

Appendix S2. Evolutionary drivers of species and community dynamics

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1 Background

In Appendix S1.5, we evaluated the ability of the statistical model to estimate the impact of trait evolution for species abundances. Trait evolution’s impact for species abundances can be modeled as a predictor in different ways. Here we derive the numerical form of the trait x_i that we hypothesize has the most impact on species abundances.

2 Model for population growth, competition, environmental change, and evolution

The growth equation for each species is:

$$N_{i,t+1} = \frac{\hat{W} e^{\frac{-[(\frac{w+(1-h^2)P}{P+w})(E-x_{i,t})]^2}{2(P+w)}}}{1 + \alpha_{ii}N_{i,t} + \alpha_{ij}N_{j,t}} N_{i,t}$$

We simulate population growth under particular parameter values, to generate population dynamics where species interactions, environment, and trait evolution all drive population dynamics.

```
set.seed(42)
# Case 1: Weaker interactions, stronger environment Simulate initial species
# population growth with environment fluctuations
N1.0 <- 10
N2.0 <- 10
alpha.11 <- 0.01
alpha.22 <- 0.01
alpha.12 <- 0.005
alpha.21 <- 0.01
E.0 <- 0.8
x1.0 <- 0.1
x2.0 <- 0.8
P <- 1
```

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```

w <- 2
Wmax <- 2
h2 <- 1
k <- (w + (1 - h2) * P)/(P + w)

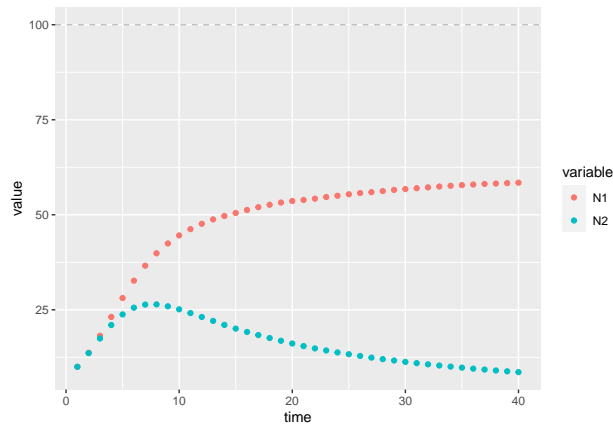
# model function
disc_LV_evol <- function(N1.0, N2.0, alpha.11, alpha.22, alpha.12, alpha.21, E, x1.0,
  x2.0, P, w, Wmax, h2) {
  What <- Wmax * sqrt(w/(P + w))
  r1 <- What * exp(-(((w + (1 - h2) * P)/(P + w)) * (E - x1.0))^2)/(2 * (P + w)))
  Nt1 <- (r1 * N1.0)/(1 + alpha.11 * N1.0 + alpha.12 * N2.0)
  r2 <- What * exp(-(((w + (1 - h2) * P)/(P + w)) * (E - x2.0))^2)/(2 * (P + w)))
  Nt2 <- (r2 * N2.0)/(1 + alpha.22 * N2.0 + alpha.21 * N1.0)
  return(c(Nt1, Nt2, r1, r2))
}

# Simulation of model for t time steps
t <- 40
N <- array(NA, dim = c(t, 2))
N <- as.data.frame(N)
colnames(N) <- c("N1", "N2")
x <- array(NA, dim = c(t, 2))
x <- as.data.frame(x)
colnames(x) <- c("x1", "x2")
r <- array(NA, dim = c(t, 2))
r <- as.data.frame(r)
colnames(r) <- c("r1", "r2")
N$N1[1] <- N1.0
N$N2[1] <- N2.0
x$x1[1] <- x1.0
x$x2[1] <- x2.0
What <- Wmax * sqrt(w/(P + w))
r$r1[1] <- What * (exp(-(((w + (1 - h2) * P)/(P + w)) * (E.0 - x1.0))^2)/(2 * (P +
  w))))
r$r2[1] <- What * (exp(-(((w + (1 - h2) * P)/(P + w)) * (E.0 - x2.0))^2)/(2 * (P +
  w))))
E <- rep(NA, t)
E[1] <- E.0
for (i in 2:t) {
  res <- disc_LV_evol(N1.0 = N[i - 1, 1], N2.0 = N[i - 1, 2], alpha.11 = alpha.11,
    alpha.22 = alpha.22, alpha.12 = alpha.12, alpha.21 = alpha.21, E = E[i -
    1], x1 = x[i - 1, 1], x2 = x[i - 1, 2], P = P, w = w, Wmax = Wmax, h2 = h2)
  N$N1[i] <- res[1]
  N$N2[i] <- res[2]
  r$r1[i] <- res[3]
  r$r2[i] <- res[4]
  # trait change
  d1 <- E[i - 1] - x[i - 1, 1]
  d2 <- E[i - 1] - x[i - 1, 2]
  d1.1 <- k * d1
  d2.1 <- k * d2
  x$x1[i] <- E[i - 1] - d1.1
  x$x2[i] <- E[i - 1] - d2.1
  # environmental change

```

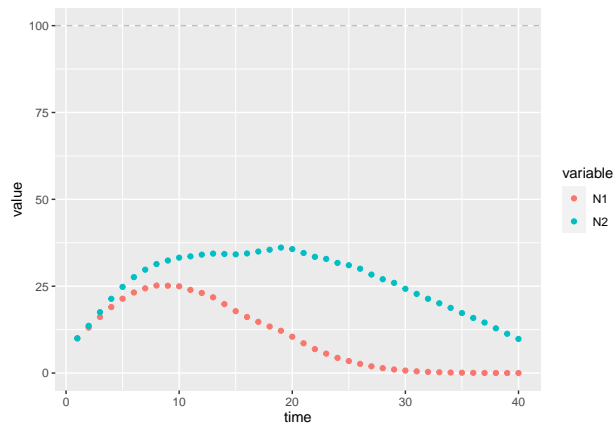
```
E[i] <- E[i - 1] + abs(rnorm(1, 0, 0.05))
}

# Plot simulation: ggplot
N$time <- 1:t
dat <- melt(N, id.vars = "time")
ggplot2::ggplot(dat, aes(time, value, col = variable)) + geom_point() + geom_hline(yintercept = (1/alpha),
  linetype = "dashed", color = "gray")
```



15

16 We can check the impact evolution had for this system by re-running dynamics with $h^2 = 0$:



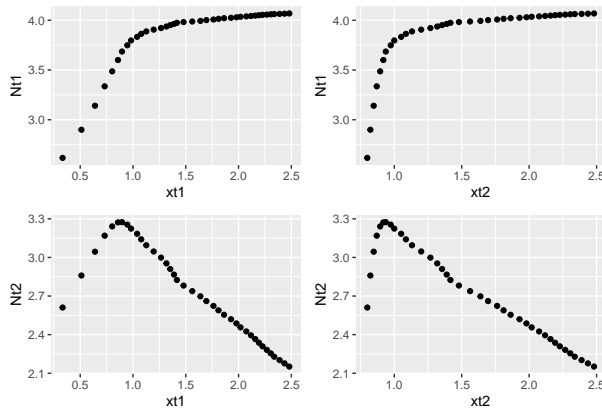
17

18 3 Hypotheses for evolution as a driver of species abundances

19 The growth equation is not linear, yet we fit abundance data to a hierarchical model with linear predictors
 20 because in most natural systems, a single theoretical growth model can't adequately capture the complexity
 21 in a given system. Linear models don't capture mechanistic relationships, but instead correlative. They
 22 are useful for inferring direction and magnitude of effect sizes. In Appendix S1, we explore how non-linear
 23 dynamics are captured when fit to a linear model. Here, we explore different forms to encode evolution in
 24 the trait x that determines the organism's fitness in this system.

3.1 H1. Raw trait value drives species abundances

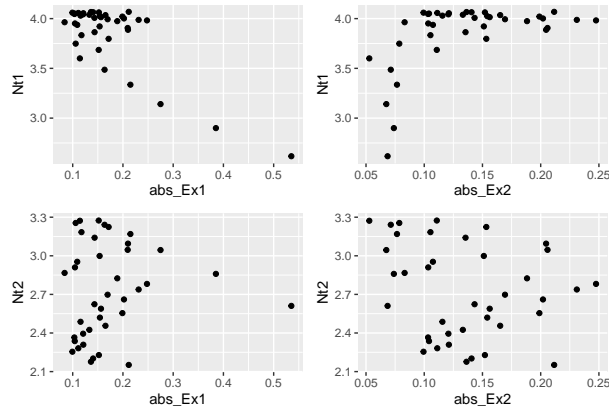
```
# Plot simulation: ggplot
dat <- as.data.frame(cbind(log(N$N1), log(N$N2), x$x1, x$x2))
colnames(dat) <- c("N1", "N2", "x1", "x2")
dat$time <- 1:t
df <- data.frame(cbind(dat$N1[2:t], dat$N2[2:t], dat$x1[2:t], dat$x2[2:t]))
colnames(df) <- c("Nt1", "Nt2", "xt1", "xt2")
# Plot
p1 <- ggplot2::ggplot(df, aes(xt1, Nt1)) + geom_point()
p2 <- ggplot2::ggplot(df, aes(xt2, Nt1)) + geom_point()
p3 <- ggplot2::ggplot(df, aes(xt1, Nt2)) + geom_point()
p4 <- ggplot2::ggplot(df, aes(xt2, Nt2)) + geom_point()
p1 + p2 + p3 + p4
```



We do not hypothesize that the raw trait value is the best predictor of species abundances, because the impacts of this evolved trait depend on the context of the environment.

3.2 H2. Absolute value of trait distance from optimum drives species abundances

```
# Plot simulation: ggplot
df$E <- E[2:t]
df$abs_Ex1 <- abs(df$E - df$xt1)
df$abs_Ex2 <- abs(df$E - df$xt2)
# Plot
p1 <- ggplot2::ggplot(df, aes(abs_Ex1, Nt1)) + geom_point()
p2 <- ggplot2::ggplot(df, aes(abs_Ex2, Nt1)) + geom_point()
p3 <- ggplot2::ggplot(df, aes(abs_Ex1, Nt2)) + geom_point()
p4 <- ggplot2::ggplot(df, aes(abs_Ex2, Nt2)) + geom_point()
p1 + p2 + p3 + p4
```



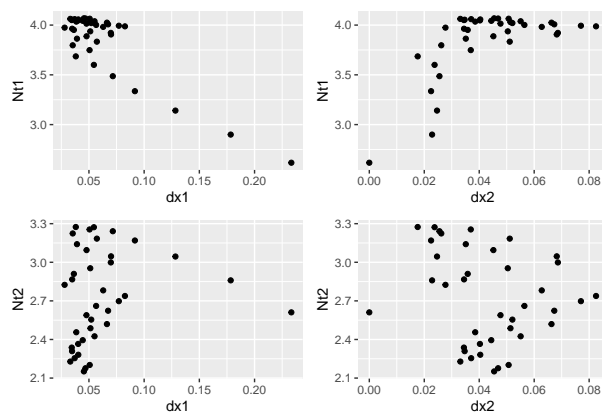
31

32 We hypothesize that this represents the true impact on species abundances, via the direct impact this measure
 33 has on fitness and consequent population growth. However, we also acknowledge this is likely difficult to
 34 measure in most empirical systems.

35 3.3 H3. Absolute value of trait change from one time to the next drives species 36 abundances

37 We now consider the magnitude of trait change from one time step to the next as a driver for species
 38 abundances.

```
df$dx1 <- abs(dat$x1[2:t] - dat$x1[1:(t - 1)])
df$dx2 <- abs(dat$x2[2:t] - dat$x2[1:(t - 1)])
# Plot
p1 <- ggplot2::ggplot(df, aes(dx1, Nt1)) + geom_point()
p2 <- ggplot2::ggplot(df, aes(dx2, Nt1)) + geom_point()
p3 <- ggplot2::ggplot(df, aes(dx1, Nt2)) + geom_point()
p4 <- ggplot2::ggplot(df, aes(dx2, Nt2)) + geom_point()
p1 + p2 + p3 + p4
```



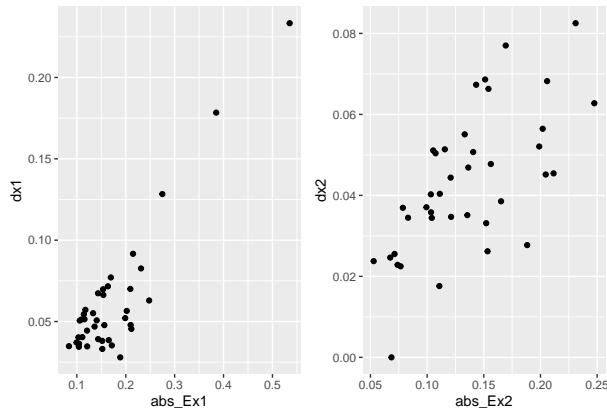
39

40 The measures correlate highly with one another, as the amount of trait change (the amount of evolution)
 41 is determined by the distance to the optimum trait value (given that P and w remain constant in the
 42 simulation).

```

cor(df$dx1, df$abs_Ex1)
#> [1] 0.8944668
cor(df$dx2, df$abs_Ex2)
#> [1] 0.6523635
# Plot
p1 <- ggplot2::ggplot(df, aes(abs_Ex1, dx1)) + geom_point()
p2 <- ggplot2::ggplot(df, aes(abs_Ex2, dx2)) + geom_point()
p1 + p2

```



43

44 4 Statistical model for evolution as a driver of species abundances

45 We hypothesize that $|E_t - x_{i,t}|$ will be the best predictor of species abundances, and that $|x_{t+1} - x_{i,t}|$ will
 46 have similar explanatory power.

```

# prepare data in HMSC format
dat <- as.data.frame(cbind(log(N$N1), log(N$N2), x$x1, x$x2))
colnames(dat) <- c("N1", "N2", "x1", "x2")
dat$time <- 1:t
df <- data.frame(cbind(dat$N1[2:t], dat$N2[2:t], dat$x1[2:t], dat$x2[2:t]))
colnames(df) <- c("Nt1", "Nt2", "xt1", "xt2")
df$dx1 <- abs(dat$x1[2:t] - dat$x1[1:(t - 1)])
df$dx2 <- abs(dat$x2[2:t] - dat$x2[1:(t - 1)])
# Y matrix
Y <- as.matrix(cbind(df$Nt1, df$Nt2))
# X matrix
XData <- data.frame(cbind(dat$N1[1:(t - 1)], dat$N2[1:(t - 1)], E[1:(t - 1)], E[1:(t - 1)]^2,
  dat$x1[1:(t - 1)], dat$x2[1:(t - 1)], abs(dat$x1[2:t] - dat$x1[1:(t - 1)]),
  abs(dat$x2[2:t] - dat$x2[1:(t - 1)]), abs(E[1:(t - 1)] - dat$x1[1:(t - 1)]),
  abs(E[1:(t - 1)] - dat$x2[1:(t - 1)]))
colnames(XData) <- c("n1", "n2", "E", "Esq", "x1", "x2", "dx1", "dx2", "dEx1", "dEx2")
# Prepare HMSC model
studyDesign = data.frame(sample = as.factor(1:(t - 1)))
rL = HmscRandomLevel(units = studyDesign$sample)
# Design and fit 4 alternative HMSC models
models = list()
for (i in 1:4) {
  XFormula = switch(i, ~1, ~E + Esq + x1 + x2, ~E + Esq + dEx1 + dEx2, ~E + Esq +

```

```

    dx1 + dx2)
  m = Hmsc(Y = Y, XData = XData, XFormula = XFormula, studyDesign = studyDesign,
    ranLevels = list(sample = rL))
  models[[i]] = m
}
# Bayesian model parameters
nChains <- 2
thin <- 5
samples <- 2000
transient <- 100 * thin
verbose <- 500 * thin
for (i in 1:4) {
  models[[i]] = sampleMcmc(models[[i]], thin = thin, sample = samples, transient = transient,
    nChains = nChains, verbose = verbose)
}
#> Computing chain 1
#> Chain 1, iteration 2500 of 10500 (sampling)
#> Chain 1, iteration 5000 of 10500 (sampling)
#> Chain 1, iteration 7500 of 10500 (sampling)
#> Chain 1, iteration 10000 of 10500 (sampling)
#> Computing chain 2
#> Chain 2, iteration 2500 of 10500 (sampling)
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#> Chain 1, iteration 10000 of 10500 (sampling)
#> Computing chain 2
#> Chain 2, iteration 2500 of 10500 (sampling)
#> Chain 2, iteration 5000 of 10500 (sampling)

```

```
#> Chain 2, iteration 7500 of 10500 (sampling)
#> Chain 2, iteration 10000 of 10500 (sampling)
```

47 We check the explanatory and predictive power of each model using cross-validation.

```
# After Ovaskainen & Abrego 2020, Chapter 7
partition = createPartition(m, nfolds = 2, column = "sample")
partition.sp = c(1, 2)
result = matrix(NA, nrow = 3, ncol = 4)
for (i in 1:4) {
  m = models[[i]]
  # Explanatory power
  preds = computePredictedValues(m)
  MF = evaluateModelFit(hM = m, predY = preds)
  result[1, i] = mean(MF$R2)
  # Predictive power based on cross-validation
  preds = computePredictedValues(m, partition = partition)
  MF = evaluateModelFit(hM = m, predY = preds)
  result[2, i] <- mean(MF$R2)
  # Predictive power based on conditional cross-validation
  preds = computePredictedValues(m, partition = partition, partition.sp = partition.sp,
    mcmcStep = 100)
  MF = evaluateModelFit(hM = m, predY = preds)
  result[3, i] = mean(MF$R2)
}
#> Cross-validation, fold 1 out of 2
#> Computing chain 1
#> Chain 1, iteration 2500 of 10500 (sampling)
#> Chain 1, iteration 5000 of 10500 (sampling)
#> Chain 1, iteration 7500 of 10500 (sampling)
#> Chain 1, iteration 10000 of 10500 (sampling)
#> Computing chain 2
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#> Cross-validation, fold 2 out of 2
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#> Computing chain 1

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#> Chain 1, iteration 2500 of 10500 (sampling)
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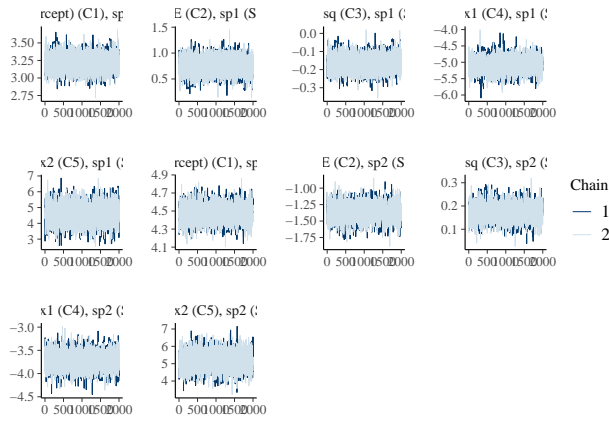
colnames(result) <- c("null", "xt", "dEx", "dx")
rownames(result) <- c("R2_expl", "R2_pred", "R2_pred_cross")
result
#>
#>      null      xt      dEx      dx
#> R2_expl  0.86789569 0.9958630 0.9958779 0.9958438
#> R2_pred -0.06763985 0.9845892 0.9840017 0.9839582
#> R2_pred_cross 0.04775282 0.9847787 0.9843090 0.9842851

```

48 An important conclusion from this analysis is that the $|E_t - x_{i,t}|$ and $|x_{t+1} - x_{i,t}|$ do have similar explanatory
49 power, and that - even though the species traits were the stronger predictor of species abundances - the

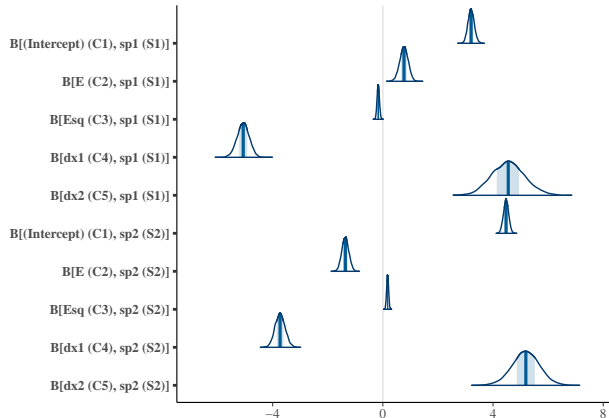
50 model with total change in trait value can be used to assess the question “How does evolution impact species
 51 abundances?”

```
m.post.hmsc <- convertToCodaObject(models[[4]])
bayesplot::mcmc_trace(m.post.hmsc$Beta)
```



52

```
bayesplot::mcmc_areas(m.post.hmsc$Beta, area_method = c("equal height"))
```

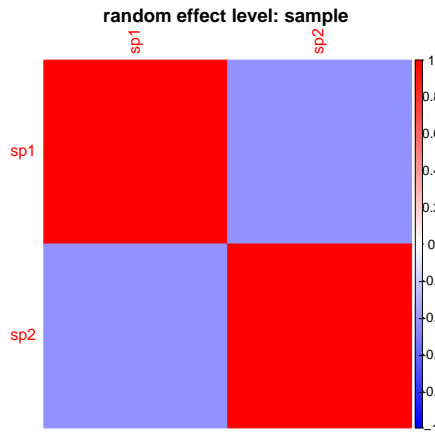


53

```
# Bayesian estimates
cbind(summary(m.post.hmsc$Beta)$statistics[1:10, 1], summary(m.post.hmsc$Beta)$quantiles[,
c(1, 5)])

#>
#> B[(Intercept) (C1), sp1 (S1)] 3.2062148 2.9907638 3.43861741
#> B[E (C2), sp1 (S1)] 0.7645592 0.4605920 1.05614131
#> B[Esq (C3), sp1 (S1)] -0.1672595 -0.2494376 -0.08281671
#> B[dx1 (C4), sp1 (S1)] -5.0723930 -5.5124602 -4.65318246
#> B[dx2 (C5), sp1 (S1)] 4.5542240 3.4173450 5.76914490
#> B[(Intercept) (C1), sp2 (S2)] 4.4729905 4.2864747 4.65632792
#> B[E (C2), sp2 (S2)] -1.3628495 -1.6049063 -1.11280451
#> B[Esq (C3), sp2 (S2)] 0.1727352 0.1028246 0.24025095
#> B[dx1 (C4), sp2 (S2)] -3.7300421 -4.0813900 -3.36742269
#> B[dx2 (C5), sp2 (S2)] 5.1890615 4.1851659 6.17434226
```

```
OmegaCor = computeAssociations(models[[4]])
supportLevel = 0.7
toPlot = ((OmegaCor[[1]]$support > supportLevel) + (OmegaCor[[1]]$support < (1 -
  supportLevel)) > 0) * OmegaCor[[1]]$mean
corrplot::corrplot(toPlot, method = "color", col = colorRampPalette(c("blue", "white",
  "red"))(200), title = paste("random effect level:", models[[4]]$rLNames[1]),
  mar = c(0, 0, 1, 0))
```

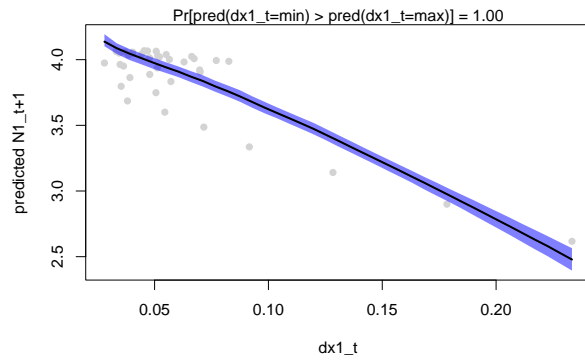


54

```
OmegaCor
#> [[1]]
#> [[1]]$mean
#>           sp1           sp2
#> sp1  1.0000000 -0.4249683
#> sp2 -0.4249683  1.0000000
#>
#> [[1]]$support
#>           sp1 sp2
#> sp1  1.00 0.24
#> sp2 0.24 1.00
```

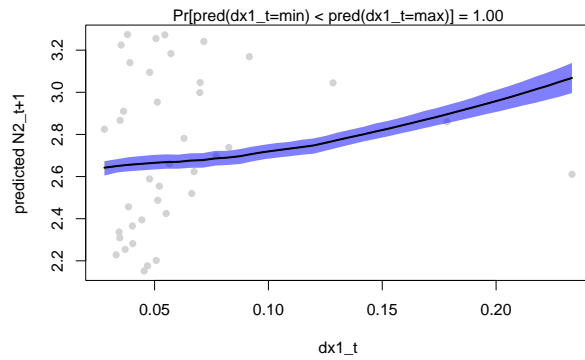
55 We can also look at a gradient plot for the effect of evolution in Species 1 for abundance in Species 1 and
 56 Species 2.

```
Gradient <- constructGradient(models[[4]], focalVariable = "dx1", non.focalVariables = list(E = list(2),
  Esq = list(2), dx2 = list(2)), ngrid = 39)
Gradient$XDataNew$Esq <- Gradient$XDataNew$E^2
predY <- predict(models[[4]], XData = Gradient$XDataNew, expected = TRUE)
a <- plotGradient(models[[4]], Gradient, pred = predY, showData = T, measure = "Y",
  index = 1, main = "", xlab = "dx1_t", ylab = "predicted N1_t+1")
```



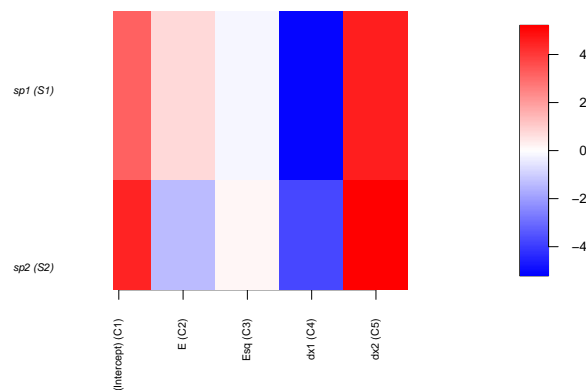
57

```
b <- plotGradient(models[[4]], Gradient, pred = predY, showData = T, measure = "Y",
  index = 2, main = "", xlab = "dx1_t", ylab = "predicted N2_t+1")
```



58

```
postBeta = getPostEstimate(models[[4]], parName = "Beta")
plotBeta(models[[4]], post = postBeta, param = "Mean", supportLevel = 0.7)
```

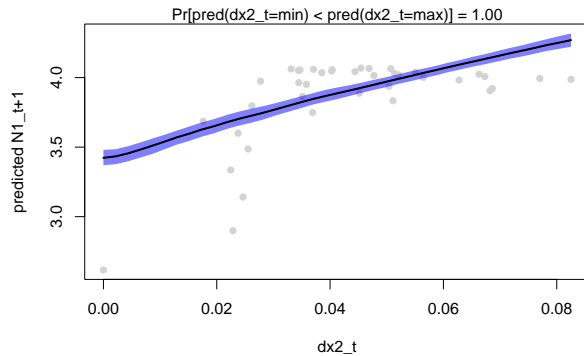


59

60 Species 2 for abundance in Species 1 and Species 2.

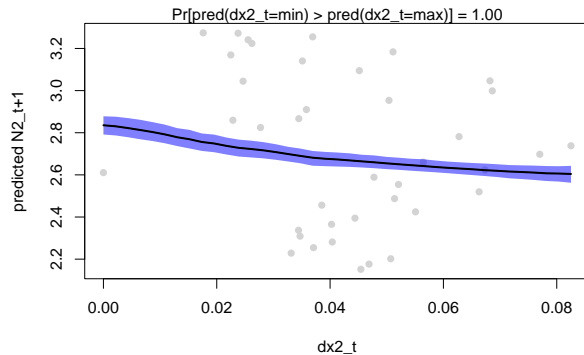
We repeat this plot for the effect of evolution in

```
Gradient <- constructGradient(models[[4]], focalVariable = "dx2", non.focalVariables = list(E = list(2)
  Esq = list(2), dx1 = list(2)), ngrid = 39)
Gradient$XDataNew$Esq <- Gradient$XDataNew$E^2
predY <- predict(models[[4]], XData = Gradient$XDataNew, expected = TRUE)
a <- plotGradient(models[[4]], Gradient, pred = predY, showData = T, measure = "Y",
  index = 1, main = "", xlab = "dx2_t", ylab = "predicted N1_t+1")
```



61

```
b <- plotGradient(models[[4]], Gradient, pred = predY, showData = T, measure = "Y",
  index = 2, main = "", xlab = "dx2_t", ylab = "predicted N2_t+1")
```

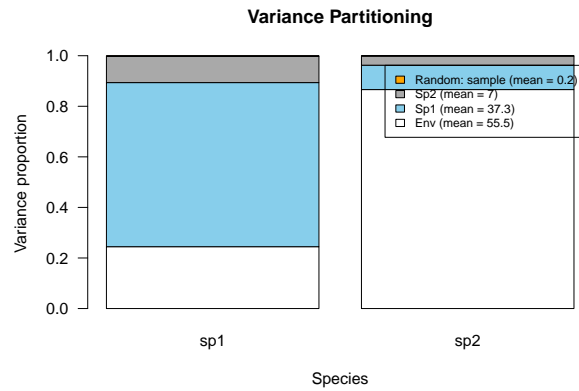


62

63 species abundances, we use variation partition:

To evaluate the relative importance of evolution for

```
VP <- computeVariancePartitioning(models[[4]], group = c(1, 1, 1, 2, 3), groupnames = c("Env",
  "Sp1", "Sp2"))
plotVariancePartitioning(models[[4]], VP, cols = c("white", "skyblue", "darkgrey",
  "orange"), args.legend = list(cex = 0.75, bg = "transparent"))
```



64 We see that trait evolution in Species 1 is an impor-
 65 tant driver of abundances in both Species 1 and Species 2.