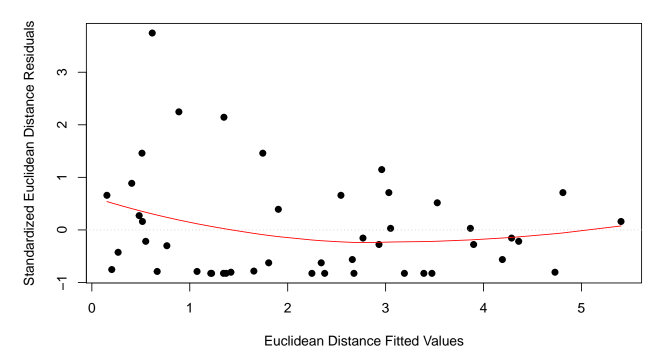


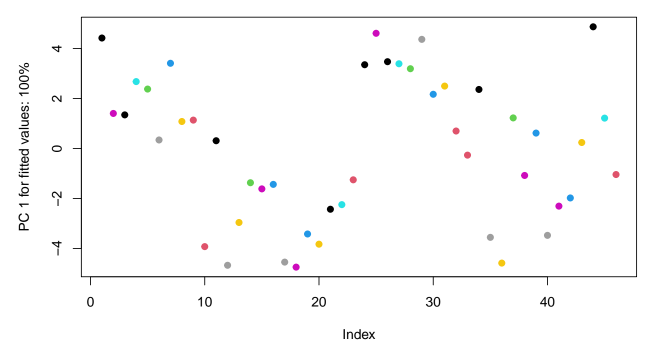


## Residuals vs. PC 1 fitted



## Residuals vs. Fitted





#### 9. Indicator Analysis

```
library(labdsv)
## High water
indicator_high <- indval(high[, 5:8], clustering = high$Fungus,
    numitr = 999, type = "long")

write.csv(cbind(indicator_high$relfrq, indicator_high$relabu,
    indicator_high$indval, data.frame(indicator_high$maxcls),
    data.frame(indicator_high$indcls), data.frame(indicator_high$pval)),
    "~/Box/Summer 2018 TX Endo Field Samples and Analysis/Statistics/Old + Young Chlorophyll/Indicator_
## Low water
indicator_low <- indval(low[, 5:8], clustering = low$Fungus,
    numitr = 999, type = "long")

write.csv(cbind(indicator_low$relfrq, indicator_low$relabu, indicator_low$indval,
    data.frame(indicator_low$maxcls), data.frame(indicator_low$indcls),
    data.frame(indicator_low$pval)), "~/Box/Summer 2018 TX Endo Field Samples and Analysis/Statistics/0</pre>
```