# experiment 1

### Owen Petchey

6/25/2021

This experiment supercedes all previous ones. It is a factorial manipulation of diversity of the three groups. It takes about 50 hours to run while using 12 cores.

## Setup

### $\mathbf{R}$

```
rm(list = ls())
knitr::opts_knit$set(progress = TRUE, verbose = FALSE, cache = TRUE)
microxanox_release <- "0.2"
#tmplib <- tempfile()</pre>
#dir.create(tmplib)
### From '?remotes::install_github`:
# auth_token
   To install from a private repo, generate a personal access token (PAT) in
    "https://github.com/settings/tokens" and supply to this argument. This is
   safer than using a password because you can easily delete a PAT without
   affecting any others. Defaults to the GITHUB_PAT environment variable.
# remotes::install_github(
  "opetchey/microxanox",
# ref = microxanox_release,
  # auth_token = "ENTER YOUR TOKEN or PROVED AS ENVIRONMENT VARIABLE",
  build_vignettes = FALSE,
  force = TRUE,
   upgrade = FALSE,
   lib = tmplib
# )
#library(microxanox, lib.loc = tmplib)
library(microxanox)
library(tidyverse)
```

```
## -- Attaching packages ------ tidyverse 1.3.1 --
## v ggplot2 3.3.5 v purrr 0.3.4
```

## Version of microxanox package used: 0.2.2

#### General simulation conditions

```
default_dynamic_model <- bushplus_dynamic_model
default_event_definition <- event_definition_1
default_event_interval <- 100
default_noise_sigma <- 0
default_minimum_abundances <- rep(1, 3)
names(default_minimum_abundances) <- c("CB", "PB", "SB")
default_sim_duration <- 80000
default_sim_sample_interval <- 100
initial_pars_from <- "bush_ssfig3"
## note that next line (log10a_series is over-ridden with getting stable states)
#default_log10a_series <- c(-2, -2, -2, -10, -10, -10, -10, -10)</pre>
```

#### Define diversity

```
num_CB_strains <- 9
num_SB_strains <- 9
num_PB_strains <- 9

## multiplier of SBPB variation

CB_var_multiplier <- 2
SBPB_var_multiplier <- 6

CB_gmax_div <- 0.015789474 * CB_var_multiplier

CB_h_div <- -0.08 * CB_var_multiplier

SB_gmax_div <- 0.015789474 * SBPB_var_multiplier

SB_h_div <- -0.323 * SBPB_var_multiplier

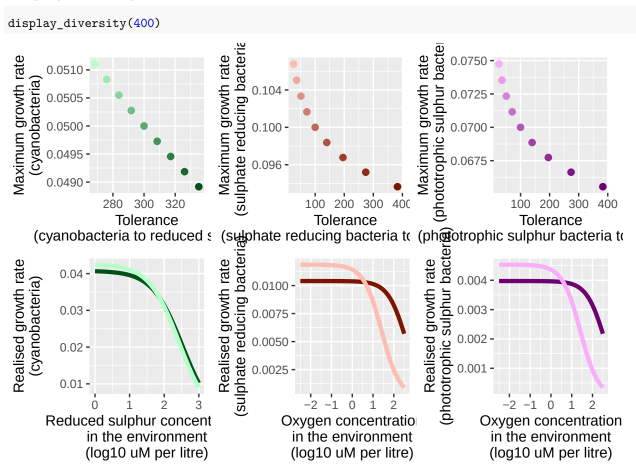
PB_gmax_div <- 0.015789474 * SBPB_var_multiplier

PB_h_div <- -0.323 * SBPB_var_multiplier</pre>
```

### Create diversity

```
var_expt <- create_diversity_factorial()</pre>
```

## Display diversity



## Temporal switching

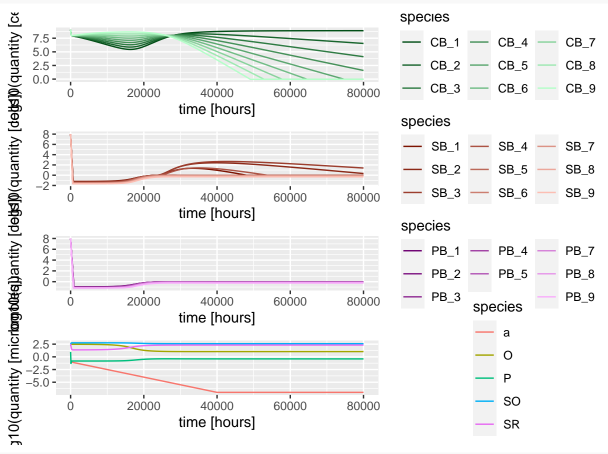
#### Oxic to anoxic

#### No diversity

```
default_sim_duration <- 80000
default_log10a_series \leftarrow c(-1, -7, -7)
initial_state <- new_initial_state(num_CB_strains,</pre>
                                          num_PB_strains,
                                          num_SB_strains,
                                          values = "bush_ssfig3")
initial_state[grep("CB_", names(initial_state))] <- 10^10/num_CB_strains
sim_res_novar1 <- run_simulation(parameter_values = var_expt$pars[[no_diversity]],</pre>
                                initial_state = initial_state)
saveRDS(sim_res_novar1, here("experiments/experiment 1/data/sim_res_novar1.RDS"))
sim_res_novar1 <- readRDS(here("experiments/experiment 1/data/sim_res_novar1.RDS"))</pre>
plot_dynamics(sim_res_novar1)
                                                                     species
310(quantity[micrloggtotes])antity[de的例(quantity[de的例)(quantity[ce
    7.5 -
                                                                           CB_1
                                                                                     CB 4
                                                                                                   CB_7
    5.0 - 2.5 - 0.0 -
                                                                           CB<sub>2</sub>
                                                                                       CB<sub>5</sub>
                                                                                                   CB_8
                     20000
                                  40000
                                                            80000
                                                                           CB_3
                                                                                       CB<sub>6</sub>
                                                                                                   CB_9
                                               60000
                               time [hours]
                                                                      species
      6 -
                                                                           SB 1 —
                                                                                       SB 4
                                                                                                   SB 7
      3 -
                                                                           SB_2
                                                                                       SB_5
                                                                                                   SB<sub>8</sub>
                     20000
                                                                           SB_3
                                                                                       SB_6
                                                                                                   SB_9
                                  40000
                                               60000
                                                            80000
           \cap
                               time [hours]
                                                                      species
      8 -
6 -
4 -
2 -
0 -
                                                                           PB_1
                                                                                       PB_4
                                                                                                   PB_7
                                                                           PB_2
                                                                                       PB_5
                                                                                                   PB_8
                     20000
                                  40000
                                                                           PB_3
                                                                                                   PB_9
                                               60000
                                                            80000
           0
                                                                                  species
                               time [hours]
     2.5 -
0.0 -
                                                                                        0
     -2.5 -
                                                                                        Р
    –5.0 -
                     20000
                                                                                        SO
                                  40000
                                               60000
                                                            80000
                               time [hours]
                                                                                        SR
#qqsave(here("simulations/expt2/figures/switching_novar.pdf"), width = 10)
```

#### Maximum diversity

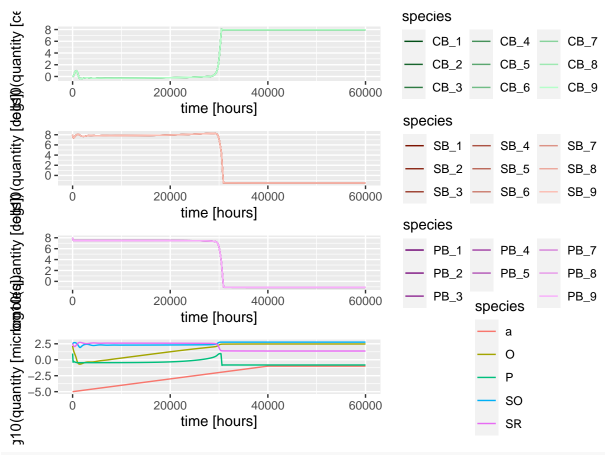




#qqsave(here("simulationsexpt2/figures/switching highvar.pdf"), width = 10)

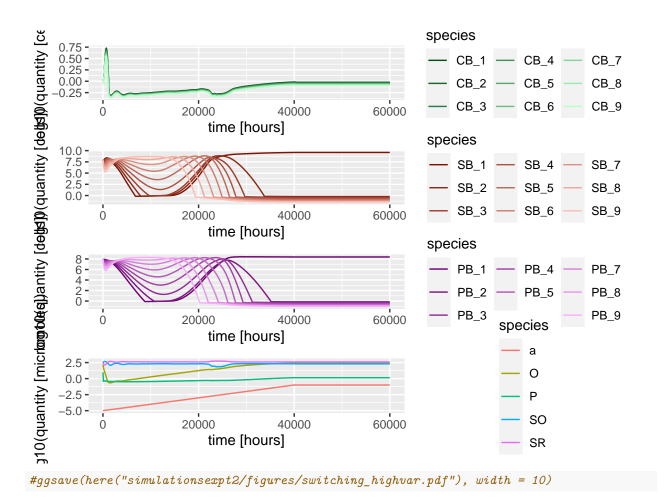
## Anoxic to oxic

#### No diversity



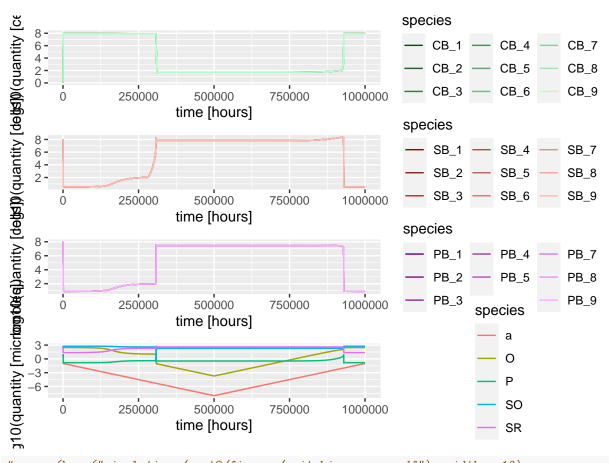
#qqsave(here("simulations/expt2/figures/switching novar.pdf"), width = 10)

### Maximum diversity



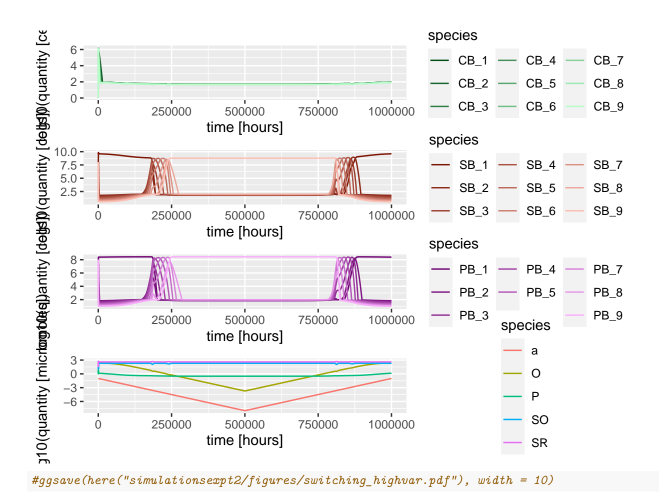
#### Anoxic to oxic to anoxic

#### No diversity



 $\#ggsave(here("simulations/expt2/figures/switching_novar.pdf"), \ width = 10)$ 

### Maximum diversity



#### Visualise

visualise\_temporal\_env\_eco()

## Stable state finding

#### **Finding**

```
options(mc.cores = 12)

default_sim_duration <- 1000000
ssfind_minimum_abundances <- rep(0, 3)
names(ssfind_minimum_abundances) <- c("CB", "PB", "SB")
ssfind_simulation_duration <- default_sim_duration
ssfind_simulation_sampling_interval <- ssfind_simulation_duration
ssfind_event_interval <- ssfind_simulation_duration
grid_num_a <- 1000 #usually 1000 ## number of a_0 values
a_0s <- 10^seq(-7, -0.5, length=grid_num_a) ## sequence of a_0 values
grid_num_N <- 2 ## number of N values
initial_CBs <- 10^seq(0, 10, length=grid_num_N) ## sequence of N values
initial_PBs <- 1e8 ## not varied
initial_SBs <- 1e8 ## not varied
# next line creates all possible combinations</pre>
```

Run stable state finding

\*Careful, this simulation takes about 600 hours on a single core

```
var_expt <- run_ss_var_experiment()
saveRDS(var_expt, here("experiments/experiment 1/data/ss_data_1e6_x2x6_factorial.RDS"))</pre>
```

Process the stable state data

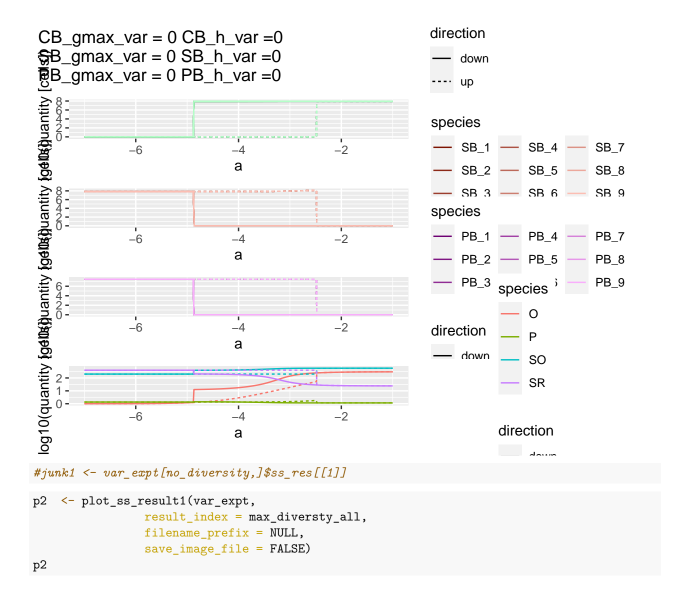
Bring in various stable state datasets

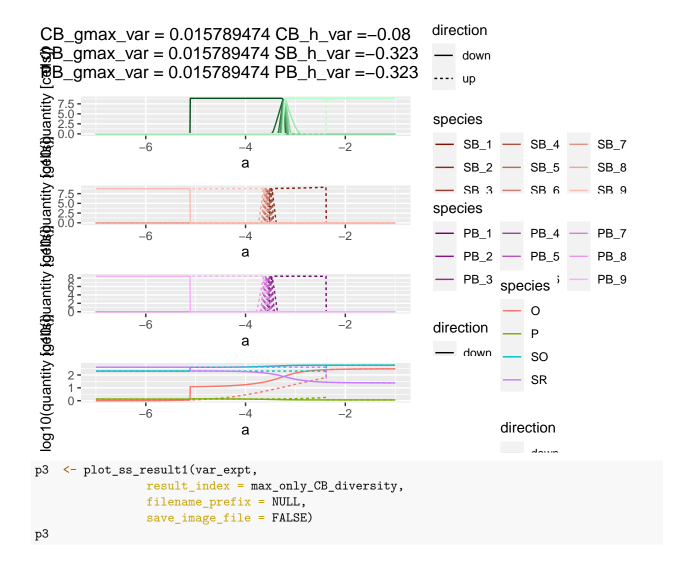
```
## sim length 80'000, 20 x 20 factorial, reference maximum diversity
var_expt1 <- readRDS(here("experiments/experiment 1/data/ss_data_80000.RDS")) %>%
  mutate(sim length = 80000)
stab_data1 <- var_expt1 %>%
  group_by(CB_var_gmax_s, CB_var_h_s,
           SB_var_gmax_s, SB_var_h_s,
           PB_var_gmax_s, PB_var_h_s, sim_length) %>%
  do(stability measures = get stability measures(.$ss res[[1]]))
stab_data1 <- unnest(stab_data1, cols = c(stability_measures))</pre>
saveRDS(stab_data1, here("experiments/experiment 1/data/stab_data_80000.RDS"))
## sim length 1'000'000, 20 x 20 factorial, reference maximum diversity
var_expt2 <- readRDS(here("experiments/experiment 1/data/ss_data_1000000_20factorial.RDS")) %>%
  mutate(sim_length = 1000000)
stab_data2 <- var_expt2 %>%
  group_by(CB_var_gmax_s, CB_var_h_s,
           SB_var_gmax_s, SB_var_h_s,
           PB_var_gmax_s, PB_var_h_s, sim_length) %>%
  do(stability_measures = get_stability_measures(.$ss_res[[1]]))
stab_data2 <- unnest(stab_data2, cols = c(stability_measures))</pre>
saveRDS(stab_data2, here("experiments/experiment 1/data/stab_data_1000000_20factorial.RDS"))
## sim length 1'000'000, 20 SBPBgrad, 5x maximum diversity
var expt3 <- readRDS(here("experiments/experiment 1/data/ss data 1e6 noCB 5xSBPB .RDS")) %>%
  mutate(sim_length = 1000000)
stab_data3 <- var_expt3 %>%
  group_by(CB_var_gmax_s, CB_var_h_s,
           SB_var_gmax_s, SB_var_h_s,
           PB_var_gmax_s, PB_var_h_s, sim_length) %>%
  do(stability_measures = get_stability_measures(.$ss_res[[1]]))
stab_data3 <- unnest(stab_data3, cols = c(stability_measures))</pre>
saveRDS(stab_data3, here("experiments/experiment 1/data/stab_data_1e6_noCB_5xSBPB_.RDS"))
## sim length 300'000, 20 SBPBgrad, reference maximum diversity
var_expt4 <- readRDS(here("experiments/experiment 1/data/ss_data_300000_small.RDS")) %>%
 mutate(sim_length = 300000)
```

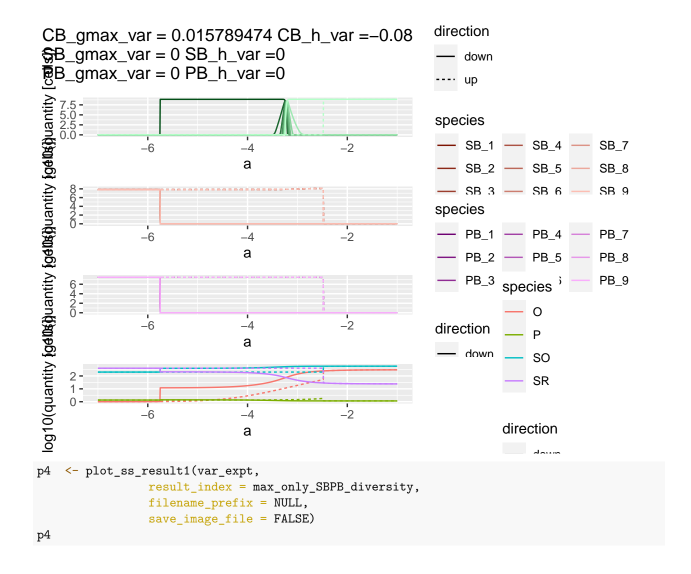
```
stab_data4 <- var_expt4 %>%
  group_by(CB_var_gmax_s, CB_var_h_s,
           SB_var_gmax_s, SB_var_h_s,
           PB_var_gmax_s, PB_var_h_s, sim_length) %>%
  do(stability_measures = get_stability_measures(.$ss_res[[1]]))
stab_data4 <- unnest(stab_data4, cols = c(stability_measures))</pre>
saveRDS(stab_data4, here("experiments/experiment 1/data/stab_data_300000.RDS"))
## sim length 1'000'000, 20 SBPBgrad, 2xCB variation, 6xSBPB variation
var_expt5 <- readRDS(here("experiments/experiment 1/data/ss_data_1e6_x2x6_factorial.RDS")) %%</pre>
 mutate(sim length = 1000000)
stab_data5 <- var_expt5 %>%
  group_by(CB_var_gmax_s, CB_var_h_s,
           SB_var_gmax_s, SB_var_h_s,
           PB_var_gmax_s, PB_var_h_s, sim_length) %>%
  do(stability_measures = get_stability_measures(.$ss_res[[1]]))
stab_data5 <- unnest(stab_data5, cols = c(stability_measures))</pre>
saveRDS(stab_data5, here("experiments/experiment 1/data/stab_data_1e6_x2x6_factorial.RDS"))
```

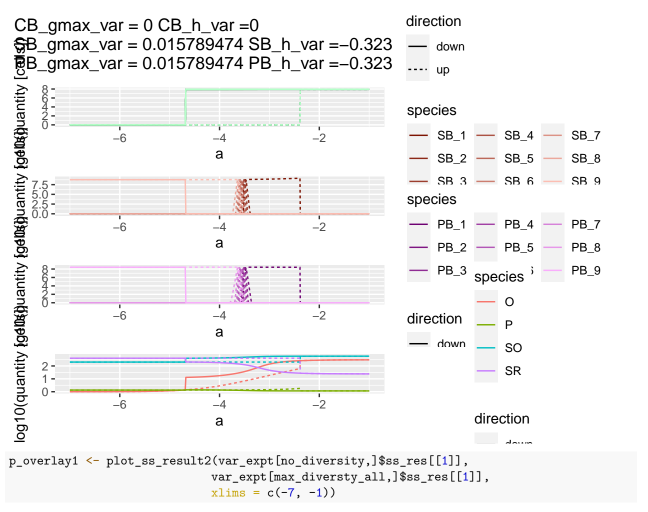
## SS, no diversity, all diversity, CB only, and SBPB only

```
## find various combinations of diversity
var_expt <- readRDS(here("experiments/experiment 1/data/ss_data_1000000_20factorial.RDS"))</pre>
var_expt_levels <- var_expt[,1:6]</pre>
no_diversity <- which(rowSums(abs(var_expt_levels))==0)</pre>
max_diversty_all <- which(max(rowSums(abs(var_expt_levels))) ==</pre>
                             rowSums(abs(var_expt_levels)))
max_only_CB_diversity <- which(max(rowSums(abs(var_expt_levels[,1:2]))) ==</pre>
                             rowSums(abs(var_expt_levels[,1:2])) &
                               rowSums(abs(var_expt_levels[,3:6]))==0)
#var_expt_levels[381,]
max_only_SBPB_diversity <- which(max(rowSums(abs(var_expt_levels[,3:6]))) ==</pre>
                             rowSums(abs(var expt levels[,3:6])) &
                               rowSums(abs(var_expt_levels[,1:2]))==0)
#var_expt_levels[20,]
p1 <- plot_ss_result1(var_expt,</pre>
                result_index = no_diversity,
                filename_prefix = NULL,
                 save image file = FALSE)
р1
```

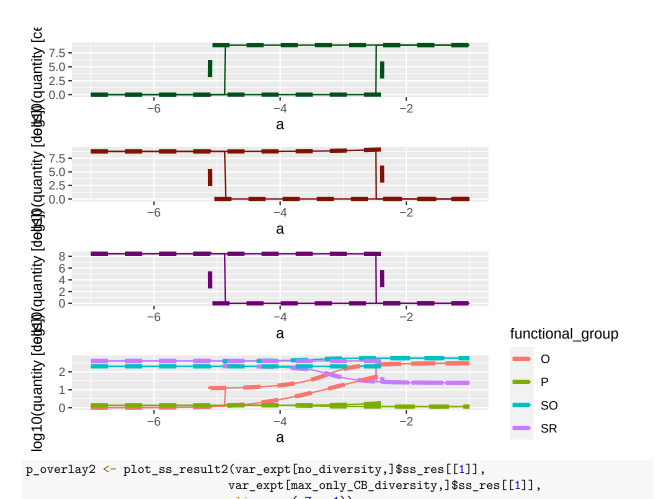




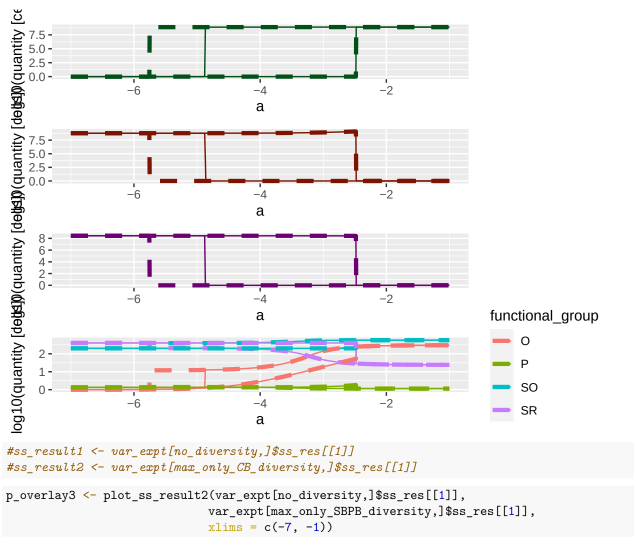




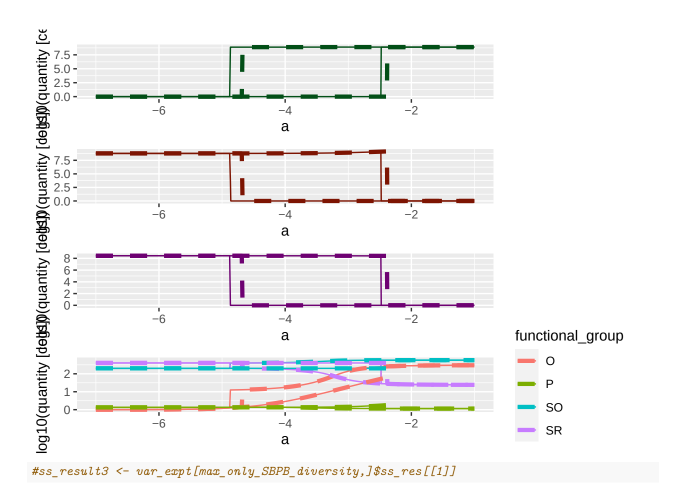
## `summarise()` has grouped output by 'a', 'direction', 'var\_type'. You can override using the `.group
## `summarise()` has grouped output by 'a', 'direction', 'var\_type'. You can override using the `.group
p\_overlay1



p\_overlay2



```
## `summarise()` has grouped output by 'a', 'direction', 'var_type'. You can override using the `.group
## `summarise()` has grouped output by 'a', 'direction', 'var_type'. You can override using the `.group
p_overlay3
```



## Look at stability measures

```
stab_data1 <- readRDS(here("experiments/experiment 1/data/stab_data_1000000_20factorial.RDS"))</pre>
stab data2 <- readRDS(here("experiments/experiment 1/data/stab data 80000.RDS"))
stab_data <- bind_rows(stab_data1, stab_data2)</pre>
CB_vars <- unique(stab_data$CB_var_gmax_s)</pre>
SB_vars <- unique(stab_data$CB_var_gmax_s)</pre>
CB_stab_data <- stab_data %>%
  filter(SB_var_gmax_s == 0) %>%
  mutate(var_treat = "CB",
         var_gmax = CB_var_gmax_s)
SBPB_stab_data <- stab_data %>%
  filter(CB_var_gmax_s == 0) %>%
  mutate(var_treat = "SB-PB",
         var_gmax = SB_var_gmax_s)
for_join <- tibble(CB_var_gmax_s = CB_vars,</pre>
                      SB_var_gmax_s = SB_vars)
CBSBPB_stab_data <- stab_data %>%
  right_join(for_join) %>%
  mutate(var_treat = "CB-SB-PB",
```

```
var_gmax = CB_var_gmax_s)

## Joining, by = c("CB_var_gmax_s", "SB_var_gmax_s")

all_stab_results <- CB_stab_data %>%
    bind_rows(SBPB_stab_data) %>%

# bind_rows(results3) %>%

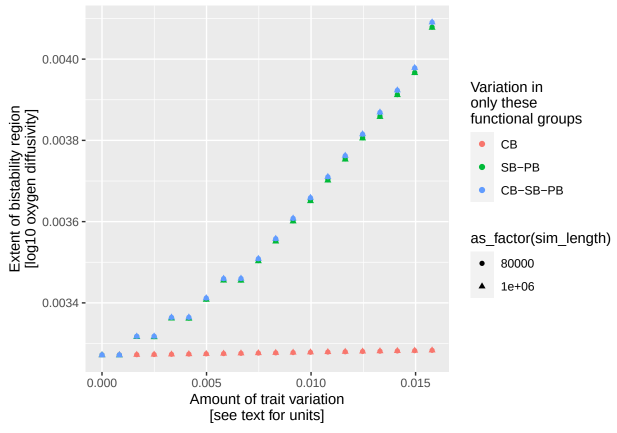
# bind_rows(results4) %>%
    bind_rows(CBSBPB_stab_data)

all_stab_results<- all_stab_results %>%
    mutate(var_treat = forcats::fct_relevel(var_treat, levels = c("CB", "SB-PB", "CB-SB-PB")))

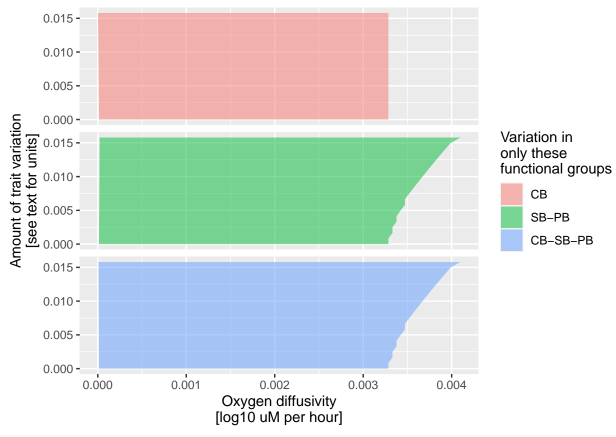
## Warning: Outer names are only allowed for unnamed scalar atomic inputs
saveRDS(all_stab_results, here("experiments/experiment 1/data/all_stab.RDS"))
```

#### Raw Stability

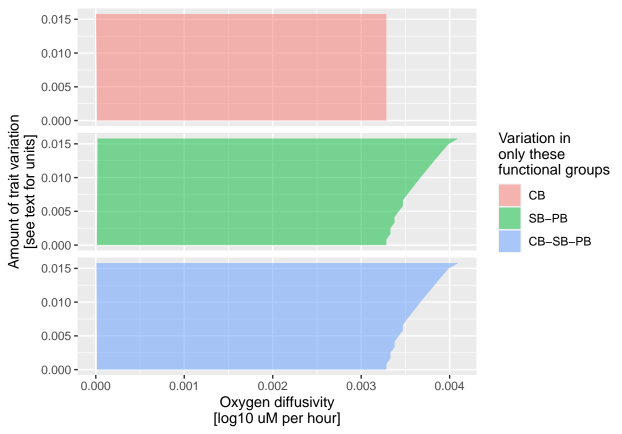
```
all_stab_results <- readRDS(here("experiments/experiment 1/data/all_stab.RDS"))
all_stab_results %>%
  filter(Species == "0") %>%
  ggplot(aes(x = var_gmax, y = hyst_range_raw, col=var_treat, shape = as_factor(sim_length))) +
  geom_point() +
  xlab("Amount of trait variation\n[see text for units]") +
  ylab("Extent of bistability region\n[log10 oxygen diffusivity]") +
  labs(col = "Variation in\nonly these\nfunctional groups")
```

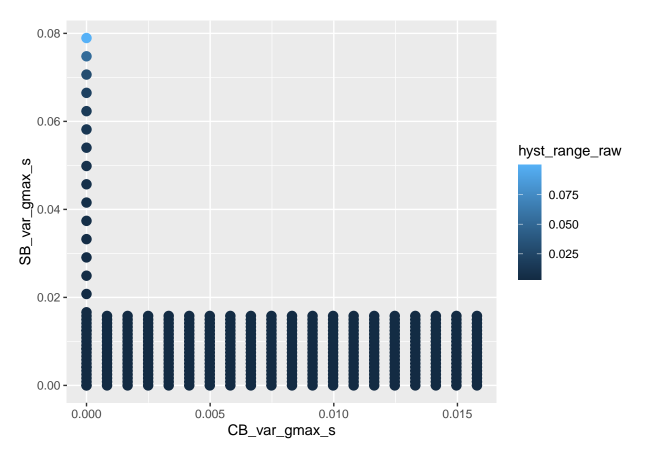


```
##ggsave("manuscript/figures/extent_of_bistab1.pdf", height = 4)
all_stab_results %>%
  #filter(var treat == "CB") %>%
  filter(Species == "0",
         sim_length == 1e6) %>%
  ggplot(aes(x = var_gmax,
             ymin = hyst_min_raw,
             ymax = hyst_max_raw,
             fill=var_treat)) +
  geom_ribbon(alpha = 0.5) +
  facet_wrap( ~ var_treat, nrow = 3) +
  xlab("Amount of trait variation\n[see text for units]") +
  ylab("Oxygen diffusivity\n[log10 uM per hour]") +
  labs(fill = "Variation in\nonly these\nfunctional groups") +
  coord_flip() +
  theme(
    strip.background = element_blank(),
    strip.text.x = element_blank()
 )
```



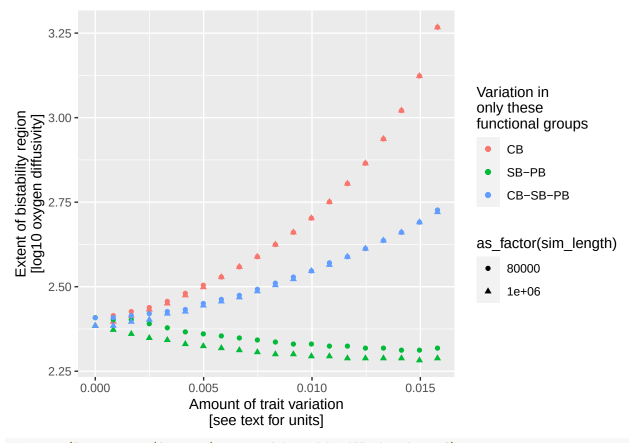
```
all_stab_results %>%
  #filter(var_treat == "CB") %>%
  filter(Species == "0",
         sim length == 8e4) %>%
  ggplot(aes(x = var_gmax,
             ymin = hyst_min_raw,
             ymax = hyst_max_raw,
             fill=var_treat)) +
  geom_ribbon(alpha = 0.5) +
  facet_wrap( ~ var_treat, nrow = 3) +
  xlab("Amount of trait variation\n[see text for units]") +
  ylab("Oxygen diffusivity\n[log10 uM per hour]") +
  labs(fill = "Variation in\nonly these\nfunctional groups") +
  coord_flip() +
  theme(
    strip.background = element_blank(),
    strip.text.x = element_blank()
```



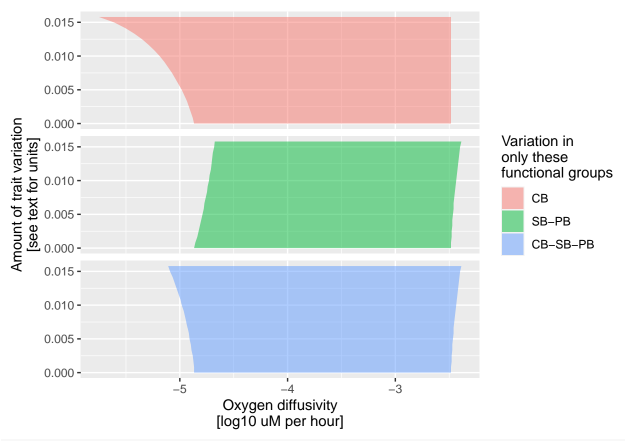


### Log Stability

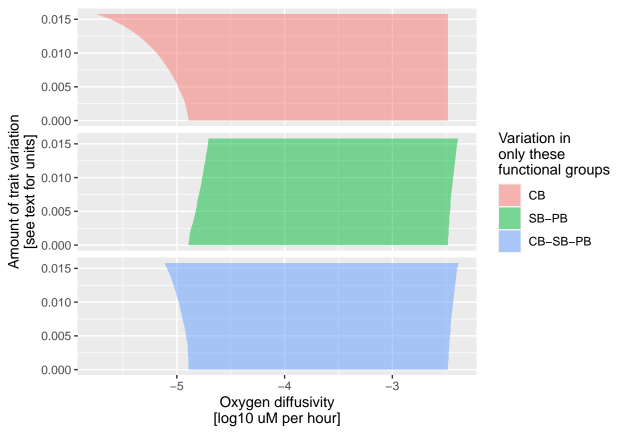
```
all_stab_results <- readRDS(here("experiments/experiment 1/data/all_stab.RDS"))
all_stab_results %>%
  filter(Species == "0") %>%
  ggplot(aes(x = var_gmax, y = hyst_range_log, col=var_treat, shape = as_factor(sim_length))) +
  geom_point() +
  xlab("Amount of trait variation\n[see text for units]") +
  ylab("Extent of bistability region\n[log10 oxygen diffusivity]") +
  labs(col = "Variation in\nonly these\nfunctional groups")
```

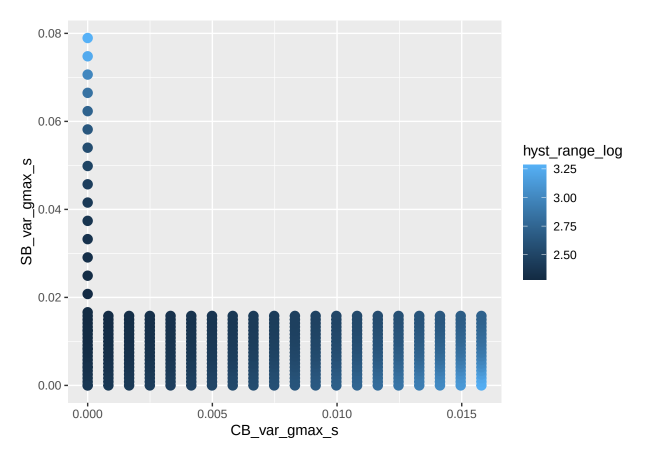


```
##ggsave("manuscript/figures/extent_of_bistab1.pdf", height = 4)
all_stab_results %>%
  #filter(var treat == "CB") %>%
  filter(Species == "0",
         sim_length == 1e6) %>%
  ggplot(aes(x = var_gmax,
             ymin = hyst_min_log,
             ymax = hyst_max_log,
             fill=var_treat)) +
  geom_ribbon(alpha = 0.5) +
  facet_wrap( ~ var_treat, nrow = 3) +
  xlab("Amount of trait variation\n[see text for units]") +
  ylab("Oxygen diffusivity\n[log10 uM per hour]") +
  labs(fill = "Variation in\nonly these\nfunctional groups") +
  coord_flip() +
  theme(
    strip.background = element_blank(),
    strip.text.x = element_blank()
 )
```



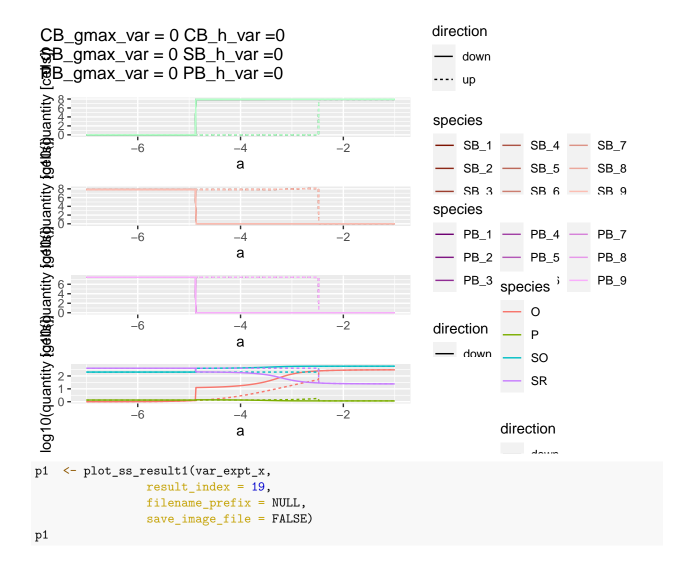
```
all_stab_results %>%
  #filter(var_treat == "CB") %>%
  filter(Species == "0",
         sim_length == 8e4) \%%
  ggplot(aes(x = var_gmax,
             ymin = hyst_min_log,
             ymax = hyst_max_log,
            fill=var_treat)) +
  geom_ribbon(alpha = 0.5) +
  facet_wrap( ~ var_treat, nrow = 3) +
  xlab("Amount of trait variation\n[see text for units]") +
  ylab("Oxygen diffusivity\n[log10 uM per hour]") +
  labs(fill = "Variation in\nonly these\nfunctional groups") +
  coord_flip() +
  theme(
    strip.background = element_blank(),
    strip.text.x = element_blank()
```





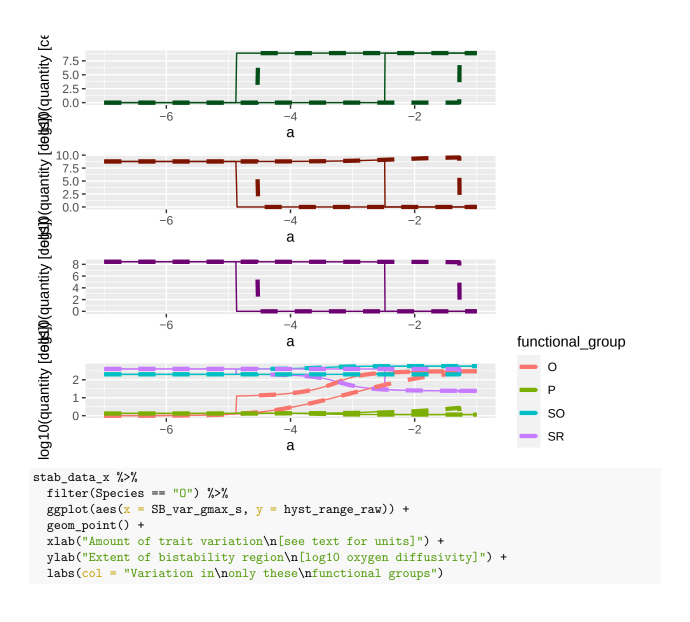
## Extra SBPB diversity

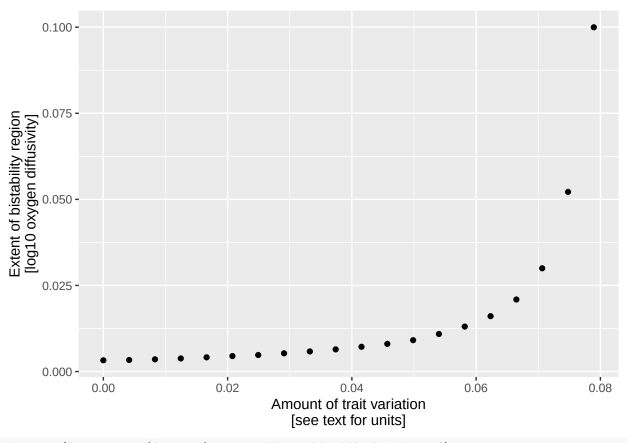
## Raw Stability

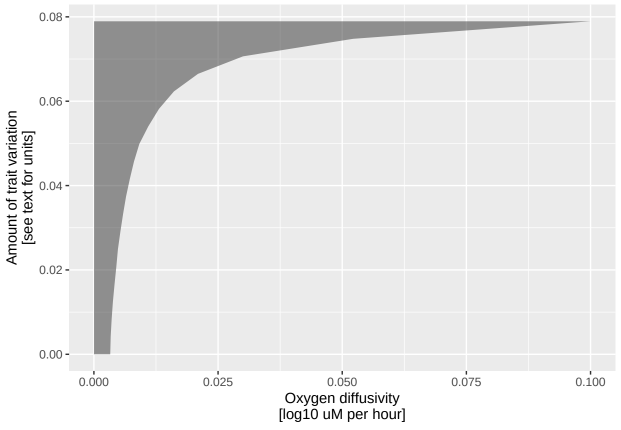


```
direction
CB_gmax_var = 0 CB_h_var =0
B_g = 0.0747922452631579 SB_h_var = -1.56wn
og10(quantity {ge伽爾hantity {ge伽爾hantity {ge伽爾hantity |
     86420
-
                                                          species
                                                              SB_1 —
                                                                        SB_4
                                                                                  SB_7
                -6
                              -4
                                             -2
                              а
                                                               SB_2
                                                                        SB_5
                                                                                  SB<sub>8</sub>
                                                               SB 3 -
                                                                        SB 6
                                                                                  SB 9
                                                          species
                                                            PB_1
                                                                        PB_4
                                                                                  PB_7
                -6
                                             -2
                              -4
                              а
                                                               PB_2
                                                                         PB_5
                                                                                  PB<sub>8</sub>
                                                                                  PB_9
                                                                    species
                <u>-</u>6
                                             <u>-</u>2
                              -4
                                                          direction
                              а
                                                            down
                                                                         SO
                                                                         SR
                                             -2
                -6
                              -4
                                                                    direction
                              а
                                                                    40.00
p_overlay1 <- plot_ss_result2(var_expt_x[1,]$ss_res[[1]],</pre>
                             var_expt_x[19,]$ss_res[[1]],
                             xlims = c(-7, -1))
## `summarise()` has grouped output by 'a', 'direction', 'var_type'. You can override using the `.group
## `summarise()` has grouped output by 'a', 'direction', 'var_type'. You can override using the `.group
```

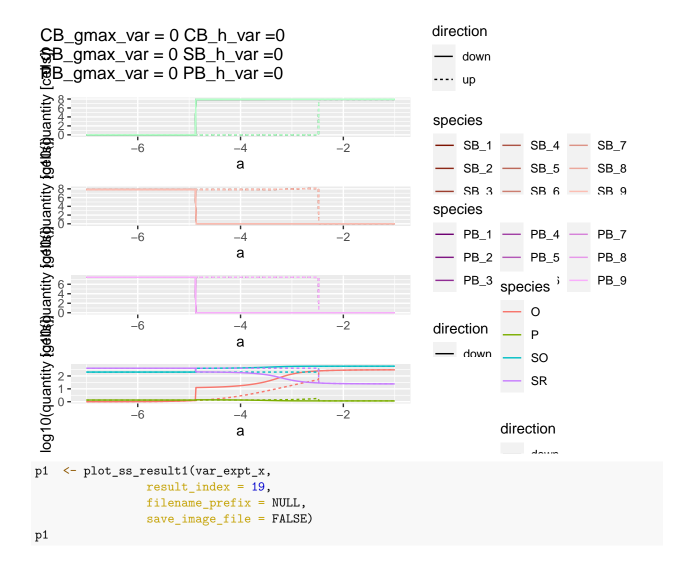
p\_overlay1





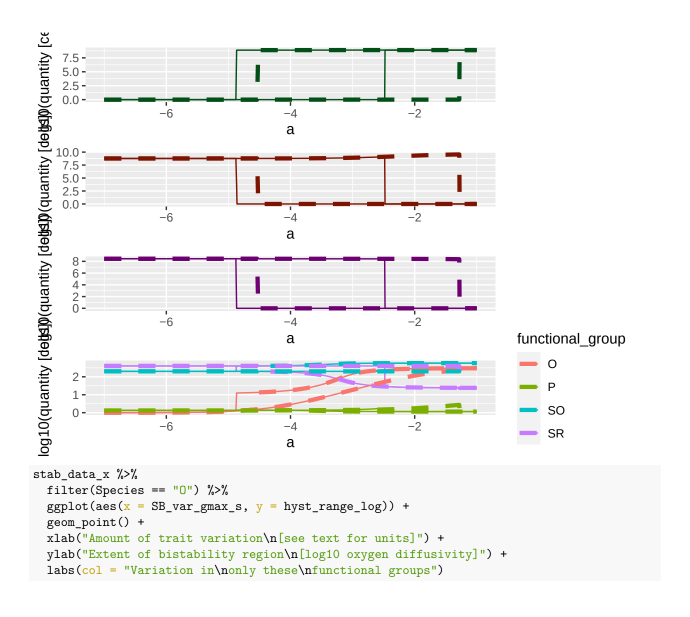


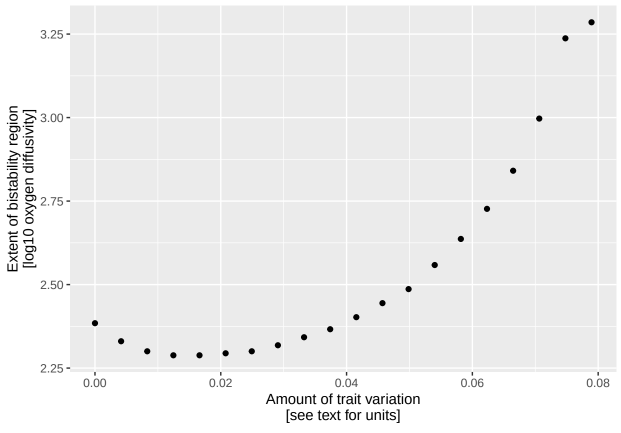
## ### Log Stability

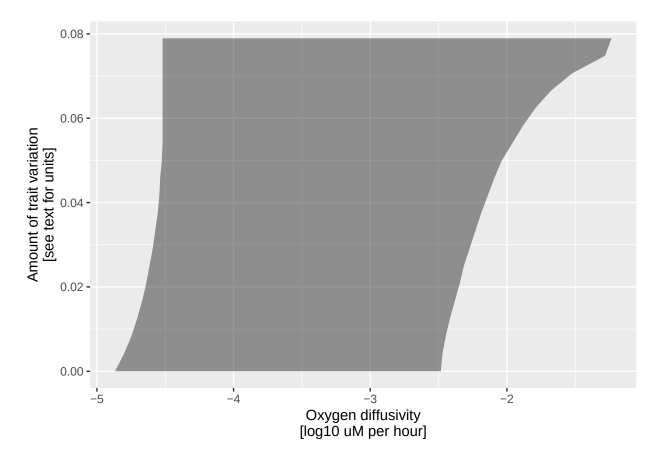


```
direction
CB_gmax_var = 0 CB_h_var =0
B_g = 0.0747922452631579 SB_h_var = -1.56wn
og10(quantity {ge伽爾hantity {ge伽爾hantity {ge伽爾hantity |
     86420
-
                                                          species
                                                              SB_1 — SB_4 -
                                                                                  SB_7
                -6
                              -4
                                             -2
                              а
                                                               SB_2
                                                                        SB_5
                                                                                  SB<sub>8</sub>
                                                               SB 3 -
                                                                        SB 6
                                                                                  SB 9
                                                          species
                                                            PB_1
                                                                        PB_4
                                                                                  PB_7
                -6
                                             -2
                              -4
                              а
                                                               PB_2
                                                                         PB_5
                                                                                  PB<sub>8</sub>
                                                                                  PB_9
                                                                    species
                <u>-</u>6
                                             <u>-</u>2
                              -4
                                                          direction
                              а
                                                            down
                                                                         SO
                                                                         SR
                                             -2
                -6
                              -4
                                                                    direction
                              а
                                                                    40.00
p_overlay1 <- plot_ss_result2(var_expt_x[1,]$ss_res[[1]],</pre>
                             var_expt_x[19,]$ss_res[[1]],
                             xlims = c(-7, -1))
## `summarise()` has grouped output by 'a', 'direction', 'var_type'. You can override using the `.group
## `summarise()` has grouped output by 'a', 'direction', 'var_type'. You can override using the `.group
```

p\_overlay1

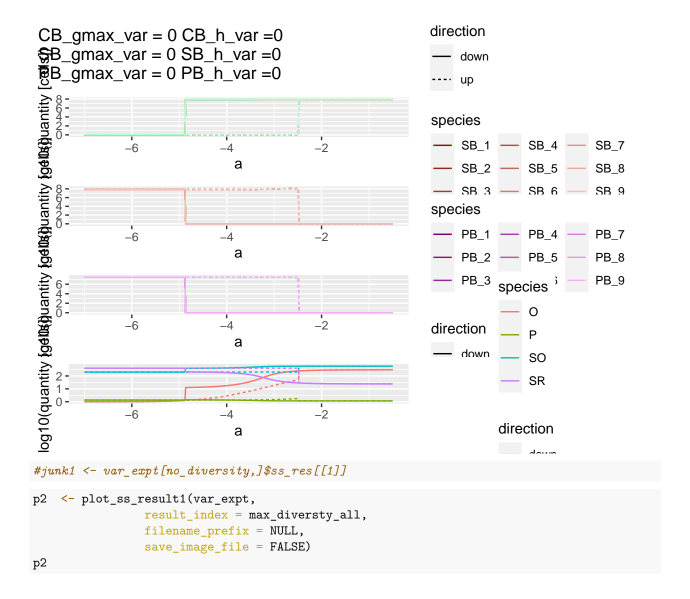


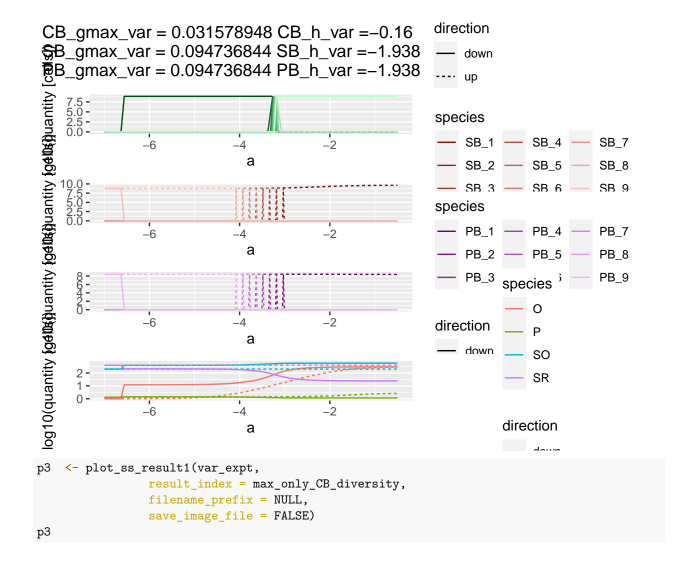


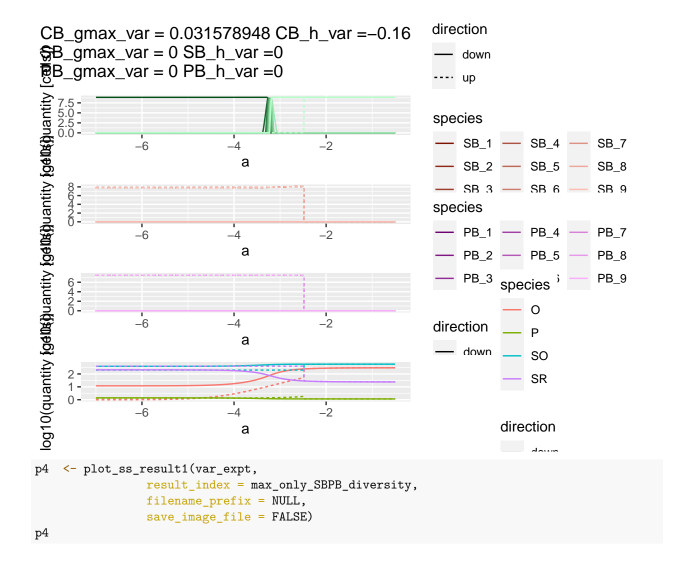


### 2x CB, 6xSBPB diversity

```
## find various combinations of diversity
var_expt <- readRDS(here("experiments/experiment 1/data/ss_data_1e6_x2x6_factorial.RDS"))</pre>
stab_data <- readRDS(here("experiments/experiment 1/data/stab_data_1e6_x2x6_factorial.RDS"))</pre>
var_expt_levels <- var_expt[,1:6]</pre>
no_diversity <- which(rowSums(abs(var_expt_levels))==0)</pre>
max_diversty_all <- which(max(rowSums(abs(var_expt_levels))) ==</pre>
                              rowSums(abs(var_expt_levels)))
max only CB diversity <- which(max(rowSums(abs(var expt levels[,1:2]))) ==</pre>
                              rowSums(abs(var_expt_levels[,1:2])) &
                                rowSums(abs(var_expt_levels[,3:6]))==0)
#var_expt_levels[381,]
max_only_SBPB_diversity <- which(max(rowSums(abs(var_expt_levels[,3:6]))) ==</pre>
                              rowSums(abs(var expt levels[,3:6])) &
                                rowSums(abs(var_expt_levels[,1:2]))==0)
#var_expt_levels[20,]
p1 <- plot_ss_result1(var_expt,</pre>
                 result_index = no_diversity,
                 filename_prefix = NULL,
                 save_image_file = FALSE)
p1
```







```
$B gmax var = 0.094736844 SB h var =−1.938
                                                                   down
     _gmax_var = 0.094736844 PB_h_var =–1.938
og10(quantity {get0k} uantity {get0k} uantity {get0k} uantity
      86420
-
                                                                 species
                                                                   — SB_1 — SB_4
                                                                                            SB 7
                 -6
                                               -2
                                  а
                                                                      SB_2
                                                                                 SB_5
                                                                                            SB<sub>8</sub>
    10.0 -
7.5 -
5.0 -
2.5 -
0.0 -
                                                                      SB 3 -
                                                                                 SB 6
                                                                                            SB 9
                                                                 species
                                                                  -- PB_1
                                                                                 PB 4
                                                                                             PB 7
                                               -2
                 -6
                                -4
                                  а
                                                                      PB_2
                                                                                  PB<sub>5</sub>
                                                                                             PB<sub>8</sub>
                                                                                             PB_9
                                                                      PB_3
                                                                            species
                                               <u>-</u>2
                 -6
                                -4
                                                                 direction
                                  а
                                                                    - down
                                                                                  SO
      2 1
                                                                                  SR
                                               <u>-2</u>
                 -6
                                -4
                                                                            direction
                                  а
                                                                             مريده
p_overlay1 <- plot_ss_result2(var_expt[no_diversity,]$ss_res[[1]],</pre>
                                 var_expt[max_diversty_all,]$ss_res[[1]],
                                 xlims = c(-7, -1))
## `summarise()` has grouped output by 'a', 'direction', 'var_type'. You can override using the `.group
## `summarise()` has grouped output by 'a', 'direction', 'var_type'. You can override using the `.group
p_overlay1
## Warning: Removed 154 row(s) containing missing values (geom_path).
## Warning: Removed 154 row(s) containing missing values (geom_path).
## Warning: Removed 154 row(s) containing missing values (geom_path).
## Warning: Removed 154 row(s) containing missing values (geom_path).
```

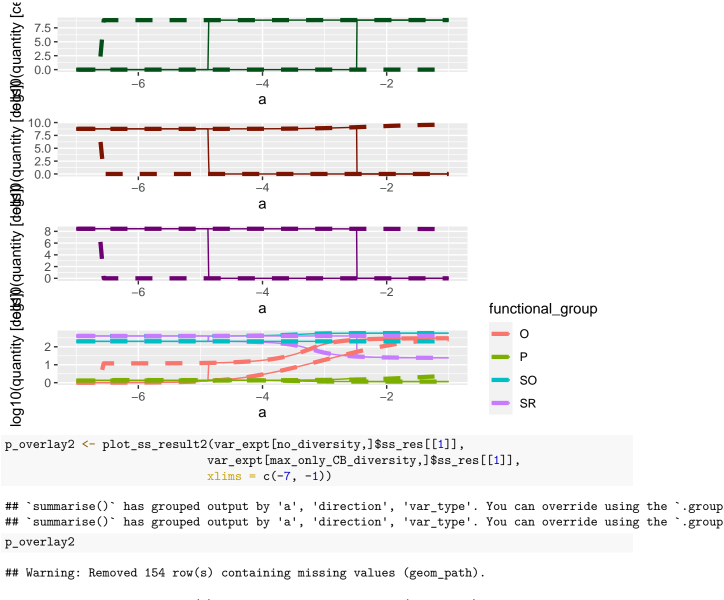
 $CB_gmax_var = 0 CB_h_var = 0$ 

direction

## Warning: Removed 154 row(s) containing missing values (geom\_path).

## Warning: Removed 154 row(s) containing missing values (geom\_path).
## Warning: Removed 616 row(s) containing missing values (geom\_path).

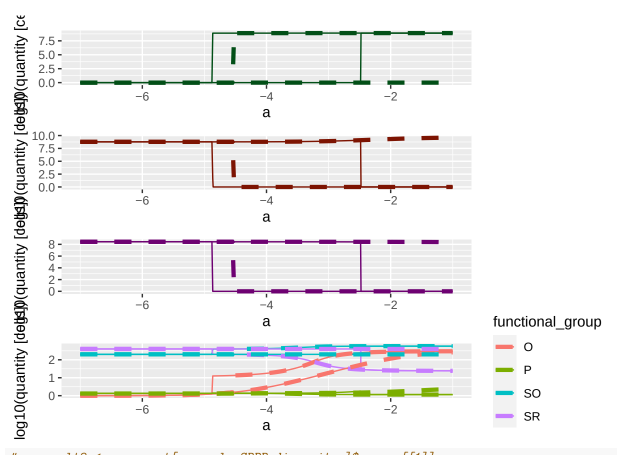
## Warning: Removed 616 row(s) containing missing values (geom\_path).



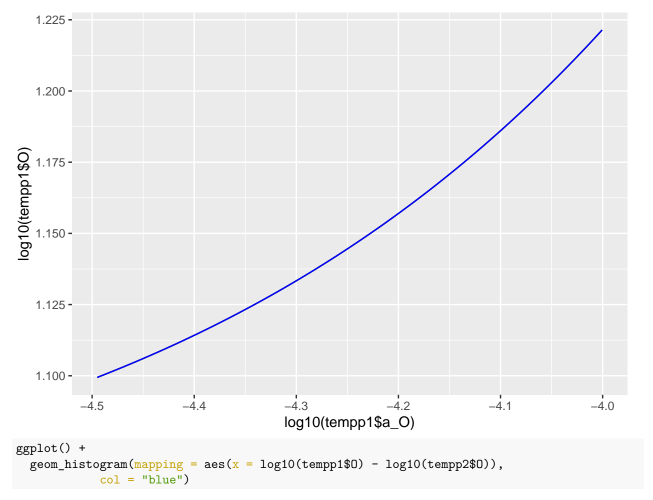
## Warning: Removed 154 row(s) containing missing values (geom\_path).
## Warning: Removed 154 row(s) containing missing values (geom\_path).
## Warning: Removed 154 row(s) containing missing values (geom\_path).
## Warning: Removed 154 row(s) containing missing values (geom\_path).
## Warning: Removed 154 row(s) containing missing values (geom\_path).
## Warning: Removed 154 row(s) containing missing values (geom\_path).
## Warning: Removed 616 row(s) containing missing values (geom\_path).
## Warning: Removed 616 row(s) containing missing values (geom\_path).

```
ce [50] (quantity [delba]) (quan
                                                     _6
                                                                                                                                                                     -2
                                                                                                             -4
                                                                                                              а
                                                                                                                                                                     <u>-</u>2
                                                                                                             -4
                                                                                                              а
                                                      -6
                                                                                                             -4
                                                                                                                                                                     -2
                                                                                                              а
                                                                                                                                                                                                                     functional_group
                                                                                                                                                                                                                                   0
                                                                                                                                                                                                                                   SO
                                                     -6
                                                                                                                                                                     -
2
                                                                                                             -4
                                                                                                                                                                                                                                   SR
                                                                                                              а
#ss_result1 <- var_expt[no_diversity,]$ss_res[[1]]</pre>
#ss_result2 <- var_expt[max_only_CB_diversity,]$ss_res[[1]]
p_overlay3 <- plot_ss_result2(var_expt[no_diversity,]$ss_res[[1]],</pre>
                                                                                         var_expt[max_only_SBPB_diversity,]$ss_res[[1]],
                                                                                        xlims = c(-7, -1))
## `summarise()` has grouped output by 'a', 'direction', 'var_type'. You can override using the `.group
## `summarise()` has grouped output by 'a', 'direction', 'var_type'. You can override using the `.group
p_overlay3
## Warning: Removed 154 row(s) containing missing values (geom_path).
## Warning: Removed 154 row(s) containing missing values (geom_path).
## Warning: Removed 154 row(s) containing missing values (geom_path).
## Warning: Removed 154 row(s) containing missing values (geom_path).
## Warning: Removed 154 row(s) containing missing values (geom_path).
## Warning: Removed 154 row(s) containing missing values (geom_path).
## Warning: Removed 616 row(s) containing missing values (geom_path).
```

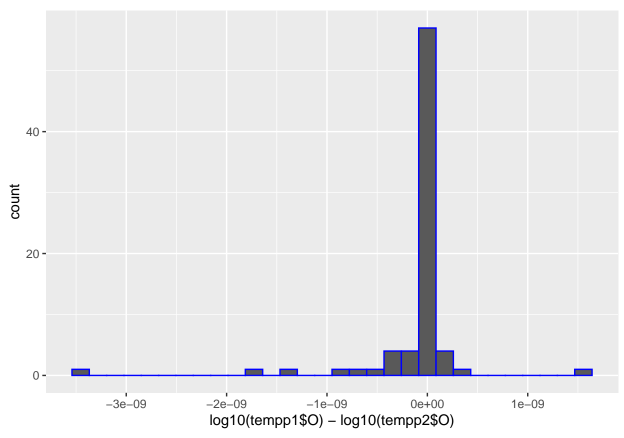
## Warning: Removed 616 row(s) containing missing values (geom\_path).



 $\#ss\_result3 \ <- \ var\_expt[max\_only\_SBPB\_diversity,] \$ss\_res[[1]]$ 



## `stat\_bin()` using `bins = 30`. Pick better value with `binwidth`.



```
ggplot() +
  geom_line(mapping = aes(x = log10(tempp1$a_0),
      y = log10(tempp1$P) - log10(tempp2$P)),
      col = "blue")
```

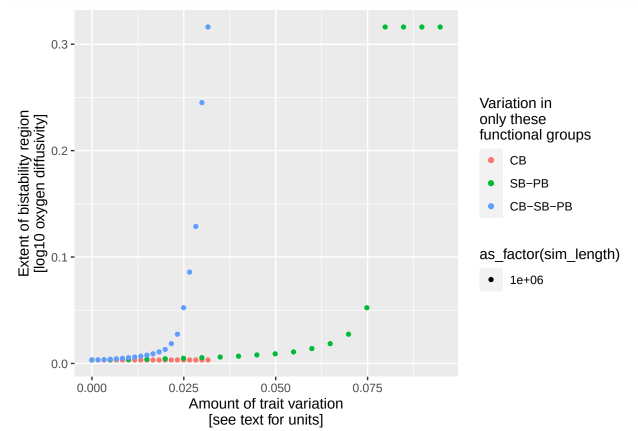
```
4e-09 -
log10(tempp1$P) - log10(tempp2$P)
   3e-09 -
   2e-09 -
    1e-09 -
   0e+00 -
                                           -4.3
                                                                           -4.1
           -4.5
                           -4.4
                                                                                           -4.0
                                           log10(tempp1$a_O)
CB_vars <- unique(stab_data$CB_var_gmax_s)</pre>
SB_vars <- unique(stab_data$SB_var_gmax_s)</pre>
CB_stab_data <- stab_data %>%
  filter(SB_var_gmax_s == 0) %>%
  mutate(var_treat = "CB",
         var_gmax = CB_var_gmax_s)
SBPB_stab_data <- stab_data %>%
  filter(CB_var_gmax_s == 0) %>%
  mutate(var_treat = "SB-PB",
         var_gmax = SB_var_gmax_s)
for_join <- tibble(CB_var_gmax_s = CB_vars,</pre>
                       SB_var_gmax_s = SB_vars)
CBSBPB_stab_data <- stab_data %>%
  right_join(for_join) %>%
  mutate(var_treat = "CB-SB-PB",
          var_gmax = CB_var_gmax_s)
## Joining, by = c("CB_var_gmax_s", "SB_var_gmax_s")
all_stab_results <- CB_stab_data %>%
  bind_rows(SBPB_stab_data) %>%
# bind_rows(results3) %>%
 # bind_rows(results4) %>%
```

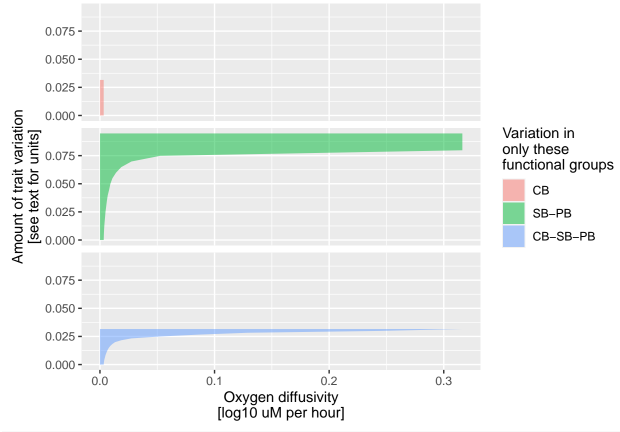
bind\_rows(CBSBPB\_stab\_data)

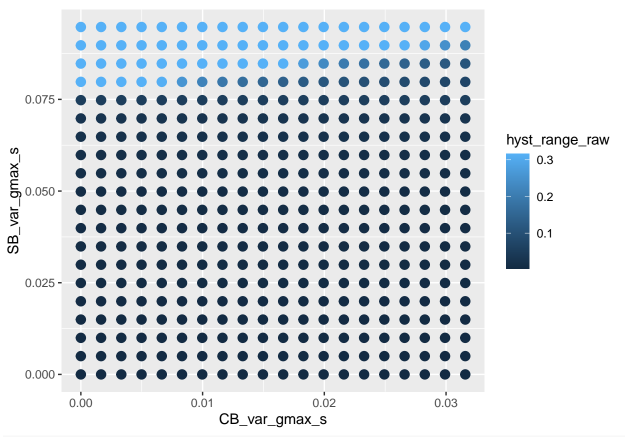
```
all_stab_results<- all_stab_results %>%
    mutate(var_treat = forcats::fct_relevel(var_treat, levels = c("CB", "SB-PB", "CB-SB-PB")))
## Warning: Outer names are only allowed for unnamed scalar atomic inputs
#saveRDS(all_stab_results, here("experiments/experiment summary/all_stab.RDS"))
#all_stab_results <- readRDS(here("experiments/experiment summary/all_stab.RDS"))</pre>
```

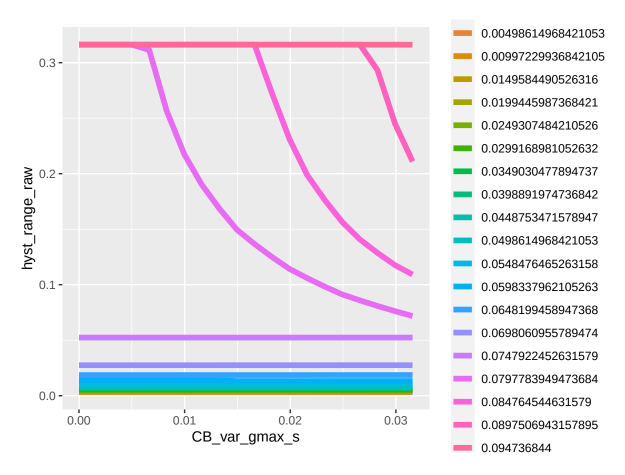
#### Raw Stability

```
all_stab_results %>%
  filter(Species == "0") %>%
  ggplot(aes(x = var_gmax, y = hyst_range_raw, col=var_treat, shape = as_factor(sim_length))) +
  geom_point() +
  xlab("Amount of trait variation\n[see text for units]") +
  ylab("Extent of bistability region\n[log10 oxygen diffusivity]") +
  labs(col = "Variation in\nonly these\nfunctional groups")
```



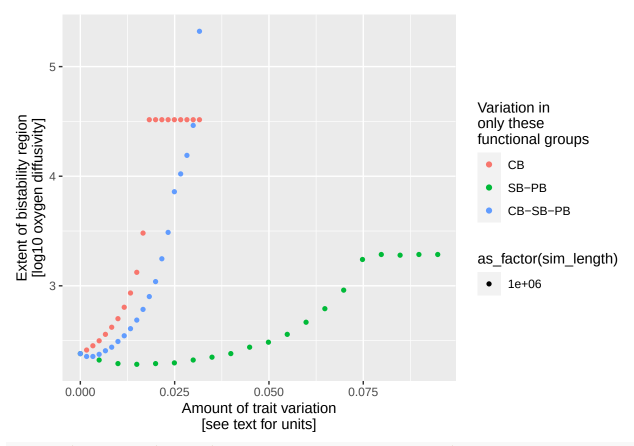




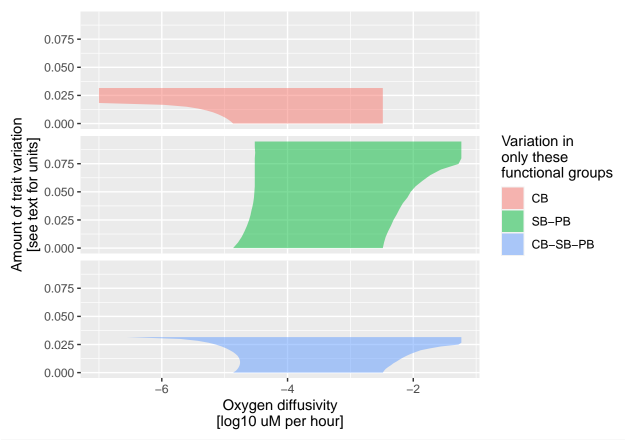


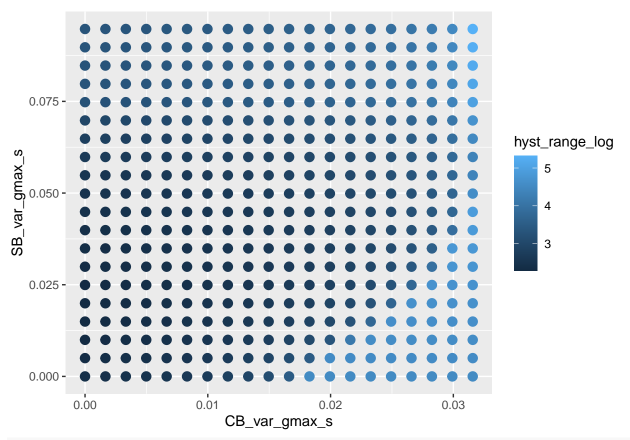
### Log Stability

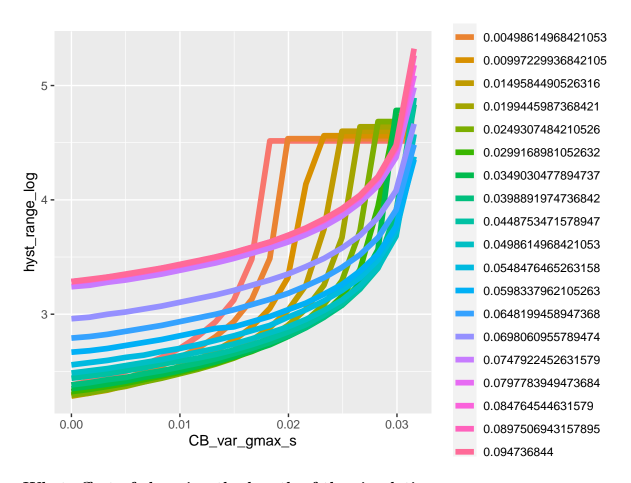
```
all_stab_results %>%
  filter(Species == "0") %>%
  ggplot(aes(x = var_gmax, y = hyst_range_log, col=var_treat, shape = as_factor(sim_length))) +
  geom_point() +
  xlab("Amount of trait variation\n[see text for units]") +
  ylab("Extent of bistability region\n[log10 oxygen diffusivity]") +
  labs(col = "Variation in\nonly these\nfunctional groups")
```



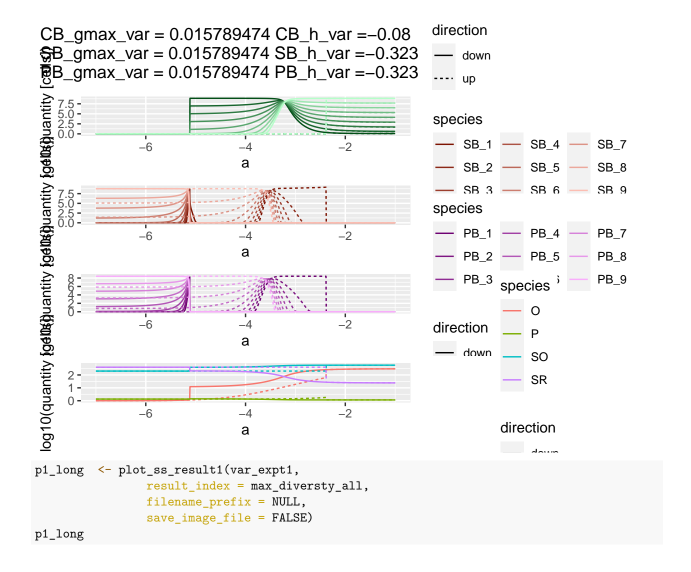
```
##ggsave("manuscript/figures/extent_of_bistab1.pdf", height = 4)
all_stab_results %>%
  #filter(var treat == "CB") %>%
  filter(Species == "0",
         sim_length == 1e6) %>%
  ggplot(aes(x = var_gmax,
             ymin = hyst_min_log,
             ymax = hyst_max_log,
             fill=var_treat)) +
  geom_ribbon(alpha = 0.5) +
  facet_wrap( ~ var_treat, nrow = 3) +
  xlab("Amount of trait variation\n[see text for units]") +
  ylab("Oxygen diffusivity\n[log10 uM per hour]") +
  labs(fill = "Variation in\nonly these\nfunctional groups") +
  coord_flip() +
  theme(
    strip.background = element_blank(),
    strip.text.x = element_blank()
 )
```

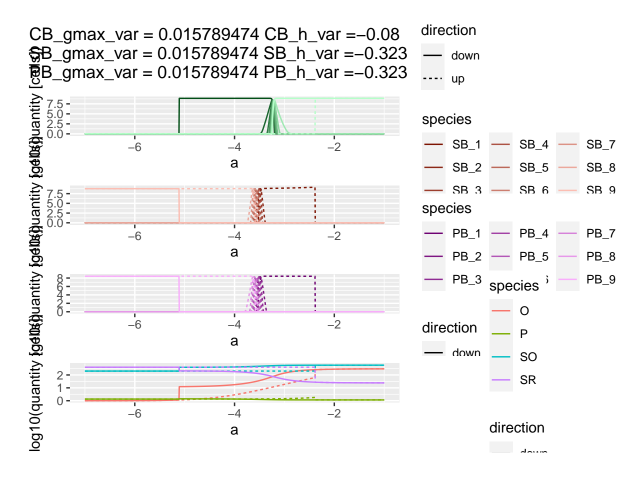






### What effect of changing the length of the simulations





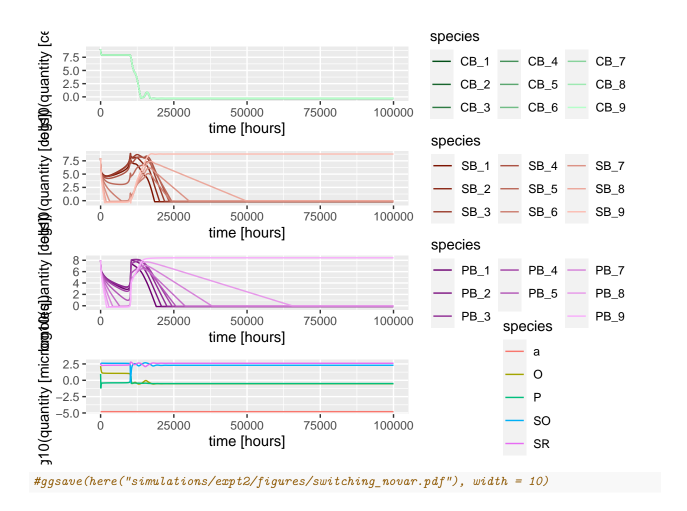
# Some puzzles

#### Puzzle 1

When the environment ameliorates for the sulphur bacteria, there is no strain replacement (in final state) along the oxygen diffusivity gradient – either the system is oxic or the least tolerant strain SB9 dominates. And yet the switch to anoxic occurs earlier than when there is no diversity, which suggests there is some role of the more tolerant strains. Indeed Uriah showed that the presence of only the most tolerant strain is sufficient to give an earlier switch, even though it is not present in the final state when less tolerant strains are present. And he showed that the presence of only the least tolerant strain creates a later switch than when there are more tolerant strains.

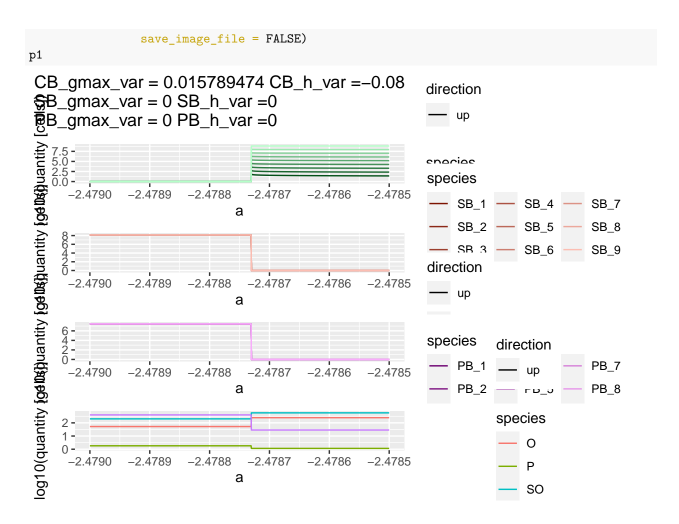
The explanation is that the most tolerant strain does play a role, but only a transient one. The following dynamics are for the system starting oxic, and with a value of oxygen diffusivity ( $log10(a_O) = -4.8$ ) for which the system remains oxic if there is no diversity, but switches to anoxic if there is diversity. There is only diversity in the sulphur bacteria. The most tolerant strain does at first grow, but is then outcompeted by less tolerant strains as the environment ameliorates (temporally).

```
max_only_SBPB_diversity <- which(max(rowSums(abs(var_expt_levels[,3:6]))) ==</pre>
                                 rowSums(abs(var expt levels[,3:6])) &
                                   rowSums(abs(var_expt_levels[,1:2]))==0)
default_sim_duration <- 100000</pre>
default_log10a_series \leftarrow c(-4.8, -4.8)
initial_state <- new_initial_state(num_CB_strains,</pre>
                                          num_PB_strains,
                                         num_SB_strains,
                                          values = "bush_ssfig3")
initial_state[grep("CB_", names(initial_state))] <- 10^10/num_CB_strains</pre>
sim_res_novar1 <- run_simulation(parameter_values = var_expt$pars[[no_diversity]],</pre>
                               initial_state = initial_state)
saveRDS(sim_res_novar1, here("experiments/experiment 1/data/puzzle1_1.RDS"))
sim_res_novar1 <- readRDS(here("experiments/experiment 1/data/puzzle1_1.RDS"))</pre>
plot_dynamics(sim_res_novar1)
310(quantity [micrlogototeq])antity [debs])(quantity [debs])(quantity [ce
                                                                    species
     9.0 -
                                                                                 — CB_4
                                                                         CB_1 -
                                                                                                CB_7
    8.7 -
     8.4 -
                                                                         CB<sub>2</sub>
                                                                                    CB_5
                                                                                                CB<sub>8</sub>
     8.1 -
                                                                         CB<sub>3</sub>
                                                                                    CB_6
                                                                                                CB_9
                     25000
                                 50000
                                                          100000
                                              75000
                              time [hours]
                                                                    species
      6 -
                                                                                    SB_4
                                                                         SB_1 -
                                                                                                SB_7
      4 -
                                                                         SB 2
                                                                                    SB 5
                                                                                                SB 8
                     25000
                                 50000
                                              75000
                                                          100000
                                                                         SB<sub>3</sub>
                                                                                    SB<sub>6</sub>
                                                                                                SB<sub>9</sub>
                              time [hours]
                                                                    species
      8 -
      6 -
                                                                         PB 1 -
                                                                                    PB_4
                                                                                                PB 7
      4 -
                                                                         PB<sub>2</sub>
                                                                                    PB_5
                                                                                                PB_8
                                                                                                PB_9
                                                                         PB<sub>3</sub>
                     25000
                                  50000
                                              75000
                                                          100000
          0
                                                                                species
                              time [hours]
                                                                                     а
     2.5 -
                                                                                     0
     0.0 -
    -2.5 -
                                                                                     Ρ
    -5.0 ·
                                                                                     SO
                     25000
                                  50000
                                              75000
                                                          100000
                              time [hours]
                                                                                     SR
#qqsave(here("simulations/expt2/figures/switching_novar.pdf"), width = 10)
sim_res_novar1 <- run_simulation(parameter_values = var_expt$pars[[max_only_SBPB_diversity]],</pre>
                               initial_state = initial_state)
saveRDS(sim_res_novar1, here("experiments/experiment 1/data/puzzle1_2.RDS"))
sim res novar1 <- readRDS(here("experiments/experiment 1/data/puzzle1 2.RDS"))</pre>
plot_dynamics(sim_res_novar1)
```



### Zoom in on SS

```
a_0s <- 10^seq(-2.479, -2.4785, length=grid_num_a) ## sequence of a_0 values
initial_CBs <- 1#10^seq(0, 0, length=grid_num_N) ## sequence of N values
initial_PBs <- 1e8 ## not varied</pre>
initial_SBs <- 1e8 ## not varied</pre>
# next line creates all possible combinations
ss_expt <- expand.grid(N_CB = initial_CBs,</pre>
                       N_PB = initial_PBs,
                       N_SB = initial_SBs,
                       a \ 0 = a \ 0s)
var expt master <- var expt</pre>
var_expt <- var_expt_master[381,]</pre>
#var_expt <- run_ss_var_experiment()</pre>
#saveRDS(var_expt, here("experiments/experiment 1/data/ss_data_zoom.RDS"))
library(here)
zoom <- readRDS(here("experiments/experiment 1/data/ss_data_zoom.RDS"))</pre>
p1 <- plot_ss_result1(zoom,</pre>
                 result_index = 1,
                 filename_prefix = NULL,
```



# Negative abundance investigation

I (Owen) found that the sampling interval had an effect on the stability of the simulation. If the sampling interval was long, then in some rare cases (see below) the odesolver failed, with negative abundances occurring. I think this is due to abundances becoming very small, and then the computer having trouble with precision. I guess that when a sample is taken, the abundance is somehow altered if it is very low, probably by some rounding.

```
var_expt$pars[[1]]
dd <- var_expt$ss_res[[1]]
dd1 <- filter(dd, PB_1<(-0.0001))
dd1$a_0

ss_expt_master <- ss_expt
ss_expt <- ss_expt_master[abs(ss_expt_master$a_0 - 1.336984e-05)<1e-10,]

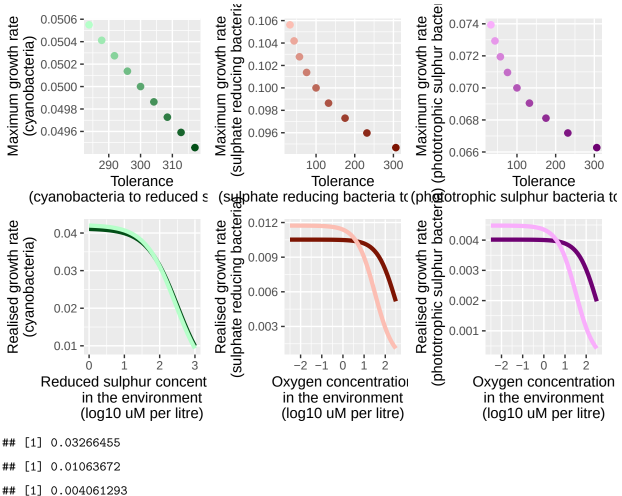
var_expt_master <- var_expt
#var_expt <- var_expt[1,]
var_expt_test <- run_ss_var_experiment()
res <- var_expt_test$ss_res[[1]]

test1 <- ss_by_a_N(ss_expt, var_expt$pars[[1]])
x <- ss_expt[2,]</pre>
```

```
param <- var_expt$pars[[1]]</pre>
get_final_states_a_N(x, param)
ssfind_parameters <- param</pre>
ssfind_simulation_sampling_interval <- 1000
## now run inside the function "get_final_states_a_N"
simres1 <- simres
ssfind_simulation_sampling_interval <- 5000
## now run inside the function "get final states a N"
simres2 <- simres # this fails</pre>
## now run inside the function "get_final_states_a_N"
plot_dynamics(simres2)
ggplot() +
  geom_line(data = simres1$result,
             mapping = aes(x = time, y = log10(PB_1))) +
  geom_point(data = simres2$result,
             mapping = aes(x = time, y = log10(PB_1))) +
  xlim(c(0, 250000))
ccc <- simres2$result
simres2$result$PB 1
simres2$result$time
log10_a <- log10(ss_expt$a_0[1]) ## very slowly goes anoxic</pre>
\#log10_a \leftarrow log10(a_0s[354]) \# very slowly goes anoxic
\#log10_a \leftarrow log10(a_0s[356]) \# very very very slowly goes anoxic
\#log10_a \leftarrow log10(a_0s[357]) \#\# does not go anoxic
default_dynamic_model <- bushplus_dynamic_model</pre>
default_event_definition <- event_definition_1</pre>
default_event_interval <- ssfind_simulation_duration</pre>
default_noise_sigma <- 0</pre>
default_minimum_abundances <- ssfind_minimum_abundances</pre>
default_sim_duration <- ssfind_simulation_duration</pre>
default_sim_sample_interval <- ssfind_simulation_duration</pre>
#initial_pars_from <- "bush_ssfig3"</pre>
default_log10a_series <- c(log10_a, log10_a)</pre>
initial_state <- new_initial_state(num_CB_strains,</pre>
                                     num_PB_strains,
                                     num_SB_strains,
                                     values = "bush_ssfig3")
initial_state[grep("CB_", names(initial_state))] <- 10^10/num_CB_strains
sim_res_novar <- run_simulation(parameter_values = var_expt$pars[[1]],</pre>
                            initial_state = initial_state)
plot_dynamics(sim_res_novar)
```

```
simulation_result <- sim_res_novar</pre>
every_n <- 1
chk <- sim_res_novar$result</pre>
sim_res_novar$result %>%
  ggplot() +
  geom_line(mapping = aes(x = time, y = PB_1))
#qqsave(here("simulations/expt2/figures/switching_novar.pdf"), width = 10)
```

## Understand about relative and absolute variation in traits



With the CB diversity multiplier set at 1 and the SB/PB multiplier set at 5 the range of realised growth rates of CB is 0.0326645, range of SB is 0.0106367, and range of PB is 0.0040613.