# Supplemental Material for Adaptive phenotypic plasticity stabilizes evolution in fluctuating environments

Alexander Lalejini, Austin J. Ferguson, Nkrumah A. Grant, and Charles Ofria

2021 - 05 - 25

# Contents

1	Intr 1.1 1.2 1.3	About our supplemental material	7
2		a availability Source code	
	2.1 2.2	Source code	
3	Con	pile and run experiments locally 11	L
	3.1	Docker	L
4	Avio	la instruction set	3
	4.1	Default instructions	3
	4.2	Custom instructions	Ĺ
5	Vali	dation experiment 17	7
	5.1	Overview	7
	5.2	Analysis dependencies	3
	5.3	Setup	3
	5.4	Evolution of phenotypic plasticity	)
6	Evo	lutionary change 23	3
	6.1	Overview	3
	6.2	Analysis dependencies	3
	6.3	Setup	l
	6.4	The evolution of phenotypic plasticity	;
	6.5	Average generation	3
	6.6	Coalescence event count	L
	6.7	Phenotypic volatility along the dominant lineage	)
	6.8	Phenotypic fidelity	í
	6.9	Mutation count	)
	6.10	Genotypic fidelity	;
	6 11		)

4 CONTENTS

	6.12	Mutational robustness	$7^{2}$
	6.13	Manuscript figures	74
7	Evo	lution and maintenance of novel traits	79
•	7.1		79
	$7.1 \\ 7.2$		80
	7.2	· · · · · · · · · · · · · · · · · · ·	81
	7.3 - 7.4	-	83
	$7.4 \\ 7.5$		
			85
	7.6		89
	7.7	* ( 0 /	92
	7.8	v (1 1 )	95
	7.9	Novel task discovery frequency (lineage)	
		Novel tasks gained (lineage)	
		Novel task loss (lineage)	
		Frequency of novel task loss (lineage)	08
	7.13	How many instances of novel trait loss co-occurred with changes	
		in base phenotype?	
		Manuscript figures	
	7.15	Combined panel	15
8	Acc	umulation of deleterious instructions 1	19
	8.1	Overview	19
	8.2	Analysis dependencies	19
	8.3	Setup	
	8.4	Evolution of phenotypic plasticity	
	8.5	Poison instruction execution	
	8.6	Characterizing mutations that increase poison instruction execution 1	
	8.7	What fraction of poison execution increases occur in unexpressed	
		phenotype (as cryptic variation)?	54
	8.8	Manuscript figures	
	0.0	1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1	
9	_	ulation in Avida 1	
	9.1	Overview	
	9.2	Analysis dependencies	
	9.3	Setup	58
	9.4	How many instructions do plastic genomes toggle depending on	
			59
	9.5	What is the distribution of toggled sequence sizes? 1	60
10	Evo	lutionary change (variable length genomes) 1	63
		Overview	
		Analysis dependencies	
		Setup	
		Evolution of phenotypic plasticity	
		Genome length	

10.6	Average generation
10.7	Coalescence event count
10.8	Phenotypic volatility along the dominant lineage 176
10.9	Mutation count (along dominant lineage) 179

6 CONTENTS

## Introduction

This is the supplemental material for our work entitled, Adaptive phenotypic plasticity stabilizes evolution in fluctuating environments. Preprint forthcoming.

## 1.1 About our supplemental material

This supplemental material is hosted on GitHub using GitHub pages. The source code and configuration files used to generate this supplemental material can be found in this GitHub repository. We compiled our data analyses and supplemental documentation into this nifty web-accessible book using bookdown.

Our supplemental material includes the following:

- Data availability (Section 2)
- Guide for running our experiments (Section 3)
- Avida instruction set (Section 4)
- Experiment analyses (including source code):
  - Validating the evolution of phenotypic plasticity (Section 5)
  - Effect of adaptive phenotypic plasticity on evolutionary change (Section 6)
    - \* Results with variable-length genomes (Section 10)
  - Effect of adaptive phenotypic plasticity on the evolution and maintenance of novel traits (Section 7)
  - Effect of adaptive phenotypic plasticity on the accumulation of deleterious instructions (Section 8)
  - Exploring how regulation is encoded in genomes in Avida (Section 9)

## 1.2 Contributing authors

• Alexander Lalejini

- Austin J. Ferguson
- Nkrumah A. Grant
- Charles Ofria

#### 1.3 Research overview

Abstract: > Fluctuating environmental conditions are ubiquitous in natural systems, and populations have evolved various strategies to cope with such fluctuations. The particular mechanisms that evolve profoundly influence subsequent evolutionary dynamics. One such mechanism is phenotypic plasticity, which is the ability of a single genotype to produce alternate phenotypes in an environmentally dependent context. Here, we use digital organisms (selfreplicating computer programs) to investigate how adaptive phenotypic plasticity alters evolutionary dynamics and influences evolutionary outcomes in cyclically changing environments. Specifically, we examined the evolutionary histories of both plastic populations and non-plastic populations to ask: (1) Does adaptive plasticity promote or constrain evolutionary change? (2) Are plastic populations better able to evolve and then maintain novel traits? And (3), how does adaptive plasticity affect the potential for maladaptive traits to accumulate in evolving genomes? We find that populations with adaptive phenotypic plasticity undergo less evolutionary change than non-plastic populations, which must rely on genetic variation from de novo mutations to continuously readapt to environmental fluctuations. Indeed, the non-plastic populations undergo more frequent selective sweeps and accumulate many more genetic changes. We find that the repeated selected sweeps in non-plastic populations drive the loss of beneficial traits via deleterious hitchhiking, whereas phenotypic plasticity can stabilize populations against environmental fluctuations. This stabilization allows plastic populations to more easily retain novel adaptive traits than their non-plastic counterparts. In general, the evolution of adaptive phenotypic plasticity shifted evolutionary dynamics to be more similar to that of populations evolving in a static environment than to non-plastic populations evolving in an identical fluctuating environment. All natural environments subject populations to some form of change; our findings suggest that the stabilizing effect of phenotypic plasticity plays an important role in subsequent adaptive evolution.

# Data availability

### 2.1 Source code

The source code for this work is publicly accessible on GitHub: https://github. com/amlalejini/evolutionary-consequences-of-plasticity

## 2.2 Experimental results

The data from our experiments are available online in our OSF repository (Lalejini and Ferguson, 2021) at https://osf.io/sav2c/.

# Compile and run experiments locally

Here, we provide a brief guide to compiling and running our experiments using our Docker image.

Please file an issue on GitHub if something is unclear or does not work.

#### 3.1 Docker

You can use the Dockerfile in our repository to build a docker image locally, or you can pull the latest docker image from DockerHub using

docker pull amlalejini/evolutionary-consequences-of-plasticity

This will pull down a docker image with:

- all of the requisite dependencies installed/downloaded
- all experiment source code
- the minimal set of raw data needed to compile the supplemental material
- a build of our supplemental material (which will also run all of our analyses)

To run the container interactively:

docker run -it --entrypoint bash amlalejini/evolutionary-consequences-of-plasticity

You can exit the container at any point with ctrl-d.

Inside the container, you should be able to navigate to /opt/evolutionary-consequences-of-plasticity:

cd /opt/evolutionary-consequences-of-plasticity

To run Avida, you'll need to cd into the avida directory and run ./build\_avida.

All of the Avida configuration files necessary for re-running our experiments can be found here:  $\frac{\text{https:}}{\text{github.com/amlalejini/evolutionary-consequences-of-plasticity/tree/master/experiments}}.$ 

For example, the configuration files for our evolutionary change experiment are here: https://github.com/amlalejini/evolutionary-consequences-of-plasticity/tree/master/experiments/2021-02-08-evo-dynamics/hpcc/config.

# Avida instruction set

#### 4.1 Default instructions

We used the following default instructions in all of our experiments:

```
# No-ops
INST nop-A
INST nop-B
INST nop-C
# Flow control operations
INST if-n-equ
INST if-less
INST if-label
INST mov-head
INST jmp-head
INST get-head
INST set-flow
# Single Argument Math
{\tt INST \ shift-r}
INST shift-l
INST inc
INST dec
INST push
INST pop
INST swap-stk
INST swap
```

# Double Argument Math

INST add
INST sub
INST nand

# Biological Operations
INST h-copy
INST h-alloc
INST h-divide

# I/O and Sensory
INST IO
INST h-search

Each of these instructions is described in the Avida documentation.

#### 4.2 Custom instructions

We implemented several custom instructions for this work:

#### • INST sense-react-NAND

 Provides sensory feedback on whether the NAND Boolean logic task is currently rewarded or punished by pushing a 1 to the organism's active stack if it is rewarded, a -1 if it is punished, and a 0 if it is neither rewarded nor punished.

#### • INST sense-react-NOT

 Provides sensory feedback on whether the NOT Boolean logic task is currently rewarded or punished by pushing a 1 to the organism's active stack if it is rewarded, a -1 if it is punished, and a 0 if it is neither rewarded nor punished.

#### • INST sense-react-AND

 Provides sensory feedback on whether the AND Boolean logic task is currently rewarded or punished by pushing a 1 to the organism's active stack if it is rewarded, a -1 if it is punished, and a 0 if it is neither rewarded nor punished.

#### • INST sense-react-ORN

 Provides sensory feedback on whether the ORN Boolean logic task is currently rewarded or punished by pushing a 1 to the organism's active stack if it is rewarded, a -1 if it is punished, and a 0 if it is neither rewarded nor punished.

#### • INST sense-react-OR

 Provides sensory feedback on whether the OR Boolean logic task is currently rewarded or punished by pushing a 1 to the organism's active stack if it is rewarded, a -1 if it is punished, and a 0 if it is neither rewarded nor punished.

#### • INST sense-react-ANDN

- Provides sensory feedback on whether the ANDN Boolean logic task

is currently rewarded or punished by pushing a 1 to the organism's active stack if it is rewarded, a -1 if it is punished, and a 0 if it is neither rewarded nor punished.

#### • INST poison

 Each time poison is executed, the organism reduces the metabolic rate of the organism by a fixed rate (specified by POISON\_PENALTY in avida.cfg).

# Validation experiment

In this experiment, we validate that (1) we observe the evolution of phenotypic plasticity in a changing environment when digital organisms have access to sensory instructions (capable of differentiating environmental states) and (2) that adaptive phenotypic plasticity does not evolve when populations lack access to sensory instructions.

#### 5.1 Overview

```
total_updates <- 200000
replicates <- 100

all_traits <- c("not", "nand", "ornot", "or", "andnot")
traits_set_a <- c("not", "and", "or")
traits_set_b <- c("nand", "ornot", "andnot")

# Relative location of data.
working_directory <- "experiments/2021-01-07-validation/analysis/"
# working_directory <- "./"
# << For bookdown
# <- For local analysis</pre>
```

We evolved populations of digital organisms under four conditions:

- 1. A fluctuating environment with access to sensory instructions
- 2. A fluctuating environment without access to sensory instructions (i.e., sensory instructions are no-operations)
- 3. A constant environment with access to sensory instructions
- 4. A constant environment without access to sensory instructions

In fluctuating environments, we alternate between rewarding and punishing different sets of computational tasks. In one environment, we reward tasks not,

and, or and punish tasks nand, ornot, andnot. In the alternative environment, we reward tasks nand, ornot, andnot and punish tasks not, and, or. In constant environments, we reward all tasks (not, nand, and, ornot, or, andnot).

For each replicate of each condition, we extract the dominant (i.e., most numerous) genotype at the end of the run to analyze further. We expect to observe the evolution of adaptive phenotypic plasticity in only the first experimental condition. In conditions without sensors, plasticity in any form should be unable to evolve.

### 5.2 Analysis dependencies

Load all required R libraries.

```
library(ggplot2)
library(tidyverse)
library(cowplot)
source("https://gist.githubusercontent.com/benmarwick/2a1bb0133ff568cbe28d/raw/fb53bd9
```

These analyses were conducted/knitted with the following computing environment:

```
print(version)
```

```
x86 64-pc-linux-gnu
## platform
## arch
                  x86_64
## os
                  linux-gnu
## system
                  x86_64, linux-gnu
## status
                  4
## major
## minor
                  1.0
                  2021
## year
## month
                  05
                  18
## day
                  80317
## svn rev
## language
## version.string R version 4.1.0 (2021-05-18)
## nickname
                  Camp Pontanezen
```

## 5.3 Setup

```
data_loc <- paste0(working_directory, "data/aggregate.csv")
data <- read.csv(data_loc, na.strings="NONE")
data$DISABLE_REACTION_SENSORS <- as.factor(data$DISABLE_REACTION_SENSORS)</pre>
```

5.3. SETUP 19

```
data$chg_env <- as.factor(data$chg_env)</pre>
data$dom_plastic_odd_even <- as.factor(data$dom_plastic_odd_even)</pre>
data$sensors <- data$DISABLE_REACTION_SENSORS == "0"</pre>
data$is_plastic <- data$dom_plastic_odd_even == "True"</pre>
env_label_fun <- function(chg_env) {</pre>
 if (chg_env) {
    return("Fluctuating")
  } else {
    return("Constant")
 }
}
sensors_label_fun <- function(has_sensors) {</pre>
  if (has_sensors) {
   return("Sensors")
 } else {
    return("No sensors")
 }
}
# Count observed plasticity for each condition (I'm sure there's a 'tidier' way to do this..)
observed_plasticity <- data.frame(</pre>
  environment=character(),
  sensors=character(),
  plastic=integer(),
  nonplastic=integer(),
  plastic_adaptive=integer(),
  plastic optimal=integer(),
  plastic_nonadaptive=integer()
for (env_chg in levels(data$chg_env)) {
  for (disabled_sensors in levels(data$DISABLE_REACTION_SENSORS)) {
    cond_data <- filter(data, chg_env == env_chg & data$DISABLE_REACTION_SENSORS == disabled_sens
    environment_label <- env_label_fun(env_chg)</pre>
    sensors_label <- sensors_label_fun(disabled_sensors == "0")</pre>
    observed_plasticity <- observed_plasticity %>% add_row(
      environment=environment_label,
      sensors=sensors_label,
      plastic=nrow(filter(cond_data, is_plastic==TRUE)),
      nonplastic=nrow(filter(cond_data, is_plastic==FALSE)),
      plastic_adaptive=nrow(filter(cond_data, dom_adaptive_plasticity=="True")),
      plastic_optimal=nrow(filter(cond_data, dom_optimal_plastic=="True")),
      plastic_nonadaptive=nrow(filter(cond_data, is_plastic==TRUE & dom_adaptive_plasticity=="Fal
```

```
}

observed_plasticity <- pivot_longer(
  observed_plasticity,
  cols=c("plastic", "plastic_adaptive", "plastic_optimal", "plastic_nonadaptive", "nongonames_to="phenotype",
  values_to="phenotype_cnt"
)

####### misc ######

# Configure our default graphing theme
theme_set(theme_cowplot())
</pre>
```

## 5.4 Evolution of phenotypic plasticity

For each experimental condition, do we observe the evolution of phenotypic plasticity? To test for phenotypic plasticity, we culture digital organisms in both environments from the fluctuating condition (including organisms evolved in a constant environment). Any plasticity that we observe from digital organisms evolved under constant conditions is cryptic variation (as these organisms were never exposed to these culturing environments).

```
ggplot(filter(observed_plasticity, phenotype %in% c("plastic", "nonplastic")), aes(x=p)
  geom_bar(
    stat="identity",
    position=position_dodge(0.9)
  geom_text(
    stat="identity",
   mapping=aes(label=phenotype_cnt),
    vjust=0.05
  ) +
  scale_fill_brewer(palette="Accent") +
  scale_x_discrete(
   name="Phenotype",
   limits=c("plastic", "nonplastic"),
   labels=c("Plastic", "Non-plastic")
  ) +
 facet_grid(sensors~environment) +
 theme(
    legend.position="none"
```



Indeed, we do not observe the evolution of phenotypic plasticity in any replicates in which digital organisms do not have access to sensory instructions. We do observe the evolution of plasticity (not necessarily adaptive plasticity) in both constant and fluctuating environments where sensors are enabled.

To what extent is the observed phenotypic plasticity adaptive?

```
ggplot(filter(observed_plasticity, environment=="Fluctuating" & sensors == "Sensors" & phenotype
  geom_bar(
   stat="identity",
   position=position_dodge(0.9)
  geom_text(
    stat="identity",
   mapping=aes(label=phenotype_cnt),
   vjust=0.05
  ) +
  scale_fill_brewer(palette="Accent") +
  scale_x_discrete(
   name="Phenotype",
   limits=c("plastic", "plastic_adaptive", "plastic_optimal", "plastic_nonadaptive"),
   labels=c("Total plastic", "Adaptive plasticity", "Optimal plasticity", "Non-adaptive plastic
  ) +
  facet_grid(sensors~environment) +
  theme(
    legend.position="none"
```



# Evolutionary change

The effect of adaptive phenotypic plasticity on evolutionary change.

#### 6.1 Overview

```
total_updates <- 200000
replicates <- 100
alpha <- 0.05

all_traits <- c("not", "nand", "ornot", "or", "andnot")
traits_set_a <- c("not", "and", "or")
traits_set_b <- c("nand", "ornot", "andnot")

# Relative location of data.
working_directory <- "experiments/2021-02-08-evo-dynamics/analysis/" # << For bookdown
# working_directory <- "./" # << For local analysis</pre>
```

## 6.2 Analysis dependencies

Load all required R libraries.

```
library(ggplot2)
library(rstatix)
library(ggsignif)
library(scales)
library(tidyverse)
library(cowplot)
library(RColorBrewer)
library(Hmisc)
```

```
library(boot)
source("https://gist.githubusercontent.com/benmarwick/2a1bb0133ff568cbe28d/raw/fb53bd9")
```

These analyses were conducted/knitted with the following computing environment:

```
print(version)
##
                  x86_64-pc-linux-gnu
## platform
## arch
                  x86_64
## os
                  linux-gnu
## system
                  x86_64, linux-gnu
## status
## major
## minor
                  1.0
## year
                  2021
## month
                  05
## day
                  18
## svn rev
                  80317
## language
                  R.
## version.string R version 4.1.0 (2021-05-18)
## nickname
                  Camp Pontanezen
```

## 6.3 Setup

```
summary_data_loc <- paste0(working_directory, "data/aggregate.csv")</pre>
summary_data <- read.csv(summary_data_loc, na.strings="NONE")</pre>
summary_data$DISABLE_REACTION_SENSORS <- as.factor(summary_data$DISABLE_REACTION_SENSO
summary_data$chg_env <- summary_data$chg_env == "True"</pre>
summary_data$dominant_plastic_odd_even <- as.factor(summary_data$dominant_plastic_odd_.
summary_data$sensors <- summary_data$DISABLE_REACTION_SENSORS == "0"</pre>
summary_data$is_plastic <- summary_data$dominant_plastic_odd_even == "True"</pre>
env_label_fun <- function(chg_env) {</pre>
  if (chg_env) {
    return("Fluctuating")
  } else {
    return("Constant")
  }
}
sensors_label_fun <- function(has_sensors) {</pre>
  if (has_sensors) {
```

6.3. SETUP 25

```
return("Sensors")
  } else {
    return("No sensors")
  }
}
# note that this labeler makes assumptions about how we set up our experiment
condition_label_fun <- function(has_sensors, env_chg) {</pre>
  if (has_sensors && env_chg) {
    return("PLASTIC")
  } else if (env_chg) {
    return("NON-PLASTIC")
  } else {
    return("STATIC")
  }
}
summary_data$env_label <- mapply(</pre>
  env_label_fun,
  summary_data$chg_env
)
summary_data$sensors_label <- mapply(</pre>
  sensors_label_fun,
  summary_data$sensors
summary_data$condition <- mapply(</pre>
  condition_label_fun,
  summary_data$sensors,
  summary_data$chg_env
condition_order = c(
  "STATIC",
  "NON-PLASTIC",
  "PLASTIC"
)
pairwise_comparisons <- list(</pre>
  c("STATIC", "NON-PLASTIC"),
  c("STATIC", "PLASTIC"),
  c("PLASTIC", "NON-PLASTIC")
p_label <- function(p_value) {</pre>
  threshold = 0.0001
  if (p_value < threshold) {</pre>
```

```
return(paste0("p < ", threshold))</pre>
  } else {
    return(paste0("p = ", p_value))
  }
}
# *really* inefficient way to identify outliers
is_outlier <- function(value, cond, data, column) {</pre>
  cond_data <- filter(data, condition==cond)</pre>
  q1 <- summary(cond_data[,column])[["1st Qu."]]
  q3 <- summary(cond_data[,column])[["3rd Qu."]]
  H <- 1.5 * IQR(cond data[,column])</pre>
  return( (value < (q1-H)) || (value > (q3+H)) )
###### misc ######
# Configure our default graphing theme
theme_set(theme_cowplot())
# Palette
cb_palette <- "Paired"
# Create a directory to store plots
dir.create(paste0(working_directory, "plots"), showWarnings=FALSE)
# Define sample mean function
samplemean <- function(x, d) {</pre>
  return(mean(x[d]))
```

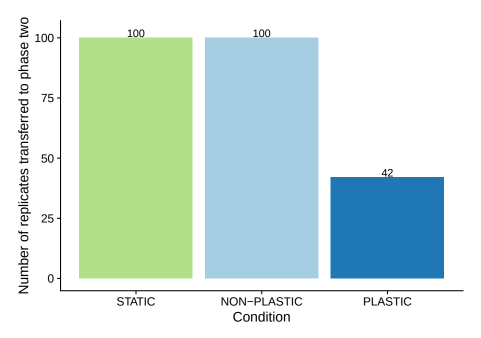
## 6.4 The evolution of phenotypic plasticity

For sensor-enabled populations in fluctuating environments, we only transfered populations containing an optimally plastic genotype to phase-two.

```
summary_data_grouped = dplyr::group_by(summary_data, condition)
summary_data_group_counts = dplyr::summarize(summary_data_grouped, n=dplyr::n())

ggplot(summary_data_group_counts, aes(x=condition, y=n, fill=condition)) +
    geom_col(position=position_dodge(0.9)) +
    geom_text(aes(label=n, y=n+2)) +
    scale_x_discrete(
        name="Condition",
        limits=condition_order
) +
    scale_fill_brewer(
        palette=cb_palette
) +
```

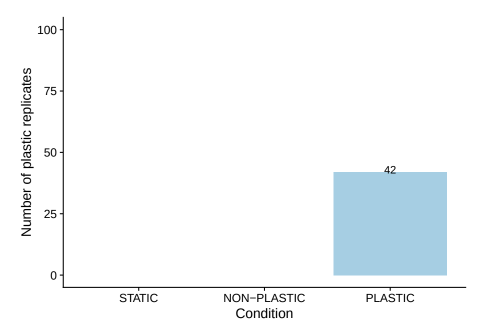
```
scale_color_brewer(
   palette=cb_palette
) +
ylab("Number of replicates transferred to phase two") +
theme(
  legend.position="none"
)
```



We can confirm our expectation that the dominant genotypes in non-plastic conditions are not phenotypically plastic.

```
summary_data_grouped = dplyr::group_by(summary_data, condition, is_plastic)
summary_data_group_counts = dplyr::summarize(summary_data_grouped, n=dplyr::n())
ggplot(filter(summary_data_group_counts, is_plastic), aes(x=condition, y=n, fill=condition)) +
geom_col(
    position=position_dodge(0.9)
) +
scale_x_discrete(
    name="Condition",
    limits=condition_order
) +
scale_fill_brewer(
    palette=cb_palette
) +
scale_color_brewer(
```

```
palette=cb_palette
) +
geom_text(aes(label=n, y=n+1)) +
ylab("Number of plastic replicates") +
ylim(0, 100) +
theme(
  legend.position="none"
)
```



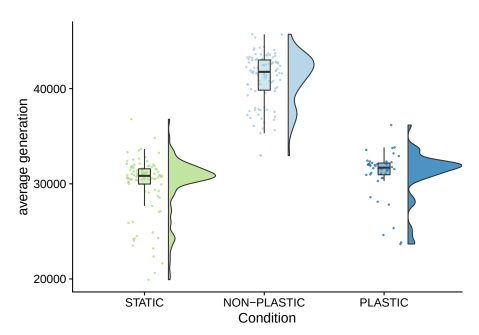
## 6.5 Average generation

How many generations elapsed in each of our treatments?

```
ggplot(summary_data, aes(x=condition, y=time_average_generation, fill=condition)) +
  geom_flat_violin(
    position = position_nudge(x = .2, y = 0),
        alpha = .8
) +
  geom_point(
    mapping=aes(color=condition),
    position = position_jitter(width = .15),
    size = .5,
    alpha = 0.8
) +
```

```
geom_boxplot(
  width = .1,
  outlier.shape = NA,
  alpha = 0.5
scale_x_discrete(
  name="Condition",
  limits=condition_order
scale_fill_brewer(
  palette=cb_palette
scale_color_brewer(
  palette=cb_palette
) +
# coord_flip() +
ylab("average generation") +
theme(
  legend.position="none"
ggsave(paste0(working_directory, "plots/", "average-generation.png"))
```

#### ## Saving $6.5 \times 4.5$ in image



```
kruskal.test(
  formula=time_average_generation~condition,
  data=summary_data
)
##
   Kruskal-Wallis rank sum test
##
##
## data: time_average_generation by condition
## Kruskal-Wallis chi-squared = 177.33, df = 2, p-value < 2.2e-16
pairwise.wilcox.test(
  x=summary_data$time_average_generation,
  g=summary_data$condition,
  p.adjust.method="bonferroni",
)
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: summary_data$time_average_generation and summary_data$condition
##
##
           NON-PLASTIC PLASTIC
## PLASTIC <2e-16
## STATIC <2e-16
                       0.004
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
    "PLASTIC median: ",
    median(filter(summary_data, condition=="PLASTIC")$time_average_generation)
  paste0(
    "STATIC median: ",
    median(filter(summary_data, condition=="STATIC")$time_average_generation)
 ),
  paste0(
    "NON-PLASTIC median: ",
    median(filter(summary_data, condition=="NON-PLASTIC")$time_average_generation)
)
```

## [1] "PLASTIC median: 31697.65; STATIC median: 30839.75; NON-PLASTIC median: 41768.6

```
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
  pair_data <- filter(summary_data, condition %in% pair)</pre>
  pair_data$condition <- as.factor(pair_data$condition)</pre>
  wt <- wilcox.test(</pre>
    formula=time_average_generation~condition,
    data=pair_data,
    exact=FALSE,
    paired=FALSE
  print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
## [1] "STATIC<-->NON-PLASTIC: W=9982"
## [1] "STATIC<-->PLASTIC: W=2818"
## [1] "PLASTIC<-->NON-PLASTIC: W=4186"
summary_data %>%
  group_by(condition) %>%
  summarise(mean=mean(time_average_generation),sd=sd(time_average_generation))
## # A tibble: 3 x 3
## condition
                 mean
                <dbl> <dbl>
##
    <chr>
## 1 NON-PLASTIC 41090. 2702.
## 2 PLASTIC 31016. 2615.
## 3 STATIC
                 30002. 3011.
```

#### 6.6 Coalescence event count

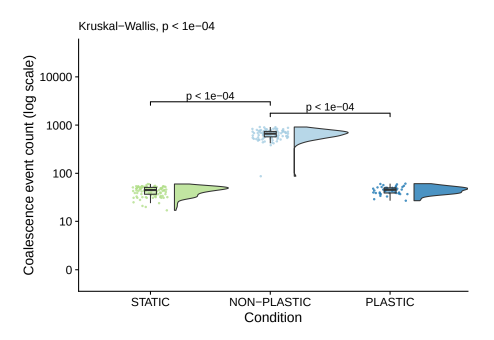
The number of times the most recent common ancestor changes gives us the number of selective sweeps that occur during the experiment.

```
# Compute manual labels for geom_signif
stat.test <- summary_data %>%
  wilcox_test(phylo_mrca_changes ~ condition) %>%
  adjust_pvalue(method = "bonferroni") %>%
  add_significance() %>%
  add_xy_position(x="condition",step.increase=1)
# Tweak y.position manually to account for scaled axis (edge case that triggers bad behavior in general stat.test$manual_position <- log10(stat.test$y.position) * c(1.0,1.0,1.03)
stat.test$label <- mapply(p_label,stat.test$p.adj)
summary_data$is_outlier <- mapply(</pre>
```

```
is_outlier,
 summary_data$phylo_mrca_changes,
 summary_data$condition,
 MoreArgs=list(data=summary_data, column="phylo_mrca_changes")
coalescence_events_fig <- ggplot(</pre>
   summary_data,
   aes(x=condition, y=phylo_mrca_changes,fill=condition)
 ) +
 geom_flat_violin(
   # data=filter(summary_data,is_outlier==FALSE),
   scale="width",
   position = position_nudge(x = .2, y = 0),
   alpha = .8
 ) +
 geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
   alpha = 0.8
 ) +
 geom_boxplot(
   width = .1,
   outlier.shape = NA,
   alpha = 0.5
 ) +
 scale_x_discrete(
   name="Condition",
   limits=condition_order,
   labels=condition_order,
   breaks=condition_order
 ) +
 scale_y_continuous(
   name="Coalescence event count (log scale)",
   trans=pseudo_log_trans(sigma = 1, base = 10),
   breaks=c(0, 10, 100, 1000, 10000),
   limits=c(-1, 35000)
 ) +
 scale_fill_brewer(
   palette=cb_palette
  scale_color_brewer(
   palette=cb_palette
```

```
labs(
  subtitle=paste0(
    "Kruskal-Wallis, ",
    p_label(signif(kruskal.test(formula=phylo_mrca_changes~condition, data=summary_data)$p.valu
) +
ggsignif::geom_signif(
  data=filter(stat.test, p.adj <= alpha),</pre>
  aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
  manual=TRUE,
  inherit.aes=FALSE
) +
# coord_flip() +
theme(
  legend.position="none"
) +
ggsave(
  pasteO(working_directory, "plots/", "selective-sweeps.pdf"),
  width=5,
  height=5
```

## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y\_position
coalescence\_events\_fig



```
kruskal.test(
  formula=phylo_mrca_changes~condition,
  data=summary_data
)
##
##
   Kruskal-Wallis rank sum test
##
## data: phylo_mrca_changes by condition
## Kruskal-Wallis chi-squared = 175.46, df = 2, p-value < 2.2e-16
pairwise.wilcox.test(
  x=summary_data$phylo_mrca_changes,
  g=summary_data$condition,
  p.adjust.method="bonferroni",
)
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: summary_data$phylo_mrca_changes and summary_data$condition
##
           NON-PLASTIC PLASTIC
##
## PLASTIC <2e-16
## STATIC <2e-16
                       1
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
    "PLASTIC median: ",
   median(filter(summary_data, condition=="PLASTIC")$phylo_mrca_changes)
  ),
  paste0(
   "STATIC median: ",
   median(filter(summary_data, condition=="STATIC")$phylo_mrca_changes)
  ),
  paste0(
    "NON-PLASTIC median: ",
    median(filter(summary_data, condition=="NON-PLASTIC")$phylo_mrca_changes)
  )
)
```

## [1] "PLASTIC median: 45.5; STATIC median: 45; NON-PLASTIC median: 663.5"

```
print("Wilcox rank sum test statistics:")

## [1] "Wilcox rank sum test statistics:"

for (pair in pairwise_comparisons) {
    pair_data <- filter(summary_data, condition %in% pair)
    pair_data$condition <- as.factor(pair_data$condition)
    wt <- wilcox.test(
        formula=phylo_mrca_changes~condition,
        data=pair_data,
        exact=FALSE,
        paired=FALSE
    )
    print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
}

## [1] "STATIC<-->NON-PLASTIC: W=10000"

## [1] "STATIC<-->PLASTIC: W=2215"

## [1] "PLASTIC<-->NON-PLASTIC: W=4200"
```

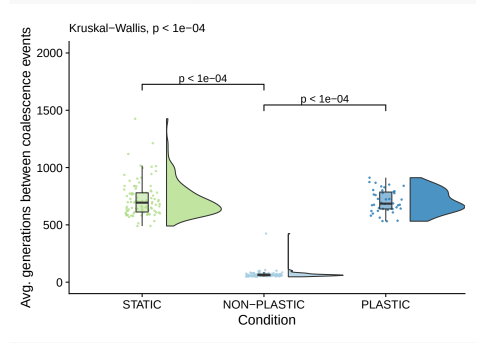
# 6.6.1 Average number of generations between coalescence events

```
# Compute frequency of coalescence events
summary_data$generations_per_mrca_change <- summary_data$time_average_generation / summary_data_average_generation / summary_data_average_gen
# Compute manual labels for geom_signif
stat.test <- summary_data %>%
      wilcox_test(generations_per_mrca_change ~ condition) %>%
      adjust_pvalue(method = "bonferroni") %>%
      add significance() %>%
      add_xy_position(x="condition")
# Tweak y.position manually to account for scaled axis (edge case that triggers bad behavior in
stat.test$manual_position <- stat.test$y.position</pre>
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
summary_data$is_outlier <- mapply(</pre>
      is_outlier,
      summary_data$generations_per_mrca_change,
      summary_data$condition,
      MoreArgs=list(data=summary_data, column="generations_per_mrca_change")
)
coalescence_events_freq_fig <- ggplot(</pre>
             summary_data,
             aes(x=condition, y=generations_per_mrca_change, fill=condition)
```

```
) +
 geom_flat_violin(
   # data=filter(summary_data,is_outlier==FALSE),
   scale="width",
   position = position_nudge(x = .2, y = 0),
   alpha = .8
 ) +
 geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
   alpha = 0.8
 ) +
 geom_boxplot(
   width = .1,
   outlier.shape = NA,
   alpha = 0.5
 scale_x_discrete(
   name="Condition",
   limits=condition_order,
   labels=condition_order
 ) +
 scale_y_continuous(
   name="Avg. generations between coalescence events",
   limits=c(0, 2000),
   breaks=seq(0, 2000, 500)
 ) +
 scale_fill_brewer(
   palette=cb_palette
 ) +
 scale_color_brewer(
   palette=cb_palette
 # coord_flip() +
 labs(
   subtitle=paste0(
     "Kruskal-Wallis, ",
     p_label(signif(kruskal.test(formula=generations_per_mrca_change~condition, data=
   )
 ggsignif::geom_signif(
   data=filter(stat.test, p.adj <= alpha),</pre>
   aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
   manual=TRUE,
```

```
inherit.aes=FALSE
) +
theme(
  legend.position="none"
) +
ggsave(
  paste0(working_directory, "plots/", "generations-between-selective-sweeps.png"),
  width=5,
  height=5
)
```

## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y\_position
coalescence\_events\_freq\_fig



```
kruskal.test(
  formula=generations_per_mrca_change~condition,
  data=summary_data
)
```

```
##
## Kruskal-Wallis rank sum test
##
## data: generations_per_mrca_change by condition
## Kruskal-Wallis chi-squared = 175.33, df = 2, p-value < 2.2e-16</pre>
```

```
pairwise.wilcox.test(
  x=summary_data$generations_per_mrca_change,
  g=summary_data$condition,
  p.adjust.method="bonferroni",
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: summary_data$generations_per_mrca_change and summary_data$condition
##
           NON-PLASTIC PLASTIC
##
## PLASTIC <2e-16
## STATIC <2e-16
##
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
    "PLASTIC median: ",
   median(filter(summary_data, condition=="PLASTIC")$generations_per_mrca_change)
  paste0(
    "STATIC median: ",
   median(filter(summary_data, condition=="STATIC")$generations_per_mrca_change)
  ),
  paste0(
    "NON-PLASTIC median: ",
    median(filter(summary_data, condition=="NON-PLASTIC")$generations_per_mrca_change)
)
## [1] "PLASTIC median: 685.001780758557; STATIC median: 693.676265008576; NON-PLASTIC
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
  pair_data <- filter(summary_data, condition %in% pair)</pre>
  pair_data$condition <- as.factor(pair_data$condition)</pre>
  wt <- wilcox.test(</pre>
    formula=generations_per_mrca_change~condition,
    data=pair_data,
    exact=FALSE,
    paired=FALSE
```

```
)
    print(pasteO(pair[1], "<-->", pair[2], ": W=",wt$statistic))
}

## [1] "STATIC<-->NON-PLASTIC: W=O"

## [1] "STATIC<-->PLASTIC: W=2151"

## [1] "PLASTIC<-->NON-PLASTIC: W=O"
```

## 6.7 Phenotypic volatility along the dominant lineage

```
# Compute manual labels for geom_signif
stat.test <- summary_data %>%
  wilcox_test(dominant_lineage_trait_volatility ~ condition) %>%
  adjust_pvalue(method = "bonferroni") %>%
  add_significance() %>%
  add_xy_position(x="condition", step.increase=1)
# Tweak y.position manually to account for scaled axis (edge case that triggers bad behavior in
stat.test$manual_position <- log10(stat.test$y.position) * c(1.0,1.0,1.03)
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
summary_data$is_outlier <- mapply(</pre>
  is outlier,
 summary_data$dominant_lineage_trait_volatility,
  summary data$condition,
 MoreArgs=list(data=summary_data, column="dominant_lineage_trait_volatility")
phenotypic_volatility_fig <- ggplot(</pre>
    summary_data,
    aes(x=condition, y=dominant_lineage_trait_volatility, fill=condition)
  geom_flat_violin(
    # data=filter(summary_data, is_outlier==FALSE),
   scale="width",
   position = position_nudge(x = .2, y = 0),
   alpha = .8
  ) +
  geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
    alpha = 0.8
```

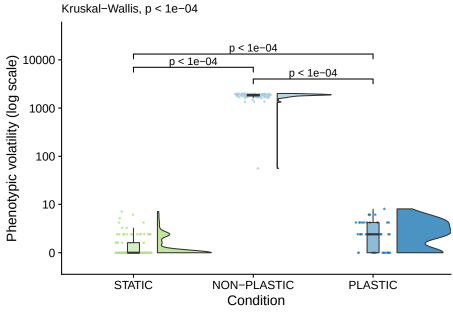
```
geom_boxplot(
  width = .1,
  outlier.shape = NA,
  alpha = 0.5
) +
scale_x_discrete(
  name="Condition",
  limits=condition_order,
  labels=condition_order
) +
scale_y_continuous(
  name="Phenotypic volatility (log scale)",
  trans=pseudo_log_trans(sigma = 1, base = 10),
  breaks=c(0, 10, 100, 1000, 10000),
  limits=c(-1, 35000)
) +
scale_fill_brewer(
  palette=cb_palette
) +
scale_color_brewer(
  palette=cb_palette
) +
labs(
  subtitle=paste0(
    "Kruskal-Wallis, ",
    p_label(signif(kruskal.test(formula=dominant_lineage_trait_volatility~condition,
  )
) +
ggsignif::geom_signif(
  data=filter(stat.test, p.adj<=alpha),</pre>
  aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
  manual=TRUE,
  inherit.aes=FALSE
) +
# coord_flip() +
theme(
  legend.position="none"
) +
ggsave(
  pasteO(working_directory, "plots/", "phenotypic-volatility.pdf"),
  width=5,
  height=5
)
```

## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y\_position

#### phenotypic\_volatility\_fig

## ##

## PLASTIC < 2e-16



```
kruskal.test(
  formula=dominant_lineage_trait_volatility~condition,
  data=summary_data
)
##
##
   Kruskal-Wallis rank sum test
## data: dominant_lineage_trait_volatility by condition
## Kruskal-Wallis chi-squared = 190.78, df = 2, p-value < 2.2e-16
pairwise.wilcox.test(
  x=summary_data$dominant_lineage_trait_volatility,
  g=summary_data$condition,
  p.adjust.method="bonferroni",
)
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
```

NON-PLASTIC PLASTIC

## data: summary\_data\$dominant\_lineage\_trait\_volatility and summary\_data\$condition

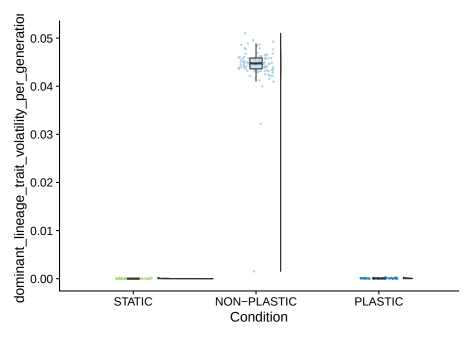
```
## STATIC < 2e-16
                       8.7e-07
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
    "PLASTIC median: ",
   median(filter(summary_data, condition=="PLASTIC")$dominant_lineage_trait_volatilit
  ),
  paste0(
    "STATIC median: ",
   median(filter(summary data, condition=="STATIC")$dominant lineage trait volatility
  ),
  paste0(
    "NON-PLASTIC median: ",
    median(filter(summary_data, condition=="NON-PLASTIC")$dominant_lineage_trait_volat
  )
## [1] "PLASTIC median: 2; STATIC median: 0; NON-PLASTIC median: 1868"
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
  pair_data <- filter(summary_data, condition %in% pair)</pre>
  pair_data$condition <- as.factor(pair_data$condition)</pre>
  wt <- wilcox.test(
    formula=dominant_lineage_trait_volatility~condition,
    data=pair_data,
    exact=FALSE,
    paired=FALSE
  )
  print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
## [1] "STATIC<-->NON-PLASTIC: W=10000"
## [1] "STATIC<-->PLASTIC: W=3116.5"
## [1] "PLASTIC<-->NON-PLASTIC: W=4200"
```

### 6.7.1 Phenotypic volatility normalized by generations elapsed

summary\_data\$dominant\_lineage\_trait\_volatility\_per\_generation <- summary\_data\$dominant</pre>

```
ggplot(summary_data, aes(x=condition, y=dominant_lineage_trait_volatility_per_generation, fill=condition)
  geom_flat_violin(
    position = position_nudge(x = .2, y = 0),
    alpha = .8
 ) +
 geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
    alpha = 0.8
  geom_boxplot(
   width = .1,
    outlier.shape = NA,
    alpha = 0.5
 scale_x_discrete(
   name="Condition",
   limits=condition_order
  ) +
  scale_fill_brewer(
   palette=cb_palette
  ) +
  scale_color_brewer(
    palette=cb_palette
  # coord_flip() +
 theme(
    legend.position="none"
```

##



```
kruskal.test(
  formula=dominant_lineage_trait_volatility_per_generation~condition,
  data=summary_data
)

##

## Kruskal-Wallis rank sum test

##

## data: dominant_lineage_trait_volatility_per_generation by condition

## Kruskal-Wallis chi-squared = 189.62, df = 2, p-value < 2.2e-16

pairwise.wilcox.test(
  x=summary_data$dominant_lineage_trait_volatility_per_generation,
  g=summary_data$condition,
  p.adjust.method="bonferroni",
)</pre>
```

```
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
    "PLASTIC median: ",
    median(filter(summary_data, condition=="PLASTIC")$dominant_lineage_trait_volatility_per_gener
  ),
  paste0(
    "STATIC median: ",
    median(filter(summary_data, condition=="STATIC")$dominant_lineage_trait_volatility_per_generate
 ),
  paste0(
    "NON-PLASTIC median: ",
    median(filter(summary_data, condition=="NON-PLASTIC")$dominant_lineage_trait_volatility_per_g
)
## [1] "PLASTIC median: 6.33339279717772e-05; STATIC median: 0; NON-PLASTIC median: 0.04474401456
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
  pair_data <- filter(summary_data, condition %in% pair)</pre>
 pair_data$condition <- as.factor(pair_data$condition)</pre>
  wt <- wilcox.test(
    formula=dominant_lineage_trait_volatility_per_generation~condition,
    data=pair_data,
    exact=FALSE,
    paired=FALSE
  print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
## [1] "STATIC<-->NON-PLASTIC: W=10000"
## [1] "STATIC<-->PLASTIC: W=3061.5"
## [1] "PLASTIC<-->NON-PLASTIC: W=4200"
```

### 6.8 Phenotypic fidelity

Frequency that an offspring's genotype is identical to a parent genotype (along the dominant lineage).

```
summary_data$dominant_lineage_trait_fidelity <- (summary_data$dominant_generation_born - summary_
# Compute manual labels for geom_signif</pre>
```

```
stat.test <- summary_data %>%
  wilcox_test(dominant_lineage_trait_fidelity ~ condition) %>%
  adjust_pvalue(method = "bonferroni") %>%
 add_significance() %>%
 add_xy_position(x="condition",step.increase=1.5)
# Tweak y.position manually to account for scaled axis (edge case that triggers bad be
stat.test$manual_position <- stat.test$y.position * c(1.0,1.0,1.0005)
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
summary_data$is_outlier <- mapply(</pre>
 is outlier,
 summary_data$dominant_lineage_trait_fidelity,
 summary data$condition,
 MoreArgs=list(data=summary_data, column="dominant_lineage_trait_fidelity")
phenotypic_fidelity_fig <- ggplot(</pre>
    summary_data,
    aes(x=condition, y=dominant_lineage_trait_fidelity, fill=condition)
 geom_flat_violin(
    data=filter(summary_data,is_outlier==FALSE),
    scale="width",
    position = position_nudge(x = .2, y = 0),
    alpha = .8
 ) +
  geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
    alpha = 0.8
  geom_boxplot(
    width = .1,
    outlier.shape = NA,
    alpha = 0.5
  ) +
  scale_x_discrete(
   name="Condition",
   limits=condition_order,
   labels=condition_order
  ) +
  scale_y_continuous(
```

```
name="Phenotypic fidelity",
  limits=c(0.94, 1.013),
  breaks=c(0.94, 0.96, 0.98, 1.0) #seq(0.94, 1.0, 0.01)
scale_fill_brewer(
  palette=cb_palette
scale_color_brewer(
  palette=cb_palette
# coord_flip() +
labs(
  subtitle=paste0(
    "Kruskal-Wallis, ",
    p_label(signif(kruskal.test(formula=dominant_lineage_trait_fidelity~condition, data=summary
  )
) +
ggsignif::geom_signif(
  data=filter(stat.test, p.adj <= alpha),</pre>
  aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
  manual=TRUE,
  inherit.aes=FALSE
) +
theme(
  legend.position="none"
ggsave(
  pasteO(working_directory, "plots/", "phenotypic-fidelity.pdf"),
  width=5,
  height=5
)
```

```
## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y_position
phenotypic_fidelity_fig
```

```
Kruskal-Wallis, p < 1e-04

1.00

1.00

1.00

1.00

STATIC

NON-PLASTIC

Condition

Cruskal test (
```

```
kruskal.test(
  formula=dominant_lineage_trait_fidelity~condition,
  data=summary_data
)

##

## Kruskal-Wallis rank sum test

##

## data: dominant_lineage_trait_fidelity by condition

## Kruskal-Wallis chi-squared = 189.62, df = 2, p-value < 2.2e-16

pairwise.wilcox.test(
  x=summary_data$dominant_lineage_trait_fidelity,
  g=summary_data$condition,
  p.adjust.method="bonferroni",
)</pre>
```

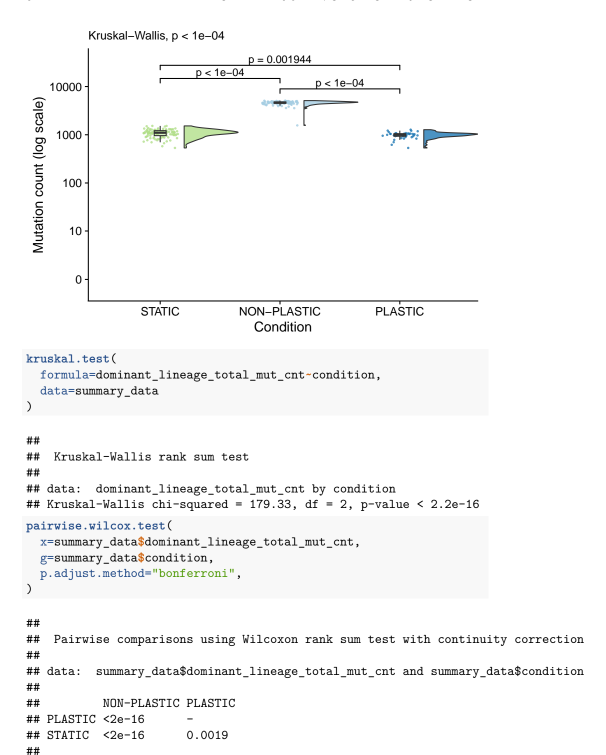
```
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
   "PLASTIC median: ",
   median(filter(summary_data, condition=="PLASTIC")$dominant_lineage_trait_fidelity)
  ),
  paste0(
    "STATIC median: ",
   median(filter(summary_data, condition=="STATIC")$dominant_lineage_trait_fidelity)
  ),
  paste0(
    "NON-PLASTIC median: ",
   median(filter(summary_data, condition=="NON-PLASTIC")$dominant_lineage_trait_fidelity)
)
## [1] "PLASTIC median: 0.999936666072028; STATIC median: 1; NON-PLASTIC median: 0.95525598543618
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
 pair_data <- filter(summary_data, condition %in% pair)</pre>
 pair_data$condition <- as.factor(pair_data$condition)</pre>
  wt <- wilcox.test(
   formula=dominant_lineage_trait_fidelity~condition,
   data=pair_data,
   exact=FALSE,
   paired=FALSE
  print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
## [1] "STATIC<-->NON-PLASTIC: W=0"
## [1] "STATIC<-->PLASTIC: W=1138.5"
## [1] "PLASTIC<-->NON-PLASTIC: W=0"
6.9
       Mutation count
# Compute manual labels for geom_signif
```

```
# Compute manual labels for geom_signif
stat.test <- summary_data %>%
  wilcox_test(dominant_lineage_total_mut_cnt ~ condition) %>%
  adjust_pvalue(method = "bonferroni") %>%
  add_significance() %>%
```

```
add_xy_position(x="condition",step.increase=1)
# Tweak y.position manually to account for scaled axis (edge case that triggers bad be
stat.test\$manual_position <- log10(stat.test\$y.position) * c(1.0,1.0,1.03) # c(1.0,1.
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
summary_data$is_outlier <- mapply(</pre>
  is_outlier,
  summary_data$dominant_lineage_total_mut_cnt,
  summary_data$condition,
 MoreArgs=list(data=summary_data, column="dominant_lineage_total_mut_cnt")
mutation_count_fig <- ggplot(</pre>
    summary_data,
    aes(x=condition, y=dominant_lineage_total_mut_cnt, fill=condition)
  ) +
  geom_flat_violin(
    # data=filter(summary_data, !is_outlier),
    scale="width",
    position = position_nudge(x = .2, y = 0),
    alpha = .8
  ) +
  geom_point(
    mapping=aes(color=condition),
    position = position_jitter(width = .15),
    size = .5,
    alpha = 0.8
  ) +
  geom_boxplot(
    width = .1,
    outlier.shape = NA,
    alpha = 0.5
  scale_x_discrete(
    name="Condition",
    limits=condition_order,
    labels=condition_order
  ) +
  scale_y_continuous(
    name="Mutation count (log scale)",
    trans=pseudo_log_trans(sigma = 1, base = 10),
    breaks=c(0, 10, 100, 1000, 10000),
    limits=c(-1, 35000)
  ) +
  scale_fill_brewer(
```

```
palette=cb_palette
scale_color_brewer(
 palette=cb_palette
) +
# coord_flip() +
labs(
  subtitle=paste0(
    "Kruskal-Wallis, ",
   p_label(signif(kruskal.test(formula=dominant_lineage_total_mut_cnt~condition, data=summary)
) +
ggsignif::geom_signif(
  data=filter(stat.test, p.adj <= alpha),</pre>
  aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
  manual=TRUE,
  inherit.aes=FALSE
) +
theme(
  legend.position="none"
ggsave(
  pasteO(working_directory, "plots/", "mutation-accumulation.pdf"),
  height=4
```

## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y\_position
mutation\_count\_fig



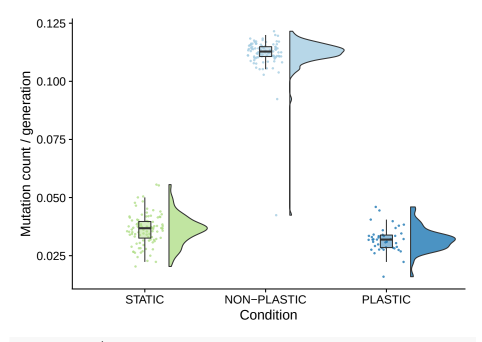
```
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
    "PLASTIC median: ",
    median(filter(summary_data, condition=="PLASTIC")$dominant_lineage_total_mut_cnt)
  ),
  paste0(
    "STATIC median: ",
    median(filter(summary_data, condition=="STATIC")$dominant_lineage_total_mut_cnt)
 ),
  paste0(
    "NON-PLASTIC median: ",
    median(filter(summary_data, condition=="NON-PLASTIC")$dominant_lineage_total_mut_cnt)
  )
)
## [1] "PLASTIC median: 998.5; STATIC median: 1100; NON-PLASTIC median: 4657.5"
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
 pair_data <- filter(summary_data, condition %in% pair)</pre>
 pair_data$condition <- as.factor(pair_data$condition)</pre>
 wt <- wilcox.test(</pre>
    formula=dominant_lineage_total_mut_cnt~condition,
    data=pair_data,
    exact=FALSE,
    paired=FALSE
  print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
## [1] "STATIC<-->NON-PLASTIC: W=10000"
## [1] "STATIC<-->PLASTIC: W=1336.5"
## [1] "PLASTIC<-->NON-PLASTIC: W=4200"
```

#### 6.9.1 Mutation count normalized by generations elapsed

```
summary_data$mutations_per_generation <- summary_data$dominant_lineage_total_mut_cnt / summary_data
ggplot(summary_data, aes(x=condition, y=mutations_per_generation, fill=condition)) +
geom_flat_violin(
   position = position_nudge(x = .2, y = 0),
   alpha = .8</pre>
```

```
) +
geom_point(
  mapping=aes(color=condition),
  position = position_jitter(width = .15),
  size = .5,
  alpha = 0.8
) +
geom_boxplot(
  width = .1,
  outlier.shape = NA,
  alpha = 0.5
) +
scale_x_discrete(
  name="Condition",
  limits=condition_order
ylab("Mutation count / generation") +
scale_fill_brewer(
 palette=cb_palette
) +
scale_color_brewer(
 palette=cb_palette
) +
# coord_flip() +
theme(
  legend.position="none"
```

##



```
kruskal.test(
  formula=mutations_per_generation~condition,
  data=summary_data
)
##
   Kruskal-Wallis rank sum test
##
## data: mutations_per_generation by condition
## Kruskal-Wallis chi-squared = 180.11, df = 2, p-value < 2.2e-16
pairwise.wilcox.test(
  x=summary_data$mutations_per_generation,
  g=summary_data$condition,
 p.adjust.method="bonferroni",
)
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: summary_data$mutations_per_generation and summary_data$condition
##
           NON-PLASTIC PLASTIC
## PLASTIC <2e-16
## STATIC <2e-16
                       2e-04
```

```
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
    "PLASTIC median: ",
    median(filter(summary_data, condition=="PLASTIC")$mutations_per_generation)
  ),
  paste0(
    "STATIC median: ",
    median(filter(summary_data, condition=="STATIC")$mutations_per_generation)
  ),
  paste0(
    "NON-PLASTIC median: ",
    median(filter(summary_data, condition=="NON-PLASTIC")$mutations_per_generation)
## [1] "PLASTIC median: 0.0319267181456982; STATIC median: 0.0368157192941933; NON-PLA
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
  pair_data <- filter(summary_data, condition %in% pair)</pre>
  pair_data$condition <- as.factor(pair_data$condition)</pre>
  wt <- wilcox.test(
    formula=mutations_per_generation~condition,
    data=pair_data,
    exact=FALSE,
    paired=FALSE
  print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
## [1] "STATIC<-->NON-PLASTIC: W=9987"
## [1] "STATIC<-->PLASTIC: W=1206"
## [1] "PLASTIC<-->NON-PLASTIC: W=4198"
```

### 6.10 Genotypic fidelity

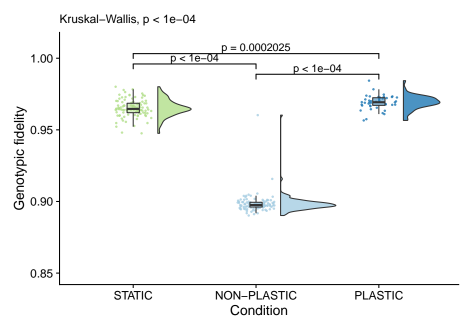
The frequency that an offspring's genotype is the same as a parent's genotype.

```
summary_data$dominant_lineage_genotypic_fidelity <- (summary_data$dominant_generation_
# Compute manual labels for geom_signif
stat.test <- summary_data %>%
```

```
wilcox_test(dominant_lineage_genotypic_fidelity ~ condition) %>%
  adjust_pvalue(method = "bonferroni") %>%
  add_significance() %>%
  add_xy_position(x="condition",step.increase=0.2)
# Tweak y.position manually to account for scaled axis (edge case that triggers bad behavior in
stat.test$manual_position <- stat.test$y.position * c(1.0,1.0,1.0)</pre>
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
genotypic_fidelity_fig <- ggplot(</pre>
    summary_data,
    aes(x=condition, y=dominant_lineage_genotypic_fidelity, fill=condition)
  geom_flat_violin(
    position = position_nudge(x = .2, y = 0),
    alpha = .8
  ) +
  geom_point(
    mapping=aes(color=condition),
    position = position_jitter(width = .15),
    size = .5,
    alpha = 0.8
  ) +
  geom_boxplot(
    width = .1,
    outlier.shape = NA,
    alpha = 0.5
  ) +
  scale_x_discrete(
    name="Condition",
    limits=condition_order,
    labels=condition_order
  ) +
  scale_y_continuous(
    name="Genotypic fidelity",
    limits=c(0.85, 1.01),
    breaks=c(0.85, 0.90, 0.95, 1.0) #seq(0.85, 1.0, 0.02)
  scale_fill_brewer(
    palette=cb_palette
  scale_color_brewer(
    palette=cb_palette
  ) +
  # coord_flip() +
  labs(
```

```
subtitle=paste0(
    "Kruskal-Wallis, ",
    p_label(signif(kruskal.test(formula=dominant_lineage_genotypic_fidelity~condition)
  )
) +
ggsignif::geom_signif(
  data=filter(stat.test, p.adj <= alpha),</pre>
  aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
  manual=TRUE,
  inherit.aes=FALSE
) +
theme(
  legend.position="none"
ggsave(
  pasteO(working_directory, "plots/", "genotypic-fidelity.pdf"),
  width=5,
  height=4
)
```

## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y\_position
genotypic\_fidelity\_fig



kruskal.test(
 formula=dominant\_lineage\_genotypic\_fidelity~condition,

```
data=summary_data
)
##
##
   Kruskal-Wallis rank sum test
## data: dominant_lineage_genotypic_fidelity by condition
## Kruskal-Wallis chi-squared = 179.86, df = 2, p-value < 2.2e-16
pairwise.wilcox.test(
  x=summary_data$dominant_lineage_genotypic_fidelity,
  g=summary_data$condition,
  p.adjust.method="bonferroni",
)
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: summary_data$dominant_lineage_genotypic_fidelity and summary_data$condition
##
           NON-PLASTIC PLASTIC
##
## PLASTIC <2e-16
                       2e-04
## STATIC <2e-16
## P value adjustment method: bonferroni
paste(
 sep="; ",
  paste0(
   "PLASTIC median: ",
   median(filter(summary data, condition=="PLASTIC")$dominant lineage genotypic fidelity)
 ),
  paste0(
   "STATIC median: ",
   median(filter(summary_data, condition=="STATIC")$dominant_lineage_genotypic_fidelity)
  ),
  paste0(
    "NON-PLASTIC median: ",
   median(filter(summary_data, condition=="NON-PLASTIC")$dominant_lineage_genotypic_fidelity)
  )
)
## [1] "PLASTIC median: 0.969286906891951; STATIC median: 0.964620594632577; NON-PLASTIC median:
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
```

```
for (pair in pairwise_comparisons) {
  pair_data <- filter(summary_data, condition %in% pair)
  pair_data$condition <- as.factor(pair_data$condition)
  wt <- wilcox.test(
    formula=dominant_lineage_genotypic_fidelity~condition,
    data=pair_data,
    exact=FALSE,
    paired=FALSE
)
  print(pasteO(pair[1], "<-->", pair[2], ": W=",wt$statistic))
}
### [1] "STATIC<-->NON-PLASTIC: W=18"
### [1] "PLASTIC<-->NON-PLASTIC: W=2992"
### [1] "PLASTIC<--->NON-PLASTIC: W=2"
```

# 6.11 Characterizing variation along dominant lineages

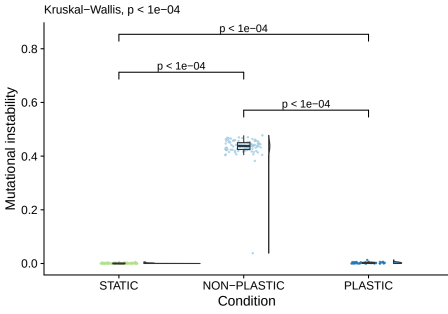
#### 6.11.1 Mutational instability

```
summary_data$frac_phenotype_changing_mut_steps <- summary_data$dominant_lineage_num_mu
summary_data$frac_phenotype_stable_mut_steps <- 1 - summary_data$frac_phenotype_changi:
# Compute manual labels for geom_signif
stat.test <- summary data %>%
 wilcox_test(frac_phenotype_changing_mut_steps ~ condition) %>%
 adjust_pvalue(method = "bonferroni") %>%
 add_significance() %>%
  add_xy_position(x="condition",step.increase=0.2)
# Tweak y.position manually to account for scaled axis (edge case that triggers bad be
stat.test\$manual_position <- stat.test\$y.position #* c(1.0,1.0,1.0)
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
ggplot(summary_data, aes(x=condition, y=frac_phenotype_changing_mut_steps, fill=condit
 geom_flat_violin(
    position = position_nudge(x = .2, y = 0),
    alpha = .8
  geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
    alpha = 0.8
```

```
) +
 geom_boxplot(
   width = .1,
   outlier.shape = NA,
   alpha = 0.5
 ) +
 scale_x_discrete(
  name="Condition",
   limits=condition_order
 ylab("Mutational instability") +
 scale fill brewer(
   palette=cb_palette
 scale_color_brewer(
   palette=cb_palette
 # coord_flip() +
 labs(
   subtitle=paste0(
     "Kruskal-Wallis, ",
     p_label(signif(kruskal.test(formula=frac_phenotype_changing_mut_steps~condition, data=summater)
   )
 ) +
 ggsignif::geom_signif(
   data=filter(stat.test, p.adj <= alpha),</pre>
   aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
   manual=TRUE,
   inherit.aes=FALSE
 ) +
 theme(
   legend.position="none"
 ggsave(paste0(working_directory, "plots/", "frac_phenotype_changing_mutational_steps.png"))
```

```
## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y_position
```

```
## Saving 6.5 x 4.5 in image
```



```
kruskal.test(
  formula=frac_phenotype_changing_mut_steps~condition,
  data=summary_data
)

##

## Kruskal-Wallis rank sum test

##

## data: frac_phenotype_changing_mut_steps by condition

## Kruskal-Wallis chi-squared = 191.23, df = 2, p-value < 2.2e-16

pairwise.wilcox.test(
  x=summary_data$frac_phenotype_changing_mut_steps,
  g=summary_data$condition,
  p.adjust.method="bonferroni",
)</pre>
```

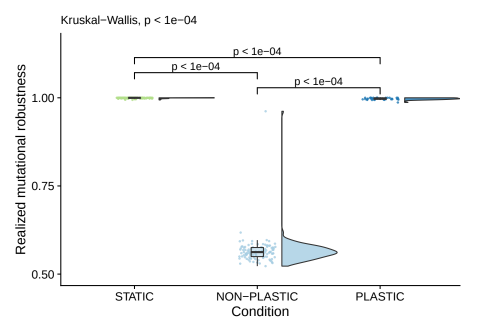
adjust\_pvalue(method = "bonferroni") %>%

```
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
   "PLASTIC median: ",
   median(filter(summary_data, condition=="PLASTIC")$frac_phenotype_changing_mut_steps)
  ),
  paste0(
    "STATIC median: ",
   median(filter(summary_data, condition=="STATIC")$frac_phenotype_changing_mut_steps)
  ),
  paste0(
    "NON-PLASTIC median: ",
   median(filter(summary_data, condition=="NON-PLASTIC")$frac_phenotype_changing_mut_steps)
)
## [1] "PLASTIC median: 0.00224941742616098; STATIC median: 0; NON-PLASTIC median: 0.437583018324
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
 pair_data <- filter(summary_data, condition %in% pair)</pre>
 pair_data$condition <- as.factor(pair_data$condition)</pre>
  wt <- wilcox.test(
   formula=frac_phenotype_changing_mut_steps~condition,
   data=pair_data,
   exact=FALSE,
   paired=FALSE
  print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
## [1] "STATIC<-->NON-PLASTIC: W=10000"
## [1] "STATIC<-->PLASTIC: W=3172"
## [1] "PLASTIC<-->NON-PLASTIC: W=4200"
6.11.2
         Mutational stability (realized mutational robust-
         ness)
# Compute manual labels for geom_signif
stat.test <- summary_data %>%
  wilcox_test(frac_phenotype_stable_mut_steps ~ condition) %>%
```

```
add_significance() %>%
  add_xy_position(x="condition", step.increase=0.75)
# Tweak y.position manually to account for scaled axis (edge case that triggers bad be
stat.test\$manual_position <- stat.test\$y.position #* c(1.0,1.0,1.0)
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
summary_data$is_outlier <- mapply(</pre>
  is_outlier,
  summary_data$dominant_lineage_trait_volatility,
  summary_data$condition,
  MoreArgs=list(data=summary_data, column="dominant_lineage_trait_volatility")
mutational_stability_fig <- ggplot(</pre>
    summary_data,
    aes(x=condition, y=frac_phenotype_stable_mut_steps, fill=condition)
  ) +
  geom_flat_violin(
    # data=filter(summary_data,is_outlier==FALSE),
    scale="width",
    position = position_nudge(x = .2, y = 0),
    alpha = .8
  ) +
  geom_point(
    mapping=aes(color=condition),
    position = position_jitter(width = .15),
    size = .5,
    alpha = 0.8
  ) +
  geom_boxplot(
    width = .1,
    outlier.shape = NA,
    alpha = 0.5
  ) +
  scale_x_discrete(
    name="Condition",
    limits=condition_order
  ) +
  scale_y_continuous(
    name="Realized mutational robustness",
    limits=c(0.5, 1.15),
    breaks=c(0.5, 0.75, 1.0)
  ) +
  scale_fill_brewer(
    palette=cb_palette
```

```
) +
scale_color_brewer(
  palette=cb_palette
) +
labs(
  subtitle=paste0(
    "Kruskal-Wallis, ",
    p_label(signif(kruskal.test(formula=frac_phenotype_stable_mut_steps~condition, data=summary
) +
ggsignif::geom_signif(
  data=filter(stat.test, p.adj <= alpha),</pre>
  aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
  manual=TRUE,
  inherit.aes=FALSE
) +
theme(
  legend.position="none"
```

## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y\_position
mutational\_stability\_fig



kruskal.test(
 formula=frac\_phenotype\_stable\_mut\_steps~condition,

```
data=summary_data
##
##
   Kruskal-Wallis rank sum test
##
## data: frac_phenotype_stable_mut_steps by condition
## Kruskal-Wallis chi-squared = 191.23, df = 2, p-value < 2.2e-16
pairwise.wilcox.test(
  x=summary_data$frac_phenotype_stable_mut_steps,
  g=summary_data$condition,
  p.adjust.method="bonferroni",
)
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
##
## data: summary_data$frac_phenotype_stable_mut_steps and summary_data$condition
##
           NON-PLASTIC PLASTIC
##
## PLASTIC < 2e-16
## STATIC < 2e-16
                       2.3e-07
##
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
    "PLASTIC median: ",
   median(filter(summary_data, condition=="PLASTIC")$frac_phenotype_stable_mut_steps)
  ),
  paste0(
    "STATIC median: ",
    median(filter(summary_data, condition=="STATIC")$frac_phenotype_stable_mut_steps)
  ),
  paste0(
    "NON-PLASTIC median: ",
    median(filter(summary_data, condition=="NON-PLASTIC")$frac_phenotype_stable_mut_st
  )
)
## [1] "PLASTIC median: 0.997750582573839; STATIC median: 1; NON-PLASTIC median: 0.562
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
```

```
for (pair in pairwise_comparisons) {
  pair_data <- filter(summary_data, condition %in% pair)
  pair_data$condition <- as.factor(pair_data$condition)
  wt <- wilcox.test(
    formula=frac_phenotype_stable_mut_steps~condition,
    data=pair_data,
    exact=FALSE,
    paired=FALSE
)
  print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
}
```

```
## [1] "STATIC<-->NON-PLASTIC: W=0"
## [1] "STATIC<-->PLASTIC: W=1028"
## [1] "PLASTIC<-->NON-PLASTIC: W=0"
```

### 6.11.3 For PLASTIC populations, what fraction of phenotype-altering mutations occurred in the

summary\_data\$frac\_unexpressed\_mut\_steps <- summary\_data\$dominant\_lineage\_num\_mut\_steps\_that\_change
summary\_data\$frac\_expressed\_mut\_steps <- summary\_data\$dominant\_lineage\_num\_mut\_steps\_that\_change
ggplot(filter(summary\_data, condition=="PLASTIC" & dominant\_lineage\_num\_mut\_steps\_that\_change\_agg
geom\_histogram(binwidth=0.1) +
scale\_x\_continuous(
 limits=c(0, 1.1),
 breaks=seq(0, 1.0, 0.1)
) +
theme(
legend.position="none"</pre>

## Warning: Removed 2 rows containing missing values (geom\_bar).

```
15
  10
count
    5
    0
                               0.4
                                     0.5
        0.0
             0.1
                   0.2
                         0.3
                                           0.6
                                                 0.7
                                                       8.0
                                                             0.9
                                                                   1.0
                          frac_unexpressed_mut_steps
```

```
print(paste0("PLASTIC - Mean with bootstrapped 95% CI"))
## [1] "PLASTIC - Mean with bootstrapped 95% CI"
bo <- boot(filter(summary_data, condition=="PLASTIC" & dominant_lineage_num_mut_steps_
print(bo)
##
## ORDINARY NONPARAMETRIC BOOTSTRAP
##
##
## Call:
## boot(data = filter(summary_data, condition == "PLASTIC" & dominant_lineage_num_mut_:
       0) $frac_unexpressed_mut_steps, statistic = samplemean, R = 10000)
##
##
##
## Bootstrap Statistics :
        original
                        bias
                                std. error
## t1* 0.8247126 -3.936782e-05
                                0.0405534
print(boot.ci(bo, conf=0.95, type="perc"))
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 10000 bootstrap replicates
##
## CALL :
```

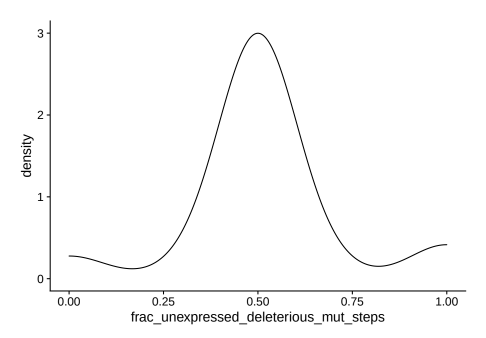
## boot.ci(boot.out = bo, conf = 0.95, type = "perc")

# 6.11.4 For PLASTIC populations, what fraction of mutations that affect the unexpressed phenotype are deleterious versus beneficial?

aggregate\_frac\_unexpressed\_deleterious\_mut\_steps <- sum(plastic\_summary\_data\$dominant\_lineage\_numaggregate\_frac\_unexpressed\_beneficial\_mut\_steps <- sum(plastic\_summary\_data\$dominant\_lineage\_num\_

#### 6.11.4.1 Deleterious mutations

```
summary_data$frac_unexpressed_deleterious_mut_steps <- summary_data$dominant_lineage_num_mut_step
ggplot(
   filter(summary_data, condition=="PLASTIC" & dominant_lineage_num_mut_steps_that_change_unexpress
   aes(x=frac_unexpressed_deleterious_mut_steps)
) +
   geom_density() +
   theme(
    legend.position="none"</pre>
```



bo <- boot(filter(summary\_data, condition=="PLASTIC" & dominant\_lineage\_num\_mut\_steps\_
print(bo)</pre>

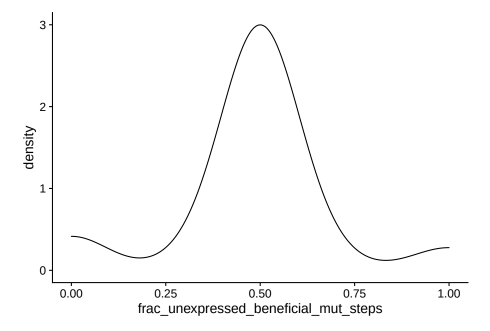
```
##
## ORDINARY NONPARAMETRIC BOOTSTRAP
##
##
## Call:
## boot(data = filter(summary_data, condition == "PLASTIC" & dominant_lineage_num_mut_
##
       0)$frac_unexpressed_deleterious_mut_steps, statistic = samplemean,
       R = 10000)
##
##
##
## Bootstrap Statistics :
        original
                        bias
                                std. error
## t1* 0.5172414 -5.229885e-05 0.03954846
print(boot.ci(bo, conf=0.95, type="perc"))
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 10000 bootstrap replicates
## boot.ci(boot.out = bo, conf = 0.95, type = "perc")
## Intervals :
```

#### 6.11. CHARACTERIZING VARIATION ALONG DOMINANT LINEAGES71

```
## Level Percentile
## 95% ( 0.4402,  0.5943 )
## Calculations and Intervals on Original Scale
```

#### 6.11.4.2 Beneficial mutations

```
ggplot(
   filter(summary_data, condition=="PLASTIC" & dominant_lineage_num_mut_steps
   aes(x=frac_unexpressed_beneficial_mut_steps)
) +
   geom_density() +
   theme(
    legend.position="none"
)
```



bo <- boot(filter(summary\_data, condition=="PLASTIC" & dominant\_lineage\_num\_mut\_steps\_that\_change
print(bo)</pre>

```
##
## ORDINARY NONPARAMETRIC BOOTSTRAP
##
##
##
##
Call:
```

```
## boot(data = filter(summary_data, condition == "PLASTIC" & dominant_lineage_num_mut_:
       0)$frac_unexpressed_beneficial_mut_steps, statistic = samplemean,
       R = 10000)
##
##
##
## Bootstrap Statistics :
       original
                      bias
                               std. error
## t1* 0.4827586 0.0004188506 0.03981129
print(boot.ci(bo, conf=0.95, type="perc"))
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 10000 bootstrap replicates
##
## boot.ci(boot.out = bo, conf = 0.95, type = "perc")
##
## Intervals :
            Percentile
## Level
## 95%
         (0.4034, 0.5586)
## Calculations and Intervals on Original Scale
```

#### 6.12 Mutational robustness

Mutational robustness measures the fraction of one-step mutations on a focal genotype that result in a phenotypic change. Here, we calculate the mutational robustness of the representative genotype from each replicate (the most abundant genotype in the final population).

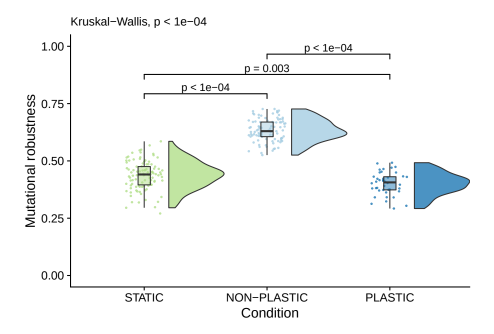
This data is located in a separate .tar.gz file on OSF, so we need to load it and wrangle the data.

```
# Load the data
df_mut = read.csv(paste0(working_directory, 'mutational_robustness/data/aggregated_mut.
# Extract the treatment for each line
df_mut$treatment = 'STATIC'
df_mut[df_mut$environment == 'chg-u100',]$treatment = 'PLASTIC'
df_mut[df_mut$environment == 'chg-u100' & df_mut$sensors == F,]$treatment = 'NON-PLAST
df_mut$treatment_factor = factor(df_mut$treatment, levels = c('STATIC', 'NON-PLASTIC',
# For compatibility with is_outlier above
df_mut$condition = df_mut$treatment_factor
# Calculate robustness (originally calculated as volatility)
df_mut$mutational_robustness = 1 - df_mut$one_step_diff_pheno_frac
```

Now we can plot mutational robustness:

```
# Compute manual labels for geom_signif
stat.test <- df_mut %>%
  wilcox_test(mutational_robustness ~ treatment_factor) %>%
  adjust_pvalue(method = "bonferroni") %>%
  add_significance() %>%
  add_xy_position(x="treatment_factor")
# Tweak y.position manually to account for scaled axis (edge case that triggers bad behavior in a
stat.test$manual_position <- stat.test$y.position * c(1.05,1.1,1.15)
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
df_mut$is_outlier <- mapply(</pre>
  is_outlier,
  df mut$mutational robustness,
  df_mut$treatment_factor,
  MoreArgs=list(data=df_mut, column="mutational_robustness")
# Remap colors so that colors map
color_map = c(
  'STATIC' = brewer.pal(3, cb_palette)[3],
  'PLASTIC' = brewer.pal(3, cb_palette)[2],
  'NON-PLASTIC' = brewer.pal(3, cb_palette)[1]
)
# Plot!
mut_robustness_fig <- ggplot(df_mut, aes(x=treatment_factor, y=mutational_robustness, fill=treatment_factor)</pre>
  geom_flat_violin( data=filter(df_mut,is_outlier==FALSE),
    scale="width", position = position_nudge(x = .2, y = 0), alpha = .8) +
  geom_point(mapping=aes(color=treatment_factor), position = position_jitter(width = .15), size =
  geom_boxplot(width = .1, outlier.shape = NA, alpha = 0.5) +
  scale_x_discrete( name="Condition") +
  scale_y_continuous( name='Mutational robustness', limits=c(0, 1)) +
  scale_fill_manual( values = color_map ) +
  scale_color_manual( values = color_map ) +
  labs( subtitle=paste0( "Kruskal-Wallis, ", p_label(signif(kruskal.test(formula=mutational_robus
  ggsignif::geom_signif( data=filter(stat.test, p.adj <= alpha), aes(xmin=group1,xmax=group2,anr
  theme( legend.position="none" )
```

```
## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y_position
mut_robustness_fig
```

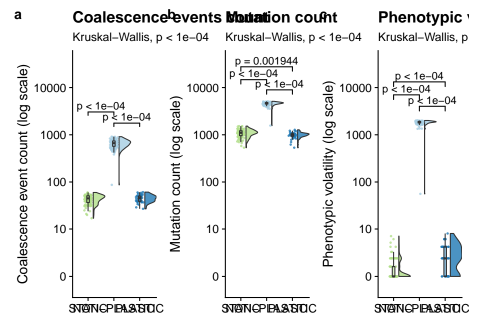


#### 6.13 Manuscript figures

Figures styled for the paper.

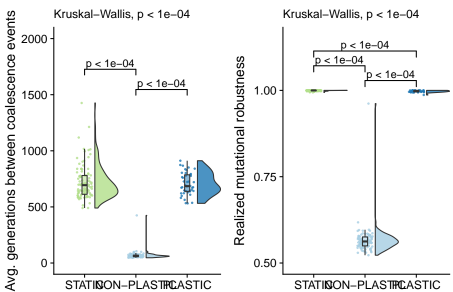
```
magnitude_grid <- plot_grid(</pre>
  coalescence_events_fig +
    theme(
      legend.position="none",
      axis.title.x=element blank()
    ggtitle("Coalescence events count"),
  mutation_count_fig +
    theme(
      legend.position="none",
      axis.title.x=element_blank()
    ) +
    ggtitle("Mutation count"),
  phenotypic_volatility_fig +
    theme(
      legend.position="none",
      axis.title.x=element_blank()
    ) +
    ggtitle("Phenotypic volatility"),
  nrow=1,
```

```
ncol=3,
align="v",
labels="auto"
)
magnitude_grid
```



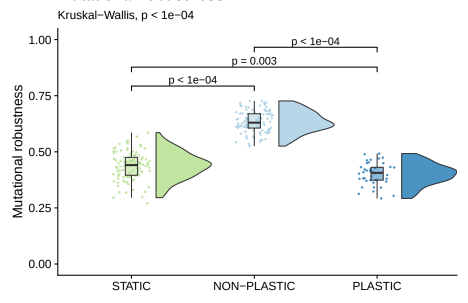
```
pace_grid <- plot_grid(
    coalescence_events_freq_fig +
        theme(
        legend.position="none",
        axis.title.x=element_blank()
    ) +
    ggtitle("Generations between coalescence events"),
    mutational_stability_fig +
        theme(
        legend.position="none",
        axis.title.x=element_blank()
    ) +
    ggtitle("Realized mutational robustness"),
    nrow=1,
    ncol=2,
    align="v",
    labels="auto"
)
pace_grid</pre>
```

#### a Generations between coalesce Realizants utational rob



```
# Even though mutational robustness is shown by itself, this ensures it is plotted ide
mut_robustness_grid = plot_grid(
    mut_robustness_fig +
        theme(
        legend.position="none",
        axis.title.x=element_blank()
    ) +
        ggtitle("Mutational robustness"),
        nrow=1,
        ncol=1,
        align="v",
        labels=""
)
mut_robustness_grid
```

#### **Mutational robustness**



```
save_plot(
  paste0(working_directory, "plots/", "evolutionary-change-magnitude-panel.pdf"),
 magnitude_grid,
  base_height=6,
  base_asp=3/1
)
save_plot(
 paste0(working_directory, "plots/", "evolutionary-change-pace-panel.pdf"),
 pace_grid,
 base_height=6,
  base_asp=2/1
save_plot(
 paste0(working_directory, "plots/", "mutational-robustness.pdf"),
 mut_robustness_grid,
 base_height=6,
  base_asp=1
```

## Chapter 7

# Evolution and maintenance of novel traits

The effect of adaptive phenotypic plasticity on the evolution and maintenance of novel tasks.

#### 7.1 Overview

```
total_updates <- 200000
replicates <- 100
alpha <- 0.05
focal_traits <- c("not", "nand", "and", "ornot", "or", "andnot")</pre>
traits_set_a <- c("not", "and", "or")</pre>
traits_set_b <- c("nand", "ornot", "andnot")</pre>
extra_traits <- c(
  "nor", "xor", "equals",
  "logic_3aa", "logic_3ab", "logic_3ac",
  "logic_3ad","logic_3ae","logic_3af",
  "logic_3ag", "logic_3ah", "logic_3ai",
  "logic_3aj", "logic_3ak", "logic_3al",
  "logic_3am", "logic_3an", "logic_3ao",
  "logic_3ap", "logic_3aq", "logic_3ar",
  "logic_3as", "logic_3at", "logic_3au",
  "logic_3av", "logic_3aw", "logic_3ax",
  "logic_3ay", "logic_3az", "logic_3ba",
  "logic_3bb", "logic_3bc", "logic_3bd",
  "logic_3be", "logic_3bf", "logic_3bg",
  "logic_3bh", "logic_3bi", "logic_3bj",
```

```
"logic_3bk","logic_3bn","logic_3bm",
  "logic_3bn","logic_3bs","logic_3bs",
  "logic_3bd","logic_3bs","logic_3bs",
  "logic_3bt","logic_3bw","logic_3bv",
  "logic_3bw","logic_3bx","logic_3by",
  "logic_3bz","logic_3ca","logic_3cb",
  "logic_3cc","logic_3cd","logic_3ce",
  "logic_3cf","logic_3cg","logic_3ch",
  "logic_3ci","logic_3cj","logic_3ck",
  "logic_3cl","logic_3cm","logic_3cn",
  "logic_3co","logic_3cp"
)

# Relative location of data.
working_directory <- "experiments/2021-01-31-complex-features/analysis/" # << For book
# working_directory <- "./"</pre>
```

#### 7.2 Analysis dependencies

Load all required R libraries.

```
library(ggplot2)
library(rstatix)
library(ggsignif)
library(scales)
library(tidyverse)
library(cowplot)
library(RColorBrewer)
library(Hmisc)
library(boot)
source("https://gist.githubusercontent.com/benmarwick/2a1bb0133ff568cbe28d/raw/fb53bd9")
```

These analyses were conducted/knitted with the following computing environment:

```
print(version)
##
## platform
                 x86_64-pc-linux-gnu
## arch
                 x86_64
## os
                 linux-gnu
## system
                 x86_64, linux-gnu
## status
## major
                 4
                 1.0
## minor
## year
                 2021
```

7.3. SETUP 81

```
## month 05

## day 18

## svn rev 80317

## language R

## version.string R version 4.1.0 (2021-05-18)

## nickname Camp Pontanezen
```

#### 7.3 Setup

```
###### summary data ######
summary_data_loc <- paste0(working_directory, "data/aggregate.csv")</pre>
summary_data <- read.csv(summary_data_loc, na.strings="NONE")</pre>
summary_data$DISABLE_REACTION_SENSORS <- as.factor(summary_data$DISABLE_REACTION_SENSORS)</pre>
summary_data$chg_env <- summary_data$chg_env == "True"</pre>
summary_data$dominant_plastic_odd_even <- as.factor(summary_data$dominant_plastic_odd_even)</pre>
summary_data$sensors <- summary_data$DISABLE_REACTION_SENSORS == "0"</pre>
summary_data$is_plastic <- summary_data$dominant_plastic_odd_even == "True"</pre>
summary_data$extra_task_value <- as.factor(summary_data$extra_task_value)
summary_data <- filter(summary_data, extra_task_value == 0.1)</pre>
env_label_fun <- function(chg_env) {</pre>
  if (chg env) {
    return("Fluctuating")
  } else {
    return("Constant")
  }
}
sensors_label_fun <- function(has_sensors) {</pre>
  if (has_sensors) {
    return("Sensors")
  } else {
    return("No sensors")
}
condition_label_fun <- function(has_sensors, env_chg) {</pre>
  if (has_sensors && env_chg) {
    return("PLASTIC")
  } else if (env_chg) {
    return("NON-PLASTIC")
  } else {
 return("STATIC")
```

```
}
}
summary_data$env_label <- mapply(</pre>
  env_label_fun,
  summary_data$chg_env
summary_data$sensors_label <- mapply(</pre>
  sensors_label_fun,
  summary_data$sensors
summary_data$condition <- mapply(</pre>
  condition_label_fun,
  summary_data$sensors,
  summary_data$chg_env
condition_order = c(
  "STATIC",
  "NON-PLASTIC",
  "PLASTIC"
pairwise_comparisons <- list(</pre>
  c("STATIC", "NON-PLASTIC"),
  c("STATIC", "PLASTIC"),
  c("PLASTIC", "NON-PLASTIC")
)
p_label <- function(p_value) {</pre>
  threshold = 0.0001
  if (p_value < threshold) {</pre>
    return(paste0("p < ", threshold))</pre>
  } else {
    return(paste0("p = ", p_value))
  }
}
# *really* inefficient way to identify outliers
is_outlier <- function(value, cond, data, column) {</pre>
  cond_data <- filter(data, condition==cond)</pre>
  q1 <- summary(cond_data[,column])[["1st Qu."]]
  q3 <- summary(cond_data[,column])[["3rd Qu."]]
  H <- 1.5 * IQR(cond_data[,column])</pre>
  return( (value < (q1-H)) | (value > (q3+H)) )
```

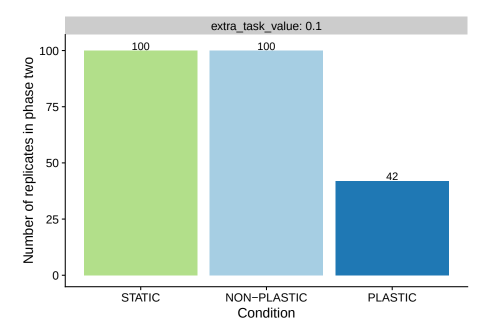
```
##### time series #####
lineage_time_series_data_loc <- paste0(working_directory, "data/lineage_series.csv")</pre>
lineage_time_series_data <- read.csv(lineage_time_series_data_loc)</pre>
lineage_time_series_data$DISABLE_REACTION_SENSORS <- as.factor(lineage_time_series_data$DISABLE_I
lineage_time_series_data$chg_env <- lineage_time_series_data$chg_env == "True"</pre>
lineage_time_series_data$sensors <- lineage_time_series_data$DISABLE_REACTION_SENSORS == "0"
lineage_time_series_data$extra_task_value <- as.factor(lineage_time_series_data$extra_task_value)
lineage_time_series_data$env_label <- mapply(</pre>
  env_label_fun,
  lineage_time_series_data$chg_env
lineage_time_series_data$sensors_label <- mapply(</pre>
  sensors_label_fun,
  lineage_time_series_data$sensors
)
lineage_time_series_data$condition <- mapply(</pre>
  condition_label_fun,
  lineage_time_series_data$sensors,
  lineage_time_series_data$chg_env
###### misc ######
# Configure our default graphing theme
theme_set(theme_cowplot())
# Palette
cb_palette <- "Paired"
# Create directory to dump plots
dir.create(paste0(working_directory, "plots"), showWarnings=FALSE)
# Sample mean function
samplemean <- function(x, d) {</pre>
  return(mean(x[d]))
```

## 7.4 The evolution of phenotypic plasticity

For sensor-enabled populations in fluctuating environments, we only transferred populations containing an optimally plastic genotype to phase two.

```
summary_data_grouped = dplyr::group_by(summary_data, sensors, env_label, condition, extra_task_vasummary_data_group_counts = dplyr::summarize(summary_data_grouped, n=dplyr::n())
ggplot(summary_data_group_counts, aes(x=condition, y=n, fill=condition)) +
```

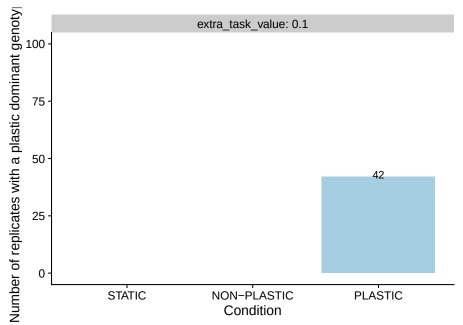
```
geom_col(position=position_dodge(0.9)) +
geom_text(aes(label=n, y=n+2)) +
scale_x_discrete(
   name="Condition",
   limits=condition_order
) +
scale_fill_brewer(
   palette=cb_palette
) +
scale_color_brewer(
   palette=cb_palette
) +
ylab("Number of replicates in phase two") +
facet_wrap(~extra_task_value, labeller=label_both) +
theme(
   legend.position="none"
)
```



We can confirm our expectation that the dominant genotypes in non-plastic conditions are not phenotypically plastic.

```
summary_data_grouped = dplyr::group_by(summary_data, condition, is_plastic, extra_task,
summary_data_group_counts = dplyr::summarize(summary_data_grouped, n=dplyr::n())
ggplot(filter(summary_data_group_counts, is_plastic), aes(x=condition, y=n, fill=condit
geom_col(position=position_dodge(0.9)) +
```

```
scale_x_discrete(
   name="Condition",
   limits=condition_order
) +
scale_fill_brewer(
   palette=cb_palette
) +
scale_color_brewer(
   palette=cb_palette
) +
ylim(0, 100) +
geom_text(aes(label=n, y=n+1)) +
ylab("Number of replicates with a plastic dominant genotype") +
facet_wrap(~extra_task_value, labeller=label_both) +
theme(
   legend.position="none"
)
```



## 7.5 Final novel task count (dominant genotype)

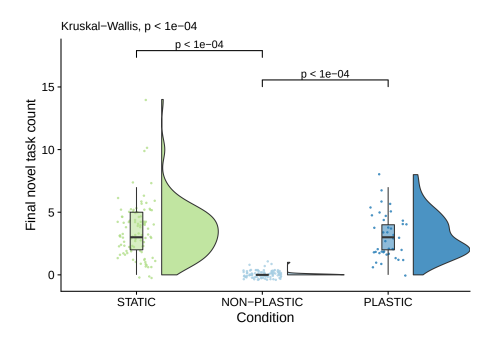
How many novel tasks do final dominant genotypes perform?

```
# Compute manual labels for geom_signif
stat.test <- summary_data %>%
```

```
wilcox_test(dominant_extra_tasks ~ condition) %>%
  adjust_pvalue(method = "bonferroni") %>%
  add_significance() %>%
  add_xy_position(x="condition") # ,step.increase=1
# Tweak y.position manually to account for scaled axis (edge case that triggers bad be
stat.test$manual_position <- stat.test$y.position #* c(1.0,1.0,1.03)
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
summary_data$is_outlier <- mapply(</pre>
  is_outlier,
  summary_data$dominant_extra_tasks,
  summary data$condition,
  MoreArgs=list(data=summary_data, column="dominant_extra_tasks")
final_novel_task_count_fig <- ggplot(</pre>
    summary_data,
    aes(x=condition, y=dominant_extra_tasks, fill=condition)
  geom_flat_violin(
    # data=filter(summary_data,is_outlier==FALSE),
    scale="width",
    position = position_nudge(x = .2, y = 0),
    alpha = .8
  ) +
  geom_point(
    mapping=aes(color=condition),
    position = position_jitter(width = .15),
    size = .5,
    alpha = 0.8
  ) +
  geom_boxplot(
    width = .1,
    outlier.shape = NA,
    alpha = 0.5
  ) +
  scale_x_discrete(
    name="Condition",
    limits=condition_order,
    labels=condition_order
  scale_y_continuous(
    name="Final novel task count"
  scale_fill_brewer(
```

```
palette=cb_palette
) +
scale_color_brewer(
  palette=cb_palette
) +
labs(
  subtitle=paste0(
    "Kruskal-Wallis, ",
    p_label(signif(kruskal.test(formula=dominant_extra_tasks~condition, data=summary_data)$p.va
  )
) +
ggsignif::geom_signif(
  data=filter(stat.test, p.adj <= alpha),</pre>
  aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
  manual=TRUE,
  inherit.aes=FALSE
) +
# coord_flip()
theme(
  legend.position="none"
```

## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y\_position
final\_novel\_task\_count\_fig



```
kruskal.test(
  formula=dominant_extra_tasks~condition,
  data=summary_data
)
##
   Kruskal-Wallis rank sum test
##
##
## data: dominant_extra_tasks by condition
## Kruskal-Wallis chi-squared = 177.17, df = 2, p-value < 2.2e-16
pairwise.wilcox.test(
  x=summary_data$dominant_extra_tasks,
  g=summary_data$condition,
  p.adjust.method="bonferroni",
  conf.int=TRUE,
  conf.level=0.95
)
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
##
## data: summary_data$dominant_extra_tasks and summary_data$condition
##
##
           NON-PLASTIC PLASTIC
## PLASTIC <2e-16
## STATIC <2e-16
                       0.9
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
    "PLASTIC median: ",
   median(filter(summary_data, condition=="PLASTIC")$dominant_extra_tasks)
  ),
  paste0(
    "STATIC median: ",
   median(filter(summary_data, condition=="STATIC")$dominant_extra_tasks)
 ),
  paste0(
    "NON-PLASTIC median: ",
    median(filter(summary_data, condition=="NON-PLASTIC")$dominant_extra_tasks)
  )
)
```

## [1] "PLASTIC median: 3; STATIC median: 3; NON-PLASTIC median: 0"

```
print("Wilcox rank sum test statistics:")

## [1] "Wilcox rank sum test statistics:"

for (pair in pairwise_comparisons) {
    pair_data <- filter(summary_data, condition %in% pair)
    pair_data$condition <- as.factor(pair_data$condition)
    wt <- wilcox.test(
        formula=dominant_extra_tasks~condition,
        data=pair_data,
        exact=FALSE,
        paired=FALSE
    )
    print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
}

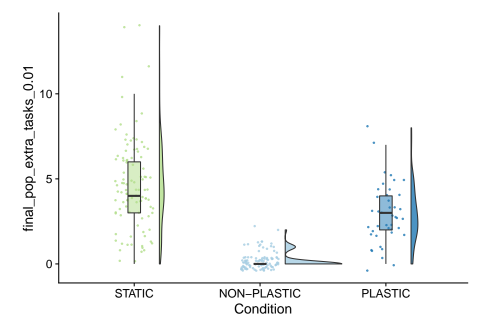
## [1] "STATIC<-->NON-PLASTIC: W=184"
## [1] "STATIC<-->PLASTIC: W=1871"
## [1] "PLASTIC<-->NON-PLASTIC: W=64"
```

### 7.6 Novel task count (final population)

How many novel tasks are performed across the final population (1% of organisms must perform to count)?

```
ggplot(summary_data, aes(x=condition, y=final_pop_extra_tasks_0.01, fill=condition)) +
  geom_flat_violin(
   position = position_nudge(x = .2, y = 0),
   alpha = .8
  ) +
  geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
   alpha = 0.8
  ) +
  geom_boxplot(
   width = .1,
   outlier.shape = NA,
   alpha = 0.5
  scale_x_discrete(
   name="Condition",
   limits=condition_order
  scale_fill_brewer(
```

```
palette=cb_palette
) +
scale_color_brewer(
  palette=cb_palette
) +
# coord_flip() +
theme(
  legend.position="none"
)
```



```
kruskal.test(
  formula=final_pop_extra_tasks_0.01~condition,
  data=summary_data
)

##

## Kruskal-Wallis rank sum test

##

## data: final_pop_extra_tasks_0.01 by condition

## Kruskal-Wallis chi-squared = 169.47, df = 2, p-value < 2.2e-16

pairwise.wilcox.test(
  x=summary_data$final_pop_extra_tasks_0.01,
  g=summary_data$condition,
  p.adjust.method="bonferroni",
  conf.int=TRUE,</pre>
```

```
conf.level=0.95
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: summary_data$final_pop_extra_tasks_0.01 and summary_data$condition
##
           NON-PLASTIC PLASTIC
## PLASTIC < 2e-16
## STATIC < 2e-16
                       0.00016
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
   "PLASTIC median: ",
   median(filter(summary_data, condition=="PLASTIC")$final_pop_extra_tasks_0.01)
 ),
  paste0(
   "STATIC median: ",
   median(filter(summary_data, condition=="STATIC")$final_pop_extra_tasks_0.01)
 ),
  paste0(
    "NON-PLASTIC median: ",
   median(filter(summary_data, condition=="NON-PLASTIC")$final_pop_extra_tasks_0.01)
)
## [1] "PLASTIC median: 3; STATIC median: 4; NON-PLASTIC median: 0"
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
 pair_data <- filter(summary_data, condition %in% pair)</pre>
 pair_data$condition <- as.factor(pair_data$condition)</pre>
  wt <- wilcox.test(
   formula=final_pop_extra_tasks_0.01~condition,
   data=pair_data,
   exact=FALSE,
   paired=FALSE
 )
 print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
```

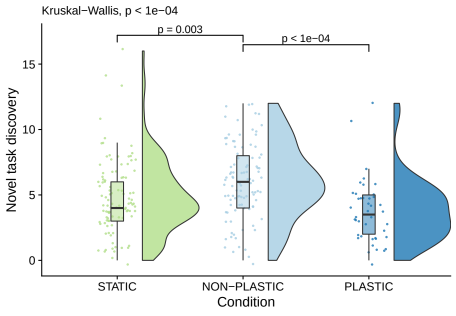
```
## [1] "STATIC<-->NON-PLASTIC: W=227.5"
## [1] "STATIC<-->PLASTIC: W=1203"
## [1] "PLASTIC<-->NON-PLASTIC: W=225.5"
```

#### 7.7 Novel task discovery (lineage)

```
# Compute manual labels for geom_signif
stat.test <- summary_data %>%
  wilcox_test(dominant_lineage_extra_traits_discovered ~ condition) %>%
  adjust_pvalue(method = "bonferroni") %>%
 add_significance() %>%
 add_xy_position(x="condition") # ,step.increase=1
# Tweak y.position manually to account for scaled axis (edge case that triggers bad be
stat.test\$manual\_position \leftarrow stat.test\$y.position #* c(1.0,1.0,1.03)
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
summary_data$is_outlier <- mapply(</pre>
  is outlier,
 summary_data$dominant_lineage_extra_traits_discovered,
 summary_data$condition,
 MoreArgs=list(data=summary_data, column="dominant_lineage_extra_traits_discovered")
lineage_novel_task_discovery_fig <- ggplot(</pre>
    summary data,
    aes(x=condition, y=dominant_lineage_extra_traits_discovered, fill=condition)
  ) +
  geom_flat_violin(
    # data=filter(summary_data, is_outlier==FALSE),
    scale="width",
    position = position_nudge(x = .2, y = 0),
    alpha = .8
  ) +
  geom_point(
    mapping=aes(color=condition),
    position = position_jitter(width = .15),
    size = .5,
    alpha = 0.8
  ) +
  geom boxplot(
   width = .1,
    outlier.shape = NA,
    alpha = 0.5
```

```
scale_x_discrete(
  name="Condition",
  limits=condition_order,
 labels=condition_order
scale_y_continuous(
  name="Novel task discovery"
) +
scale_fill_brewer(
  palette=cb_palette
scale_color_brewer(
  palette=cb_palette
) +
labs(
  subtitle=paste0(
    "Kruskal-Wallis, ",
   p_label(signif(kruskal.test(formula=dominant_lineage_extra_traits_discovered~condition, dat
  )
) +
ggsignif::geom_signif(
  data=filter(stat.test, p.adj <= alpha),</pre>
  aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
  manual=TRUE,
  inherit.aes=FALSE
) +
# coord_flip()
theme(
  legend.position="none"
```

## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y\_position
lineage\_novel\_task\_discovery\_fig



```
kruskal.test(
  formula=dominant_lineage_extra_traits_discovered~condition,
  data=summary_data
)
##
   Kruskal-Wallis rank sum test
##
##
## data: dominant_lineage_extra_traits_discovered by condition
## Kruskal-Wallis chi-squared = 24.099, df = 2, p-value = 5.846e-06
pairwise.wilcox.test(
  x=summary_data$dominant_lineage_extra_traits_discovered,
  g=summary_data$condition,
 p.adjust.method="bonferroni",
 conf.int=TRUE,
  conf.level=0.95
)
```

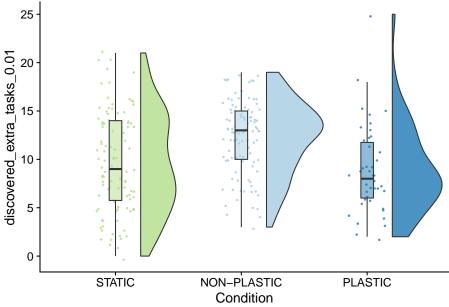
```
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: summary_data$dominant_lineage_extra_traits_discovered and summary_data$condi
##
## NON-PLASTIC PLASTIC
## PLASTIC 1.7e-05 -
```

```
## STATIC 0.0035
                       0.0561
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
    "PLASTIC median: ",
   median(filter(summary_data, condition=="PLASTIC")$dominant_lineage_extra_traits_discovered)
  ),
  paste0(
   "STATIC median: ",
   median(filter(summary_data, condition=="STATIC")$dominant_lineage_extra_traits_discovered)
 ),
  paste0(
   "NON-PLASTIC median: ",
   median(filter(summary_data, condition=="NON-PLASTIC")$dominant_lineage_extra_traits_discovered
  )
)
## [1] "PLASTIC median: 3.5; STATIC median: 4; NON-PLASTIC median: 6"
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
  pair_data <- filter(summary_data, condition %in% pair)</pre>
 pair_data$condition <- as.factor(pair_data$condition)</pre>
  wt <- wilcox.test(
   formula=dominant_lineage_extra_traits_discovered~condition,
    data=pair data,
   exact=FALSE,
   paired=FALSE
  print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
## [1] "STATIC<-->NON-PLASTIC: W=6319.5"
## [1] "STATIC<-->PLASTIC: W=1578"
## [1] "PLASTIC<-->NON-PLASTIC: W=3110.5"
```

## 7.8 Novel task discovery (population)

```
ggplot(
    summary_data,
    aes(x=condition, y=discovered_extra_tasks_0.01, fill=condition)
```

```
) +
  geom_flat_violin(
    position = position_nudge(x = .2, y = 0),
    alpha = .8
  ) +
  geom_point(
    mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
    alpha = 0.8
  geom_boxplot(
   width = .1,
   outlier.shape = NA,
   alpha = 0.5
  ) +
  scale_x_discrete(
    name="Condition",
   limits=condition_order
  scale_fill_brewer(
   palette=cb_palette
  ) +
  scale_color_brewer(
   palette=cb_palette
  # coord_flip() +
  theme(
    legend.position="none"
```



```
kruskal.test(
  formula=discovered_extra_tasks_0.01~condition,
  data=summary_data
)
##
   Kruskal-Wallis rank sum test
##
##
## data: discovered_extra_tasks_0.01 by condition
## Kruskal-Wallis chi-squared = 24.271, df = 2, p-value = 5.365e-06
pairwise.wilcox.test(
  x=summary_data$discovered_extra_tasks_0.01,
  g=summary_data$condition,
 p.adjust.method="bonferroni",
  conf.int=TRUE,
  conf.level=0.95
)
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: summary_data$discovered_extra_tasks_0.01 and summary_data$condition
##
```

##

## PLASTIC 2.4e-05

NON-PLASTIC PLASTIC

```
## STATIC 0.00035
                       1.00000
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
    "PLASTIC median: ",
   median(filter(summary_data, condition=="PLASTIC")$discovered_extra_tasks_0.01)
  ),
  paste0(
    "STATIC median: ",
    median(filter(summary data, condition=="STATIC")$discovered extra tasks 0.01)
  ),
  paste0(
    "NON-PLASTIC median: ",
    median(filter(summary_data, condition=="NON-PLASTIC")$discovered_extra_tasks_0.01)
  )
## [1] "PLASTIC median: 8; STATIC median: 9; NON-PLASTIC median: 13"
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
  pair_data <- filter(summary_data, condition %in% pair)</pre>
  pair_data$condition <- as.factor(pair_data$condition)</pre>
  wt <- wilcox.test(
    formula=discovered_extra_tasks_0.01~condition,
    data=pair_data,
    exact=FALSE,
    paired=FALSE
  print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
## [1] "STATIC<-->NON-PLASTIC: W=6573.5"
## [1] "STATIC<-->PLASTIC: W=1918.5"
## [1] "PLASTIC<-->NON-PLASTIC: W=3096"
```

### 7.9 Novel task discovery frequency (lineage)

```
summary_data$dominant_lineage_extra_traits_discovered_per_generation <- summary_data$dominant_lineage_extra_traits_generations_per_discovery <- summary_data$dominant_lineage_extra_traits_generations_per_discovery <- summary_data$dominant_lineage_extra_traits_generations_per_discovery <- summary_data$dominant_lineage_extra_traits_generations_per_discovery <- summary_data$dominant_lineage_extra_traits_discovered_per_generation</pre>
```

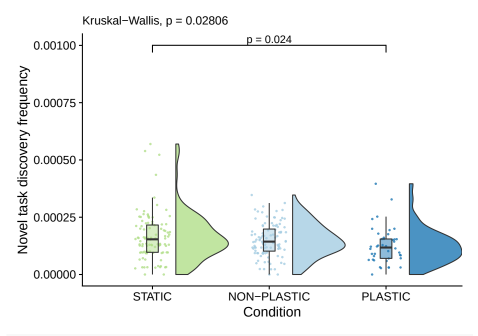
```
# Compute manual labels for geom_signif
# stat.test <- filter(summary_data, dominant_lineage_extra_traits_discovered > 0) %>%
# wilcox_test(dominant_lineage_extra_traits_generations_per_discovery ~ condition) %>%
# adjust_pvalue(method = "bonferroni") %>%
# add_significance() %>%
# add_xy_position(x="condition") # ,step.increase=1
stat.test <- summary data %>%
 wilcox_test(dominant_lineage_extra_traits_discovered_per_generation ~ condition) %>%
  adjust_pvalue(method = "bonferroni") %>%
  add_significance() %>%
  add_xy_position(x="condition", step.increase=0.0001) # ,step.increase=1
# Tweak y.position manually to account for scaled axis (edge case that triggers bad behavior in
stat.testmanual_position \leftarrow stat.test_y.position #* <math>c(1.0,1.0,1.03)
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
summary_data$is_outlier <- mapply(</pre>
  is_outlier,
  summary_data$dominant_lineage_extra_traits_discovered_per_generation,
  summary_data$condition,
  MoreArgs=list(data=summary_data, column="dominant_lineage_extra_traits_discovered_per_generation)
lineage_novel_task_discovery_freq_fig <- ggplot(</pre>
    # filter(summary data, dominant lineage extra traits discovered > 0),
    summary_data,
    aes(x=condition, y=dominant_lineage_extra_traits_discovered_per_generation, fill=condition)
  ) +
  geom_flat_violin(
    # data=filter(summary_data, is_outlier==FALSE),
    scale="width",
   position = position_nudge(x = .2, y = 0),
   alpha = .8
  ) +
  geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
   alpha = 0.8
  ) +
  geom_boxplot(
   width = .1,
   outlier.shape = NA,
    alpha = 0.5
  ) +
  scale_x_discrete(
```

#### 100CHAPTER 7. EVOLUTION AND MAINTENANCE OF NOVEL TRAITS

```
name="Condition",
  limits=condition_order
) +
ylab("Novel task discovery frequency") +
scale_fill_brewer(
  palette=cb_palette
scale_color_brewer(
 palette=cb_palette
) +
labs(
  subtitle=paste0(
    "Kruskal-Wallis, ",
    p_label(signif(kruskal.test(formula=dominant_lineage_extra_traits_discovered_per_
) +
ggsignif::geom_signif(
  data=filter(stat.test, p.adj <= alpha),</pre>
  aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
  manual=TRUE,
  inherit.aes=FALSE
) +
# coord_flip() +
theme(
  legend.position="none"
```

## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y\_position
lineage\_novel\_task\_discovery\_freq\_fig

## PLASTIC 0.092

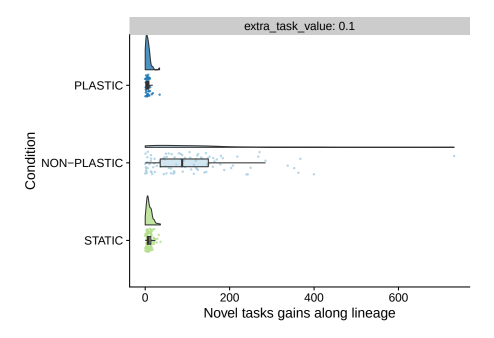


```
kruskal.test(
  formula=dominant_lineage_extra_traits_discovered_per_generation~condition,
  data=summary_data
)
##
##
   Kruskal-Wallis rank sum test
##
## data: dominant_lineage_extra_traits_discovered_per_generation by condition
## Kruskal-Wallis chi-squared = 7.1465, df = 2, p-value = 0.02806
pairwise.wilcox.test(
  x=summary_data$dominant_lineage_extra_traits_discovered_per_generation,
  g=summary_data$condition,
 p.adjust.method="bonferroni",
  conf.int=TRUE,
  conf.level=0.95
)
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: summary_data$dominant_lineage_extra_traits_discovered_per_generation and summary_data$0
##
           NON-PLASTIC PLASTIC
##
```

```
## STATIC 1.000
                       0.025
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
    "PLASTIC median: ",
   median(filter(summary_data, condition=="PLASTIC")$dominant_lineage_extra_traits_di
  ),
  paste0(
    "STATIC median: ",
   median(filter(summary data, condition=="STATIC")$dominant lineage extra traits dis
  ),
  paste0(
    "NON-PLASTIC median: ",
   median(filter(summary_data, condition=="NON-PLASTIC")$dominant_lineage_extra_traits
  )
## [1] "PLASTIC median: 0.000117695011124939; STATIC median: 0.00015363220504867; NON-
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
  pair_data <- filter(summary_data, condition %in% pair)</pre>
  pair_data$condition <- as.factor(pair_data$condition)</pre>
  wt <- wilcox.test(
    formula=dominant_lineage_extra_traits_discovered_per_generation~condition,
    data=pair data,
    exact=FALSE,
    paired=FALSE
  print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
## [1] "STATIC<-->NON-PLASTIC: W=4751"
## [1] "STATIC<-->PLASTIC: W=1510.5"
## [1] "PLASTIC<-->NON-PLASTIC: W=2584"
        Novel tasks gained (lineage)
ggplot(
```

```
ggplot(
    summary_data,
    aes(x=condition, y=dominant_lineage_extra_traits_gained, fill=condition)
```

```
) +
 geom_flat_violin(
   position = position_nudge(x = .2, y = 0),
   alpha = .8
 ) +
 geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
   alpha = 0.8
 geom_boxplot(
   width = .1,
   outlier.shape = NA,
   alpha = 0.5
 scale_x_discrete(
   name="Condition",
   limits=condition_order
 ylab("Novel tasks gains along lineage") +
 scale_fill_brewer(
   palette=cb_palette
 ) +
 scale_color_brewer(
   palette=cb_palette
 ) +
 coord_flip() +
 facet_wrap(
   ~extra_task_value,
   labeller=label_both
 ) +
 theme(
   legend.position="none"
 ggsave(
   paste0(working_directory, "plots/dominant-lineage-extra-tasks-gained.pdf"),
   width=15,
   height=10
```



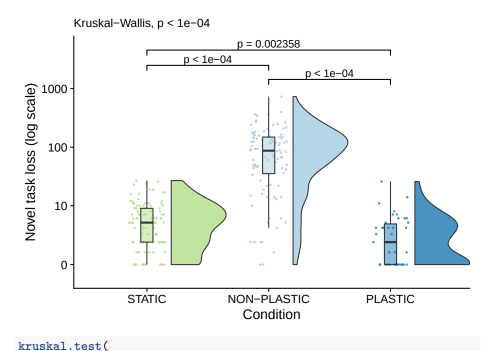
## 7.11 Novel task loss (lineage)

```
# Compute manual labels for geom_signif
stat.test <- summary_data %>%
  wilcox_test(dominant_lineage_extra_traits_lost ~ condition) %>%
  adjust_pvalue(method = "bonferroni") %>%
  add_significance() %>%
  add_xy_position(x="condition", step.increase=1)
# Tweak y.position manually to account for scaled axis (edge case that triggers bad be
stat.test$manual_position <- log10(stat.test$y.position) * c(1.0,1.0,1.03)
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
summary_data$is_outlier <- mapply(</pre>
  is_outlier,
  summary_data$dominant_lineage_extra_traits_lost,
  summary_data$condition,
  MoreArgs=list(data=summary_data, column="dominant_lineage_extra_traits_lost")
lineage_novel_task_loss_fig <- ggplot(</pre>
    summary_data,
    aes(x=condition, y=dominant_lineage_extra_traits_lost, fill=condition)
```

```
geom_flat_violin(
  # data=filter(summary_data,is_outlier==FALSE),
 scale="width",
 position = position_nudge(x = .2, y = 0),
 alpha = .8
) +
geom_point(
 mapping=aes(color=condition),
 position = position_jitter(width = .15),
 size = .5,
 alpha = 0.8
) +
geom_boxplot(
 width = .1,
 outlier.shape = NA,
 alpha = 0.5
) +
scale_x_discrete(
 name="Condition",
 limits=condition_order,
 labels=condition_order
) +
scale_y_continuous(
 name="Novel task loss (log scale)",
 trans=pseudo_log_trans(sigma=1, base=10),
 breaks=c(0,10,100,1000),
 limits=c(-1,5000)
) +
scale fill brewer(
 palette=cb_palette
) +
scale_color_brewer(
 palette=cb_palette
) +
labs(
  subtitle=paste0(
    "Kruskal-Wallis, ",
    p_label(signif(kruskal.test(formula=dominant_lineage_extra_traits_lost~condition, data=summer)
) +
ggsignif::geom_signif(
  data=filter(stat.test, p.adj<=alpha),</pre>
  aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
 manual=TRUE,
 inherit.aes=FALSE
```

```
) +
# coord_flip()
theme(
  legend.position="none"
)
```

## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y\_position
lineage\_novel\_task\_loss\_fig



```
formula=dominant_lineage_extra_traits_lost~condition,
  data=summary_data
)

##

## Kruskal-Wallis rank sum test

##

## data: dominant_lineage_extra_traits_lost by condition

## Kruskal-Wallis chi-squared = 129.06, df = 2, p-value < 2.2e-16

pairwise.wilcox.test(
    x=summary_data$dominant_lineage_extra_traits_lost,
    g=summary_data$condition,
    p.adjust.method="bonferroni",
    conf.int=TRUE,</pre>
```

```
conf.level=0.95
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: summary_data$dominant_lineage_extra_traits_lost and summary_data$condition
##
##
           NON-PLASTIC PLASTIC
## PLASTIC 2.7e-16
## STATIC < 2e-16
                       0.0024
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
   "PLASTIC median: ",
   median(filter(summary_data, condition=="PLASTIC")$dominant_lineage_extra_traits_lost)
  ),
  paste0(
    "STATIC median: ",
   median(filter(summary_data, condition=="STATIC")$dominant_lineage_extra_traits_lost)
 ),
  paste0(
    "NON-PLASTIC median: ",
   median(filter(summary_data, condition=="NON-PLASTIC")$dominant_lineage_extra_traits_lost)
)
## [1] "PLASTIC median: 2; STATIC median: 5; NON-PLASTIC median: 87.5"
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
 pair_data <- filter(summary_data, condition %in% pair)</pre>
 pair_data$condition <- as.factor(pair_data$condition)</pre>
  wt <- wilcox.test(
   formula=dominant_lineage_extra_traits_lost~condition,
   data=pair_data,
   exact=FALSE,
   paired=FALSE
 )
 print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
```

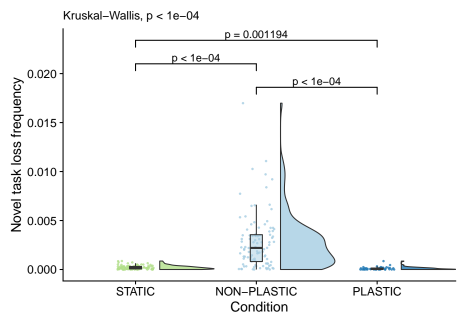
```
## [1] "STATIC<-->NON-PLASTIC: W=9105"
## [1] "STATIC<-->PLASTIC: W=1353.5"
## [1] "PLASTIC<-->NON-PLASTIC: W=3959"
```

#### 7.12 Frequency of novel task loss (lineage)

```
summary_data$dominant_lineage_extra_traits_lost_per_generation <- summary_data$dominan
summary_data$dominant_lineage_extra_traits_generations_per_loss <- summary_data$dominat
# Compute manual labels for geom_signif
# stat.test <- filter(summary_data, dominant_lineage_extra_traits_lost > 0) %>%
   wilcox_test(dominant_lineage_extra_traits_generations_per_loss ~ condition) %>%
   adjust_pvalue(method = "bonferroni") %>%
   add_significance() %>%
    add_xy_position(x="condition", step.increase=1)
stat.test <- summary_data %>%
  wilcox_test(dominant_lineage_extra_traits_lost_per_generation ~ condition) %>%
  adjust_pvalue(method = "bonferroni") %>%
  add_significance() %>%
 add_xy_position(x="condition", step.increase=.1)
# Tweak y.position manually to account for scaled axis (edge case that triggers bad be
\text{stat.test\$manual\_position} \leftarrow \text{stat.test\$y.position} \# * c(1.0,1.0,1.03)
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
summary_data$is_outlier <- mapply(</pre>
  is outlier,
  summary_data$dominant_lineage_extra_traits_lost_per_generation,
 summary_data$condition,
 MoreArgs=list(data=summary_data, column="dominant_lineage_extra_traits_lost_per_gene:
lineage_novel_task_loss_freq_fig <- ggplot(</pre>
    # filter(summary_data, dominant_lineage_extra_traits_lost > 0),
    summary_data,
    aes(x=condition, y=dominant_lineage_extra_traits_lost_per_generation, fill=condition)
  geom_flat_violin(
    # data=filter(summary_data,is_outlier==FALSE),
    scale="width",
    position = position_nudge(x = .2, y = 0),
   alpha = .8
  ) +
 geom_point(
```

```
mapping=aes(color=condition),
  position = position_jitter(width = .15),
  size = .5,
  alpha = 0.8
) +
geom_boxplot(
  width = .1,
  outlier.shape = NA,
  alpha = 0.5
) +
scale_x_discrete(
 name="Condition",
  limits=condition_order
ylab("Novel task loss frequency") +
scale_fill_brewer(
  palette=cb_palette
scale_color_brewer(
  palette=cb_palette
) +
labs(
  subtitle=paste0(
    "Kruskal-Wallis, ",
    p_label(signif(kruskal.test(formula=dominant_lineage_extra_traits_lost_per_generation~cond
) +
ggsignif::geom_signif(
  data=filter(stat.test, p.adj<=alpha),</pre>
  aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
  manual=TRUE,
  inherit.aes=FALSE
) +
theme(
  legend.position="none"
```

```
## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y_position
lineage_novel_task_loss_freq_fig
```



```
kruskal.test(
  formula=dominant_lineage_extra_traits_lost_per_generation~condition,
  data=summary_data
)
##
##
   Kruskal-Wallis rank sum test
##
## data: dominant_lineage_extra_traits_lost_per_generation by condition
## Kruskal-Wallis chi-squared = 121.41, df = 2, p-value < 2.2e-16
pairwise.wilcox.test(
  x=summary_data$dominant_lineage_extra_traits_lost_per_generation,
  g=summary_data$condition,
 p.adjust.method="bonferroni",
 conf.int=TRUE,
  conf.level=0.95
```

```
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: summary_data$dominant_lineage_extra_traits_lost_per_generation and summary_d
##
## NON-PLASTIC PLASTIC
## PLASTIC 1.1e-15 -
```

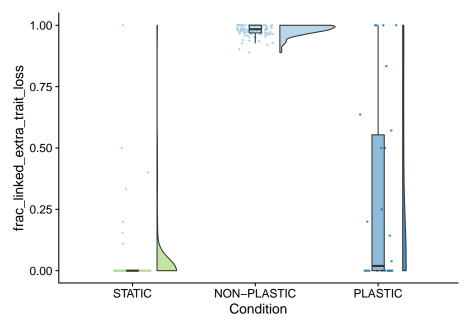
```
## STATIC < 2e-16
                        0.0012
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
    "PLASTIC median: ",
    median(filter(summary_data, condition=="PLASTIC")$dominant_lineage_extra_traits_lost_per_general
  ),
  paste0(
    "STATIC median: ",
    median(filter(summary_data, condition=="STATIC")$dominant_lineage_extra_traits_lost_per_gener
 ),
  paste0(
    "NON-PLASTIC median: ",
    median(filter(summary_data, condition=="NON-PLASTIC")$dominant_lineage_extra_traits_lost_per_
  )
)
## [1] "PLASTIC median: 6.25141973661864e-05; STATIC median: 0.000161396283669756; NON-PLASTIC median: 0.000161396283669756;
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
  pair_data <- filter(summary_data, condition %in% pair)</pre>
  pair_data$condition <- as.factor(pair_data$condition)</pre>
  wt <- wilcox.test(
    formula=dominant_lineage_extra_traits_lost_per_generation~condition,
    data=pair_data,
    exact=FALSE,
    paired=FALSE
 )
 print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
## [1] "STATIC<-->NON-PLASTIC: W=8940"
## [1] "STATIC<-->PLASTIC: W=1311"
## [1] "PLASTIC<-->NON-PLASTIC: W=3922"
```

#### 7.13 How many instances of novel trait loss cooccurred with changes in base phenotype?

Task loss linked with primary trait changes.

```
lost_traits_summary_data <- filter(summary_data, extra_task_value==0.1 & dominant_line lost_traits_summary_data$frac_linked_extra_trait_loss <- lost_traits_summary_data$dominates.
```

```
ggplot(lost_traits_summary_data, aes(x=condition, y=frac_linked_extra_trait_loss, fill=
 geom_flat_violin(
   position = position_nudge(x = .2, y = 0),
   alpha = .8
 ) +
 geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
   alpha = 0.8
 ) +
 geom_boxplot(
   width = .1,
   outlier.shape = NA,
   alpha = 0.5
 ) +
 scale_x_discrete(
   name="Condition",
   limits=condition_order
 ) +
  scale_fill_brewer(
   palette=cb_palette
 ) +
 scale_color_brewer(
   palette=cb_palette
  # coord_flip() +
 theme(
    legend.position="none"
```



```
kruskal.test(
  formula=frac_linked_extra_trait_loss~condition,
  data=lost_traits_summary_data
)
##
##
   Kruskal-Wallis rank sum test
##
## data: frac_linked_extra_trait_loss by condition
## Kruskal-Wallis chi-squared = 153.68, df = 2, p-value < 2.2e-16
pairwise.wilcox.test(
  x=lost_traits_summary_data$frac_linked_extra_trait_loss,
  g=lost_traits_summary_data$condition,
 p.adjust.method="bonferroni",
 conf.int=TRUE,
  conf.level=0.95
)
```

##

## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: lost\_traits\_summary\_data\$frac\_linked\_extra\_trait\_loss and lost\_traits\_summary\_data\$cond
##
## NON-PLASTIC PLASTIC
## PLASTIC 1.9e-08 -

```
## STATIC < 2e-16
                       1.8e-06
## P value adjustment method: bonferroni
paste(
  sep="; ",
  paste0(
    "PLASTIC median: ",
    median(filter(lost_traits_summary_data, condition=="PLASTIC")$frac_linked_extra_traits_summary_data
  ),
  paste0(
    "STATIC median: ",
    median(filter(lost_traits_summary_data, condition=="STATIC")$frac_linked_extra_tra
  ),
  paste0(
    "NON-PLASTIC median: ",
    median(filter(lost_traits_summary_data, condition=="NON-PLASTIC") $frac_linked_extra
  )
## [1] "PLASTIC median: 0.0192307692307692; STATIC median: 0; NON-PLASTIC median: 0.98
print("Wilcox rank sum test statistics:")
## [1] "Wilcox rank sum test statistics:"
for (pair in pairwise_comparisons) {
  pair_data <- filter(lost_traits_summary_data, condition %in% pair)</pre>
  pair_data$condition <- as.factor(pair_data$condition)</pre>
  wt <- wilcox.test(
    formula=frac_linked_extra_trait_loss~condition,
    data=pair_data,
    exact=FALSE,
    paired=FALSE
  )
  print(paste0(pair[1], "<-->", pair[2], ": W=",wt$statistic))
## [1] "STATIC<-->NON-PLASTIC: W=8344"
## [1] "STATIC<-->PLASTIC: W=1602"
## [1] "PLASTIC<-->NON-PLASTIC: W=2212"
sum(filter(lost_traits_summary_data, condition=="NON-PLASTIC")$dominant_lineage_extra_
## [1] 10998
sum(filter(lost_traits_summary_data, condition=="NON-PLASTIC")$dominant_lineage_extra_
## [1] 11229
```

```
aggregate_frac_linked_extra_trait_loss_nonplastic <- sum(filter(lost_traits_summary_data, condition))
aggregate_frac_linked_extra_trait_loss_nonplastic
## [1] 0.9794283
sum(filter(lost_traits_summary_data, condition=="PLASTIC")$dominant_lineage_extra_traits_lost_lineage
## [1] 29
sum(filter(lost_traits_summary_data, condition=="PLASTIC")$dominant_lineage_extra_traits_lost)
## [1] 142
aggregate_frac_linked_extra_trait_loss_plastic <- sum(filter(lost_traits_summary_data, condition=
aggregate_frac_linked_extra_trait_loss_plastic
## [1] 0.2042254
sum(filter(lost_traits_summary_data, condition=="STATIC")$dominant_lineage_extra_traits_lost_link
## [1] 13
sum(filter(lost_traits_summary_data, condition=="STATIC")$dominant_lineage_extra_traits_lost)
## [1] 631
aggregate_frac_linked_extra_trait_loss_nonplastic <- sum(filter(lost_traits_summary_data, condition))
aggregate_frac_linked_extra_trait_loss_nonplastic
## [1] 0.02060222
```

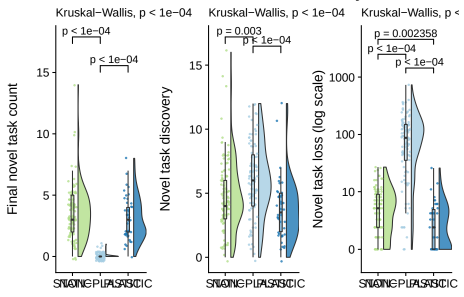
#### 7.14 Manuscript figures

#### 7.15 Combined panel

```
magnitude_grid <- plot_grid(
  final_novel_task_count_fig +
    theme(
      axis.title.x=element_blank()
    ) +
    ggtitle("Final novel task count"),
  lineage_novel_task_discovery_fig +
    theme(
      axis.title.x=element_blank()
    ) +
    ggtitle("Novel task discovery"),
  lineage_novel_task_loss_fig +
    theme(</pre>
```

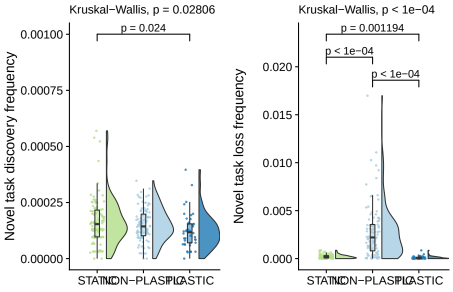
```
axis.title.x=element_blank()
) +
    ggtitle("Novel task loss"),
    nrow=1,
    align="v",
    labels="auto"
)
magnitude_grid
```

#### a Final novel task countovel task discoveryNovel task lo



```
pace_grid <- plot_grid(
  lineage_novel_task_discovery_freq_fig +
    theme(
      axis.title.x=element_blank()
  ) +
    ggtitle("Novel task discovery frequency"),
  lineage_novel_task_loss_freq_fig +
    theme(
      axis.title.x=element_blank()
  ) +
    ggtitle("Novel task loss frequency"),
    nrow=1,
    align="v",
    labels="auto"
)
pace_grid</pre>
```

#### a Novel task discovery Prequency Novel task loss frequ



```
save_plot(
   paste0(working_directory, "plots/", "complex-traits-magnitude-panel.pdf"),
   magnitude_grid,
   base_height=6,
   base_asp=3/1
)

save_plot(
   paste0(working_directory, "plots/", "complex-traits-pace-panel.pdf"),
   pace_grid,
   base_height=6,
   base_asp=2/1
)
```

#### 118CHAPTER 7. EVOLUTION AND MAINTENANCE OF NOVEL TRAITS

#### Chapter 8

# Accumulation of deleterious instructions

The effect of adaptive phenotypic plasticity on the accumulation of deleterious genes.

#### 8.1 Overview

```
total_updates <- 200000
replicates <- 100
alpha <- 0.05
focal_poison_penalty <- 0.1

focal_traits <- c("not", "nand", "ornot", "or", "andnot")
traits_set_a <- c("not", "and", "or")
traits_set_b <- c("nand", "ornot", "andnot")

# Relative location of data.
working_directory <- "experiments/2021-02-05-hitchhiking/analysis/" # << For bookdown
# working_directory <- "./"</pre>
```

#### 8.2 Analysis dependencies

Load all required R libraries.

```
library(RColorBrewer)
library(ggplot2)
library(rstatix)
```

```
library(ggsignif)
library(scales)
library(tidyverse)
library(cowplot)
library(Hmisc)
library(fmsb)
library(knitr)
source("https://gist.githubusercontent.com/benmarwick/2a1bb0133ff568cbe28d/raw/fb53bd9)
These analyses were conducted/knitted with the following computing environment:
print(version)
```

```
## platform
                 x86_64-pc-linux-gnu
## arch
                 x86_64
## os
                 linux-gnu
## system
                 x86_64, linux-gnu
## status
## major
## minor
                 1.0
                 2021
## year
## month
                05
## day
                 18
                80317
## svn rev
## language
## version.string R version 4.1.0 (2021-05-18)
## nickname
               Camp Pontanezen
```

#### 8.3 Setup

```
###### summary data ######
summary_data_loc <- pasteO(working_directory, "data/aggregate.csv")
summary_data <- read.csv(summary_data_loc, na.strings="NONE")

summary_data$DISABLE_REACTION_SENSORS <- as.factor(summary_data$DISABLE_REACTION_SENSORS
summary_data$chg_env <- summary_data$chg_env == "True"
summary_data$dominant_plastic_odd_even <- as.factor(summary_data$dominant_plastic_odd_even summary_data$sensors <- summary_data$DISABLE_REACTION_SENSORS == "0"
summary_data$is_plastic <- summary_data$dominant_plastic_odd_even == "True"
summary_data$POISON_PENALTY <- as.factor(summary_data$POISON_PENALTY)

summary_data$dominant_lineage_num_times_hitchhike_inst_exec_increases_per_generation <</pre>
```

8.3. SETUP 121

summary\_data\$frac\_hitchhiking\_linked\_trait\_change <- summary\_data\$dominant\_lineage\_num\_times\_hitchhiker summary\_data\$frac\_unexpressed\_hitchhiker\_inc <- summary\_data\$dominant\_lineage\_num\_times\_hitchhike summary\_data\$frac\_expressed\_hitchiker\_inc <- summary\_data\$dominant\_lineage\_num\_times\_hitchhike\_inc

```
env_label_fun <- function(chg_env) {</pre>
  if (chg_env) {
    return("Fluctuating")
  } else {
    return("Constant")
  }
}
sensors_label_fun <- function(has_sensors) {</pre>
  if (has_sensors) {
    return("Sensors")
  } else {
    return("No sensors")
}
condition_label_fun <- function(has_sensors, env_chg) {</pre>
  if (has_sensors && env_chg) {
    return("PLASTIC")
  } else if (env_chg) {
    return("NON-PLASTIC")
  } else {
    return("STATIC")
  }
}
summary_data$env_label <- mapply(</pre>
  env_label_fun,
  summary_data$chg_env
summary_data$sensors_label <- mapply(</pre>
  sensors_label_fun,
  summary_data$sensors
)
summary_data$condition <- mapply(</pre>
  condition_label_fun,
  summary_data$sensors,
  summary_data$chg_env
)
condition_order = c(
```

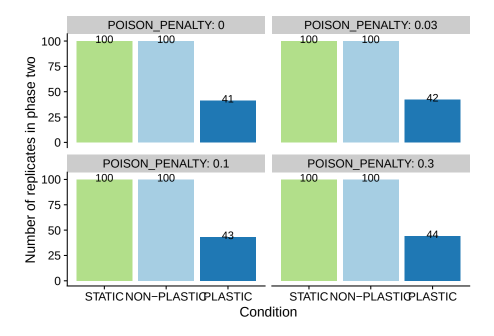
```
"STATIC",
 "NON-PLASTIC",
 "PLASTIC"
pairwise_comparisons <- list(</pre>
  c("STATIC", "NON-PLASTIC"),
  c("STATIC", "PLASTIC"),
  c("PLASTIC", "NON-PLASTIC")
)
p_label <- function(p_value) {</pre>
  threshold = 0.0001
  if (p_value < threshold) {</pre>
    return(paste0("p < ", threshold))</pre>
  } else {
    return(paste0("p = ", p_value))
  }
poison_penalties <- levels(summary_data$POISON_PENALTY)</pre>
##### time series #####
lineage time series data loc <- pasteO(working directory, "data/lineage series.csv")
lineage_time_series_data <- read.csv(lineage_time_series_data_loc)</pre>
lineage_time_series_data$DISABLE_REACTION_SENSORS <- as.factor(lineage_time_series_data)</pre>
lineage_time_series_data$chg_env <- lineage_time_series_data$chg_env == "True"</pre>
lineage_time_series_data$sensors <- lineage_time_series_data$DISABLE_REACTION_SENSORS</pre>
lineage_time_series_data$POISON_PENALTY <- as.factor(lineage_time_series_data$POISON_V.
lineage_time_series_data$env_label <- mapply(</pre>
  env_label_fun,
  lineage_time_series_data$chg_env
lineage_time_series_data$sensors_label <- mapply(</pre>
  sensors_label_fun,
  lineage_time_series_data$sensors
lineage_time_series_data$condition <- mapply(</pre>
  condition label fun,
  lineage_time_series_data$sensors,
  lineage_time_series_data$chg_env
)
```

```
####### misc ######
# Configure our default graphing theme
focal_summary_data <- filter(summary_data, POISON_PENALTY==focal_poison_penalty)
theme_set(theme_cowplot())
cb_palette <- "Paired"
dir.create(pasteO(working_directory, "plots"), showWarnings=FALSE)
samplemean <- function(x, d) {
   return(mean(x[d]))
}</pre>
```

#### 8.4 Evolution of phenotypic plasticity

For sensor-enabled populations in fluctuating environments, we only transferred populations containing an optimally plastic genotype to phase-two.

```
summary_data_grouped = dplyr::group_by(summary_data, sensors, env_label, condition, POISON_PENALT
summary_data_group_counts = dplyr::summarize(summary_data_grouped, n=dplyr::n())
ggplot(summary_data_group_counts, aes(x=condition, y=n, fill=condition)) +
  geom_col(position=position_dodge(0.9)) +
  geom_text(aes(label=n, y=n+2)) +
  scale_x_discrete(
   name="Condition",
   limits=condition_order
  scale_fill_brewer(
   palette=cb_palette
  scale_color_brewer(
   palette=cb_palette
  ylab("Number of replicates in phase two") +
  facet_wrap(~POISON_PENALTY, labeller=label_both) +
  theme(
   legend.position="none"
```

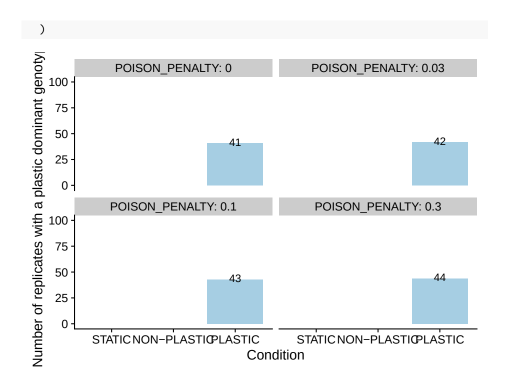


We can confirm our expectation that the dominant genotypes in non-plastic conditions are not phenotypically plastic.

```
summary_data_grouped = dplyr::group_by(summary_data, condition, is_plastic, POISON_PEN.
summary_data_group_counts = dplyr::summarize(summary_data_grouped, n=dplyr::n())
```

## `summarise()` has grouped output by 'condition', 'is\_plastic'. You can override using ggplot(filter(summary\_data\_group\_counts, is\_plastic), aes(x=condition, y=n, fill=condition, y=n, fill=conditi

```
geom_col(position=position_dodge(0.9)) +
scale_x_discrete(
   name="Condition",
   limits=condition_order
) +
geom_text(aes(label=n, y=n+1)) +
scale_fill_brewer(
   palette=cb_palette
) +
scale_color_brewer(
   palette=cb_palette
) +
ylab("Number of replicates with a plastic dominant genotype") +
ylim(0, 100) +
facet_wrap(~POISON_PENALTY, labeller=label_both) +
theme(
   legend.position="none"
```



#### 8.5 Poison instruction execution

# 8.5.1 Number of replicates where final dominant genotype executes the poison instruction

```
for (penalty in poison_penalties) {
  occurrences <- c(
    length(filter(summary_data, POISON_PENALTY==penalty & condition=="NON-PLASTIC" & dominant_timelength(filter(summary_data, POISON_PENALTY==penalty & condition=="PLASTIC" & dominant_times_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_pointies_poin
```

```
names(occurrences) <- c(</pre>
    "NON-PLASTIC",
    "PLASTIC",
    "STATIC"
  poison_exec_table <- data.frame(</pre>
    executes.poison=occurrences,
    replicates=trials
  cat(paste0("#### Penalty: ", penalty, "\n"))
  cat(print(kable(poison_exec_table)))
  cat("\n")
  ft <- pairwise.fisher.test(x=occurrences, n=trials, p.adjust.method="bonferroni")</pre>
  print(ft)
  cat("\n\n")
## #### Penalty: 0
##
## \begin{tabular}{||r|r}
## \hline
## & executes.poison & replicates\\
## \hline
## NON-PLASTIC & 86 & 100\\
## \hline
## PLASTIC & 27 & 41\\
## \hline
## STATIC & 85 & 100\\
## \hline
## \end{tabular}
##
##
## Pairwise comparisons using Pairwise comparison of proportions (Fisher)
##
## data: occurrences out of trials
           NON-PLASTIC PLASTIC
## PLASTIC 0.03
## STATIC 1.00
                       0.06
## P value adjustment method: bonferroni
##
##
## #### Penalty: 0.03
##
```

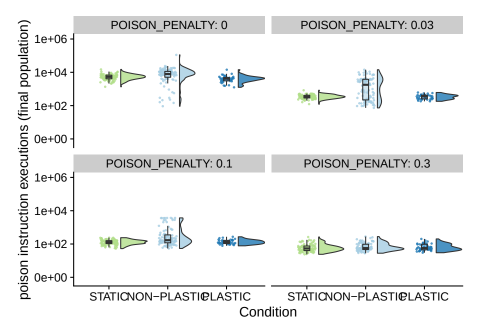
```
## \begin{tabular}{l|r|r}
## \hline
## & executes.poison & replicates\\
## \hline
## NON-PLASTIC & 46 & 100\\
## \hline
## PLASTIC & 1 & 42\\
## \hline
## STATIC & 1 & 100\\
## \hline
## \end{tabular}
##
## Pairwise comparisons using Pairwise comparison of proportions (Fisher)
## data: occurrences out of trials
##
          NON-PLASTIC PLASTIC
## PLASTIC 1.2e-07
## STATIC 2.9e-15
                   1
## P value adjustment method: bonferroni
##
## #### Penalty: 0.1
## \begin{tabular}{l|r|r}
## \hline
## & executes.poison & replicates\\
## \hline
## NON-PLASTIC & 14 & 100\\
## \hline
## PLASTIC & 0 & 43\\
## \hline
## STATIC & 0 & 100\\
## \hline
## \end{tabular}
##
##
## Pairwise comparisons using Pairwise comparison of proportions (Fisher)
##
## data: occurrences out of trials
##
         NON-PLASTIC PLASTIC
## PLASTIC 0.03212 -
## STATIC 0.00022 1.00000
```

```
##
## P value adjustment method: bonferroni
##
##
## #### Penalty: 0.3
##
## \begin{tabular}{||r|r}
## \hline
    & executes.poison & replicates\\
## \hline
## NON-PLASTIC & 0 & 100\\
## \hline
## PLASTIC & 0 & 44\\
## \hline
## STATIC & 0 & 100\\
## \hline
## \end{tabular}
##
##
##
   Pairwise comparisons using Pairwise comparison of proportions (Fisher)
##
## data: occurrences out of trials
          NON-PLASTIC PLASTIC
## PLASTIC 1
## STATIC 1
## P value adjustment method: bonferroni
```

#### 8.5.2 Poison instruction execution (final population)

```
ggplot(summary_data, aes(x=condition, y=final_population_poison, fill=condition)) +
  geom_flat_violin(
    position = position_nudge(x = .2, y = 0),
        alpha = .8
) +
    geom_point(
        mapping=aes(color=condition),
        position = position_jitter(width = .15),
        size = .5,
        alpha = 0.8
) +
    geom_boxplot(
    width = .1,
    outlier.shape = NA,
```

```
alpha = 0.5
) +
scale_x_discrete(
 name="Condition",
  limits=condition_order
) +
scale_y_continuous(
  name="poison instruction executions (final population)",
  trans=pseudo_log_trans(sigma=1,base=10),
  breaks=c(0,100,10000,1000000),
  limits=c(-1,1000000)
) +
scale_fill_brewer(
  palette=cb_palette
) +
scale_color_brewer(
  palette=cb_palette
) +
facet_wrap(
  ~POISON_PENALTY,
  labeller=label_both
) +
# coord_flip() +
theme(
  legend.position="none"
ggsave(
  paste0(working_directory, "plots/final-population-poison-log.pdf"),
  width=15,
  height=10
)
```



```
for (penalty in poison_penalties) {
  stat_data <- filter(summary_data, POISON_PENALTY==penalty)</pre>
  print(
    paste0(
      "PENALTY: ", penalty
    )
  )
  kt <- kruskal.test(</pre>
      formula=final_population_poison~condition,
      data=stat_data
    )
  print(
    kt
  )
  if (is.na(kt$p.value)) { next }
  if (kt$p.value > 0.05) { next }
  print(
    pairwise.wilcox.test(
      x=stat_data$final_population_poison,
      g=stat_data$condition,
      p.adjust.method="bonferroni"
    )
  )
}
```

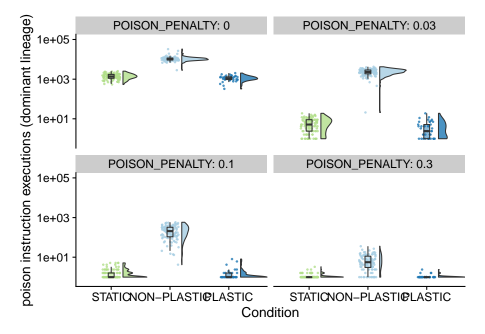
```
## [1] "PENALTY: O"
##
## Kruskal-Wallis rank sum test
## data: final_population_poison by condition
## Kruskal-Wallis chi-squared = 43.589, df = 2, p-value = 3.426e-10
##
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: stat_data$final_population_poison and stat_data$condition
          NON-PLASTIC PLASTIC
## PLASTIC 8.7e-07
## STATIC 9.8e-07
                      0.00074
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.03"
##
## Kruskal-Wallis rank sum test
## data: final_population_poison by condition
## Kruskal-Wallis chi-squared = 20.74, df = 2, p-value = 3.136e-05
##
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: stat_data$final_population_poison and stat_data$condition
##
          NON-PLASTIC PLASTIC
##
## PLASTIC 0.003
## STATIC 1e-04
                      1.000
##
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.1"
## Kruskal-Wallis rank sum test
## data: final_population_poison by condition
## Kruskal-Wallis chi-squared = 20.608, df = 2, p-value = 3.35e-05
##
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: stat_data$final_population_poison and stat_data$condition
##
```

```
## NON-PLASTIC PLASTIC
## PLASTIC 0.0093    -
## STATIC 4.9e-05    1.0000
##
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.3"
##
## Kruskal-Wallis rank sum test
##
## data: final_population_poison by condition
## Kruskal-Wallis chi-squared = 3.3994, df = 2, p-value = 0.1827
```

# 8.5.3 Cummulative poison instruction execution along final dominant lineages

```
ggplot(summary_data, aes(x=condition, y=dominant_lineage_times_poison_executed, fill=c
 geom_flat_violin(
   position = position_nudge(x = .2, y = 0),
   alpha = .8
 ) +
 geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
   alpha = 0.8
 ) +
 geom_boxplot(
   width = .1,
   outlier.shape = NA,
   alpha = 0.5
 scale_x_discrete(
   name="Condition",
   limits=condition_order
 ) +
 scale_y_continuous(
   name="poison instruction executions (dominant lineage)",
   trans=pseudo_log_trans(sigma = 1, base = 10),
   breaks=c(10,1000,100000),
   limits=c(-1,100000)
 ) +
 facet_wrap(
   ~POISON PENALTY,
   labeller=label_both
```

```
scale_fill_brewer(
   palette=cb_palette
) +
scale_color_brewer(
   palette=cb_palette
) +
theme(
   legend.position="none"
) +
ggsave(
   paste0(working_directory, "plots/final-dominant-lineage-poison-log.pdf"),
   width=15,
   height=10
)
```



```
for (penalty in poison_penalties) {
  stat_data <- filter(summary_data, POISON_PENALTY==penalty)
  print(
    paste0(
        "PENALTY: ", penalty
    )
  )
  kt <- kruskal.test(
        formula=dominant_lineage_times_poison_executed~condition,
        data=stat_data</pre>
```

```
)
 print(
   kt
  )
 if (is.na(kt$p.value)) { next }
  if (kt$p.value > 0.05) { next }
 print(
   pairwise.wilcox.test(
     x=stat_data$dominant_lineage_times_poison_executed,
      g=stat_data$condition,
      p.adjust.method="bonferroni"
    )
  )
## [1] "PENALTY: O"
##
##
   Kruskal-Wallis rank sum test
##
## data: dominant_lineage_times_poison_executed by condition
## Kruskal-Wallis chi-squared = 178.84, df = 2, p-value < 2.2e-16
##
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: stat_data$dominant_lineage_times_poison_executed and stat_data$condition
##
          NON-PLASTIC PLASTIC
##
## PLASTIC <2e-16
## STATIC <2e-16
                       0.0018
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.03"
##
##
   Kruskal-Wallis rank sum test
##
## data: dominant_lineage_times_poison_executed by condition
## Kruskal-Wallis chi-squared = 178.62, df = 2, p-value < 2.2e-16
##
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: stat_data$dominant_lineage_times_poison_executed and stat_data$condition
##
```

##

NON-PLASTIC PLASTIC

```
## PLASTIC <2e-16
## STATIC <2e-16
                      0.011
##
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.1"
## Kruskal-Wallis rank sum test
##
## data: dominant_lineage_times_poison_executed by condition
## Kruskal-Wallis chi-squared = 184.83, df = 2, p-value < 2.2e-16
##
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: stat_data$dominant_lineage_times_poison_executed and stat_data$condition
##
##
          NON-PLASTIC PLASTIC
## PLASTIC <2e-16
## STATIC <2e-16
                      0.21
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.3"
## Kruskal-Wallis rank sum test
## data: dominant_lineage_times_poison_executed by condition
## Kruskal-Wallis chi-squared = 149.48, df = 2, p-value < 2.2e-16
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: stat_data$dominant_lineage_times_poison_executed and stat_data$condition
##
          NON-PLASTIC PLASTIC
## PLASTIC 4.4e-16
## STATIC < 2e-16
                      0.84
## P value adjustment method: bonferroni
```

# 8.6 Characterizing mutations that increase poison instruction execution

```
ggplot(summary_data, aes(x=condition, y=dominant_lineage_num_times_hitchhike_inst_exec
 geom_flat_violin(
   position = position_nudge(x = .2, y = 0),
    alpha = .8
  geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
   alpha = 0.8
  geom_boxplot(
   width = .1,
   outlier.shape = NA,
   alpha = 0.5
 ) +
 scale_x_discrete(
   name="Condition",
   limits=condition_order
 ) +
 facet_wrap(
    ~POISON_PENALTY,
   labeller=label_both
 ) +
 scale_fill_brewer(
   palette=cb_palette
 ) +
 scale_color_brewer(
   palette=cb_palette
 ) +
 theme(
   legend.position="none"
```

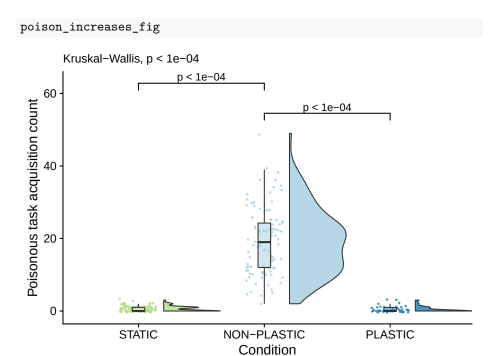
```
ninant_lineage_num_times_hitchhike_inst_exec_incr
              POISON_PENALTY: 0
                                               POISON_PENALTY: 0.03
   200
   150
    100
    50
     0 -
             POISON_PENALTY: 0.1
                                               POISON_PENALTY: 0.3
   200
   150
   100
    50
     0 -
          STATICNON-PLASTICPLASTIC
                                            STATICNON-PLASTICPLASTIC
                                    Condition
for (penalty in poison_penalties) {
  stat_data <- filter(summary_data, POISON_PENALTY==penalty)</pre>
  print(
    paste0(
       "PENALTY: ", penalty
  )
  kt <- kruskal.test(</pre>
      formula=dominant_lineage_num_times_hitchhike_inst_exec_increases~condition,
      data=stat_data
    )
  print(
    kt
  )
  if (is.na(kt$p.value)) { next }
  if (kt$p.value > 0.05) { next }
  print(
    pairwise.wilcox.test(
      x=stat_data$dominant_lineage_num_times_hitchhike_inst_exec_increases,
       g=stat_data$condition,
      p.adjust.method="bonferroni"
    )
  )
```

```
## [1] "PENALTY: O"
##
##
   Kruskal-Wallis rank sum test
##
## data: dominant_lineage_num_times_hitchhike_inst_exec_increases by condition
## Kruskal-Wallis chi-squared = 179.79, df = 2, p-value < 2.2e-16
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: stat_data$dominant_lineage_num_times_hitchhike_inst_exec_increases and stat_
##
          NON-PLASTIC PLASTIC
## PLASTIC < 2e-16
## STATIC < 2e-16
                      0.00046
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.03"
## Kruskal-Wallis rank sum test
##
## data: dominant_lineage_num_times_hitchhike_inst_exec_increases by condition
## Kruskal-Wallis chi-squared = 179.35, df = 2, p-value < 2.2e-16
##
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: stat_data$dominant_lineage_num_times_hitchhike_inst_exec_increases and stat_
##
##
          NON-PLASTIC PLASTIC
## PLASTIC <2e-16
## STATIC <2e-16
                      0.03
##
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.1"
##
## Kruskal-Wallis rank sum test
##
## data: dominant_lineage_num_times_hitchhike_inst_exec_increases by condition
## Kruskal-Wallis chi-squared = 185.34, df = 2, p-value < 2.2e-16
##
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: stat_data$dominant_lineage_num_times_hitchhike_inst_exec_increases and stat_
##
```

```
##
                          NON-PLASTIC PLASTIC
## PLASTIC <2e-16
## STATIC <2e-16
                                                       0.27
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.3"
##
## Kruskal-Wallis rank sum test
## data: dominant_lineage_num_times_hitchhike_inst_exec_increases by condition
## Kruskal-Wallis chi-squared = 146.35, df = 2, p-value < 2.2e-16
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: stat_data$dominant_lineage_num_times_hitchhike_inst_exec_increases and stat_data$condit
##
                          NON-PLASTIC PLASTIC
## PLASTIC 7.8e-16
## STATIC < 2e-16
                                                       0.86
## P value adjustment method: bonferroni
# sum(filter(summary_data, condition=="NON-PLASTIC" & POISON_PENALTY==0.1)$dominant_lineage_num_
\# \ sum(filter(summary\_data, \ condition == "PLASTIC" \ \& \ POISON\_PENALTY == 0.1) \\ \$ dominant\_lineage\_num\_times = 0.1 
# sum(filter(summary_data, condition=="STATIC" & POISON_PENALTY==0.1)$dominant_lineage_num_times_
Focal figure for the manuscript:
# Compute manual labels for geom_signif
stat.test <- focal_summary_data %>%
    wilcox_test(dominant_lineage_num_times_hitchhike_inst_exec_increases ~ condition) %>%
    adjust_pvalue(method = "bonferroni") %>%
    add_significance() %>%
    add_xy_position(x="condition")
# Tweak y.position manually to account for scaled axis (edge case that triggers bad behavior in g
stat.test$manual_position <- stat.test$y.position</pre>
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
poison_increases_fig <- ggplot(</pre>
         focal_summary_data,
         aes(x=condition, y=dominant_lineage_num_times_hitchhike_inst_exec_increases, fill=condition)
    geom_flat_violin(
         scale="width",
         position = position_nudge(x = .2, y = 0),
         alpha = .8
```

```
) +
  geom_point(
    mapping=aes(color=condition),
    position = position_jitter(width = .15),
    size = .5,
    alpha = 0.8
  ) +
  geom_boxplot(
    width = .1,
    outlier.shape = NA,
    alpha = 0.5
 ) +
 scale_x_discrete(
    name="Condition",
   limits=condition_order,
   labels=condition_order
 ) +
 scale_y_continuous(
   name="Poisonous task acquisition count",
 ) +
  scale_fill_brewer(
   palette=cb_palette
 ) +
  scale_color_brewer(
    palette=cb_palette
  # coord_flip()
 labs(
    subtitle=paste0(
      "Kruskal-Wallis, ",
     p_label(signif(kruskal.test(formula=dominant_lineage_num_times_hitchhike_inst_ex
    )
  ) +
  ggsignif::geom_signif(
    data=filter(stat.test, p.adj <= alpha),</pre>
    aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
   manual=TRUE,
    inherit.aes=FALSE
 ) +
 theme(
    legend.position="none"
```

 $\hbox{\tt \#\# Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y\_position}$ 



# 8.6.2 Frequency of increases in poison instruction execution (lineage)

```
ggplot(summary_data, aes(x=condition, y=dominant_lineage_num_times_hitchhike_inst_exec_increases_
  geom_flat_violin(
   position = position_nudge(x = .2, y = 0),
   alpha = .8
  geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
   alpha = 0.8
  geom_boxplot(
   width = .1,
   outlier.shape = NA,
   alpha = 0.5
  scale_x_discrete(
   name="Condition",
   limits=condition_order
```

```
) +
 scale_fill_brewer(
   palette=cb_palette
 ) +
 scale_color_brewer(
   palette=cb_palette
 ) +
 facet_wrap(
   ~POISON_PENALTY,
   labeller=label_both,
   scales="free_y"
 ) +
 # coord_flip() +
 theme(
   legend.position="none"
 ) +
 ggsave(
   paste0(working_directory, "plots/final-dominant-lineage-poison-increase-per-general
   width=15,
   height=10
 )
neage_num_times_hitchhike_inst_exec_increases_p
                                                 POISON_PENALTY: 0.03
              POISON_PENALTY: 0
                                       0.005
  0.005
                                       0.004
   0.004
  0.003
                                       0.003
   0.002
                                       0.002
   0.001
                                       0.001
   0.000 -
                                       0.000 -
             POISON_PENALTY: 0.1
                                                 POISON_PENALTY: 0.3
                                       0.005
   0.005
   0.004
                                       0.004
   0.003
                                       0.003
  0.002
                                       0.002
  0.001
                                       0.001
   0.000
                                       0.000
           STATIGION-PLASTIC
                                               STATICION-PLASTICLASTIC
                                     Condition
```

for (penalty in poison\_penalties) {

print(

stat\_data <- filter(summary\_data, POISON\_PENALTY==penalty)</pre>

```
paste0(
      "PENALTY: ", penalty
  )
  kt <- kruskal.test(</pre>
      formula=dominant_lineage_num_times_hitchhike_inst_exec_increases_per_generation~condition,
      data=stat_data
   )
 print(
   kt
  if (is.na(kt$p.value)) { next }
  if (kt$p.value > 0.05) { next }
  print(
   pairwise.wilcox.test(
     x=stat_data$dominant_lineage_num_times_hitchhike_inst_exec_increases_per_generation,
      g=stat_data$condition,
      p.adjust.method="bonferroni"
   )
 )
}
## [1] "PENALTY: O"
##
## Kruskal-Wallis rank sum test
## data: dominant_lineage_num_times_hitchhike_inst_exec_increases_per_generation by condition
## Kruskal-Wallis chi-squared = 180.05, df = 2, p-value < 2.2e-16
##
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: stat_data$dominant_lineage_num_times_hitchhike_inst_exec_increases_per_generation and s
##
          NON-PLASTIC PLASTIC
## PLASTIC < 2e-16
## STATIC < 2e-16
                      7.8e-05
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.03"
##
## Kruskal-Wallis rank sum test
## data: dominant_lineage_num_times_hitchhike_inst_exec_increases_per_generation by condition
## Kruskal-Wallis chi-squared = 176.25, df = 2, p-value < 2.2e-16
```

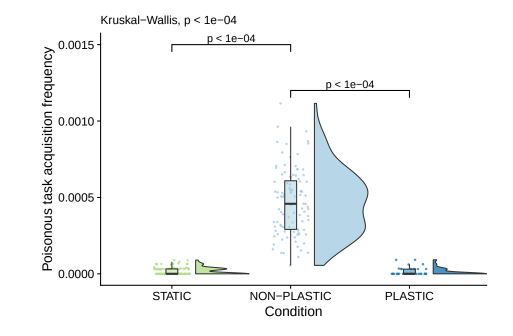
```
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
##
## data: stat_data$dominant_lineage_num_times_hitchhike_inst_exec_increases_per_gener
##
          NON-PLASTIC PLASTIC
##
## PLASTIC <2e-16
## STATIC <2e-16
                       0.019
##
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.1"
##
##
   Kruskal-Wallis rank sum test
##
## data: dominant_lineage_num_times_hitchhike_inst_exec_increases_per_generation by c
## Kruskal-Wallis chi-squared = 184.17, df = 2, p-value < 2.2e-16
##
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: stat_data$dominant_lineage_num_times_hitchhike_inst_exec_increases_per_gener
##
          NON-PLASTIC PLASTIC
##
## PLASTIC <2e-16
## STATIC <2e-16
                       0.2
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.3"
##
##
   Kruskal-Wallis rank sum test
##
## data: dominant_lineage_num_times_hitchhike_inst_exec_increases_per_generation by c
## Kruskal-Wallis chi-squared = 140.99, df = 2, p-value < 2.2e-16
##
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: stat_data$dominant_lineage_num_times_hitchhike_inst_exec_increases_per_gener
##
           NON-PLASTIC PLASTIC
## PLASTIC 2.2e-15
## STATIC < 2e-16
                       0.79
## P value adjustment method: bonferroni
```

Figure for the manuscript:

```
# Compute manual labels for geom_signif
stat.test <- focal_summary_data %>%
  wilcox_test(dominant_lineage_num_times_hitchhike_inst_exec_increases_per_generation ~ condition
  adjust_pvalue(method = "bonferroni") %>%
 add_significance() %>%
 add_xy_position(x="condition", step.increase=0.2)
# Tweak y.position manually to account for scaled axis (edge case that triggers bad behavior in
stat.test$manual_position <- stat.test$y.position</pre>
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
poison_increases_per_gen_fig <- ggplot(</pre>
    focal_summary_data,
    aes(x=condition, y=dominant_lineage_num_times_hitchhike_inst_exec_increases_per_generation, 1
  ) +
  geom_flat_violin(
    scale="width",
    position = position_nudge(x = .2, y = 0),
    alpha = .8
  ) +
  geom_point(
    mapping=aes(color=condition),
    position = position_jitter(width = .15),
    size = .5,
    alpha = 0.8
  ) +
  geom_boxplot(
    width = .1,
    outlier.shape = NA,
    alpha = 0.5
  ) +
  scale_x_discrete(
    name="Condition",
    limits=condition_order,
    labels=condition_order
  ) +
  scale_y_continuous(
    name="Poisonous task acquisition frequency",
  ) +
  scale_fill_brewer(
    palette=cb_palette
  scale_color_brewer(
    palette=cb_palette
```

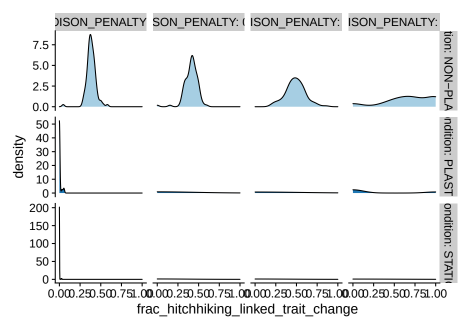
```
# coord_flip()
labs(
    subtitle=paste0(
        "Kruskal-Wallis, ",
        p_label(signif(kruskal.test(formula=dominant_lineage_num_times_hitchhike_inst_ex.)
)
) +
ggsignif::geom_signif(
    data=filter(stat.test, p.adj <= alpha),
    aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
    manual=TRUE,
    inherit.aes=FALSE
) +
theme(
    legend.position="none"
)</pre>
```

## Warning: Ignoring unknown aesthetics: xmin, xmax, annotations, y\_position
poison\_increases\_per\_gen\_fig



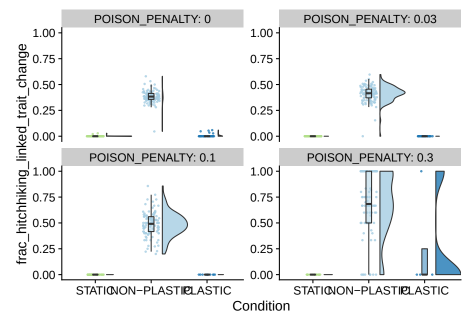
8.6.3 What fraction of mutations that increase poison instruction execution co-occur with base trait changes?

```
ggplot(filter(summary_data, dominant_lineage_num_times_hitchhike_inst_exec_increases>0), aes(x=filter)
 geom_density() +
 facet_grid(
   condition~POISON_PENALTY,
   labeller=label_both,
   scales="free_y"
  scale_fill_brewer(
   palette=cb_palette
  scale_color_brewer(
   palette=cb_palette
  ) +
 theme(
   legend.position="none"
  ggsave(
   paste0(working_directory, "plots/dominant-lineage-frac_hitchhiking_linked_trait_change.png");
   width=15,
   height=10
```



```
ggplot(filter(summary_data, dominant_lineage_num_times_hitchhike_inst_exec_increases>0
  geom_flat_violin(
    position = position_nudge(x = .2, y = 0),
    alpha = .8
  geom_point(
    mapping=aes(color=condition),
   position = position_jitter(width = .15),
    size = .5,
    alpha = 0.8
  ) +
  geom_boxplot(
   width = .1,
    outlier.shape = NA,
    alpha = 0.5
  ) +
  scale_x_discrete(
    name="Condition",
   limits=condition_order
  scale_fill_brewer(
   palette=cb_palette
 ) +
  scale_color_brewer(
    palette=cb_palette
```

```
facet_wrap(
    ~POISON_PENALTY,
    labeller=label_both,
    scales="free_y"
) +
# coord_flip() +
theme(
    legend.position="none"
)
```



print(
 kt

```
for (penalty in poison_penalties) {
   stat_data <- filter(summary_data, POISON_PENALTY==penalty & dominant_lineage_num_times_hitchhil
   print(
     paste0(
        "PENALTY: ", penalty
     )
   )
   kt <- kruskal.test(
        formula=frac_hitchhiking_linked_trait_change~condition,
        data=stat_data
   )</pre>
```

```
if (is.na(kt$p.value)) { next }
  if (kt$p.value > 0.05) { next }
  print(
    pairwise.wilcox.test(
      x=stat_data$frac_hitchhiking_linked_trait_change,
      g=stat_data$condition,
     p.adjust.method="bonferroni",
      exact=FALSE
    )
  )
}
## [1] "PENALTY: O"
##
##
   Kruskal-Wallis rank sum test
##
## data: frac_hitchhiking_linked_trait_change by condition
## Kruskal-Wallis chi-squared = 211.29, df = 2, p-value < 2.2e-16
##
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: stat_data$frac_hitchhiking_linked_trait_change and stat_data$condition
##
           NON-PLASTIC PLASTIC
##
## PLASTIC <2e-16
## STATIC <2e-16
                       0.031
##
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.03"
##
##
   Kruskal-Wallis rank sum test
##
## data: frac_hitchhiking_linked_trait_change by condition
## Kruskal-Wallis chi-squared = 186.88, df = 2, p-value < 2.2e-16
##
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: stat_data$frac_hitchhiking_linked_trait_change and stat_data$condition
##
           NON-PLASTIC PLASTIC
## PLASTIC 2.9e-16
## STATIC < 2e-16
```

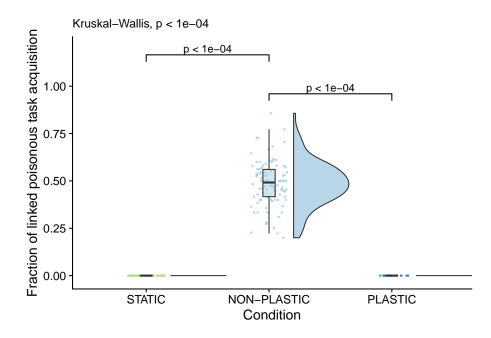
##

```
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.1"
##
## Kruskal-Wallis rank sum test
##
## data: frac_hitchhiking_linked_trait_change by condition
## Kruskal-Wallis chi-squared = 113.72, df = 2, p-value < 2.2e-16
##
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: stat_data$frac_hitchhiking_linked_trait_change and stat_data$condition
##
          NON-PLASTIC PLASTIC
##
## PLASTIC 3.3e-08
## STATIC < 2e-16
## P value adjustment method: bonferroni
## [1] "PENALTY: 0.3"
##
## Kruskal-Wallis rank sum test
##
## data: frac_hitchhiking_linked_trait_change by condition
## Kruskal-Wallis chi-squared = 34.791, df = 2, p-value = 2.788e-08
##
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: stat_data$frac_hitchhiking_linked_trait_change and stat_data$condition
          NON-PLASTIC PLASTIC
##
## PLASTIC 0.26
## STATIC 2.4e-08
                       0.18
## P value adjustment method: bonferroni
denom <- sum(filter(summary_data, condition=="NON-PLASTIC" & POISON_PENALTY==0.1)$dominant_lineag
num <- sum(filter(summary_data, condition=="NON-PLASTIC" & POISON_PENALTY==0.1)$dominant_lineage_
paste0("NON-PLASTIC (0.1 penalty): ", num/denom, "(", num, "/", denom, ")")
## [1] "NON-PLASTIC (0.1 penalty): 0.498956158663883(956/1916)"
denom <- sum(filter(summary_data, condition=="PLASTIC" & POISON_PENALTY==0.1)$dominant_lineage_nu
num <- sum(filter(summary_data, condition=="PLASTIC" & POISON_PENALTY==0.1)$dominant_lineage_num_
paste0("PLASTIC (0.1 penalty): ", num/denom, " (", num, "/", denom, ")")
## [1] "PLASTIC (0.1 penalty): 0 (0/18)"
```

```
denom <- sum(filter(summary_data, condition=="STATIC" & POISON_PENALTY==0.1)$dominant_
num <- sum(filter(summary_data, condition=="STATIC" & POISON_PENALTY==0.1)$dominant_li</pre>
paste0("STATIC (0.1 penalty): ", num/denom, " (", num, "/", denom, ")")
## [1] "STATIC (0.1 penalty): 0 (0/58)"
Focal figure for the manuscript:
# Compute manual labels for geom_signif
stat.test <-filter(focal_summary_data,dominant_lineage_num_times_hitchhike_inst_exec_i:
  wilcox_test(frac_hitchhiking_linked_trait_change ~ condition, comparisons=list(c("PL
 adjust_pvalue(method = "bonferroni") %>%
 add_significance() %>%
 add xy position(x="condition")
# Tweak y.position manually to account for scaled axis (edge case that triggers bad be
stat.test$manual_position <- stat.test$y.position</pre>
stat.test$label <- mapply(p_label,stat.test$p.adj)</pre>
linked_trait_change_fig <- ggplot(</pre>
    filter(focal_summary_data, dominant_lineage_num_times_hitchhike_inst_exec_increase
    aes(x=condition, y=frac_hitchhiking_linked_trait_change, fill=condition)
  ) +
  geom_flat_violin(
    scale="width",
    position = position_nudge(x = .2, y = 0),
    alpha = .8
 ) +
 geom_point(
    mapping=aes(color=condition),
    position = position_jitter(width = .15),
    size = .5,
    alpha = 0.8
  ) +
  geom_boxplot(
    width = .1,
    outlier.shape = NA,
    alpha = 0.5
 ) +
  scale_x_discrete(
    name="Condition",
    limits=condition_order,
   labels=condition_order
  scale_y_continuous(
   name="Fraction of linked poisonous task acquisition",
    limits=c(-0.01, 1.2),
```

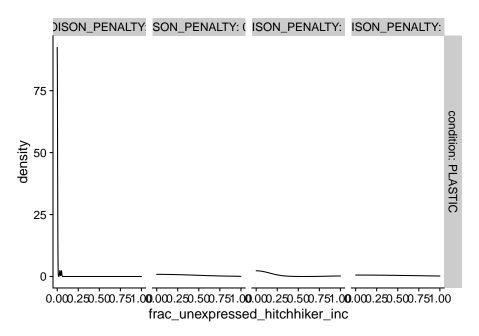
### 8.6. CHARACTERIZING MUTATIONS THAT INCREASE POISON INSTRUCTION EXECUTION153

```
breaks=c(0, 0.25, 0.50, 0.75, 1.0)
) +
scale_fill_brewer(
        palette=cb_palette
scale_color_brewer(
        palette=cb_palette
) +
labs(
          subtitle=paste0(
                    "Kruskal-Wallis, ",
                    p_label(signif(kruskal.test(formula=frac_hitchhiking_linked_trait_change~condition, data=formula=frac_hitchhiking_linked_trait_change~condition, data=formula=frac_hitchhiking_linked_trait_change~condition, data=formula=frac_hitchhiking_linked_trait_change~condition, data=formula=frac_hitchhiking_linked_trait_change~condition, data=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=formula=f
          )
) +
ggsignif::geom_signif(
          data=filter(stat.test, p.adj <= alpha),</pre>
          aes(xmin=group1,xmax=group2,annotations=label,y_position=manual_position),
          manual=TRUE,
          inherit.aes=FALSE
) +
theme(
          legend.position="none"
```



# 8.7 What fraction of poison execution increases

```
ggplot(filter(summary_data, dominant_lineage_num_times_hitchhike_inst_exec_increases>0
   geom_density() +
   facet_grid(
      condition~POISON_PENALTY,
      labeller=label_both,
      scales="free_y"
) +
   theme(
   legend.position="none"
)
```



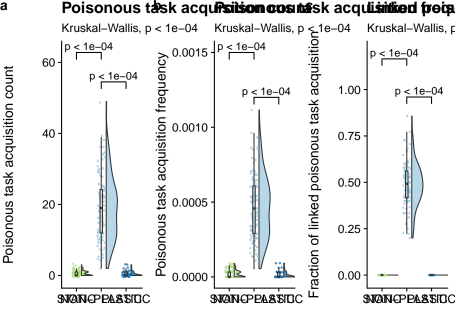
```
denom <- sum(filter(summary_data, condition=="PLASTIC" & POISON_PENALTY==0.1)$dominant_lineage_nu
num <- sum(filter(summary_data, condition=="PLASTIC" & POISON_PENALTY==0.1)$dominant_lineage_num
paste0("PLASTIC: ", num/denom, " (", num, "/", denom, ")")
## [1] "PLASTIC: 0.05555555555555556 (1/18)"</pre>
```

# 8.8 Manuscript figures

```
grid <- plot_grid(
  poison_increases_fig +
    theme(
       axis.title.x=element_blank()
    ) +
    ggtitle("Poisonous task acquisition count"),
  poison_increases_per_gen_fig +
    theme(
       axis.title.x=element_blank()
    ) +
    ggtitle("Poisonous task acquisition frequency"),
  linked_trait_change_fig +
    theme(
       axis.title.x=element_blank()
    ) +
    ggtitle("Linked poisonous task acquisition"),</pre>
```

```
nrow=1,
  align="v",
 labels="auto"
save_plot(
  pasteO(working_directory, "plots/", "poison-accumulation-panel.pdf"),
  base_height=6,
   base_asp=3/1
grid
```

#### Poisonous takk acquiraitisom cosutakk acquiraitised freig а



# Chapter 9

# Regulation in Avida

### 9.1 Overview

```
total_updates <- 200000
replicates <- 100

all_traits <- c("not", "nand", "ornot", "or", "andnot")
traits_set_a <- c("not", "and", "or")
traits_set_b <- c("nand", "ornot", "andnot")

# Relative location of data.
working_directory <- "experiments/2021-02-08-evo-dynamics/analysis/" # << For bookdown
# working_directory <- "./"
# << For local analysis</pre>
```

# 9.2 Analysis dependencies

Load all required R libraries.

```
library(ggplot2)
library(tidyverse)
library(cowplot)
library(RColorBrewer)
library(Hmisc)
library(boot)
source("https://gist.githubusercontent.com/benmarwick/2a1bb0133ff568cbe28d/raw/fb53bd97121f7f9ce9
```

These analyses were conducted/knitted with the following computing environment:

```
print(version)
##
## platform
                  x86_64-pc-linux-gnu
## arch
                  x86 64
## os
                  linux-gnu
## system
                  x86_64, linux-gnu
## status
## major
                  1.0
## minor
## year
                  2021
                  05
## month
## day
                  18
                  80317
## svn rev
## language
                  R
## version.string R version 4.1.0 (2021-05-18)
## nickname
                  Camp Pontanezen
```

## 9.3 Setup

```
trace_summary_data_loc <- paste0(working_directory, "data/trace_summary.csv")</pre>
trace_summary_data <- read.csv(trace_summary_data_loc, na.strings="NONE")</pre>
trace_summary_data$DISABLE_REACTION_SENSORS <- as.factor(trace_summary_data$DISABLE_RE.
trace_summary_data$chg_env <- trace_summary_data$chg_env == "True"</pre>
trace_summary_data$sensors <- trace_summary_data$DISABLE_REACTION_SENSORS == "0"</pre>
env_label_fun <- function(chg_env) {</pre>
  if (chg_env) {
    return("Fluctuating")
  } else {
    return("Constant")
  }
}
sensors_label_fun <- function(has_sensors) {</pre>
  if (has_sensors) {
    return("Sensors")
  } else {
    return("No sensors")
  }
}
```

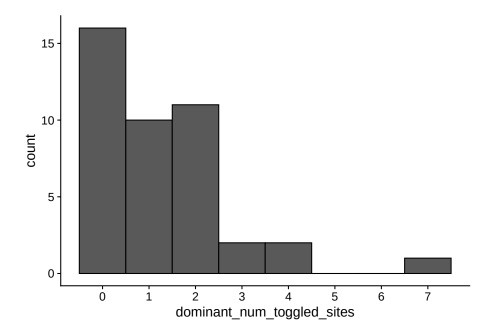
```
# note that this labeler makes assumptions about how we set up our experiment
condition_label_fun <- function(has_sensors, env_chg) {</pre>
  if (has_sensors && env_chg) {
    return("PLASTIC")
 } else if (env_chg) {
    return("NON-PLASTIC")
  } else {
    return("STATIC")
 }
}
trace_summary_data$env_label <- mapply(</pre>
  env_label_fun,
  trace_summary_data$chg_env
trace_summary_data$sensors_label <- mapply(</pre>
  sensors_label_fun,
  trace_summary_data$sensors
trace_summary_data$condition <- mapply(</pre>
  condition_label_fun,
  trace_summary_data$sensors,
  trace_summary_data$chg_env
###### misc ######
# Configure our default graphing theme
theme_set(theme_cowplot())
dir.create(paste0(working_directory, "plots"), showWarnings=FALSE)
```

# 9.4 How many instructions do plastic genomes toggle depending on environmental context?

```
ggplot(trace_summary_data, aes(x=dominant_num_toggled_sites)) +
    geom_histogram(
        binwidth=1,
        color="black"
) +
    scale_fill_brewer(
        palette="Paired"
) +
    scale_color_brewer(
        palette="Paired"
```

```
) +
scale_x_continuous(
   breaks=seq(0, max(trace_summary_data$dominant_num_toggled_sites)+1)
) +
theme(
  legend.position="none"
) +
ggsave(paste0(working_directory, "plots/", "toggled-sites.png"))
```

## Saving 6.5 x 4.5 in image



# 9.5 What is the distribution of toggled sequence sizes?

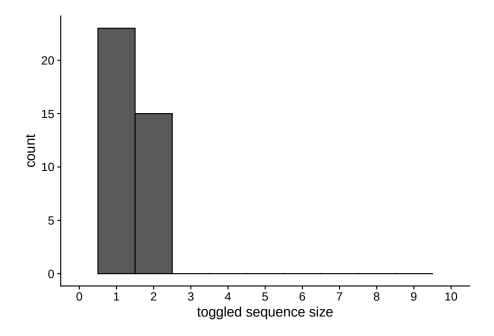
```
chunk_sizes <- data.frame(
    size=integer()
)
for (sizes in trace_summary_data$dominant_toggled_chunk_sizes) {
    if (sizes == "") { next }
        sizes <- unlist(lapply(str_split(sizes, ';'), as.integer))
        chunk_sizes <- rbind(chunk_sizes, data.frame(size=c(sizes)))
}</pre>
```

### 9.5. WHAT IS THE DISTRUBUTION OF TOGGLED SEQUENCE SIZES?161

```
ggplot(chunk_sizes, aes(x=size)) +
   geom_histogram(
     binwidth=1,
     color="black"
   scale_fill_brewer(
     palette="Paired"
   ) +
   scale_color_brewer(
     palette="Paired"
   scale_x_continuous(
     name="toggled sequence size",
     breaks=seq(0, 10),
     limits=c(0, 10)
   ) +
   theme(
     legend.position="none"
   ggsave(paste0(working_directory, "plots/", "toggled-chunk-sizes.png"))
```

```
## Saving 6.5 \times 4.5 in image
```

```
## Warning: Removed 2 rows containing missing values (geom_bar).
## Warning: Removed 2 rows containing missing values (geom_bar).
```



# Chapter 10

# Evolutionary change (variable length genomes)

### 10.1 Overview

```
total_updates <- 200000
replicates <- 100

all_traits <- c("not", "nand", "ornot", "or", "andnot")
traits_set_a <- c("not", "and", "or")
traits_set_b <- c("nand", "ornot", "andnot")

# Relative location of data.
working_directory <- "experiments/2021-01-30-evo-dynamics/analysis/" # << For bookdown
# working_directory <- "./"
# << For local analysis</pre>
```

# 10.2 Analysis dependencies

Load all required R libraries.

```
library(ggplot2)
library(tidyverse)
library(cowplot)
library(RColorBrewer)
library(Hmisc)
library(boot)
source("https://gist.githubusercontent.com/benmarwick/2a1bb0133ff568cbe28d/raw/fb53bd97121f7f9ce8
```

These analyses were conducted/knitted with the following computing environ-

ment:

```
print(version)
##
                  x86_64-pc-linux-gnu
## platform
## arch
                  x86_64
## os
                  linux-gnu
                  x86_64, linux-gnu
## system
## status
## major
                  4
## minor
                  1.0
                  2021
## year
## month
                  05
                  18
## day
## svn rev
                  80317
## language
                  R
## version.string R version 4.1.0 (2021-05-18)
                  Camp Pontanezen
## nickname
```

# 10.3 Setup

```
summary_data_loc <- paste0(working_directory, "data/aggregate.csv")</pre>
summary_data <- read.csv(summary_data_loc, na.strings="NONE")</pre>
summary_data$DISABLE_REACTION_SENSORS <- as.factor(summary_data$DISABLE_REACTION_SENSO
summary_data$chg_env <- summary_data$chg_env == "True"</pre>
summary_data$dominant_plastic_odd_even <- as.factor(summary_data$dominant_plastic_odd_.
summary_data$sensors <- summary_data$DISABLE_REACTION_SENSORS == "0"</pre>
summary_data$is_plastic <- summary_data$dominant_plastic_odd_even == "True"
env_label_fun <- function(chg_env) {</pre>
  if (chg_env) {
   return("Fluctuating")
 } else {
    return("Constant")
  }
sensors_label_fun <- function(has_sensors) {</pre>
  if (has sensors) {
   return("Sensors")
 } else {
    return("No sensors")
```

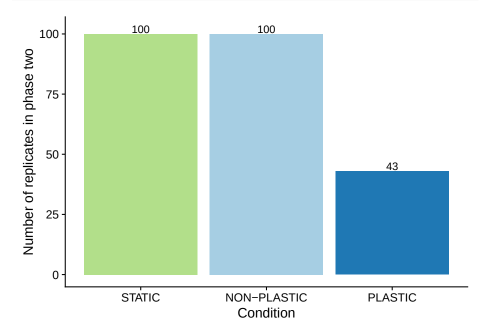
```
# note that this labeler makes assumptions about how we set up our experiment
condition_label_fun <- function(has_sensors, env_chg) {</pre>
  if (has_sensors && env_chg) {
    return("PLASTIC")
  } else if (env_chg) {
    return("NON-PLASTIC")
  } else {
    return("STATIC")
 }
}
summary_data$env_label <- mapply(</pre>
  env_label_fun,
  summary_data$chg_env
)
summary_data$sensors_label <- mapply(</pre>
  sensors_label_fun,
  summary_data$sensors
)
summary_data$condition <- mapply(</pre>
  condition_label_fun,
  summary_data$sensors,
  summary_data$chg_env
condition_order = c(
  "STATIC",
  "NON-PLASTIC",
  "PLASTIC"
)
###### misc ######
# Configure our default graphing theme
theme_set(theme_cowplot())
dir.create(paste0(working_directory, "plots"), showWarnings=FALSE)
```

# 10.4 Evolution of phenotypic plasticity

For sensor-enabled populations in fluctuating environments, we only transfered populations containing an optimally plastic genotype to phase two.

```
summary_data_grouped = dplyr::group_by(summary_data, sensors, env_label, condition)
summary_data_group_counts = dplyr::summarize(summary_data_grouped, n=dplyr::n())
```

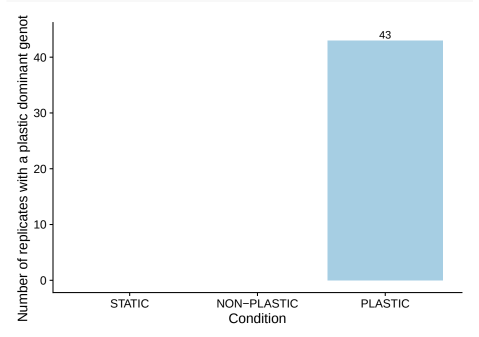
```
## `summarise()` has grouped output by 'sensors', 'env_label'. You can override using 'ggplot(summary_data_group_counts, aes(x=condition, y=n, fill=condition)) +
    geom_col(position=position_dodge(0.9)) +
    geom_text(aes(label=n, y=n+2)) +
    scale_x_discrete(
        name="Condition",
        limits=condition_order
    ) +
    scale_fill_brewer(
    palette="Paired"
    ) +
    scale_color_brewer(
    palette="Paired"
    ) +
    ylab("Number of replicates in phase two") +
    theme(
    legend.position="none"
    )
}
```



We can confirm our expectation that the dominant genotypes in non-plastic conditions are not phenotypically plastic.

```
summary_data_grouped = dplyr::group_by(summary_data, condition, is_plastic)
summary_data_group_counts = dplyr::summarize(summary_data_grouped, n=dplyr::n())
```

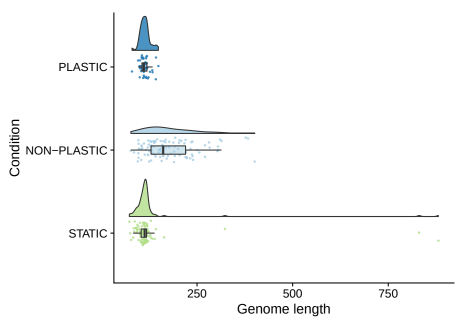
```
ggplot(filter(summary_data_group_counts, is_plastic), aes(x=condition, y=n, fill=condition)) +
  geom_col(position=position_dodge(0.9)) +
  scale_x_discrete(
   name="Condition",
   limits=condition_order
  ) +
  scale_fill_brewer(
   palette="Paired"
  scale_color_brewer(
   palette="Paired"
  ) +
  geom_text(aes(label=n, y=n+1)) +
  ylab("Number of replicates with a plastic dominant genotype") +
  theme(
   legend.position="none"
 )
```



# 10.5 Genome length

Single-instruction insertions and deletions were possible for this experiment, so genome size also evolved.

```
ggplot(summary_data, aes(x=condition, y=dominant_genome_length, fill=condition)) +
 geom_flat_violin(
    position = position_nudge(x = .2, y = 0),
    alpha = .8
 ) +
 geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
   alpha = 0.8
  geom_boxplot(
   width = .1,
   outlier.shape = NA,
   alpha = 0.5
 ) +
 scale_x_discrete(
   name="Condition",
   limits=condition_order
  scale_fill_brewer(
   palette="Paired"
 ) +
  scale_color_brewer(
   palette="Paired"
  coord_flip() +
 ylab("Genome length") +
 theme(
    legend.position="none"
```



```
kruskal.test(
 formula=dominant_genome_length~condition,
  data=summary_data
)
##
   Kruskal-Wallis rank sum test
##
##
## data: dominant_genome_length by condition
## Kruskal-Wallis chi-squared = 82.798, df = 2, p-value < 2.2e-16
pairwise.wilcox.test(
 x=summary_data$dominant_genome_length,
  g=summary_data$condition,
 p.adjust.method="bonferroni",
)
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: summary_data$dominant_genome_length and summary_data$condition
##
           NON-PLASTIC PLASTIC
## PLASTIC 1.8e-10
## STATIC < 2e-16
##
```

```
## P value adjustment method: bonferroni
median(filter(summary_data, condition=="PLASTIC")$phylo_mrca_changes)

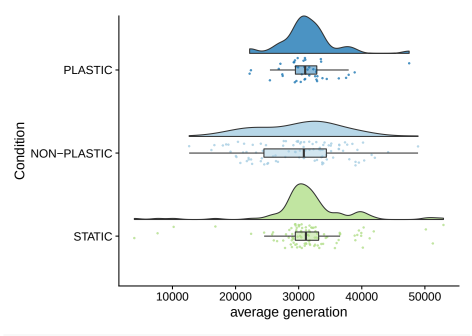
## [1] 45
median(filter(summary_data, condition=="STATIC")$phylo_mrca_changes)

## [1] 47
median(filter(summary_data, condition=="NON-PLASTIC")$phylo_mrca_changes)

## [1] 393
```

## 10.6 Average generation

```
ggplot(summary_data, aes(x=condition, y=time_average_generation, fill=condition)) +
  geom_flat_violin(
    position = position_nudge(x = .2, y = 0),
   alpha = .8
  ) +
 geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
   alpha = 0.8
  ) +
  geom_boxplot(
   width = .1,
   outlier.shape = NA,
   alpha = 0.5
  scale_x_discrete(
   name="Condition",
   limits=condition_order
  scale_fill_brewer(
   palette="Paired"
 scale_color_brewer(
   palette="Paired"
  coord flip() +
 ylab("average generation") +
 theme(
    legend.position="none"
```

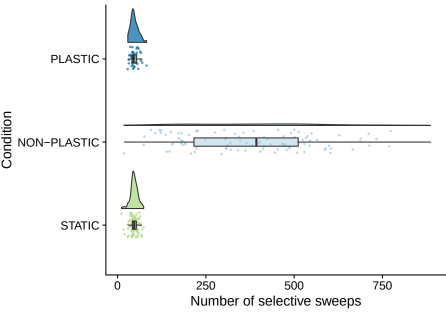


```
median(filter(summary_data, condition=="PLASTIC")$time_average_generation)
## [1] 31028.6
median(filter(summary_data, condition=="STATIC")$time_average_generation)
## [1] 31147.5
median(filter(summary_data, condition=="NON-PLASTIC")$time_average_generation)
## [1] 30817.95
kruskal.test(
  formula=time_average_generation~condition,
  data=summary_data
)
##
##
   Kruskal-Wallis rank sum test
##
## data: time_average_generation by condition
## Kruskal-Wallis chi-squared = 1.3804, df = 2, p-value = 0.5015
```

## 10.7 Coalescence event count

The number of times the most recent common ancestor changes gives us the number of selective sweeps that occur during the experiment.

```
ggplot(summary_data, aes(x=condition, y=phylo_mrca_changes, fill=condition)) +
 geom_flat_violin(
    position = position_nudge(x = .2, y = 0),
    alpha = .8
 ) +
 geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
   alpha = 0.8
 geom_boxplot(
   width = .1,
   outlier.shape = NA,
   alpha = 0.5
  scale_fill_brewer(
   palette="Paired"
  scale_color_brewer(
    palette="Paired"
  ) +
  coord_flip() +
 scale_x_discrete(
   name="Condition",
   limits=condition_order
 ylab("Number of selective sweeps") +
 theme(
    legend.position="none"
```



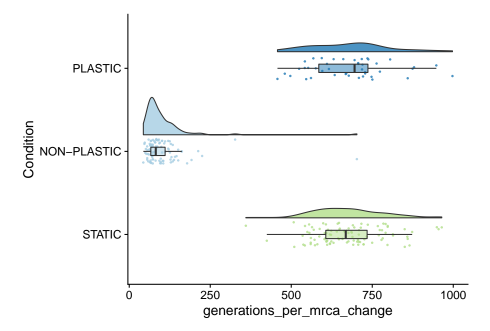
```
paste0(
 "PLASTIC: ",
  median(filter(summary_data, condition=="PLASTIC")$phylo_mrca_changes)
## [1] "PLASTIC: 45"
paste0(
  "STATIC: ",
  median(filter(summary_data, condition=="STATIC")$phylo_mrca_changes)
)
## [1] "STATIC: 47"
paste0(
  "NON-PLASTIC: ",
  median(filter(summary_data, condition=="NON-PLASTIC")$phylo_mrca_changes)
## [1] "NON-PLASTIC: 393"
kruskal.test(
  formula=phylo_mrca_changes~condition,
  data=summary_data
)
##
## Kruskal-Wallis rank sum test
```

```
##
## data: phylo_mrca_changes by condition
## Kruskal-Wallis chi-squared = 168.89, df = 2, p-value < 2.2e-16
pairwise.wilcox.test(
  x=summary_data$phylo_mrca_changes,
  g=summary_data$condition,
  p.adjust.method="bonferroni",
)
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: summary data$phylo mrca changes and summary data$condition
##
##
           NON-PLASTIC PLASTIC
## PLASTIC <2e-16
## STATIC <2e-16
## P value adjustment method: bonferroni
```

# 10.7.1 Average number of generations between coalescence events

```
summary data$generations per mrca change <- summary data$time average generation / sum
ggplot(summary_data, aes(x=condition, y=generations_per_mrca_change, fill=condition))
 geom_flat_violin(
   position = position_nudge(x = .2, y = 0),
   alpha = .8
  ) +
  geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
    alpha = 0.8
  ) +
  geom_boxplot(
   width = .1,
    outlier.shape = NA,
   alpha = 0.5
 ) +
  scale_x_discrete(
   name="Condition",
    limits=condition_order
```

```
scale_fill_brewer(
   palette="Paired"
) +
scale_color_brewer(
   palette="Paired"
) +
coord_flip() +
theme(
   legend.position="none"
)
```



```
paste0(
   "PLASTIC: ",
   median(filter(summary_data, condition=="PLASTIC")$generations_per_mrca_change)
)

## [1] "PLASTIC: 695.504761904762"

paste0(
   "STATIC: ",
   median(filter(summary_data, condition=="STATIC")$generations_per_mrca_change)
)
```

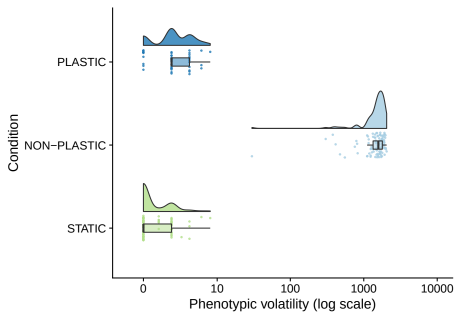
## [1] "STATIC: 668.25523255814"

```
paste0(
  "NON-PLASTIC: ",
  median(filter(summary_data, condition=="NON-PLASTIC")$generations_per_mrca_change)
## [1] "NON-PLASTIC: 81.9208459944751"
kruskal.test(
  formula=generations_per_mrca_change~condition,
  data=summary_data
##
##
   Kruskal-Wallis rank sum test
## data: generations_per_mrca_change by condition
## Kruskal-Wallis chi-squared = 171.73, df = 2, p-value < 2.2e-16
pairwise.wilcox.test(
  x=summary_data$generations_per_mrca_change,
  g=summary_data$condition,
  p.adjust.method="bonferroni",
)
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: summary_data$generations_per_mrca_change and summary_data$condition
##
           NON-PLASTIC PLASTIC
## PLASTIC <2e-16
## STATIC <2e-16
## P value adjustment method: bonferroni
```

# 10.8 Phenotypic volatility along the dominant lineage

```
ggplot(summary_data, aes(x=condition, y=dominant_lineage_trait_volatility, fill=condit
  geom_flat_violin(
    position = position_nudge(x = .2, y = 0),
    alpha = .8
) +
  geom_point(
    mapping=aes(color=condition),
```

```
position = position_jitter(width = .15),
  size = .5,
  alpha = 0.8
) +
geom_boxplot(
 width = .1,
  outlier.shape = NA,
 alpha = 0.5
scale_x_discrete(
 name="Condition",
  limits=condition_order
) +
scale_y_continuous(
 name="Phenotypic volatility (log scale)",
 trans="pseudo_log",
 breaks=c(0, 10, 100, 1000, 10000),
 limits=c(-1,10000)
) +
scale_fill_brewer(
  palette="Paired"
scale_color_brewer(
 palette="Paired"
) +
coord_flip() +
theme(
  legend.position="none"
```



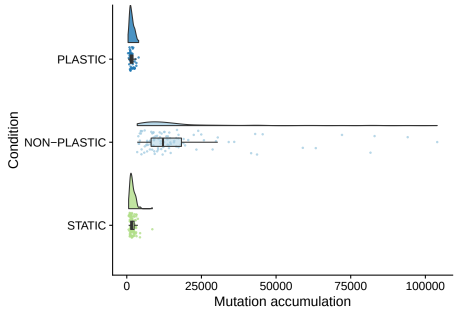
```
paste0(
  "PLASTIC: ",
  median(filter(summary_data, condition=="PLASTIC")$dominant_lineage_trait_volatility)
## [1] "PLASTIC: 2"
paste0(
  "STATIC: ",
  median(filter(summary_data, condition=="STATIC")$dominant_lineage_trait_volatility)
## [1] "STATIC: O"
paste0(
  "NON-PLASTIC: ",
  median(filter(summary_data, condition=="NON-PLASTIC")$dominant_lineage_trait_volatil
## [1] "NON-PLASTIC: 1580"
kruskal.test(
  formula=dominant_lineage_trait_volatility~condition,
  data=summary_data
)
##
## Kruskal-Wallis rank sum test
```

```
##
## data: dominant_lineage_trait_volatility by condition
## Kruskal-Wallis chi-squared = 191.98, df = 2, p-value < 2.2e-16
pairwise.wilcox.test(
  x=summary_data$dominant_lineage_trait_volatility,
  g=summary_data$condition,
  p.adjust.method="bonferroni",
)
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: summary_data$dominant_lineage_trait_volatility and summary_data$condition
##
           NON-PLASTIC PLASTIC
##
## PLASTIC < 2e-16
## STATIC < 2e-16
                       5.2e-08
## P value adjustment method: bonferroni
```

## 10.9 Mutation count (along dominant lineage)

```
ggplot(summary_data, aes(x=condition, y=dominant_lineage_total_mut_cnt, fill=condition)) +
  geom_flat_violin(
   position = position_nudge(x = .2, y = 0),
   alpha = .8
  ) +
  geom_point(
   mapping=aes(color=condition),
   position = position_jitter(width = .15),
   size = .5,
   alpha = 0.8
  ) +
  geom_boxplot(
   width = .1,
   outlier.shape = NA,
   alpha = 0.5
  ) +
  ylab("Mutation accumulation") +
  scale_x_discrete(
   name="Condition",
   limits=condition_order
  scale_fill_brewer(
```

```
palette="Paired"
) +
scale_color_brewer(
  palette="Paired"
) +
coord_flip() +
theme(
  legend.position="none"
)
```



```
paste0(
   "PLASTIC: ",
   median(filter(summary_data, condition=="PLASTIC")$dominant_lineage_total_mut_cnt)
)
## [1] "PLASTIC: 1552"

paste0(
   "STATIC: ",
   median(filter(summary_data, condition=="STATIC")$dominant_lineage_total_mut_cnt)
)
## [1] "STATIC: 1724.5"

paste0(
   "NON-PLASTIC: ",
   median(filter(summary_data, condition=="NON-PLASTIC")$dominant_lineage_total_mut_cnt
```

```
## [1] "NON-PLASTIC: 12123"
kruskal.test(
  formula=dominant_lineage_total_mut_cnt~condition,
 data=summary_data
)
##
##
   Kruskal-Wallis rank sum test
##
## data: dominant_lineage_total_mut_cnt by condition
## Kruskal-Wallis chi-squared = 174.38, df = 2, p-value < 2.2e-16
pairwise.wilcox.test(
 x=summary_data$dominant_lineage_total_mut_cnt,
  g=summary_data$condition,
 p.adjust.method="bonferroni",
)
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: summary_data$dominant_lineage_total_mut_cnt and summary_data$condition
##
          NON-PLASTIC PLASTIC
## PLASTIC <2e-16
## STATIC <2e-16
                      0.57
## P value adjustment method: bonferroni
```

 $182 CHAPTER\ 10.\ EVOLUTIONARY\ CHANGE\ (VARIABLE\ LENGTH\ GENOMES)$ 

# Bibliography

Lalejini, A. M. and Ferguson, A. J. (2021). Data for evolutionary consequences of phenotypic plasticity.