# Supplemental Material: Selection Scheme Parameter Sweep Base Diagnostics

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# Chapter 1

# Introduction

This is the supplemental material for selection scheme parameter sweep experiments with basic diagnostics.

### 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.

This supplemental material includes the following selection schemes:

- Truncation (Section 2)
- Tournament (Section 3)
- Genotypic fitness sharing (Section 4)
- Phenotypic fitness sharing (Section 5)
- Nondominated sorting (Section 6)
- Novelty search (Section 7)

### 1.2 Contributing authors

- Jose Guadalupe Hernandez
- Alexander Lalejini
- Charles Ofria

### 1.3 Computer Setup

These analyses were conducted in 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
                  3.1
                  2023
## year
## month
                  06
## day
                  16
## svn rev
                  84548
## language
                  R
## version.string R version 4.3.1 (2023-06-16)
## nickname
                Beagle Scouts
```

### 1.4 Experimental setup

Setting up required variables variables.

```
# libraries we are using
library(ggplot2)
library(cowplot)
library(dplyr)
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
library(PupillometryR)
## Loading required package: rlang
# data diractory for gh-pages
DATA_DIR = '/opt/ECJ-2023-Suite-Of-Diagnostic-Metrics-For-Characterizing-Selection-School
# data diractory for local testing
\# DATA_DIR = '~\Desktop/Repositories/ECJ-2023-Suite-Of-Diagnostic-Metrics-For-Characte
```

```
# graph variables
SHAPE = c(5,3,1,2,6,0,4,20,8)
cb_palette <- c('#332288','#88CCEE','#EE7733','#EE3377','#117733','#882255','#44AA99','#CCBB44',
TSIZE = 26
p_theme <- theme(</pre>
 text = element_text(size = 28),
  plot.title = element text( face = "bold", size = 22, hjust=0.5),
 panel.border = element_blank(),
 panel.grid.minor = element_blank(),
 legend.title=element_text(size=22),
 legend.text=element_text(size=23),
  axis.title = element_text(size=23),
  axis.text = element_text(size=22),
  legend.position="bottom",
  panel.background = element_rect(fill = "#f1f2f5",
                                   colour = "white",
                                   size = 0.5, linetype = "solid")
## Warning: The `size` argument of `element_rect()` is deprecated as of ggplot2 3.4.0.
## i Please use the `linewidth` argument instead.
## This warning is displayed once every 8 hours.
## Call `lifecycle::last_lifecycle_warnings()` to see where this warning was
## generated.
# default variables
REPLICATES = 50
DIMENSIONALITY = 100
GENERATIONS = 50000
# selection scheme params exploring
TR_LIST = c('1', '2', '4', '8', '16', '32', '64', '128', '256')
TS_LIST = c('2', '4', '8', '16', '32', '64', '128', '256', '512')
FS_LIST = c('0', '0.1', '0.3', '0.6', '1.2', '2.5', '5')
ND_LIST = c('0', '0.1', '0.3', '0.6', '1.2', '2.5', '5')
NS_LIST = c('1', '2', '4', '8', '15', '30')
```

# Chapter 2

# Truncation selection

Results for the truncation selection parameter sweep on the diagnostics with no valleys.

### 2.1 Data setup

```
over_time_df <- read.csv(paste(DATA_DIR,'OVER-TIME/tru.csv', sep = "", collapse = NULL), header =
over_time_df$T <- factor(over_time_df$T, levels = TR_LIST)

best_df <- read.csv(paste(DATA_DIR,'BEST/tru.csv', sep = "", collapse = NULL), header = TRUE, str
best_df$T <- factor(best_df$T, levels = TR_LIST)

sati_df <- read.csv(paste(DATA_DIR,'SOL-FND/tru.csv', sep = "", collapse = NULL), header = TRUE,
sati_df$T <- factor(sati_df$T, levels = TR_LIST)</pre>
```

### 2.2 Exploitation rate results

Here we present the results for **best performances** found by each selection scheme parameter on the exploitation rate diagnostic. 50 replicates are conducted for each scheme explored.

#### 2.2.1 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'exp') %>%
group_by(T, gen) %>%
```

```
dplyr::summarise(
  min = min(pop_fit_max) / DIMENSIONALITY,
  mean = mean(pop_fit_max) / DIMENSIONALITY,
  max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'T'. You can override using the `.groups`
## argument.
over_time_plot = ggplot(lines, aes(x=gen, y=mean, group = T, fill = T, color = T, shape
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale_y_continuous(
   name="Average trait score",
   limits=c(0, 100),
   breaks=seq(0,100, 20),
   labels=c("0", "20", "40", "60", "80", "100")
 ) +
  scale x continuous(
   name="Generations",
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
 ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
 guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
    fill=guide_legend(nrow=2, title.position = "bottom")
over_time_plot
```



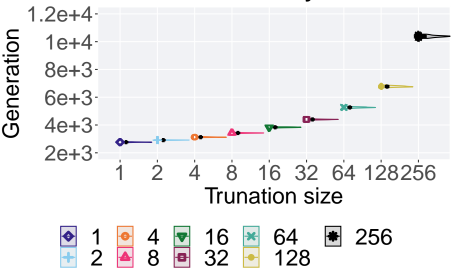
### 2.2.2 Generation satisfactory solution found

First generation a satisfactory solution is found throughout the  $50,\!000$  generations.

```
plot = filter(sati_df, acro == 'exp') %>%
  ggplot(., aes(x = T, y = gen , color = T, fill = T, shape = T)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Generation",
   limits=c(2000, 12000),
   breaks=c(2000, 4000, 6000, 8000, 10000, 12000),
   labels=c("2e+3", "4e+3", "6e+3", "8e+3", "1e+4", "1.2e+4")
  ) +
  scale_x_discrete(
   name="Trunation size"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Generation satisfactory solution found')+
  p_theme + theme(legend.title=element_blank())
```

```
plot_grid(
  plot +
    theme(legend.position="none"),
  legend,
  nrow=2,
  rel_heights = c(3,1)
```

## **Generation satisfactory solution foun**





#### 2.2.2.1 Stats

```
ssf = filter(sati_df, gen <= GENERATIONS & acro == 'exp')</pre>
ssf$acro = factor(ssf$acro, levels = TR_LIST)
ssf %>%
  group_by(T) %>%
 dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(gen)),
   min = min(gen, na.rm = TRUE),
   median = median(gen, na.rm = TRUE),
   mean = mean(gen, na.rm = TRUE),
   max = max(gen, na.rm = TRUE),
    IQR = IQR(gen, na.rm = TRUE)
```

```
## # A tibble: 9 x 8
    Τ
          count na_cnt
                         min median
                                      mean
                                             max
                                                   IQR
     <fct> <int> <int> <int> <dbl>
                                     <dbl> <int> <dbl>
## 1 1
             50
                     0 2734 2765
                                     2766.
                                            2795
                                                  17.8
## 2 2
                                            2952
                                                  18.5
             50
                     0 2889 2914.
                                     2914.
## 3 4
             50
                     0
                        3093 3124.
                                     3127.
                                            3167
## 4 8
             50
                        3385 3426.
                                     3425.
                                            3473
                                                  21.2
                     0
## 5 16
             50
                     0
                        3786 3836
                                     3835.
                                            3869
                                                  34
## 6 32
                     0 4361 4402.
             50
                                     4400.
                                            4450
                                                  26.5
## 7 64
             50
                     0 5201 5264
                                     5266.
                                            5337
                                                  44.5
## 8 128
             50
                     0 6667 6766. 6772.
                                            6905
                                                  64.2
## 9 256
             50
                     0 10236 10387 10382. 10538
                                                  86.8
```

Kruskal-Wallis test illustrates evidence of statistical differences.

```
kruskal.test(gen ~ T, data = ssf)
```

```
##
## Kruskal-Wallis rank sum test
##
## data: gen by T
## Kruskal-Wallis chi-squared = 443.46, df = 8, p-value < 2.2e-16</pre>
```

Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.

```
##
  Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: ssf$gen and ssf$T
##
##
             2
                   4
                          8
                                       32
                                                    128
      1
                                 16
                                              64
## 2
      <2e-16 -
## 4
      <2e-16 <2e-16 -
      <2e-16 <2e-16 <2e-16 -
## 16 <2e-16 <2e-16 <2e-16 -
## 32
      <2e-16 <2e-16 <2e-16 <2e-16 -
## 64 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16 <
## 128 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16 <
## 256 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16
##
## P value adjustment method: bonferroni
```

### 2.3 Ordered exploitation results

Here we present the results for **best performances** found by each selection scheme parameter on the exploitation rate diagnostic. 50 replicates are conducted for each scheme explored.

#### 2.3.1 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'ord') %>%
group_by(T, gen) %>%
dplyr::summarise(
   min = min(pop_fit_max) / DIMENSIONALITY,
   mean = mean(pop_fit_max) / DIMENSIONALITY,
   max = max(pop_fit_max) / DIMENSIONALITY
)
```

## `summarise()` has grouped output by 'T'. You can override using the `.groups`
## argument.

```
ggplot(lines, aes(x=gen, y=mean, group = T, fill = T, color = T, shape = T)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale_y_continuous(
    name="Average trait score",
   limits=c(0, 100),
    breaks=seq(0,100, 20),
    labels=c("0", "20", "40", "60", "80", "100")
  scale_x_continuous(
    name="Generations",
    limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
    labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Performance over time')+
 p theme + theme(legend.title=element blank()) +
  guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
```

```
color=guide_legend(nrow=2, title.position = "bottom"),
fill=guide_legend(nrow=2, title.position = "bottom")
)
```



#### 2.3.2 Generation satisfactory solution found

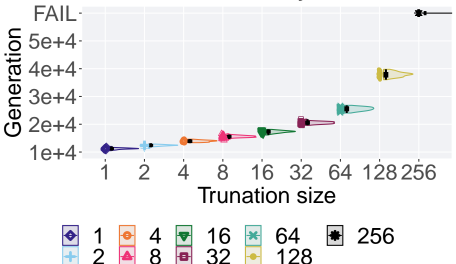
First generation a satisfactory solution is found throughout the 50,000 generations.

```
plot = filter(sati_df, acro == 'ord') %>%
    ggplot(., aes(x = T, y = gen , color = T, fill = T, shape = T)) +
    geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
    geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
    geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
    scale_y_continuous(
        name="Generation",
        limits=c(10000, 60000),
        breaks=c(10000, 20000, 30000, 40000,50000,60000),
        labels=c("1e+4","2e+4","3e+4","4e+4","5e+4","FAIL")
) +
    scale_x_discrete(
        name="Trunation size"
)+
    scale_shape_manual(values=SHAPE)+
    scale_colour_manual(values = cb_palette) +
```

```
scale_fill_manual(values = cb_palette) +
   ggtitle('Generation satisfactory solution found')+
   p_theme + theme(legend.title=element_blank())

plot_grid(
   plot +
        theme(legend.position="none"),
   legend,
   nrow=2,
   rel_heights = c(3,1)
)
```

### Generation satisfactory solution found



#### 2.3.2.1 Stats

```
ssf = filter(sati_df, gen <= GENERATIONS & acro == 'ord')
ssf$acro = factor(ssf$acro, levels = TR_LIST)
ssf %>%
  group_by(T) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(gen)),
    min = min(gen, na.rm = TRUE),
    median = median(gen, na.rm = TRUE),
```

```
mean = mean(gen, na.rm = TRUE),
   max = max(gen, na.rm = TRUE),
   IQR = IQR(gen, na.rm = TRUE)
)
## # A tibble: 8 x 8
    Т
          count na_cnt    min median
                                     mean
                                             max
                                                    IQR
     <fct> <int> <int> <dbl> <dbl> <int> <dbl>
                     0 10494 11246. 11226. 12014 316
## 1 1
             50
## 2 2
             50
                     0 11332 12438 12389. 12862
## 3 4
             50
                     0 13379 13941 13950. 14630 529.
## 4 8
             50
                     0 14261 15563 15567, 16591 476.
                     0 16147 17385 17307. 18144
## 5 16
             50
## 6 32
              50
                     0 19612 20528. 20543. 21845
## 7 64
              50
                      0 24048 25548. 25513. 26807 1075
## 8 128
              50
                      0 36034 37956 37965. 39783 1251.
Kruskal–Wallis test illustrates evidence of statistical differences.
kruskal.test(gen ~ T, data = ssf)
##
## Kruskal-Wallis rank sum test
##
## data: gen by T
## Kruskal-Wallis chi-squared = 392.52, df = 7, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = ssf$gen, g = ssf$T, p.adjust.method = "bonferroni",
                     paired = FALSE, conf.int = FALSE, alternative = 'g')
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: ssf$gen and ssf$T
##
##
              2
                       4
                              8
                                       16
                                               32
                                                       64
## 2
       3.1e-16 -
       < 2e-16 < 2e-16 -
       < 2e-16 < 2e-16 < 2e-16 -
## 16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 -
## 32 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 -
## 64 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 <
## 128 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16
## P value adjustment method: bonferroni
```

### 2.4 Contradictory objectives results

Here we present the results for activation gene coverage and satisfactory trait coverage found by each selection scheme parameter on the contradictory objectives diagnostic. 50 replicates are conducted for each scheme parameters explored.

#### 2.4.1 Activation gene coverage over time

Activation gene coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations. Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'con') %>%
group_by(T, gen) %>%
dplyr::summarise(
  min = min(uni_str_pos),
  mean = mean(uni_str_pos),
  max = max(uni_str_pos)
)
```

## `summarise()` has grouped output by 'T'. You can override using the `.groups`
## argument.

```
ggplot(lines, aes(x=gen, y=mean, group = T, fill = T, color = T, shape = T)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
 geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale y continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks=seq(0,100, 20),
   labels=c("0", "20", "40", "60", "80", "100")
  ) +
  scale_x_continuous(
    name="Generations",
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene coverage over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
```

```
shape=guide_legend(nrow=2, title.position = "bottom"),
color=guide_legend(nrow=2, title.position = "bottom"),
fill=guide_legend(nrow=2, title.position = "bottom")
)
```



#### 2.4.2 Final activation gene coverage

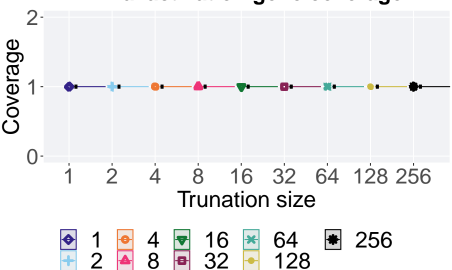
Activation gene coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'con') %>%
    ggplot(., aes(x = T, y = uni_str_pos, color = T, fill = T, shape = T)) +
    geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
    geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
    geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
    scale_y_continuous(
    name="Coverage",
    limits=c(0, 2),
    breaks=c(0,1,2)
) +
    scale_x_discrete(
    name="Trunation size"
)+
    scale_shape_manual(values=SHAPE)+
    scale_colour_manual(values = cb_palette, ) +
    scale_fill_manual(values = cb_palette) +
```

```
ggtitle('Final activation gene coverage')+
p_theme + theme(legend.title=element_blank())

plot_grid(
  plot +
    theme(legend.position="none"),
  legend,
  nrow=2,
  rel_heights = c(3,1)
)
```

### Final activation gene coverage



#### 2.4.2.1 Stats

```
act_coverage = filter(over_time_df, gen == 50000 & acro == 'con')
act_coverage$acro = factor(act_coverage$acro, levels = TR_LIST)
act_coverage %>%
  group_by(T) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(uni_str_pos)),
    min = min(uni_str_pos, na.rm = TRUE),
    median = median(uni_str_pos, na.rm = TRUE),
    mean = mean(uni_str_pos, na.rm = TRUE),
```

```
max = max(uni_str_pos, na.rm = TRUE),
    IQR = IQR(uni_str_pos, na.rm = TRUE)
 )
## # A tibble: 9 x 8
          count na_cnt
                          min median mean
                                                   IQR
   <fct> <int> <int> <dbl> <dbl> <int> <dbl>
## 1 1
            50
                      0
                            1
                                   1
                                         1
                                               1
## 2 2
             50
                      0
                            1
                                   1
                                         1
                                               1
## 3 4
             50
                      0
                            1
                                   1
                                         1
                                               1
                                                     0
              50
## 4 8
                      0
                                                     0
                            1
                                   1
                                         1
                                               1
              50
                      0
## 5 16
                            1
                                   1
                                         1
                                               1
## 6 32
              50
                     0
                            1
                                   1
                                         1
                                               1
## 7 64
              50
                      0
                            1
                                   1
                                         1
                                                     0
                                               1
## 8 128
              50
                      0
                            1
                                   1
                                         1
                                               1
                                                     0
## 9 256
              50
                            1
                                   1
                                         1
```

#### 2.4.3 Satisfactory trait coverage over time

name="Generations",
limits=c(0, 50000),

Satisfactory trait coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations. Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'con') %>%
group_by(T, gen) %>%
dplyr::summarise(
  min = min(pop_uni_obj),
  mean = mean(pop_uni_obj),
  max = max(pop_uni_obj)
)
```

breaks=c(0, 10000, 20000, 30000, 40000, 50000),

## `summarise()` has grouped output by 'T'. You can override using the `.groups`
## argument.

ggplot(lines, aes(x=gen, y=mean, group = T, fill = T, color = T, shape = T)) +
 geom\_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
 geom\_line(size = 0.5) +
 geom\_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2.0, alpha =
 scale\_y\_continuous(
 name="Coverage",
 limits=c(0, 2),
 breaks=c(0,1,2)
 ) +
 scale\_x\_continuous(

```
labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
) +
scale_shape_manual(values=SHAPE)+
scale_colour_manual(values = cb_palette) +
scale_fill_manual(values = cb_palette) +
ggtitle('Satisfactory trait coverage over time')+
p_theme + theme(legend.title=element_blank()) +
guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
    fill=guide_legend(nrow=2, title.position = "bottom"))
```

### Satisfactory trait coverage over time



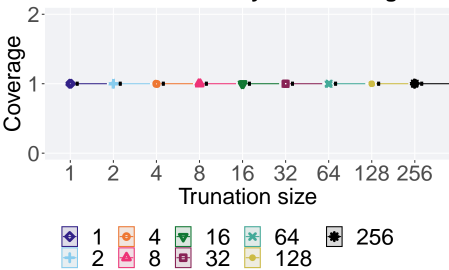
#### 2.4.4 Final satisfactory trait coverage

Satisfactory trait coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'con') %>%
    ggplot(., aes(x = T, y = pop_uni_obj, color = T, fill = T, shape = T)) +
    geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1
    geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha
    scale_y_continuous(
    name="Coverage",
```

```
limits=c(0, 2),
   breaks=c(0,1,2)
  ) +
  scale_x_discrete(
   name="Trunation size"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final satisfactory trait coverage')+
 p_theme + theme(legend.title=element_blank())
plot_grid(
 plot +
    theme(legend.position="none"),
 legend,
 nrow=2,
  rel_heights = c(3,1)
```





#### 2.4.4.1 Stats

```
sat_coverage = filter(over_time_df, gen == 50000 & acro == 'con')
sat_coverage$acro = factor(sat_coverage$acro, levels = TR_LIST)
sat_coverage %>%
  group_by(T) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(pop_uni_obj)),
    min = min(pop_uni_obj, na.rm = TRUE),
    median = median(pop_uni_obj, na.rm = TRUE),
    mean = mean(pop_uni_obj, na.rm = TRUE),
    max = max(pop_uni_obj, na.rm = TRUE),
    IQR = IQR(pop_uni_obj, na.rm = TRUE)
)
```

```
## # A tibble: 9 x 8
##
     Τ
                                                          IQR
            count na_cnt
                             min median mean
                                                   max
     <fct> <int>
                    <int> <int>
                                   <dbl> <dbl> <int>
                                                        <dbl>
## 1 1
                         0
                                                            0
                50
                                1
                                        1
                                               1
                                                     1
## 2 2
                50
                         0
                                1
                                                     1
                                                            0
                                        1
                                               1
## 3 4
                50
                         0
                                1
                                        1
                                               1
                                                     1
                                                            0
## 4 8
               50
                         0
                                                            0
                                1
                                               1
                                                     1
                                        1
## 5 16
               50
                         0
                                1
                                        1
                                               1
                                                     1
                                                            0
## 6 32
               50
                         0
                                1
                                        1
                                              1
                                                     1
                                                            0
                                                            0
## 7 64
               50
                         0
                                1
                                               1
                                                     1
## 8 128
                         0
                                                            0
               50
                                1
                                               1
                                                     1
                                        1
## 9 256
               50
                         0
                                1
                                               1
                                                            0
```

### 2.5 Multi-path exploration results

Here we present the results for **best performances** and **activation gene coverage** found by each selection scheme parameter on the multi-path exploration diagnostic. 50 replicates are conducted for each scheme parameter explored.

#### 2.5.1 Activation gene coverage over time

Activation gene coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations. Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'mpe') %>%
group_by(T, gen) %>%
dplyr::summarise(
  min = min(uni_str_pos),
  mean = mean(uni_str_pos),
  max = max(uni_str_pos)
```

```
)
```

```
## `summarise()` has grouped output by 'T'. You can override using the `.groups`
## argument.
ggplot(lines, aes(x=gen, y=mean, group = T, fill = T, color = T, shape = T)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2.0, alpha =
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks=seq(0,100, 20),
   labels=c("0", "20", "40", "60", "80", "100")
  ) +
  scale_x_continuous(
   name="Generations".
   limits=c(0, 50000),
   breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene coverage over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
   shape=guide_legend(nrow=2, title.position = "bottom"),
   color=guide_legend(nrow=2, title.position = "bottom"),
   fill=guide_legend(nrow=2, title.position = "bottom")
```



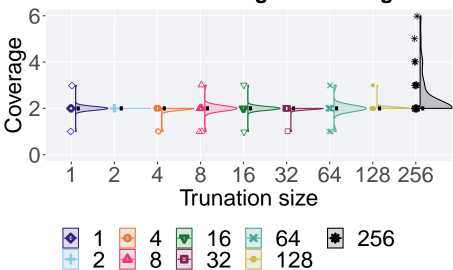
### 2.5.2 Final activation gene coverage

Activation gene coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'mpe') %>%
  ggplot(., aes(x = T, y = uni_str_pos, color = T, fill = T, shape = T)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = '
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha
  scale_y_continuous(
    name="Coverage",
    limits=c(0, 6),
    breaks=c(0,2,4,6)
  ) +
  scale_x_discrete(
    name="Trunation size"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final activation gene coverage')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
```

```
theme(legend.position="none"),
legend,
nrow=2,
rel_heights = c(3,1)
)
```

# Final activation gene coverage



### 2.5.2.1 Stats

```
act_coverage = filter(over_time_df, gen == 50000 & acro == 'mpe')
act_coverage$acro = factor(act_coverage$acro, levels = TR_LIST)
act_coverage %>%
group_by(T) %>%
dplyr::summarise(
   count = n(),
   na_cnt = sum(is.na(uni_str_pos)),
   min = min(uni_str_pos, na.rm = TRUE),
   median = median(uni_str_pos, na.rm = TRUE),
   mean = mean(uni_str_pos, na.rm = TRUE),
   max = max(uni_str_pos, na.rm = TRUE),
   IQR = IQR(uni_str_pos, na.rm = TRUE)
)
```

```
##
     <fct> <int>
                    <int> <int>
                                   <dbl> <dbl> <int> <dbl>
                                       2
                                           2
## 1 1
                50
                         0
                               1
                                                     3
## 2 2
                               2
                                           2
                                                     2
                                                            0
                50
                         0
                                       2
## 3 4
                                                     2
                50
                         0
                               1
                                       2
                                           1.96
                                                            0
## 4 8
                                                     3
                50
                         0
                               1
                                       2
                                           2
                                                            0
## 5 16
                50
                         0
                               1
                                       2
                                           2
                                                     3
                                                            0
## 6 32
                         0
                                       2
                                          1.98
                                                     2
                                                            0
                50
                               1
## 7 64
                50
                         0
                               1
                                       2
                                           2
                                                     3
                                                            0
## 8 128
                               2
                                           2.02
                                                     3
                50
                         0
                                       2
                                                            0
## 9 256
                50
                         0
                                2
                                       2 2.36
                                                     6
                                                            0
```

Kruskal-Wallis test illustrates evidence of statistical differences.

```
kruskal.test(uni_str_pos ~ T, data = act_coverage)
##
##
   Kruskal-Wallis rank sum test
##
## data: uni_str_pos by T
## Kruskal-Wallis chi-squared = 32.719, df = 8, p-value = 6.92e-05
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = act_coverage$uni_str_pos, g = act_coverage$T, p.adjust.method
                     paired = FALSE, conf.int = FALSE, alternative = 't')
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: act_coverage$uni_str_pos and act_coverage$T
##
##
                                                 128
       1
             2
                   4
                         8
                               16
                                     32
                                           64
## 2
      1.000 -
## 4
       1.000 1.000 -
## 8
       1.000 1.000 1.000 -
## 16
      1.000 1.000 1.000 1.000 -
      1.000 1.000 1.000 1.000 -
      1.000 1.000 1.000 1.000 1.000 -
## 128 1.000 1.000 1.000 1.000 1.000 1.000 -
## 256 0.092 0.034 0.015 0.187 0.092 0.022 0.320 0.142
## P value adjustment method: bonferroni
```

### 2.5.3 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'mpe') %>%
group_by(T, gen) %>%
dplyr::summarise(
  min = min(pop_fit_max) / DIMENSIONALITY,
  mean = mean(pop_fit_max) / DIMENSIONALITY,
  max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'T'. You can override using the `.groups`
## argument.
ggplot(lines, aes(x=gen, y=mean, group = T, fill = T, color = T, shape = T)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2.0, alpha =
  scale_y_continuous(
   name="Average trait score",
   limits=c(0, 100),
   breaks=seq(0,100, 20),
   labels=c("0", "20", "40", "60", "80", "100")
  ) +
  scale_x_continuous(
   name="Generations".
   limits=c(0, 50000),
   breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
   shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
   fill=guide_legend(nrow=2, title.position = "bottom")
```

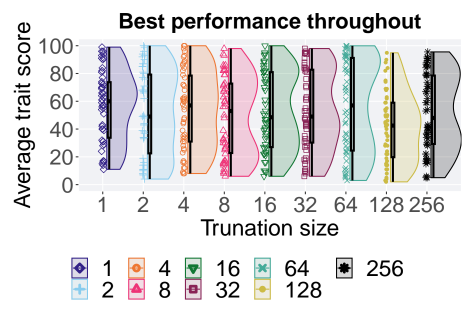


### 2.5.4 Best performance throughout

Best performance reached throughout 50,000 generations in a population.

```
plot = filter(best_df, var == 'pop_fit_max' & acro == 'mpe') %>%
  ggplot(., aes(x = T, y = val / DIMENSIONALITY, color = T, fill = T, shape = T)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = '
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha
  scale_y_continuous(
    name="Average trait score",
    limits=c(0, 100),
    breaks=seq(0,100, 20),
    labels=c("0", "20", "40", "60", "80", "100")
  scale_x_discrete(
    name="Trunation size"
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
```

```
plot +
    theme(legend.position="none"),
legend,
nrow=2,
rel_heights = c(3,1)
)
```



#### 2.5.4.1 Stats

Summary statistics for the best performance.

```
performance = filter(best_df, var == 'pop_fit_max' & acro == 'mpe')
performance %>%
  group_by(T) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(val)),
    min = min(val / DIMENSIONALITY, na.rm = TRUE),
    median = median(val / DIMENSIONALITY, na.rm = TRUE),
    mean = mean(val / DIMENSIONALITY, na.rm = TRUE),
    max = max(val / DIMENSIONALITY, na.rm = TRUE),
    IQR = IQR(val / DIMENSIONALITY, na.rm = TRUE)
)
```

```
## # A tibble: 9 x 8
## T count na_cnt min median mean max IQR
```

```
## <fct> <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
## 1 1
            50
                    0 11
                             60.0 56.6 99.0 40.0
## 2 2
            50
                    0 4
                             49.0 51.6 100.
                                              56.7
## 3 4
            50
                    0 8.00
                             57.0 53.9 100.
                                              47.5
## 4 8
            50
                    0 6
                             53.0 48.9 98.0 50.0
## 5 16
            50
                    0 6
                             48.5 52.7 100.
                                              54.0
## 6 32
            50
                    0 6
                             49.0 53.3 98.0 52.5
## 7 64
            50
                    0 3
                             57.0 55.8 99.9 66.7
## 8 128
                    0 2
            50
                             42.5 41.7 94.9 39.2
## 9 256
            50
                    0 5
                             48.0 51.6 95.5 49.3
```

Kruskal–Wallis test illustrates evidence of no statistical differences.

```
kruskal.test(val ~ T, data = performance)
```

```
##
## Kruskal-Wallis rank sum test
##
## data: val by T
## Kruskal-Wallis chi-squared = 9.7113, df = 8, p-value = 0.2859
```

# Chapter 3

# Tournament selection

Results for the tournament selection parameter sweep on the diagnostics with no valleys.

### 3.1 Data setup

```
over_time_df <- read.csv(paste(DATA_DIR,'OVER-TIME/tor.csv', sep = "", collapse = NULL), header =
over_time_df$T <- factor(over_time_df$T, levels = TS_LIST)

best_df <- read.csv(paste(DATA_DIR,'BEST/tor.csv', sep = "", collapse = NULL), header = TRUE, str
best_df$T <- factor(best_df$T, levels = TS_LIST)

sati_df <- read.csv(paste(DATA_DIR,'SOL-FND/tor.csv', sep = "", collapse = NULL), header = TRUE,
sati_df$T <- factor(sati_df$T, levels = TS_LIST)</pre>
```

### 3.2 Exploitation rate results

Here we present the results for **best performances** found by each selection scheme parameter on the exploitation rate diagnostic. 50 replicates are conducted for each scheme explored.

#### 3.2.1 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'exp') %>%
group_by(T, gen) %>%
```

```
dplyr::summarise(
   min = min(pop_fit_max) / DIMENSIONALITY,
   mean = mean(pop_fit_max) / DIMENSIONALITY,
   max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'T'. You can override using the `.groups`
## argument.
over_time_plot = ggplot(lines, aes(x=gen, y=mean, group = T, fill = T, color = T, shape
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale_y_continuous(
   name="Average trait score",
   limits=c(0, 100),
   breaks=seq(0,100, 20),
   labels=c("0", "20", "40", "60", "80", "100")
 ) +
  scale x continuous(
   name="Generations",
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
 ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
 guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
    fill=guide_legend(nrow=2, title.position = "bottom")
over_time_plot
```



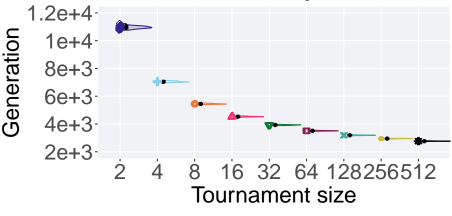
### 3.2.2 Generation satisfactory solution found

First generation a satisfactory solution is found throughout the  $50,\!000$  generations.

```
plot = filter(sati_df, acro == 'exp') %>%
  ggplot(., aes(x = T, y = gen , color = T, fill = T, shape = T)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Generation",
   limits=c(2000, 12000),
   breaks=c(2000, 4000, 6000, 8000, 10000, 12000),
   labels=c("2e+3", "4e+3", "6e+3", "8e+3", "1e+4", "1.2e+4")
  ) +
  scale_x_discrete(
   name="Tournament size"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Generation satisfactory solution found')+
  p_theme + theme(legend.title=element_blank())
```

```
plot_grid(
  plot +
    theme(legend.position="none"),
  legend,
  nrow=2,
  rel_heights = c(3,1)
)
```







#### 3.2.2.1 Stats

```
ssf = filter(sati_df, gen <= GENERATIONS & acro == 'exp')
ssf$acro = factor(ssf$acro, levels = TS_LIST)
ssf %>%
  group_by(T) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(gen)),
    min = min(gen, na.rm = TRUE),
    median = median(gen, na.rm = TRUE),
    mean = mean(gen, na.rm = TRUE),
    max = max(gen, na.rm = TRUE),
    IQR = IQR(gen, na.rm = TRUE)
)
```

```
## # A tibble: 9 x 8
    Τ
          count na_cnt
                         min median
                                      mean
                                             max
                                                   IQR
     <fct> <int> <int> <int> <dbl>
                                     <dbl> <int> <dbl>
## 1 2
             50
                     0 10756 10958. 10960. 11232 140
## 2 4
             50
                     0 6959 7040
                                     7049.
                                            7141
                                                  66
## 3 8
             50
                     0
                        5387 5442
                                     5449.
                                            5518
                                                  45.5
## 4 16
             50
                     0 4455 4528
                                     4532.
                                            4592
                                                  32.5
## 5 32
             50
                     0 3888 3930.
                                     3929.
                                            3974
                                                  30.8
## 6 64
                                            3545
             50
                     0
                        3468
                              3509
                                     3510.
                                                  23
## 7 128
             50
                     0 3156 3189
                                     3191.
                                            3234
                                                  22.5
## 8 256
             50
                     0 2908 2949
                                     2948.
                                            2985
                                                 19.5
## 9 512
             50
                     0 2718 2764.
                                     2766.
                                            2801 16.8
```

Kruskal–Wallis test illustrates evidence of statistical differences.

```
kruskal.test(gen ~ T, data = ssf)
```

```
##
## Kruskal-Wallis rank sum test
##
## data: gen by T
## Kruskal-Wallis chi-squared = 443.46, df = 8, p-value < 2.2e-16</pre>
```

Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.

```
##
  Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: ssf$gen and ssf$T
##
##
      2
                   8
                          16
                                              128
                                                    256
                                32
                                       64
## 4
      <2e-16 -
## 8
      <2e-16 <2e-16 -
## 16 <2e-16 <2e-16 -
## 32 <2e-16 <2e-16 <2e-16 -
## 64 <2e-16 <2e-16 <2e-16 <2e-16 -
## 128 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16 <
## 256 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16 <
## 512 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16
## P value adjustment method: bonferroni
```

# 3.3 Ordered exploitation results

Here we present the results for **best performances** found by each selection scheme parameter on the exploitation rate diagnostic. 50 replicates are conducted for each scheme explored.

#### 3.3.1 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'ord') %>%
group_by(T, gen) %>%
dplyr::summarise(
  min = min(pop_fit_max) / DIMENSIONALITY,
  mean = mean(pop_fit_max) / DIMENSIONALITY,
  max = max(pop_fit_max) / DIMENSIONALITY
)
```

## `summarise()` has grouped output by 'T'. You can override using the `.groups`
## argument.

```
ggplot(lines, aes(x=gen, y=mean, group = T, fill = T, color = T, shape = T)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale_y_continuous(
    name="Average trait score",
   limits=c(0, 100),
    breaks=seq(0,100, 20),
    labels=c("0", "20", "40", "60", "80", "100")
  scale_x_continuous(
    name="Generations",
    limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
    labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Performance over time')+
 p theme + theme(legend.title=element blank()) +
  guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
```

```
color=guide_legend(nrow=2, title.position = "bottom"),
fill=guide_legend(nrow=2, title.position = "bottom")
)
```



# 3.3.2 Generation satisfactory solution found

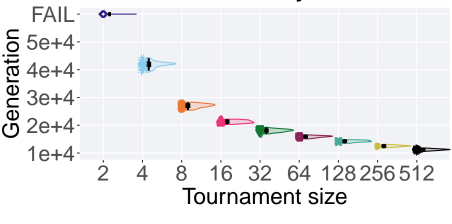
First generation a satisfactory solution is found throughout the 50,000 generations.

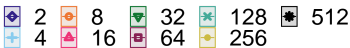
```
plot = filter(sati_df, acro == 'ord') %>%
    ggplot(., aes(x = T, y = gen , color = T, fill = T, shape = T)) +
    geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
    geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
    geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
    scale_y_continuous(
    name="Generation",
    limits=c(10000, 60000),
    breaks=c(10000, 20000, 30000, 40000,50000,60000),
    labels=c("1e+4","2e+4","3e+4","4e+4","5e+4","FAIL")
) +
    scale_x_discrete(
    name="Tournament size"
)+
    scale_shape_manual(values=SHAPE)+
    scale_colour_manual(values = cb_palette) +
```

```
scale_fill_manual(values = cb_palette) +
ggtitle('Generation satisfactory solution found')+
p_theme + theme(legend.title=element_blank())

plot_grid(
  plot +
    theme(legend.position="none"),
  legend,
  nrow=2,
  rel_heights = c(3,1)
)
```

# Generation satisfactory solution found





#### 3.3.2.1 Stats

Summary statistics for the generation a satisfactory solution is found.

```
ssf = filter(sati_df, gen <= GENERATIONS & acro == 'ord')
ssf$acro = factor(ssf$acro, levels = TS_LIST)
ssf %>%
  group_by(T) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(gen)),
    min = min(gen, na.rm = TRUE),
    median = median(gen, na.rm = TRUE),
```

```
mean = mean(gen, na.rm = TRUE),
   max = max(gen, na.rm = TRUE),
   IQR = IQR(gen, na.rm = TRUE)
)
## # A tibble: 8 x 8
    Т
          count na_cnt    min median
                                     mean
                                             max
                                                   IQR
    <fct> <int> <int> <dbl> <dbl> <int> <dbl>
                     0 39102 42086 41858. 44378 1207.
## 1 4
             50
## 2 8
             50
                     0 25443 27089 27014. 28293
## 3 16
             50
                     0 20292 21306. 21277. 22188
## 4 32
             50
                     0 16868 18107 18085. 19256
                     0 15114 15949 15885. 16540
## 5 64
             50
                                                  488
## 6 128
             50
                     0 13487 14228. 14238. 14789
## 7 256
             50
                     0 11756 12532. 12520. 13078 412.
## 8 512
             50
                     0 10311 11221 11209. 11823
Kruskal–Wallis test illustrates evidence of statistical differences.
kruskal.test(gen ~ T, data = ssf)
##
## Kruskal-Wallis rank sum test
##
## data: gen by T
## Kruskal-Wallis chi-squared = 392.76, df = 7, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = ssf$gen, g = ssf$T, p.adjust.method = "bonferroni",
                    paired = FALSE, conf.int = FALSE, alternative = '1')
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: ssf$gen and ssf$T
##
##
             8
                    16
                           32
                                  64
                                         128
                                                256
## 8
      <2e-16 -
## 16 <2e-16 <2e-16 -
## 32 <2e-16 <2e-16 -
## 64 <2e-16 <2e-16 <2e-16 -
## 128 <2e-16 <2e-16 <2e-16 <2e-16 -
## 256 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16 <
## 512 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16 <2e-16
## P value adjustment method: bonferroni
```

# 3.4 Contradictory objectives results

Here we present the results for activation gene coverage and satisfactory trait coverage found by each selection scheme parameter on the contradictory objectives diagnostic. 50 replicates are conducted for each scheme parameters explored.

# 3.4.1 Activation gene coverage over time

Activation gene coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations. Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'con') %>%
group_by(T, gen) %>%
dplyr::summarise(
  min = min(uni_str_pos),
  mean = mean(uni_str_pos),
  max = max(uni_str_pos)
)
```

## `summarise()` has grouped output by 'T'. You can override using the `.groups`
## argument.

```
ggplot(lines, aes(x=gen, y=mean, group = T, fill = T, color = T, shape = T)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
 geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale y continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks=seq(0,100, 20),
   labels=c("0", "20", "40", "60", "80", "100")
  ) +
  scale_x_continuous(
    name="Generations",
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene coverage over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
```

```
shape=guide_legend(nrow=2, title.position = "bottom"),
color=guide_legend(nrow=2, title.position = "bottom"),
fill=guide_legend(nrow=2, title.position = "bottom")
)
```



## 3.4.2 Final activation gene coverage

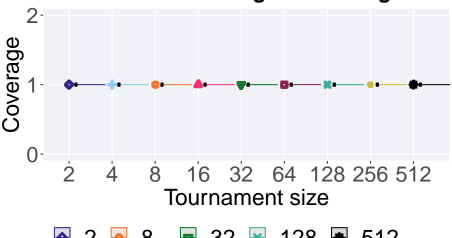
Activation gene coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'con') %>%
    ggplot(., aes(x = T, y = uni_str_pos, color = T, fill = T, shape = T)) +
    geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
    geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
    geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
    scale_y_continuous(
    name="Coverage",
    limits=c(0, 2),
    breaks=c(0,1,2)
) +
    scale_x_discrete(
    name="Tournament size"
)+
    scale_shape_manual(values=SHAPE)+
    scale_colour_manual(values = cb_palette, ) +
    scale_fill_manual(values = cb_palette) +
```

```
ggtitle('Final activation gene coverage')+
p_theme + theme(legend.title=element_blank())

plot_grid(
  plot +
    theme(legend.position="none"),
  legend,
  nrow=2,
  rel_heights = c(3,1)
)
```

# Final activation gene coverage





#### 3.4.2.1 Stats

Summary statistics for the generation a satisfactory solution is found.

```
act_coverage = filter(over_time_df, gen == 50000 & acro == 'con')
act_coverage$acro = factor(act_coverage$acro, levels = TS_LIST)
act_coverage %>%
group_by(T) %>%
dplyr::summarise(
   count = n(),
   na_cnt = sum(is.na(uni_str_pos)),
   min = min(uni_str_pos, na.rm = TRUE),
   median = median(uni_str_pos, na.rm = TRUE),
   mean = mean(uni_str_pos, na.rm = TRUE),
```

```
max = max(uni_str_pos, na.rm = TRUE),
    IQR = IQR(uni_str_pos, na.rm = TRUE)
 )
## # A tibble: 9 x 8
          count na_cnt
                          min median mean
                                                   IQR
   <fct> <int> <int> <dbl> <dbl> <int> <dbl>
## 1 2
            50
                      0
                            1
                                   1
                                         1
                                               1
## 2 4
             50
                      0
                            1
                                   1
                                         1
                                               1
## 3 8
             50
                      0
                            1
                                   1
                                         1
                                               1
                                                     0
              50
## 4 16
                      0
                                                     0
                            1
                                   1
                                         1
                                               1
## 5 32
              50
                      0
                            1
                                   1
                                         1
                                               1
## 6 64
              50
                      0
                            1
                                   1
                                         1
                                               1
## 7 128
              50
                      0
                            1
                                   1
                                         1
                                                     0
                                               1
## 8 256
              50
                      0
                            1
                                   1
                                         1
                                               1
                                                     0
## 9 512
              50
                            1
                                   1
                                         1
```

# 3.4.3 Satisfactory trait coverage over time

scale\_x\_continuous(
 name="Generations",
 limits=c(0, 50000),

Satisfactory trait coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations. Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'con') %>%
group_by(T, gen) %>%
dplyr::summarise(
  min = min(pop_uni_obj),
  mean = mean(pop_uni_obj),
  max = max(pop_uni_obj)
)
```

breaks=c(0, 10000, 20000, 30000, 40000, 50000),

## `summarise()` has grouped output by 'T'. You can override using the `.groups`

## argument.
ggplot(lines, aes(x=gen, y=mean, group = T, fill = T, color = T, shape = T)) +
 geom\_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
 geom\_line(size = 0.5) +
 geom\_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2.0, alpha =
 scale\_y\_continuous(
 name="Coverage",
 limits=c(0, 2),
 breaks=c(0,1,2)
) +

```
labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
) +
scale_shape_manual(values=SHAPE)+
scale_colour_manual(values = cb_palette) +
scale_fill_manual(values = cb_palette) +
ggtitle('Satisfactory trait coverage over time')+
p_theme + theme(legend.title=element_blank()) +
guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
    fill=guide_legend(nrow=2, title.position = "bottom"))
```

# Satisfactory trait coverage over time



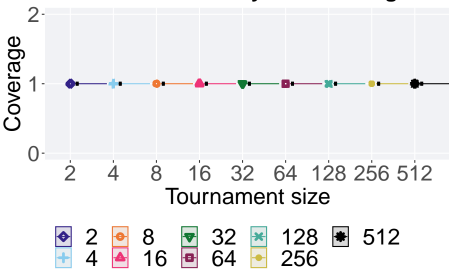
## 3.4.4 Final satisfactory trait coverage

Satisfactory trait coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'con') %>%
    ggplot(., aes(x = T, y = pop_uni_obj, color = T, fill = T, shape = T)) +
    geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1
    geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha
    scale_y_continuous(
    name="Coverage",
```

```
limits=c(0, 2),
   breaks=c(0,1,2)
  ) +
  scale_x_discrete(
   name="Tournament size"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final satisfactory trait coverage')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
 plot +
    theme(legend.position="none"),
 legend,
 nrow=2,
  rel_heights = c(3,1)
```





#### 3.4.4.1 Stats

Summary statistics for the generation a satisfactory solution is found.

```
sat_coverage = filter(over_time_df, gen == 50000 & acro == 'con')
sat_coverage$acro = factor(sat_coverage$acro, levels = TS_LIST)
sat_coverage %>%
  group_by(T) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(pop_uni_obj)),
    min = min(pop_uni_obj, na.rm = TRUE),
    median = median(pop_uni_obj, na.rm = TRUE),
    mean = mean(pop_uni_obj, na.rm = TRUE),
    max = max(pop_uni_obj, na.rm = TRUE),
    IQR = IQR(pop_uni_obj, na.rm = TRUE)
)
```

```
## # A tibble: 9 x 8
##
     Τ
                                                          IQR
            count na_cnt
                             min median mean
                                                   max
     <fct> <int>
                    <int> <int>
                                   <dbl> <dbl> <int>
                                                       <dbl>
## 1 2
                         0
                                                            0
                50
                               1
                                       1
                                              1
                                                     1
## 2 4
                50
                         0
                               1
                                                     1
                                                            0
                                       1
                                              1
## 3 8
                50
                         0
                               1
                                       1
                                              1
                                                     1
                                                            0
## 4 16
               50
                         0
                                                            0
                               1
                                              1
                                                     1
                                       1
## 5 32
               50
                         0
                               1
                                       1
                                              1
                                                     1
                                                            0
## 6 64
               50
                         0
                               1
                                       1
                                              1
                                                     1
                                                            0
                                                            0
## 7 128
               50
                         0
                               1
                                              1
## 8 256
                         0
                                                            0
               50
                               1
                                              1
                                                     1
                                       1
## 9 512
               50
                         0
                                1
                                              1
                                                            0
```

# 3.5 Multi-path exploration results

Here we present the results for **best performances** and **activation gene coverage** found by each selection scheme parameter on the multi-path exploration diagnostic. 50 replicates are conducted for each scheme parameter explored.

## 3.5.1 Activation gene coverage over time

Activation gene coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations. Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'mpe') %>%
group_by(T, gen) %>%
dplyr::summarise(
  min = min(uni_str_pos),
  mean = mean(uni_str_pos),
  max = max(uni_str_pos)
```

```
)
```

```
## `summarise()` has grouped output by 'T'. You can override using the `.groups`
## argument.
ggplot(lines, aes(x=gen, y=mean, group = T, fill = T, color = T, shape = T)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2.0, alpha =
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks=seq(0,100, 20),
   labels=c("0", "20", "40", "60", "80", "100")
  ) +
  scale_x_continuous(
   name="Generations".
   limits=c(0, 50000),
   breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene coverage over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
   shape=guide_legend(nrow=2, title.position = "bottom"),
   color=guide_legend(nrow=2, title.position = "bottom"),
   fill=guide_legend(nrow=2, title.position = "bottom")
```



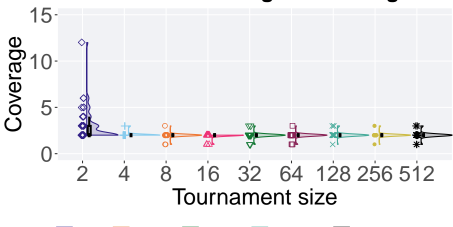
# 3.5.2 Final activation gene coverage

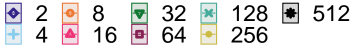
Activation gene coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'mpe') %>%
  ggplot(., aes(x = T, y = uni_str_pos, color = T, fill = T, shape = T)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = '
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha
  scale_y_continuous(
    name="Coverage",
    limits=c(0, 15),
    breaks=c(0,5,10,15)
  ) +
  scale_x_discrete(
    name="Tournament size"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final activation gene coverage')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
```

```
theme(legend.position="none"),
legend,
nrow=2,
rel_heights = c(3,1)
)
```

# Final activation gene coverage





## 3.5.2.1 Stats

Summary statistics for the generation a satisfactory solution is found.

```
act_coverage = filter(over_time_df, gen == 50000 & acro == 'mpe')
act_coverage$acro = factor(act_coverage$acro, levels = TS_LIST)
act_coverage %>%
group_by(T) %>%
dplyr::summarise(
   count = n(),
   na_cnt = sum(is.na(uni_str_pos)),
   min = min(uni_str_pos, na.rm = TRUE),
   median = median(uni_str_pos, na.rm = TRUE),
   mean = mean(uni_str_pos, na.rm = TRUE),
   max = max(uni_str_pos, na.rm = TRUE),
   IQR = IQR(uni_str_pos, na.rm = TRUE)
)
```

```
## # A tibble: 9 x 8
## T count na_cnt min median mean max IQR
```

```
##
     <fct> <int>
                    <int> <int>
                                   <dbl> <dbl> <int> <dbl>
                               2
                                          2.92
## 1 2
               50
                         0
                                       2
                                                    12
## 2 4
                               2
                                                           0
               50
                         0
                                       2
                                           2.06
                                                     3
## 3 8
               50
                        0
                               1
                                       2
                                          1.98
                                                     3
                                                           0
## 4 16
               50
                        0
                               1
                                       2
                                           1.94
                                                     2
                                                           0
## 5 32
               50
                        0
                               1
                                       2
                                           2
                                                     3
                                                           0
## 6 64
                        0
                                                     3
                                                           0
               50
                               1
                                       2
                                          1.96
## 7 128
               50
                        0
                               1
                                       2
                                          2.04
                                                     3
                                                           0
## 8 256
                                                     3
               50
                        0
                               1
                                       2
                                          2.02
                                                           0
## 9 512
               50
                         0
                               1
                                       2 2.02
                                                     3
                                                           0
```

Kruskal-Wallis test illustrates evidence of statistical differences.

```
## data: act_coverage$uni_str_pos and act_coverage$T
##
##
      2
              4
                                                            256
                     8
                             16
                                     32
                                            64
                                                    128
## 4
      0.00033 -
      1.6e-05 1.00000 -
## 8
      3.0e-06 0.27265 1.00000 -
      5.6e-05 1.00000 1.00000 1.00000 -
      1.2e-05 1.00000 1.00000 1.00000 -
## 128 0.00024 1.00000 1.00000 1.00000 1.00000 -
## 256 7.5e-05 1.00000 1.00000 1.00000 1.00000 1.00000 -
## 512 0.00018 1.00000 1.00000 1.00000 1.00000 1.00000 1.00000
## P value adjustment method: bonferroni
```

# 3.5.3 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'mpe') %>%
group_by(T, gen) %>%
dplyr::summarise(
  min = min(pop_fit_max) / DIMENSIONALITY,
  mean = mean(pop_fit_max) / DIMENSIONALITY,
  max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'T'. You can override using the `.groups`
## argument.
ggplot(lines, aes(x=gen, y=mean, group = T, fill = T, color = T, shape = T)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2.0, alpha =
  scale_y_continuous(
   name="Average trait score",
   limits=c(0, 100),
   breaks=seq(0,100, 20),
   labels=c("0", "20", "40", "60", "80", "100")
  ) +
  scale_x_continuous(
   name="Generations".
   limits=c(0, 50000),
   breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
   shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
   fill=guide_legend(nrow=2, title.position = "bottom")
```

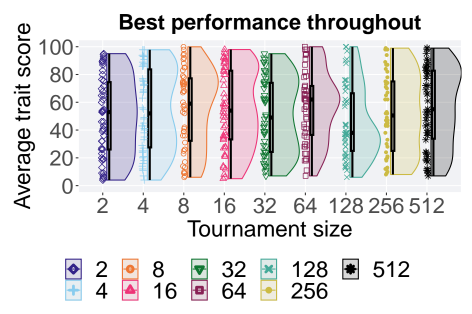


# 3.5.4 Best performance throughout

Best performance reached throughout 50,000 generations in a population.

```
plot = filter(best_df, var == 'pop_fit_max' & acro == 'mpe') %>%
  ggplot(., aes(x = T, y = val / DIMENSIONALITY, color = T, fill = T, shape = T)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = '
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1
 geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha
  scale_y_continuous(
    name="Average trait score",
    limits=c(0, 100),
    breaks=seq(0,100, 20),
    labels=c("0", "20", "40", "60", "80", "100")
  scale_x_discrete(
    name="Tournament size"
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
 p_theme + theme(legend.title=element_blank())
plot_grid(
```

```
plot +
    theme(legend.position="none"),
legend,
nrow=2,
rel_heights = c(3,1)
)
```



#### 3.5.4.1 Stats

Summary statistics for the best performance.

```
performance = filter(best_df, var == 'pop_fit_max' & acro == 'mpe')
performance %>%
  group_by(T) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(val)),
    min = min(val / DIMENSIONALITY, na.rm = TRUE),
    median = median(val / DIMENSIONALITY, na.rm = TRUE),
    mean = mean(val / DIMENSIONALITY, na.rm = TRUE),
    max = max(val / DIMENSIONALITY, na.rm = TRUE),
    IQR = IQR(val / DIMENSIONALITY, na.rm = TRUE)
)
```

```
## # A tibble: 9 x 8
## T count na_cnt min median mean max IQR
```

```
## <fct> <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
## 1 2
            50
                   0 4
                             53.0 49.7 95.0 48.5
## 2 4
            50
                    0 4
                             52.0 54.4 97.8 56.2
## 3 8
            50
                    0 6
                             59.0 55.5 99.9 45.0
## 4 16
            50
                    0 5
                             54.0 54.7 98.0 49.5
## 5 32
            50
                    0 7.00
                             49.0 50.4 95.0 49.7
## 6 64
            50
                    0 7
                             62.0 57.2 100.
                                             35.2
## 7 128
            50
                   0 6
                             38.0 45.7 100.
                                             41.5
## 8 256
                    0 8
            50
                             50.5 52.6 99.0 50.0
## 9 512
            50
                    0 7.00
                             55.0 55.9 99.0 49.0
```

Kruskal–Wallis test illustrates evidence of no statistical differences.

```
kruskal.test(val ~ T, data = performance)
```

```
##
## Kruskal-Wallis rank sum test
##
## data: val by T
## Kruskal-Wallis chi-squared = 6.8162, df = 8, p-value = 0.5566
```

# Chapter 4

# Genotypic fitness sharing

Results for the genotypic fitness sharing parameter sweep on the diagnostics with no valleys.

# 4.1 Data setup

```
over_time_df <- read.csv(paste(DATA_DIR,'OVER-TIME/gfs.csv', sep = "", collapse = NULL), header =
over_time_df$Sigma <- factor(over_time_df$Sigma, levels = FS_LIST)

best_df <- read.csv(paste(DATA_DIR,'BEST/gfs.csv', sep = "", collapse = NULL), header = TRUE, str
best_df$Sigma <- factor(best_df$Sigma, levels = FS_LIST)

sati_df <- read.csv(paste(DATA_DIR,'SOL-FND/gfs.csv', sep = "", collapse = NULL), header = TRUE,
sati_df$Sigma <- factor(sati_df$Sigma, levels = FS_LIST)</pre>
```

# 4.2 Exploitation rate results

Here we present the results for **best performances** found by each selection scheme parameter on the exploitation rate diagnostic. 50 replicates are conducted for each scheme explored.

#### 4.2.1 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'exp') %>%
group_by(Sigma, gen) %>%
```

```
dplyr::summarise(
   min = min(pop_fit_max) / DIMENSIONALITY,
   mean = mean(pop_fit_max) / DIMENSIONALITY,
   max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
over_time_plot = ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color =
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
 scale_y_continuous(
   name="Average trait score"
 ) +
  scale x continuous(
   name="Generations",
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
 ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
 scale_fill_manual(values = cb_palette) +
 ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
 guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
    fill=guide_legend(nrow=2, title.position = "bottom")
over_time_plot
```

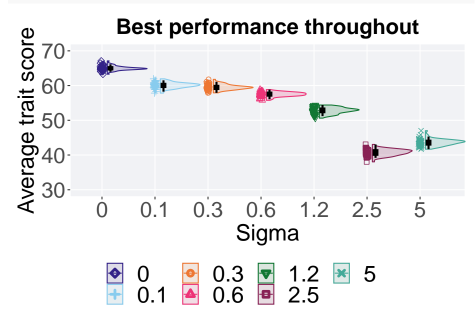


# 4.2.2 Best performance throughout

Best performance reached throughout 50,000 generations in a population.

```
plot = filter(best_df, var == 'pop_fit_max' & acro == 'exp') %>%
  ggplot(., aes(x = Sigma, y = val / DIMENSIONALITY, color = Sigma, fill = Sigma, shape = Sigma)
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Average trait score",
   limits = c(30,70)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
   theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



# 4.2.2.1 Stats

Summary statistics for the best performance.

```
performance = filter(best_df, var == 'pop_fit_max' & acro == 'exp')
performance$Sigma = factor(performance$Sigma, levels = c('0','0.1','0.3','0.6','1.2',')

performance %>%
  group_by(Sigma) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(val)),
    min = min(val / DIMENSIONALITY, na.rm = TRUE),
    median = median(val / DIMENSIONALITY, na.rm = TRUE),
    mean = mean(val / DIMENSIONALITY, na.rm = TRUE),
    max = max(val / DIMENSIONALITY, na.rm = TRUE),
    IQR = IQR(val / DIMENSIONALITY, na.rm = TRUE)
)
```

## # A tibble: 7 x 8
## Sigma count na\_cnt min median mean max IQR

```
##
     <fct> <int>
                 <int> <dbl>
                              <dbl> <dbl> <dbl> <dbl>
## 1 0
                     0 63.3
             50
                               64.9
                                    65.0 67.1 0.649
             50
## 2 0.1
                        57.8
                               60.1
                                    60.1 61.9 0.921
## 3 0.3
             50
                     0
                        57.9
                               59.5 59.5 61.8 0.900
## 4 0.6
             50
                     0
                        55.8
                               57.5 57.5 58.8 0.880
## 5 1.2
             50
                     0 50.6
                               53.0 52.8 54.4 1.06
## 6 5
             50
                     0 41.7
                               43.7 43.6 47.0 1.12
## 7 2.5
             50
                     0 38.0
                               40.7 40.7 43.0 1.33
```

Kruskal-Wallis test illustrates evidence of statistical differences.

```
kruskal.test(val ~ Sigma, data = performance)
```

```
##
##
   Kruskal-Wallis rank sum test
##
## data: val by Sigma
## Kruskal-Wallis chi-squared = 335.66, df = 6, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = performance$val, g = performance$Sigma, p.adjust.method = "bonferroni",
                     paired = FALSE, conf.int = FALSE, alternative = '1')
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: performance$val and performance$Sigma
##
##
                       0.3
       0
               0.1
                                0.6
                                        1.2
                                                5
## 0.1 < 2e-16 -
## 0.3 < 2e-16 0.0084
## 0.6 < 2e-16 2.9e-16 6.7e-16 -
## 1.2 < 2e-16 < 2e-16 < 2e-16 < 2e-16 -
       < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 -
## 2.5 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16
## P value adjustment method: bonferroni
```

# 4.3 Ordered exploitation results

Here we present the results for **best performances** found by each selection scheme parameter on the exploitation rate diagnostic. 50 replicates are conducted for each scheme explored.

#### 4.3.1 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'ord') %>%
group_by(Sigma, gen) %>%
dplyr::summarise(
   min = min(pop_fit_max) / DIMENSIONALITY,
   mean = mean(pop_fit_max) / DIMENSIONALITY,
   max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = S
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom line(size = 0.5) +
 geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale y continuous(
    name="Average trait score"
  ) +
  scale_x_continuous(
   name="Generations",
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides (
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
    fill=guide_legend(nrow=2, title.position = "bottom")
```

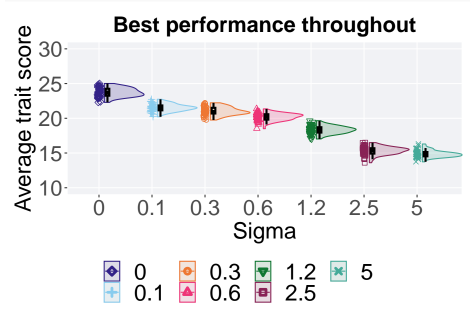


# 4.3.2 Best performance throughout

Best performance reached throughout 50,000 generations in a population.

```
plot = filter(best_df, var == 'pop_fit_max' & acro == 'ord') %>%
  ggplot(., aes(x = Sigma, y = val / DIMENSIONALITY, color = Sigma, fill = Sigma, shape = Sigma)
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Average trait score",
   limits = c(10,30)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



# 4.3.2.1 Stats

Summary statistics for the best performance.

```
performance = filter(best_df, var == 'pop_fit_max' & acro == 'ord')
performance %>%
  group_by(Sigma) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(val)),
    min = min(val / DIMENSIONALITY, na.rm = TRUE),
    median = median(val / DIMENSIONALITY, na.rm = TRUE),
    mean = mean(val / DIMENSIONALITY, na.rm = TRUE),
    max = max(val / DIMENSIONALITY, na.rm = TRUE),
    IQR = IQR(val / DIMENSIONALITY, na.rm = TRUE)
)
```

```
## # A tibble: 7 x 8
## Sigma count na_cnt min median mean max IQR
## <fct> <int> <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> > 1.00
## 1 0 50 0 22.3 23.7 23.7 25.0 1.00
```

```
## 2 0.1
              50
                        20.2
                               21.5
                                     21.5 22.7 0.705
## 3 0.3
              50
                        19.8
                               20.9
                                     21.0
                                           22.2 0.821
## 4 0.6
              50
                        18.5
                               20.3
                                     20.2
                                           21.4 0.776
## 5 1.2
              50
                        17.0
                               18.4 18.3 19.7 0.769
## 6 2.5
              50
                     0
                        13.6
                               15.4
                                     15.3
                                           16.5 0.774
## 7 5
              50
                        13.7
                               14.8 14.9 16.3 0.615
```

Kruskal–Wallis test illustrates evidence of statistical differences.

```
kruskal.test(val ~ Sigma, data = performance)
##
##
   Kruskal-Wallis rank sum test
##
## data: val by Sigma
## Kruskal-Wallis chi-squared = 326.67, df = 6, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = performance$val, g = performance$Sigma, p.adjust.method = "bonferroni",
                     paired = FALSE, conf.int = FALSE, alternative = '1')
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: performance$val and performance$Sigma
##
##
               0.1
                       0.3
                                0.6
                                        1.2
                                                2.5
       0
## 0.1 < 2e-16 -
## 0.3 < 2e-16 0.00102 -
## 0.6 < 2e-16 8.3e-15 1.6e-08 -
## 1.2 < 2e-16 < 2e-16 < 2e-16 4.4e-16 -
## 2.5 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 -
       < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 0.00019
##
## P value adjustment method: bonferroni
```

# 4.4 Contradictory objectives results

Here we present the results for activation gene coverage and satisfactory trait coverage found by each selection scheme parameter on the contradictory objectives diagnostic. 50 replicates are conducted for each scheme parameters explored.

# 4.4.1 Activation gene coverage over time

Activation gene coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations.

Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'con') %>%
group_by(Sigma, gen) %>%
dplyr::summarise(
   min = min(uni_str_pos),
   mean = mean(uni_str_pos),
   max = max(uni_str_pos)
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = S
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks = seq(0, 100, 20),
   labels=c("0", "20", "40", "60", "80", "100")
  ) +
  scale_x_continuous(
    name="Generations".
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene coverage over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
    fill=guide_legend(nrow=2, title.position = "bottom")
```



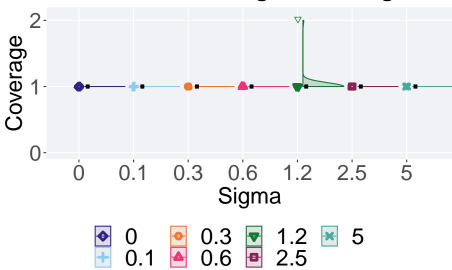
# 4.4.2 Final activation gene coverage

Activation gene coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'con') %>%
  ggplot(., aes(x = Sigma, y = uni_str_pos, color = Sigma, fill = Sigma, shape = Sigma)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 2.1),
   breaks=c(0,1,2)
  ) +
  scale_x_discrete(
   name="Sigma"
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final activation gene coverage')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
 plot +
```

```
theme(legend.position="none"),
legend,
nrow=2,
rel_heights = c(3,1)
)
```

# Final activation gene coverage



## 4.4.2.1 Stats

Summary statistics for the generation a satisfactory solution is found.

```
act_coverage = filter(over_time_df, gen == 50000 & acro == 'con')
act_coverage$Sigma = factor(act_coverage$Sigma, levels = FS_LIST)
act_coverage %>%
group_by(Sigma) %>%
dplyr::summarise(
   count = n(),
   na_cnt = sum(is.na(uni_str_pos)),
   min = min(uni_str_pos, na.rm = TRUE),
   median = median(uni_str_pos, na.rm = TRUE),
   mean = mean(uni_str_pos, na.rm = TRUE),
   max = max(uni_str_pos, na.rm = TRUE),
   IQR = IQR(uni_str_pos, na.rm = TRUE)
)
```

```
## # A tibble: 7 x 8
## Sigma count na_cnt min median mean max IQR
```

```
<fct> <int>
                   <int> <int>
                                <dbl> <dbl> <int> <dbl>
## 1 0
               50
                       0
                             1
                                     1
                                        1
                                                  1
               50
## 2 0.1
                       0
                                                        0
                             1
                                     1
                                        1
                                                  1
## 3 0.3
               50
                       0
                             1
                                     1
                                        1
                                                  1
                                                        0
## 4 0.6
               50
                       0
                             1
                                     1
                                        1
                                                  1
                                                        0
## 5 1.2
               50
                       0
                             1
                                     1
                                        1.02
                                                  2
                                                        0
## 6 2.5
               50
                       0
                                                        0
                             1
                                     1
                                        1
                                                  1
## 7 5
               50
                             1
                                     1
                                        1
                                                  1
                                                        0
```

Kruskal–Wallis test illustrates evidence of no statistical differences.

```
### Kruskal-Wallis rank sum test
## data: uni_str_pos by Sigma
## Kruskal-Wallis chi-squared = 6, df = 6, p-value = 0.4232
```

## 4.4.3 Satisfactory trait coverage over time

Satisfactory trait coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations. Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'con') %>%
  group_by(Sigma, gen) %>%
  dplyr::summarise(
    min = min(pop_uni_obj),
    mean = mean(pop_uni_obj),
    max = max(pop_uni_obj)
)
```

## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.

```
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = Sigma)) +
    geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
    geom_line(size = 0.5) +
    geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2.0, alpha =
    scale_y_continuous(
    name="Coverage",
    limits=c(0, 2.1),
    breaks=c(0,1,2)
) +
    scale_x_continuous(
    name="Generations",
    limits=c(0, 50000),
```

```
breaks=c(0, 10000, 20000, 30000, 40000, 50000),
  labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")

) +
scale_shape_manual(values=SHAPE)+
scale_colour_manual(values = cb_palette) +
scale_fill_manual(values = cb_palette) +
ggtitle('Satisfactory trait coverage over time')+
p_theme + theme(legend.title=element_blank()) +
guides(
  shape=guide_legend(nrow=2, title.position = "bottom"),
  color=guide_legend(nrow=2, title.position = "bottom"),
  fill=guide_legend(nrow=2, title.position = "bottom")
)
```

# Satisfactory trait coverage over time



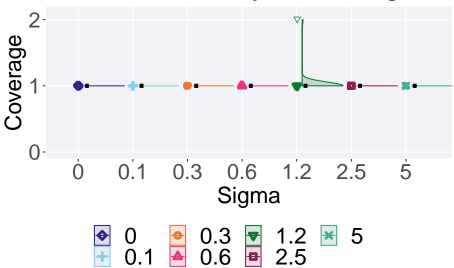
## 4.4.4 Final satisfactory trait coverage

Satisfactory trait coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'con') %>%
    ggplot(., aes(x = Sigma, y = pop_uni_obj, color = Sigma, fill = Sigma, shape = Sigma
    geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1
    geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha
    scale_y_continuous(
```

```
name="Coverage",
   limits=c(0, 2.1),
   breaks=c(0,1,2)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final satisfactory trait coverage')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
 plot +
    theme(legend.position="none"),
 legend,
  nrow=2,
  rel_heights = c(3,1)
)
```

# Final satisfactory trait coverage



## 4.4.4.1 Stats

Summary statistics for the generation a satisfactory solution is found.

```
sat_coverage = filter(over_time_df, gen == 50000 & acro == 'con')
sat_coverage$Sigma = factor(sat_coverage$Sigma, levels = FS_LIST)
sat_coverage %>%
  group_by(Sigma) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(pop_uni_obj)),
    min = min(pop_uni_obj, na.rm = TRUE),
    median = median(pop_uni_obj, na.rm = TRUE),
    mean = mean(pop_uni_obj, na.rm = TRUE),
    max = max(pop_uni_obj, na.rm = TRUE),
    IQR = IQR(pop_uni_obj, na.rm = TRUE)
)
```

```
## # A tibble: 7 x 8
##
                                                   IOR
    Sigma count na_cnt
                         min median mean
                                            max
     <fct> <int> <int> <dbl> <dbl> <int> <dbl>
                                  1 1
## 1 0
             50
                     0
                                              1
                                                    0
                           1
## 2 0.1
             50
                     0
                           1
                                  1
                                     1
                                                    0
## 3 0.3
             50
                     0
                           1
                                  1 1
                                              1
                                                    0
## 4 0.6
             50
                     0
                           1
                                  1 1
                                              1
                                                    0
## 5 1.2
             50
                     0
                                              2
                                                    0
                           1
                                  1 1.02
## 6 2.5
             50
                     0
                                                    0
                           1
                                  1 1
                                              1
## 7 5
             50
                           1
                                  1 1
                                              1
                                                    0
```

Kruskal–Wallis test illustrates evidence of no statistical differences.

```
kruskal.test(pop_uni_obj ~ Sigma, data = sat_coverage)
```

```
##
## Kruskal-Wallis rank sum test
##
## data: pop_uni_obj by Sigma
## Kruskal-Wallis chi-squared = 6, df = 6, p-value = 0.4232
```

# 4.5 Multi-path exploration results

Here we present the results for **best performances** and **activation gene coverage** found by each selection scheme parameter on the multi-path exploration diagnostic. 50 replicates are conducted for each scheme parameter explored.

## 4.5.1 Activation gene coverage over time

Activation gene coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations. Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'mpe') %>%
group_by(Sigma, gen) %>%
dplyr::summarise(
  min = min(uni_str_pos),
  mean = mean(uni_str_pos),
  max = max(uni_str_pos)
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = Sigma)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2.0, alpha =
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks=seq(0,100, 20),
   labels=c("0", "20", "40", "60", "80", "100")
  ) +
  scale_x_continuous(
   name="Generations".
   limits=c(0, 50000),
   breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene coverage over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
   shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
   fill=guide_legend(nrow=2, title.position = "bottom")
```



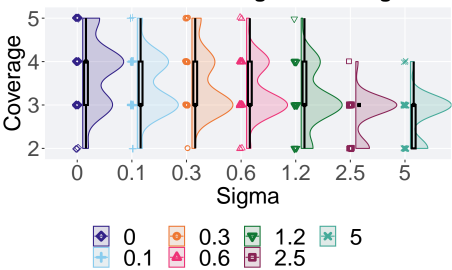
#### 4.5.2 Final activation gene coverage

Activation gene coverage found in the final population at 50,000 generations.

```
plot = filter(over time df, gen == 50000 & acro == 'mpe') %>%
  ggplot(., aes(x = Sigma, y = uni_str_pos, color = Sigma, fill = Sigma, shape = Sigma
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = '
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha
  scale_y_continuous(
    name="Coverage",
    limits=c(1.9, 5.1)
  scale_x_discrete(
    name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final activation gene coverage')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
 plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```

## Final activation gene coverage



#### 4.5.2.1 Stats

Summary statistics for the generation a satisfactory solution is found.

```
act_coverage = filter(over_time_df, gen == 50000 & acro == 'mpe')
act_coverage$Sigma = factor(act_coverage$Sigma, levels = FS_LIST)
act_coverage %>%
group_by(Sigma) %>%
dplyr::summarise(
   count = n(),
   na_cnt = sum(is.na(uni_str_pos)),
   min = min(uni_str_pos, na.rm = TRUE),
   median = median(uni_str_pos, na.rm = TRUE),
   mean = mean(uni_str_pos, na.rm = TRUE),
   max = max(uni_str_pos, na.rm = TRUE),
   IQR = IQR(uni_str_pos, na.rm = TRUE)
)
```

```
## # A tibble: 7 x 8
## Sigma count na_cnt min median mean max IQR
## <fct> <int> <int> <int> <dbl> <dbl> <int> <dbl>
```

```
## 1 0
               50
                                         3.74
                                                    5
                                                          1
## 2 0.1
               50
                              2
                                                    5
                        0
                                      3
                                         3.44
                                                          1
                              2
                                                    5
## 3 0.3
               50
                        0
                                        3.5
                                                          1
                                      3
## 4 0.6
                              2
               50
                        0
                                      3 3.34
                                                    5
                                                          1
## 5 1.2
               50
                        0
                              2
                                      3 3.14
                                                   5
                                                          1
## 6 2.5
               50
                        0
                              2
                                      3
                                         2.8
                                                   4
                                                          0
## 7 5
               50
                        0
                              2
                                      3 2.78
                                                    4
                                                          1
```

Kruskal-Wallis test illustrates evidence of statistical differences.
kruskal.test(uni\_str\_pos ~ Sigma, data = act\_coverage)

```
##
## Kruskal-Wallis rank sum test
##
## data: uni_str_pos by Sigma
## Kruskal-Wallis chi-squared = 69.982, df = 6, p-value = 4.123e-13
**Proof: The form the William of the first proof: The Proof: T
```

pairwise.wilcox.test(x = act\_coverage\$uni\_str\_pos, g = act\_coverage\$Sigma, p.adjust.me

Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.

```
paired = FALSE, conf.int = FALSE, alternative = '1')
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
##
## data: act_coverage$uni_str_pos and act_coverage$Sigma
##
##
       0
               0.1
                       0.3
                               0.6
                                        1.2
                                                2.5
## 0.1 0.52684 -
## 0.3 0.98117 1.00000 -
## 0.6 0.08439 1.00000 1.00000 -
## 1.2 0.00273 0.43402 0.29939 1.00000 -
## 2.5 5.9e-08 2.3e-05 1.3e-05 0.00023 0.09112 -
       8.8e-08 2.9e-05 1.5e-05 0.00025 0.07467 1.00000
## 5
```

#### 4.5.3 Performance over time

## P value adjustment method: bonferroni

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'mpe') %>%
group_by(Sigma, gen) %>%
dplyr::summarise(
   min = min(pop_fit_max) / DIMENSIONALITY,
   mean = mean(pop_fit_max) / DIMENSIONALITY,
```

```
max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = Sigma)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom line(size = 0.5) +
  geom_point(data = filter(lines, gen 1000 == 0 & gen != 0), size = 1.5, stroke = 2.0, alpha =
  scale_y_continuous(
   name="Average trait score"
  ) +
  scale_x_continuous(
   name="Generations",
   limits=c(0, 50000),
   breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
   shape=guide_legend(nrow=2, title.position = "bottom"),
   color=guide_legend(nrow=2, title.position = "bottom"),
   fill=guide_legend(nrow=2, title.position = "bottom")
```

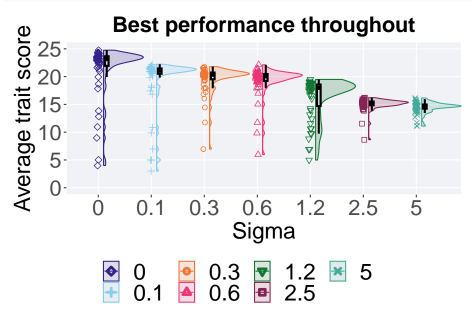


#### 4.5.4 Best performance throughout

Best performance reached throughout 50,000 generations in a population.

```
plot = filter(best_df, var == 'pop_fit_max' & acro == 'mpe') %>%
  ggplot(., aes(x = Sigma, y = val / DIMENSIONALITY, color = Sigma, fill = Sigma, shap
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = '
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha
  scale_y_continuous(
    name="Average trait score",
    limits=c(0, 25)
  ) +
  scale_x_discrete(
    name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



#### 4.5.4.1 Stats

Summary statistics for the best performance.

```
performance = filter(best_df, var == 'pop_fit_max' & acro == 'mpe')
performance %>%
  group_by(Sigma) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(val)),
    min = min(val / DIMENSIONALITY, na.rm = TRUE),
    median = median(val / DIMENSIONALITY, na.rm = TRUE),
    mean = mean(val / DIMENSIONALITY, na.rm = TRUE),
    max = max(val / DIMENSIONALITY, na.rm = TRUE),
    IQR = IQR(val / DIMENSIONALITY, na.rm = TRUE)
)
```

```
## # A tibble: 7 x 8
## Sigma count na_cnt min median mean max IQR
## <fct> <int> <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> 1.72
## 1 0 50 0 4.00 23.2 21.3 24.8 1.72
```

```
## 2 0.1
             50
                     0 3.00
                               21.1 19.2 22.2 0.956
## 3 0.3
             50
                     0 6.96
                               20.4 19.5
                                          21.8 1.21
## 4 0.6
             50
                     0 5.97
                               20.0 19.2 22.1 1.23
## 5 1.2
             50
                     0 4.98
                               18.0 16.1 19.5 3.87
## 6 2.5
             50
                     0 8.65
                               15.3 15.0 16.2 0.675
## 7 5
             50
                     0 11.1
                               14.7 14.5 16.2 0.754
```

Kruskal–Wallis test illustrates evidence of statistical differences.

## P value adjustment method: bonferroni

```
kruskal.test(val ~ Sigma, data = performance)
##
## Kruskal-Wallis rank sum test
##
## data: val by Sigma
## Kruskal-Wallis chi-squared = 187.09, df = 6, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = performance$val, g = performance$Sigma, p.adjust.method = "box
                     paired = FALSE, conf.int = FALSE, alternative = '1')
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: performance$val and performance$Sigma
##
##
       0
               0.1
                       0.3
                               0.6
                                        1.2
                                                2.5
## 0.1 3.9e-07 -
## 0.3 1.4e-07 0.00651 -
## 0.6 8.1e-08 0.00029 0.76146 -
## 1.2 6.8e-10 3.7e-08 1.6e-09 2.4e-10 -
## 2.5 4.7e-10 5.7e-10 2.8e-13 1.5e-12 0.00022 -
      3.2e-10 6.2e-10 1.4e-13 5.6e-13 0.00014 0.00038
##
```

## Chapter 5

# Phenotypic fitness sharing

Results for the phenotypic fitness sharing parameter sweep on the diagnostics with no valleys.

#### 5.1 Data setup

```
over_time_df <- read.csv(paste(DATA_DIR,'OVER-TIME/pfs.csv', sep = "", collapse = NULL), header =
over_time_df$Sigma <- factor(over_time_df$Sigma, levels = FS_LIST)

best_df <- read.csv(paste(DATA_DIR,'BEST/pfs.csv', sep = "", collapse = NULL), header = TRUE, str
best_df$Sigma <- factor(best_df$Sigma, levels = FS_LIST)

sati_df <- read.csv(paste(DATA_DIR,'SOL-FND/pfs.csv', sep = "", collapse = NULL), header = TRUE,
sati_df$Sigma <- factor(sati_df$Sigma, levels = FS_LIST)</pre>
```

## 5.2 Exploitation rate results

Here we present the results for **best performances** found by each selection scheme parameter on the exploitation rate diagnostic. 50 replicates are conducted for each scheme explored.

#### 5.2.1 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'exp') %>%
group_by(Sigma, gen) %>%
```

```
dplyr::summarise(
   min = min(pop_fit_max) / DIMENSIONALITY,
   mean = mean(pop_fit_max) / DIMENSIONALITY,
   max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
over_time_plot = ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color =
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
 scale_y_continuous(
   name="Average trait score"
 ) +
  scale x continuous(
   name="Generations",
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
 ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
 scale_fill_manual(values = cb_palette) +
 ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
 guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
    fill=guide_legend(nrow=2, title.position = "bottom")
over_time_plot
```

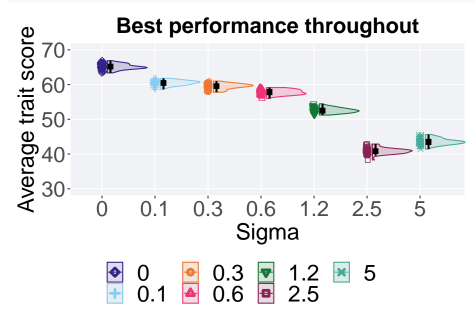


#### 5.2.2 Best performance throughout

Best performance reached throughout 50,000 generations in a population.

```
plot = filter(best_df, var == 'pop_fit_max' & acro == 'exp') %>%
  ggplot(., aes(x = Sigma, y = val / DIMENSIONALITY, color = Sigma, fill = Sigma, shape = Sigma)
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Average trait score",
   limits = c(30,70)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



#### 5.2.2.1 Stats

Summary statistics for the best performance.

Sigma count na\_cnt

```
performance = filter(best_df, var == 'pop_fit_max' & acro == 'exp')
performance$Sigma = factor(performance$Sigma, levels = c('0','0.1','0.3','0.6','1.2',')

performance %>%
    group_by(Sigma) %>%
    dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(val)),
    min = min(val / DIMENSIONALITY, na.rm = TRUE),
    median = median(val / DIMENSIONALITY, na.rm = TRUE),
    mean = mean(val / DIMENSIONALITY, na.rm = TRUE),
    max = max(val / DIMENSIONALITY, na.rm = TRUE),
    IQR = IQR(val / DIMENSIONALITY, na.rm = TRUE)
)

## # A tibble: 7 x 8
```

max

IQR

min median mean

```
##
     <fct> <int>
                 <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
## 1 0
                     0 63.4
             50
                               65.0
                                     65.1
                                           66.9 1.07
             50
## 2 0.1
                        58.6
                               60.5
                                     60.5
                                           61.8 1.06
## 3 0.3
             50
                     0
                        57.8
                               59.6
                                     59.5
                                           61.0 1.03
## 4 0.6
             50
                     0
                        56.0
                               57.8 57.8 59.1 1.24
## 5 1.2
             50
                     0 51.2
                               52.5
                                     52.6 54.5 1.10
## 6 5
             50
                     0 41.4
                               43.5 43.6 45.6 1.15
## 7 2.5
             50
                     0 38.3
                               40.9 40.9 42.9 1.04
```

Kruskal-Wallis test illustrates evidence of statistical differences.

kruskal.test(val ~ Sigma, data = performance)

##

```
##
## Kruskal-Wallis rank sum test
##
```

## data: val by Sigma
## Kruskal-Wallis chi-squared = 335.67, df = 6, p-value < 2.2e-16</pre>

Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.

```
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: performance$val and performance$Sigma
##
##
                       0.3
       0
               0.1
                               0.6
                                       1.2
                                               5
## 0.1 < 2e-16 -
## 0.3 < 2e-16 6.4e-07 -
## 0.6 < 2e-16 < 2e-16 1.7e-12 -
## 1.2 < 2e-16 < 2e-16 < 2e-16 < 2e-16 -
       < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 -
## 2.5 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 5.9e-16
## P value adjustment method: bonferroni
```

## 5.3 Ordered exploitation results

Here we present the results for **best performances** found by each selection scheme parameter on the exploitation rate diagnostic. 50 replicates are conducted for each scheme explored.

#### 5.3.1 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'ord') %>%
group_by(Sigma, gen) %>%
dplyr::summarise(
  min = min(pop_fit_max) / DIMENSIONALITY,
  mean = mean(pop_fit_max) / DIMENSIONALITY,
  max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = S
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom line(size = 0.5) +
 geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale y continuous(
    name="Average trait score"
  ) +
  scale_x_continuous(
   name="Generations".
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
    labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides (
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
    fill=guide_legend(nrow=2, title.position = "bottom")
```

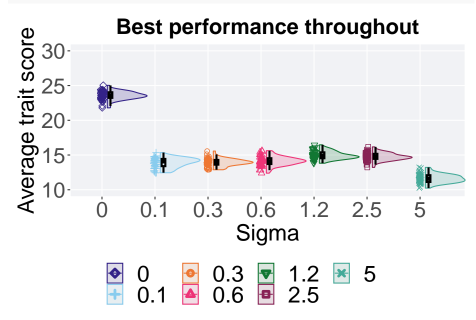


#### 5.3.2 Best performance throughout

Best performance reached throughout 50,000 generations in a population.

```
plot = filter(best df, var == 'pop fit max' & acro == 'ord') %>%
  ggplot(., aes(x = Sigma, y = val / DIMENSIONALITY, color = Sigma, fill = Sigma, shape = Sigma)
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Average trait score",
   limits = c(10,30)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



#### **5.3.2.1** Stats

Summary statistics for the best performance.

Sigma count na\_cnt

```
performance = filter(best_df, var == 'pop_fit_max' & acro == 'ord')
performance$Sigma = factor(performance$Sigma, levels = c('0','1.2','2.5','0.6','0.3',')
performance %>%
    group_by(Sigma) %>%
    dplyr::summarise(
        count = n(),
        na_cnt = sum(is.na(val)),
        min = min(val / DIMENSIONALITY, na.rm = TRUE),
        median = median(val / DIMENSIONALITY, na.rm = TRUE),
        mean = mean(val / DIMENSIONALITY, na.rm = TRUE),
        max = max(val / DIMENSIONALITY, na.rm = TRUE),
        IQR = IQR(val / DIMENSIONALITY, na.rm = TRUE)
)
```

min median mean

<fct> <int> <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>

```
## 1 0
              50
                         21.8
                                23.6
                                      23.6
                                            25.0 0.807
## 2 1.2
              50
                         13.8
                                14.9
                                      15.0
                                            16.5 0.851
## 3 2.5
              50
                         13.2
                                      14.7
                                            16.2 0.727
                                14.7
## 4 0.6
                                      14.1
              50
                      0
                         12.4
                                14.1
                                            15.6 0.834
## 5 0.3
              50
                      0 12.9
                                14.0
                                      13.9
                                            15.6 0.680
## 6 0.1
              50
                      0
                         12.4
                                14.1
                                      13.9
                                            15.4 1.04
## 7 5
                         10.2
                                11.6
                                     11.6 13.2 0.973
              50
```

Kruskal–Wallis test illustrates evidence of statistical differences.

kruskal.test(val ~ Sigma, data = performance)

##

```
##
## Kruskal-Wallis rank sum test
##
## data: val by Sigma
## Kruskal-Wallis chi-squared = 265.47, df = 6, p-value < 2.2e-16</pre>
```

Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.

```
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: performance$val and performance$Sigma
##
##
       0
                       2.5
                               0.6
                                       0.3
                                                0.1
## 1.2 < 2e-16 -
## 2.5 < 2e-16 0.90
## 0.6 < 2e-16 1.3e-07 3.7e-05 -
## 0.3 < 2e-16 1.3e-11 4.7e-09 0.95
## 0.1 < 2e-16 2.8e-10 1.2e-07 1.00
                                       1.00
       < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 3.3e-16
##
## P value adjustment method: bonferroni
```

### 5.4 Contradictory objectives results

Here we present the results for activation gene coverage and satisfactory trait coverage found by each selection scheme parameter on the contradictory objectives diagnostic. 50 replicates are conducted for each scheme parameters explored.

#### 5.4.1 Activation gene coverage over time

p\_theme + theme(legend.title=element\_blank()) +

shape=guide\_legend(nrow=2, title.position = "bottom"),
color=guide\_legend(nrow=2, title.position = "bottom"),
fill=guide\_legend(nrow=2, title.position = "bottom")

guides (

Activation gene coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations. Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'con') %>%
group_by(Sigma, gen) %>%
dplyr::summarise(
  min = min(uni_str_pos),
  mean = mean(uni_str_pos),
  max = max(uni_str_pos)
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = S
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks=seq(0,100, 20),
    labels=c("0", "20", "40", "60", "80", "100")
  ) +
  scale_x_continuous(
    name="Generations".
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
    labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene coverage over time')+
```

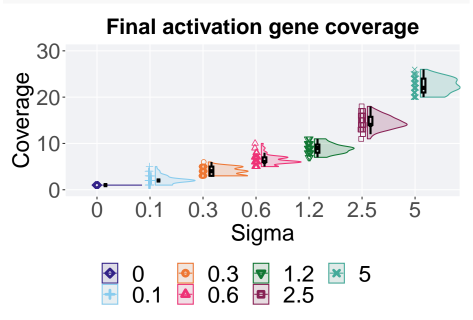


#### 5.4.2 Final activation gene coverage

Activation gene coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'con') %>%
  ggplot(., aes(x = Sigma, y = uni_str_pos, color = Sigma, fill = Sigma, shape = Sigma)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 30)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final activation gene coverage')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



#### 5.4.2.1 Stats

Summary statistics for the generation a satisfactory solution is found.

```
act_coverage = filter(over_time_df, gen == 50000 & acro == 'con')
act_coverage$Sigma = factor(act_coverage$Sigma, levels = FS_LIST)
act_coverage %>%
  group_by(Sigma) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(uni_str_pos)),
    min = min(uni_str_pos, na.rm = TRUE),
    median = median(uni_str_pos, na.rm = TRUE),
    mean = mean(uni_str_pos, na.rm = TRUE),
    max = max(uni_str_pos, na.rm = TRUE),
    IQR = IQR(uni_str_pos, na.rm = TRUE)
)
```

```
## # A tibble: 7 x 8
## Sigma count na_cnt min median mean max IQR
## <fct> <int> <int> <int> <dbl> <dbl> <int> <dbl>
```

```
## 1 0
              50
                      0
                                                   0
                             1
                                    1
                                       1
                                                1
## 2 0.1
              50
                                    2
                      0
                             1
                                       2.12
                                                5
## 3 0.3
              50
                      0
                             3
                                    4
                                      4
                                                6
                                                   2
## 4 0.6
              50
                      0
                             5
                                    6 6.42
                                               10 1
                            7
                                               11 1.75
## 5 1.2
              50
                      0
                                    9 8.88
## 6 2.5
              50
                      0
                            11
                                   14 14.6
                                               18 1.75
## 7 5
              50
                            20
                                   22 22.5
                                               26 3
                      0
```

Kruskal-Wallis test illustrates evidence of statistical differences.
kruskal.test(uni\_str\_pos ~ Sigma, data = act\_coverage)

```
##
## Kruskal-Wallis rank sum test
##
## data: uni_str_pos by Sigma
## Kruskal-Wallis chi-squared = 337.59, df = 6, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = act_coverage$uni_str_pos, g = act_coverage$Sigma, p.adjust.method = "bor paired = FALSE, conf.int = FALSE, alternative = 'g')</pre>
```

```
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
##
## data: act_coverage$uni_str_pos and act_coverage$Sigma
##
##
               0.1
                       0.3
                               0.6
                                       1.2
                                                2.5
## 0.1 1.0e-13 -
## 0.3 < 2e-16 5.9e-13 -
## 0.6 < 2e-16 < 2e-16 3.3e-15 -
## 1.2 < 2e-16 < 2e-16 < 2e-16 3.9e-13 -
## 2.5 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 -
## 5
       < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16
##
## P value adjustment method: bonferroni
```

#### 5.4.3 Satisfactory trait coverage over time

Satisfactory trait coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations. Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'con') %>%
group_by(Sigma, gen) %>%
dplyr::summarise(
   min = min(pop_uni_obj),
   mean = mean(pop_uni_obj),
```

```
max = max(pop_uni_obj)
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = S
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom point(data = filter(lines, gen \( \frac{1}{2} \) 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale_y_continuous(
    name="Coverage",
   limits=c(0, 30)
  ) +
  scale_x_continuous(
   name="Generations",
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
 ) +
 scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Satisfactory trait coverage over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
   fill=guide_legend(nrow=2, title.position = "bottom")
```

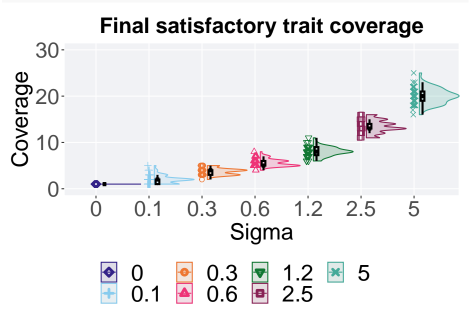


#### 5.4.4 Final satisfactory trait coverage

Satisfactory trait coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'con') %>%
  ggplot(., aes(x = Sigma, y = pop_uni_obj, color = Sigma, fill = Sigma, shape = Sigma)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 30)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final satisfactory trait coverage')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



#### 5.4.4.1 Stats

Summary statistics for the generation a satisfactory solution is found.

```
sat_coverage = filter(over_time_df, gen == 50000 & acro == 'con')
sat_coverage$Sigma = factor(sat_coverage$Sigma, levels = FS_LIST)
sat_coverage %>%
  group_by(Sigma) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(pop_uni_obj)),
    min = min(pop_uni_obj, na.rm = TRUE),
    median = median(pop_uni_obj, na.rm = TRUE),
    mean = mean(pop_uni_obj, na.rm = TRUE),
    max = max(pop_uni_obj, na.rm = TRUE),
    IQR = IQR(pop_uni_obj, na.rm = TRUE)
)
```

```
## # A tibble: 7 x 8
## Sigma count na_cnt min median mean max IQR
## <fct> <int> <int> <int> <dbl> <dbl> <int> <dbl>
```

```
## 1 0
               50
                        0
                                                       0
                               1
                                       1
                                          1
                                                    1
## 2 0.1
               50
                                       2
                                          2
                               1
                                                    5
## 3 0.3
               50
                        0
                               2
                                          3.72
                                                    5
                                                       1
## 4 0.6
               50
                        0
                               4
                                       5
                                          5.6
                                                    8 1
## 5 1.2
               50
                        0
                               6
                                       8
                                         8.04
                                                   11 1.75
## 6 2.5
               50
                        0
                              11
                                      13 13.4
                                                   16
                                                       1
## 7 5
               50
                                      20 20.0
                                                       2
                        0
                              16
                                                   25
```

Kruskal-Wallis test illustrates evidence of statistical differences.

```
kruskal.test(pop_uni_obj ~ Sigma, data = sat_coverage)
##
##
    Kruskal-Wallis rank sum test
##
## data: pop_uni_obj by Sigma
## Kruskal-Wallis chi-squared = 337.1, df = 6, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = sat_coverage$pop_uni_obj, g = sat_coverage$Sigma, p.adjust.method = "bor
                     paired = FALSE, conf.int = FALSE, alternative = 'g')
##
    Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
##
## data: sat_coverage$pop_uni_obj and sat_coverage$Sigma
##
##
       0
               0.1
                       0.3
                                0.6
                                        1.2
                                                2.5
## 0.1 9.2e-12 -
## 0.3 < 2e-16 1.4e-12 -
## 0.6 < 2e-16 < 2e-16 1.2e-14 -
## 1.2 < 2e-16 < 2e-16 < 2e-16 1.1e-14 -
## 2.5 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 -
## 5
       < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16
```

### 5.5 Multi-path exploration results

## P value adjustment method: bonferroni

##

Here we present the results for **best performances** and **activation gene coverage** found by each selection scheme parameter on the multi-path exploration diagnostic. 50 replicates are conducted for each scheme parameter explored.

#### 5.5.1 Activation gene coverage over time

Activation gene coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations.

Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'mpe') %>%
group_by(Sigma, gen) %>%
dplyr::summarise(
   min = min(uni_str_pos),
   mean = mean(uni_str_pos),
   max = max(uni_str_pos)
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = S
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks = seq(0, 100, 20),
   labels=c("0", "20", "40", "60", "80", "100")
  ) +
  scale_x_continuous(
    name="Generations".
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene coverage over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
    fill=guide_legend(nrow=2, title.position = "bottom")
```



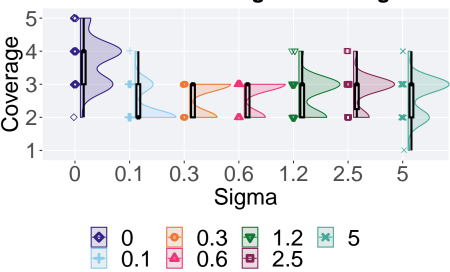
#### 5.5.2 Final activation gene coverage

Activation gene coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'mpe') %>%
  ggplot(., aes(x = Sigma, y = uni_str_pos, color = Sigma, fill = Sigma, shape = Sigma)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Coverage",
   limits=c(0.9, 5.1)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final activation gene coverage')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```

## Final activation gene coverage



#### 5.5.2.1 Stats

Summary statistics for the generation a satisfactory solution is found.

```
act_coverage = filter(over_time_df, gen == 50000 & acro == 'mpe')
act_coverage$Sigma = factor(act_coverage$Sigma, levels = FS_LIST)
act_coverage %>%
  group_by(Sigma) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(uni_str_pos)),
    min = min(uni_str_pos, na.rm = TRUE),
    median = median(uni_str_pos, na.rm = TRUE),
    mean = mean(uni_str_pos, na.rm = TRUE),
    max = max(uni_str_pos, na.rm = TRUE),
    IQR = IQR(uni_str_pos, na.rm = TRUE)
)
```

```
## # A tibble: 7 x 8
## Sigma count na_cnt min median mean max IQR
## <fct> <int> <int> <int> <dbl> <dbl> <int> <dbl>
```

```
## 1 0
              50
                     0
                           2
                                  4 3.7
                                              5 1
## 2 0.1
              50
                           2
                                  2
                                     2.34
                      0
                                               4
                                                 1
                           2
## 3 0.3
              50
                     0
                                  3 2.6
                                               3 1
## 4 0.6
                           2
                                  3 2.66
              50
                     0
                                              3 1
## 5 1.2
                           2
                                  3 2.68
              50
                     0
                                              4 1
## 6 2.5
              50
                     0
                           2
                                  3
                                     2.8
                                              4 0.75
## 7 5
              50
                     0
                                  3 2.68
                           1
                                               4 1
```

Kruskal-Wallis test illustrates evidence of statistical differences.
kruskal.test(uni\_str\_pos ~ Sigma, data = act\_coverage)

```
##
## Kruskal-Wallis rank sum test
##
## data: uni str pos by Sigma
## Kruskal-Wallis chi-squared = 98.878, df = 6, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = act_coverage$uni_str_pos, g = act_coverage$Sigma, p.adjust.method = "bor
                     paired = FALSE, conf.int = FALSE, alternative = '1')
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
##
## data: act_coverage$uni_str_pos and act_coverage$Sigma
##
               0.1 0.3 0.6 1.2 2.5
##
## 0.1 3.9e-13 -
## 0.3 6.5e-11 1
## 0.6 1.8e-10 1
                   1
```

#### 5.5.3 Performance over time

1

1

1

1

1

1

## P value adjustment method: bonferroni

## 1.2 3.4e-09 1

## 2.5 2.3e-08 1

## 5 2.1e-09 1

##

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'mpe') %>%
group_by(Sigma, gen) %>%
dplyr::summarise(
  min = min(pop_fit_max) / DIMENSIONALITY,
  mean = mean(pop_fit_max) / DIMENSIONALITY,
```

```
max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = S
 geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom line(size = 0.5) +
 geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale_y_continuous(
   name="Average trait score"
  ) +
  scale_x_continuous(
   name="Generations",
   limits=c(0, 50000),
   breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
 ) +
 scale_shape_manual(values=SHAPE)+
 scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
 ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
 guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
   fill=guide_legend(nrow=2, title.position = "bottom")
```

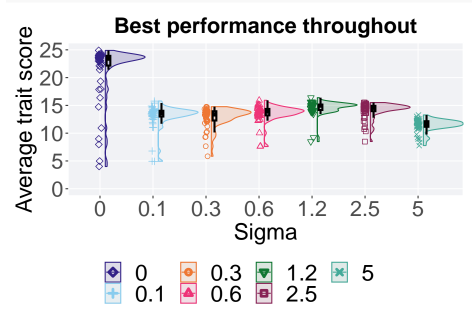


#### 5.5.4 Best performance throughout

Best performance reached throughout 50,000 generations in a population.

```
plot = filter(best_df, var == 'pop_fit_max' & acro == 'mpe') %>%
  ggplot(., aes(x = Sigma, y = val / DIMENSIONALITY, color = Sigma, fill = Sigma, shape = Sigma)
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Average trait score",
   limits=c(0, 25)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



#### 5.5.4.1 Stats

Summary statistics for the best performance.

Sigma count na\_cnt

```
performance = filter(best_df, var == 'pop_fit_max' & acro == 'mpe')
performance$Sigma = factor(performance$Sigma, levels = c('0','2.5','1.2','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.3','0.6','0.6','0.3','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6
```

max

IQR

min median mean

## P value adjustment method: bonferroni

```
<fct> <int> <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
## 1 0
              50
                      0 4.00
                                23.4 21.3 24.9 1.74
              50
                      0 8.50
## 2 2.5
                                14.6 14.1 15.5 0.949
              50
## 3 1.2
                      0 8.49
                                15.0 14.6 16.4 0.967
## 4 0.6
                      0 7.60
                                13.9 13.6 15.9 1.15
              50
## 5 0.3
              50
                      0 5.85
                                13.4 12.8 14.8 1.66
## 6 0.1
              50
                      0 4.95
                                13.6 13.1 15.8 1.03
## 7 5
              50
                      0 7.77
                                11.6 11.4 13.3 0.997
Kruskal-Wallis test illustrates evidence of statistical differences.
```

kruskal.test(val ~ Sigma, data = performance) ## ## Kruskal-Wallis rank sum test ## ## data: val by Sigma ## Kruskal-Wallis chi-squared = 183.09, df = 6, p-value < 2.2e-16 Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction. pairwise.wilcox.test(x = performance\$val, g = performance\$Sigma, p.adjust.method = "bonferroni", paired = FALSE, conf.int = FALSE, alternative = '1') ## ## Pairwise comparisons using Wilcoxon rank sum test with continuity correction ## ## data: performance\$val and performance\$Sigma ## ## 2.5 0 1.2 0.6 0.3 0.1 ## 2.5 1.1e-10 -## 1.2 2.2e-10 1.00000 -## 0.6 5.6e-11 0.01153 5.2e-06 -## 0.3 3.8e-11 1.9e-06 2.7e-10 0.07612 -## 0.1 4.2e-11 0.00013 1.7e-08 0.85825 1.00000 -3.3e-11 6.6e-12 1.7e-14 1.4e-12 1.1e-07 5.2e-10 ##

## Chapter 6

# Nondominated sorting

Results for the nondominated sorting parameter sweep on the diagnostics with no valleys.

#### 6.1 Data setup

```
over_time_df <- read.csv(paste(DATA_DIR,'OVER-TIME/nds.csv', sep = "", collapse = NULL), header =
over_time_df$Sigma <- factor(over_time_df$Sigma, levels = ND_LIST)

best_df <- read.csv(paste(DATA_DIR,'BEST/nds.csv', sep = "", collapse = NULL), header = TRUE, str
best_df$Sigma <- factor(best_df$Sigma, levels = ND_LIST)

sati_df <- read.csv(paste(DATA_DIR,'SOL-FND/nds.csv', sep = "", collapse = NULL), header = TRUE,
sati_df$Sigma <- factor(sati_df$Sigma, levels = ND_LIST)</pre>
```

## 6.2 Exploitation rate results

Here we present the results for **best performances** found by each selection scheme parameter on the exploitation rate diagnostic. 50 replicates are conducted for each scheme explored.

#### 6.2.1 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'exp') %>%
group_by(Sigma, gen) %>%
```

```
dplyr::summarise(
   min = min(pop_fit_max) / DIMENSIONALITY,
   mean = mean(pop_fit_max) / DIMENSIONALITY,
   max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
over_time_plot = ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color =
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
 scale_y_continuous(
   name="Average trait score"
 ) +
  scale x continuous(
   name="Generations",
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
 ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
 scale_fill_manual(values = cb_palette) +
 ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
 guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
    fill=guide_legend(nrow=2, title.position = "bottom")
over_time_plot
```

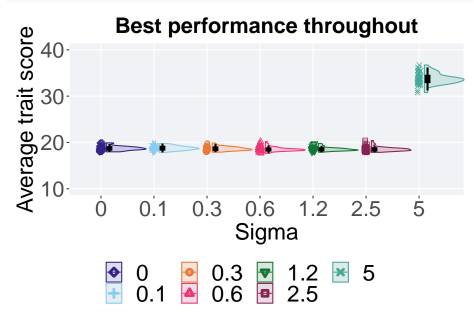


#### 6.2.2 Best performance throughout

Best performance reached throughout 50,000 generations in a population.

```
plot = filter(best_df, var == 'pop_fit_max' & acro == 'exp') %>%
  ggplot(., aes(x = Sigma, y = val / DIMENSIONALITY, color = Sigma, fill = Sigma, shape = Sigma)
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Average trait score",
   limits = c(10,40)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



#### **6.2.2.1** Stats

Summary statistics for the best performance.

```
performance = filter(best_df, var == 'pop_fit_max' & acro == 'exp')
performance %>%
  group_by(Sigma) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(val)),
    min = min(val / DIMENSIONALITY, na.rm = TRUE),
    median = median(val / DIMENSIONALITY, na.rm = TRUE),
    mean = mean(val / DIMENSIONALITY, na.rm = TRUE),
    max = max(val / DIMENSIONALITY, na.rm = TRUE),
    IQR = IQR(val / DIMENSIONALITY, na.rm = TRUE)
)
```

```
## # A tibble: 7 x 8
## Sigma count na_cnt min median mean max IQR
## <fct> <int> <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> > 0 18.0 18.7 18.7 19.9 0.535
```

```
## 2 0.1
              50
                        17.8
                                18.7
                                      18.8 19.8 0.613
## 3 0.3
              50
                         17.9
                                18.6
                                      18.7
                                            19.9 0.581
## 4 0.6
              50
                        17.7
                                18.5
                                      18.5
                                            20.2 0.442
## 5 1.2
              50
                        17.9
                                18.5 18.6
                                            19.8 0.428
## 6 2.5
              50
                      0
                        17.9
                                18.5 18.6
                                            20.4 0.480
## 7 5
              50
                         30.8
                                33.7
                                      33.8 36.7 1.35
```

Kruskal–Wallis test illustrates evidence of statistical differences.

```
kruskal.test(val ~ Sigma, data = performance)
##
##
   Kruskal-Wallis rank sum test
##
## data: val by Sigma
## Kruskal-Wallis chi-squared = 139.66, df = 6, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = performance$val, g = performance$Sigma, p.adjust.method = "bonferroni",
                     paired = FALSE, conf.int = FALSE, alternative = 'g')
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: performance$val and performance$Sigma
##
##
       0
              0.1
                     0.3
                                           2.5
                            0.6
                                    1.2
## 0.1 1
## 0.3 1
## 0.6 1
              1
                     1
## 1.2 1
                     1
                             1
## 2.5 1
                     1
                            1
                                    1
              1
## 5
       <2e-16 <2e-16 <2e-16 <2e-16 <2e-16
##
## P value adjustment method: bonferroni
```

## 6.3 Ordered exploitation results

Here we present the results for **best performances** found by each selection scheme parameter on the exploitation rate diagnostic. 50 replicates are conducted for each scheme explored.

#### 6.3.1 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'ord') %>%
group_by(Sigma, gen) %>%
dplyr::summarise(
  min = min(pop_fit_max) / DIMENSIONALITY,
  mean = mean(pop_fit_max) / DIMENSIONALITY,
  max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = S
 geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom line(size = 0.5) +
 geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale_y_continuous(
   name="Average trait score"
  ) +
  scale_x_continuous(
   name="Generations",
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
 ) +
 scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
   fill=guide_legend(nrow=2, title.position = "bottom")
```

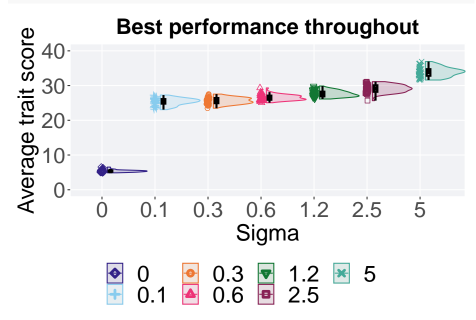


#### 6.3.2 Best performance throughout

Best performance reached throughout 50,000 generations in a population.

```
plot = filter(best_df, var == 'pop_fit_max' & acro == 'ord') %>%
  ggplot(., aes(x = Sigma, y = val / DIMENSIONALITY, color = Sigma, fill = Sigma, shape = Sigma)
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Average trait score",
   limits = c(0,40)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



#### 6.3.2.1 Stats

Summary statistics for the best performance.

```
performance = filter(best_df, var == 'pop_fit_max' & acro == 'ord')
performance %>%
  group_by(Sigma) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(val)),
    min = min(val / DIMENSIONALITY, na.rm = TRUE),
    median = median(val / DIMENSIONALITY, na.rm = TRUE),
    mean = mean(val / DIMENSIONALITY, na.rm = TRUE),
    max = max(val / DIMENSIONALITY, na.rm = TRUE),
    IQR = IQR(val / DIMENSIONALITY, na.rm = TRUE)
)
```

```
## # A tibble: 7 x 8
## Sigma count na_cnt min median mean max IQR
## <fct> <int> <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> =# 6.52 0.316
```

```
## 2 0.1
              50
                      0 23.1
                               25.5
                                     25.4
                                           27.3 1.31
## 3 0.3
              50
                      0 23.5
                                           27.5
                               25.6
                                     25.6
                                                 1.48
## 4 0.6
              50
                      0 25.1
                               26.5
                                     26.6
                                           29.3 1.18
## 5 1.2
              50
                      0 26.1
                               27.6 27.7
                                           29.9 1.34
## 6 2.5
              50
                      0 25.7
                               29.1 29.1
                                          31.1 1.87
## 7 5
              50
                      0 31.6
                               33.9 33.9 36.9
                                                1.89
```

Kruskal–Wallis test illustrates evidence of statistical differences.

```
kruskal.test(val ~ Sigma, data = performance)
##
##
   Kruskal-Wallis rank sum test
##
## data: val by Sigma
## Kruskal-Wallis chi-squared = 307.06, df = 6, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = performance$val, g = performance$Sigma, p.adjust.method = "bonferroni",
                     paired = FALSE, conf.int = FALSE, alternative = 'g')
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: performance$val and performance$Sigma
##
##
               0.1
                       0.3
                                0.6
                                        1.2
                                                2.5
       0
## 0.1 < 2e-16 -
## 0.3 < 2e-16 1
## 0.6 < 2e-16 3.6e-07 7.9e-05 -
## 1.2 < 2e-16 5.9e-15 8.1e-14 3.1e-07 -
## 2.5 < 2e-16 2.8e-16 3.5e-16 1.8e-14 3.3e-08 -
       < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16
##
## P value adjustment method: bonferroni
```

## 6.4 Contradictory objectives results

Here we present the results for activation gene coverage and satisfactory trait coverage found by each selection scheme parameter on the contradictory objectives diagnostic. 50 replicates are conducted for each scheme parameters explored.

#### 6.4.1 Activation gene coverage over time

Activation gene coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations.

Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'con') %>%
group_by(Sigma, gen) %>%
dplyr::summarise(
  min = min(uni_str_pos),
  mean = mean(uni_str_pos),
  max = max(uni_str_pos)
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = S
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks = seq(0, 100, 20),
   labels=c("0", "20", "40", "60", "80", "100")
  ) +
  scale_x_continuous(
    name="Generations".
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene coverage over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
    fill=guide_legend(nrow=2, title.position = "bottom")
```

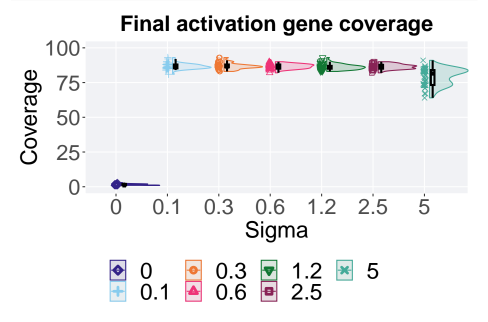


#### 6.4.2 Final activation gene coverage

Activation gene coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'con') %>%
  ggplot(., aes(x = Sigma, y = uni_str_pos, color = Sigma, fill = Sigma, shape = Sigma)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final activation gene coverage')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



#### 6.4.2.1 Stats

Summary statistics for the generation a satisfactory solution is found.

```
act_coverage = filter(over_time_df, gen == 50000 & acro == 'con')
act_coverage$Sigma = factor(act_coverage$Sigma, levels = c('2.5','1.2','0.6','0.3','0.
act_coverage %>%
    group_by(Sigma) %>%
    dplyr::summarise(
        count = n(),
        na_cnt = sum(is.na(uni_str_pos)),
        min = min(uni_str_pos, na.rm = TRUE),
        median = median(uni_str_pos, na.rm = TRUE),
        mean = mean(uni_str_pos, na.rm = TRUE),
        max = max(uni_str_pos, na.rm = TRUE),
        IQR = IQR(uni_str_pos, na.rm = TRUE)
)
```

```
## # A tibble: 7 x 8
## Sigma count na_cnt min median mean max IQR
## <fct> <int> <int> <int> <dbl> <dbl> <int> <dbl>
```

```
## 1 2.5
              50
                       0
                            82
                                 87
                                       86.5
                                                90 2.75
## 2 1.2
              50
                                       86.3
                       0
                            83
                                 86
                                                93
                                                    2
## 3 0.6
              50
                            82
                                 87
                                       86.6
                                                90 3
                       0
## 4 0.3
              50
                       0
                            83
                                 86.5 86.8
                                                93 2
## 5 0.1
              50
                       0
                            81
                                 86
                                       86.7
                                                93 2.75
## 6 5
              50
                       0
                            64
                                 81.5 79.2
                                                91 10.8
## 7 0
              50
                             1
                       0
                                  1
                                        1.44
                                                 3 1
```

Kruskal-Wallis test illustrates evidence of statistical differences.
kruskal.test(uni\_str\_pos ~ Sigma, data = act\_coverage)

```
##
## Kruskal-Wallis rank sum test
##
## data: uni_str_pos by Sigma
## Kruskal-Wallis chi-squared = 193.36, df = 6, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = act_coverage$uni_str_pos, g = act_coverage$Sigma, p.adjust.method = "bor paired = FALSE, conf.int = FALSE, alternative = 'l')
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##</pre>
```

```
## data: act_coverage$uni_str_pos and act_coverage$Sigma
##
##
       2.5
               1.2
                       0.6
                                0.3
                                        0.1
                                                5
## 1.2 1
               _
## 0.6 1
               1
## 0.3 1
               1
                       1
## 0.1 1
               1
                       1
                                1
       1.9e-11 9.3e-11 4.9e-12 1.2e-11 3.4e-12 -
## 5
## 0
       < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16
##
## P value adjustment method: bonferroni
```

#### 6.4.3 Satisfactory trait coverage over time

Satisfactory trait coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations. Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'con') %>%
group_by(Sigma, gen) %>%
dplyr::summarise(
  min = min(pop_uni_obj),
  mean = mean(pop_uni_obj),
```

```
max = max(pop_uni_obj)
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = S
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
 scale_y_continuous(
    name="Coverage",
   limits=c(0, 100)
  ) +
  scale_x_continuous(
   name="Generations",
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
 ) +
 scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Satisfactory trait coverage over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
   fill=guide_legend(nrow=2, title.position = "bottom")
```

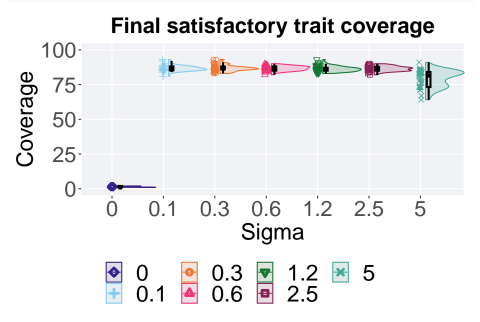


#### 6.4.4 Final satisfactory trait coverage

Satisfactory trait coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'con') %>%
  ggplot(., aes(x = Sigma, y = pop_uni_obj, color = Sigma, fill = Sigma, shape = Sigma)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final satisfactory trait coverage')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



#### 6.4.4.1 Stats

Summary statistics for the generation a satisfactory solution is found.

```
sat_coverage = filter(over_time_df, gen == 50000 & acro == 'con')
sat_coverage$Sigma = factor(sat_coverage$Sigma, levels = c('0.1','0.3','0.6','1.2','2.')
sat_coverage %>%
group_by(Sigma) %>%
dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(pop_uni_obj)),
    min = min(pop_uni_obj, na.rm = TRUE),
    median = median(pop_uni_obj, na.rm = TRUE),
    mean = mean(pop_uni_obj, na.rm = TRUE),
    max = max(pop_uni_obj, na.rm = TRUE),
    IQR = IQR(pop_uni_obj, na.rm = TRUE)
)
```

```
## # A tibble: 7 x 8
## Sigma count na_cnt min median mean max IQR
## <fct> <int> <int> <int> <dbl> <dbl> <int> <dbl>
```

```
## 1 0.1
               50
                       0
                             81
                                  86
                                        86.7
                                                      2.75
                                                 93
## 2 0.3
               50
                                                      2
                        0
                             83
                                  86.5 86.8
                                                 93
## 3 0.6
               50
                                  87
                                        86.6
                                                 90
                                                      3
                       0
                             82
## 4 1.2
               50
                       0
                             83
                                  86
                                        86.3
                                                 93
                                                      2
## 5 2.5
               50
                       0
                             82
                                  87
                                        86.5
                                                 90 2.75
## 6 5
               50
                       0
                             64
                                  81.5 79.2
                                                 91 10.8
## 7 0
               50
                              1
                                         1.38
                       0
                                   1
                                                   2 1
```

Kruskal-Wallis test illustrates evidence of statistical differences.

```
kruskal.test(pop_uni_obj ~ Sigma, data = sat_coverage)
##
##
    Kruskal-Wallis rank sum test
##
## data: pop_uni_obj by Sigma
## Kruskal-Wallis chi-squared = 193.38, df = 6, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = sat_coverage$pop_uni_obj, g = sat_coverage$Sigma, p.adjust.method = "bor
                     paired = FALSE, conf.int = FALSE, alternative = '1')
##
    Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
##
## data: sat_coverage$pop_uni_obj and sat_coverage$Sigma
##
##
       0.1
               0.3
                        0.6
                                1.2
                                        2.5
                                                 5
## 0.3 1
## 0.6 1
               1
## 1.2 1
                        1
## 2.5 1
                                1
               1
                        1
## 5
       3.4e-12 1.2e-11 4.9e-12 9.3e-11 1.9e-11 -
## 0
       < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16
##
```

## 6.5 Multi-path exploration results

## P value adjustment method: bonferroni

Here we present the results for **best performances** and **activation gene coverage** found by each selection scheme parameter on the multi-path exploration diagnostic. 50 replicates are conducted for each scheme parameter explored.

#### 6.5.1 Activation gene coverage over time

Activation gene coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations.

Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'mpe') %>%
group_by(Sigma, gen) %>%
dplyr::summarise(
   min = min(uni_str_pos),
   mean = mean(uni_str_pos),
   max = max(uni_str_pos)
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = S
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks = seq(0, 100, 20),
   labels=c("0", "20", "40", "60", "80", "100")
  ) +
  scale_x_continuous(
    name="Generations".
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene coverage over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
    fill=guide_legend(nrow=2, title.position = "bottom")
```

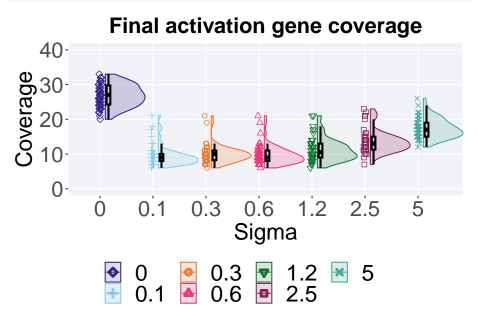


#### 6.5.2 Final activation gene coverage

Activation gene coverage found in the final population at 50,000 generations.

```
plot = filter(over time df, gen == 50000 & acro == 'mpe') %>%
  ggplot(., aes(x = Sigma, y = uni_str_pos, color = Sigma, fill = Sigma, shape = Sigma)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 40)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final activation gene coverage')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



#### 6.5.2.1 Stats

Sigma count na\_cnt

Summary statistics for activation gene coverage found in the final population at 50,000 generations.

min median mean

```
act_coverage = filter(over_time_df, gen == 50000 & acro == 'mpe')
act_coverage$Sigma = factor(act_coverage$Sigma, levels = c('0','5','2.5','1.2','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6','0.6',
```

max

IQR

```
<fct> <int>
                   <int> <int>
                                 <dbl> <dbl> <int> <dbl>
## 1 0
              50
                       0
                            20
                                 27
                                       26.8
                                                33 5.5
## 2 5
              50
                            12
                                       17.4
                                                26 4
                       0
                                 17
## 3 2.5
              50
                       0
                             7
                                 13
                                       13.6
                                                23 3.75
## 4 1.2
              50
                       0
                             6
                                 10.5 11.3
                                                21 4
## 5 0.6
              50
                       0
                             6
                                  9
                                        9.76
                                                21
                                                    3
## 6 0.3
                             6
              50
                       0
                                 10
                                        9.94
                                                21 2.75
## 7 0.1
              50
                       0
                             6
                                   9
                                        9.48
                                                21 2
```

Kruskal-Wallis test illustrates evidence of statistical differences.
kruskal.test(uni\_str\_pos ~ Sigma, data = act\_coverage)

```
##
##
   Kruskal-Wallis rank sum test
##
## data: uni_str_pos by Sigma
## Kruskal-Wallis chi-squared = 232.29, df = 6, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = act_coverage$uni_str_pos, g = act_coverage$Sigma, p.adjust.method = "bor
                     paired = FALSE, conf.int = FALSE, alternative = '1')
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: act_coverage$uni_str_pos and act_coverage$Sigma
##
##
                       2.5
       0
               5
                                1.2
                                       0.6
                                              0.3
## 5
       9.5e-16 -
## 2.5 < 2e-16 5.7e-08 -
```

#### 6.5.3 Performance over time

## 1.2 < 2e-16 8.9e-12 0.0013

##

## 0.6 < 2e-16 2.6e-14 5.3e-09 0.0582 -

## P value adjustment method: bonferroni

## 0.3 < 2e-16 8.7e-15 2.3e-09 0.2651 1.0000 - ## 0.1 < 2e-16 4.4e-14 1.1e-09 0.0062 1.0000 0.7100

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'mpe') %>%
group_by(Sigma, gen) %>%
dplyr::summarise(
   min = min(pop_fit_max) / DIMENSIONALITY,
```

```
mean = mean(pop_fit_max) / DIMENSIONALITY,
max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'Sigma'. You can override using the
## `.groups` argument.
ggplot(lines, aes(x=gen, y=mean, group = Sigma, fill = Sigma, color = Sigma, shape = S
 geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom line(size = 0.5) +
 geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale_y_continuous(
   name="Average trait score"
  ) +
  scale_x_continuous(
   name="Generations",
   limits=c(0, 50000),
   breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
 ) +
 scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
 guides(
    shape=guide_legend(nrow=2, title.position = "bottom"),
    color=guide_legend(nrow=2, title.position = "bottom"),
   fill=guide_legend(nrow=2, title.position = "bottom")
```

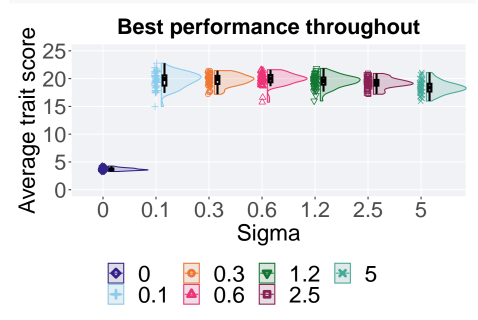


#### 6.5.4 Best performance throughout

Best performance reached throughout 50,000 generations in a population.

```
plot = filter(best_df, var == 'pop_fit_max' & acro == 'mpe') %>%
  ggplot(., aes(x = Sigma, y = val / DIMENSIONALITY, color = Sigma, fill = Sigma, shape = Sigma)
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Average trait score",
   limits=c(0, 25)
  scale_x_discrete(
   name="Sigma"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



#### 6.5.4.1 Stats

Summary statistics for the best performance.

Sigma count na\_cnt

```
performance = filter(best_df, var == 'pop_fit_max' & acro == 'mpe')
performance$Sigma = factor(performance$Sigma, levels = rev(c('0','5','2.5','1.2','0.6')

performance %>%
    group_by(Sigma) %>%
    dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(val)),
    min = min(val / DIMENSIONALITY, na.rm = TRUE),
    median = median(val / DIMENSIONALITY, na.rm = TRUE),
    mean = mean(val / DIMENSIONALITY, na.rm = TRUE),
    max = max(val / DIMENSIONALITY, na.rm = TRUE),
    IQR = IQR(val / DIMENSIONALITY, na.rm = TRUE)
)

## # A tibble: 7 x 8
```

max

IQR

min median mean

## P value adjustment method: bonferroni

```
<fct> <int> <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
## 1 0.1
             50
                     0 15.0
                             19.9 19.8 22.8 1.82
             50
## 2 0.3
                     0 17.2
                             19.9 19.7 21.4 1.43
## 3 0.6
             50
                     0 15.8
                             20.0 19.9 21.6 1.30
                             19.6 19.5 21.8 1.27
## 4 1.2
             50
                    0 16.0
## 5 2.5
             50
                    0 17.2
                             19.2 19.2 20.9 1.03
## 6 5
             50
                     0 16.0
                             18.4 18.4 21.1 1.38
## 7 0
                             3.63 3.69 4.26 0.307
             50
                     0 3.28
```

Kruskal-Wallis test illustrates evidence of statistical differences.

```
kruskal.test(val ~ Sigma, data = performance)
##
## Kruskal-Wallis rank sum test
##
## data: val by Sigma
## Kruskal-Wallis chi-squared = 166, df = 6, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = performance$val, g = performance$Sigma, p.adjust.method = "bonferroni",
                     paired = FALSE, conf.int = FALSE, alternative = '1')
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: performance$val and performance$Sigma
##
##
               0.3
       0.1
                       0.6
                               1.2
                                       2.5
                                               5
## 0.3 1.0000 -
## 0.6 1.0000 1.0000
## 1.2 1.0000 1.0000 0.7056
## 2.5 0.1012 0.0631 0.0019 0.6949
      1.8e-05 2.4e-06 6.1e-08 6.1e-05 0.0057
## 0
       < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16 < 2e-16
##
```

## Chapter 7

# Novelty search

Results for the novelty search parameter sweep on the diagnostics with no valleys.

### 7.1 Data setup

```
over_time_df <- read.csv(paste(DATA_DIR,'OVER-TIME/nov.csv', sep = "", collapse = NULL), header =
over_time_df$uni_str_pos = over_time_df$uni_str_pos + over_time_df$arc_acti_gene - over_time_df$c
over_time_df$K <- factor(over_time_df$K, levels = NS_LIST)

best_df <- read.csv(paste(DATA_DIR,'BEST/nov.csv', sep = "", collapse = NULL), header = TRUE, str
best_df$K <- factor(best_df$K, levels = NS_LIST)</pre>
```

## 7.2 Exploitation rate results

Here we present the results for **best performances** found by each selection scheme parameter on the exploitation rate diagnostic. 50 replicates are conducted for each scheme explored.

#### 7.2.1 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'exp') %>%
group_by(K, gen) %>%
dplyr::summarise(
  min = min(pop_fit_max) / DIMENSIONALITY,
  mean = mean(pop_fit_max) / DIMENSIONALITY,
```

over\_time\_plot

```
max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'K'. You can override using the `.groups`
## argument.
over_time_plot = ggplot(lines, aes(x=gen, y=mean, group = K, fill = K, color = K, shape
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
 scale_y_continuous(
   name="Average trait score"
 ) +
  scale_x_continuous(
   name="Generations",
   limits=c(0, 50000),
   breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
 ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
 scale_fill_manual(values = cb_palette) +
 ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
 guides(
    shape=guide_legend(nrow=1, title.position = "bottom"),
    color=guide_legend(nrow=1, title.position = "bottom"),
    fill=guide_legend(nrow=1, title.position = "bottom")
```

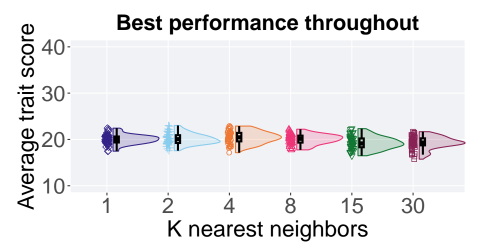


#### 7.2.2 Best performance throughout

Best performance reached throughout 50,000 generations in a population.

```
plot = filter(best_df, var == 'pop_fit_max' & acro == 'exp') %>%
  ggplot(., aes(x = K, y = val / DIMENSIONALITY, color = K, fill = K, shape = K)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Average trait score",
   limits = c(10,40)
  scale_x_discrete(
   name="K nearest neighbors"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



◆ 1 ★ 2 ◆ 4 ★ 8 ▼ 15 ★ 30

#### 7.2.2.1 Stats

Summary statistics for the best performance.

```
performance = filter(best_df, var == 'pop_fit_max' & acro == 'exp')
performance$K = factor(performance$K, levels = c('1','2','4','8','30','15'))
performance %>%
  group_by(K) %>%
  dplyr::summarise(
    count = n(),
   na_cnt = sum(is.na(val)),
   min = min(val / DIMENSIONALITY, na.rm = TRUE),
   median = median(val / DIMENSIONALITY, na.rm = TRUE),
   mean = mean(val / DIMENSIONALITY, na.rm = TRUE),
    max = max(val / DIMENSIONALITY, na.rm = TRUE),
    IQR = IQR(val / DIMENSIONALITY, na.rm = TRUE)
  )
## # A tibble: 6 x 8
    K
##
          count na_cnt
                         min median mean
                                             max
                                                   IQR
```

```
##
     <fct> <int>
                  <int> <dbl>
                                <dbl> <dbl> <dbl> <dbl> <dbl>
## 1 1
                         17.5
              50
                      0
                                 20.0
                                       20.0
                                             22.4 1.29
## 2 2
                                 20.1
              50
                         17.6
                                       20.1
                                             23.0 1.68
## 3 4
              50
                         17.1
                                 20.5
                                       20.4
                                             22.9 1.83
## 4 8
              50
                      0
                         17.7
                                 20.1
                                       20.0
                                             22.2 1.50
## 5 30
              50
                      0
                         15.7
                                 19.3
                                       19.3
                                             21.7
                                                   1.51
## 6 15
              50
                         16.5
                                       19.3
                                             22.3 1.90
                                 19.1
```

Kruskal-Wallis test illustrates evidence of statistical differences.

```
kruskal.test(val ~ K, data = performance)
##
##
   Kruskal-Wallis rank sum test
##
## data: val by K
## Kruskal-Wallis chi-squared = 28.774, df = 5, p-value = 2.568e-05
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = performance$val, g = performance$K, p.adjust.method = "bonferroni",
                     paired = FALSE, conf.int = FALSE, alternative = '1')
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
##
## data:
          performance$val and performance$K
##
##
                               8
                                       30
      1
## 2 1.00000 -
     1.00000 1.00000 -
## 8 1.00000 1.00000 0.98993 -
## 30 0.04143 0.03971 0.00158 0.04701 -
## 15 0.01794 0.01297 0.00089 0.02200 1.00000
##
## P value adjustment method: bonferroni
```

## 7.3 Ordered exploitation results

Here we present the results for **best performances** found by each selection scheme parameter on the exploitation rate diagnostic. 50 replicates are conducted for each scheme explored.

#### 7.3.1 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'ord') %>%
group_by(K, gen) %>%
dplyr::summarise(
  min = min(pop_fit_max) / DIMENSIONALITY,
  mean = mean(pop_fit_max) / DIMENSIONALITY,
  max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'K'. You can override using the `.groups`
## argument.
ggplot(lines, aes(x=gen, y=mean, group = K, fill = K, color = K, shape = K)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom line(size = 0.5) +
 geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale y continuous(
   name="Average trait score"
  ) +
  scale_x_continuous(
   name="Generations".
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale shape manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
    shape=guide_legend(nrow=1, title.position = "bottom"),
    color=guide_legend(nrow=1, title.position = "bottom"),
   fill=guide_legend(nrow=1, title.position = "bottom")
```



#### 7.3.2 Best performance throughout

Best performance reached throughout 50,000 generations in a population.

```
plot = filter(best df, var == 'pop fit max' & acro == 'ord') %%
  ggplot(., aes(x = K, y = val / DIMENSIONALITY, color = K, fill = K, shape = K)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Average trait score",
   limits = c(0,10)
  scale_x_discrete(
   name="K nearest neighbors"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



🚱 1 🛨 2 💽 4 📤 8 🔻 15 🖪 30

#### 7.3.2.1 Stats

Summary statistics for the best performance.

```
performance = filter(best_df, var == 'pop_fit_max' & acro == 'ord')
performance$K = factor(performance$K, levels = c('1','2','4','8','15','30'))
performance %>%
  group_by(K) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(val)),
    min = min(val / DIMENSIONALITY, na.rm = TRUE),
    median = median(val / DIMENSIONALITY, na.rm = TRUE),
    mean = mean(val / DIMENSIONALITY, na.rm = TRUE),
    max = max(val / DIMENSIONALITY, na.rm = TRUE),
    IQR = IQR(val / DIMENSIONALITY, na.rm = TRUE)
)

### # A tibble: 6 x 8

### # A tibble: 6 x 8
```

## # A tibble: 6 x 8
## K count na\_cnt min median mean max IQR
## <fct> <int> <int> <dbl> <dbl> <dbl> <dbl> <dbl> </dbl>

```
## 1 1
              50
                      0
                         2.44
                                4.13 4.09 6.06 0.873
## 2 2
              50
                      0
                         3.19
                                4.85
                                      4.90
                                            6.24 0.832
## 3 4
              50
                         3.66
                                5.07
                                      5.16
                                            6.55 1.00
## 4 8
              50
                      0
                        3.13
                                4.71
                                      4.64
                                            6.42 1.08
## 5 15
              50
                      0
                         2.11
                                3.72
                                      3.82 5.64 0.710
## 6 30
              50
                         2.98
                                4.78 4.81 6.36 1.03
```

Kruskal–Wallis test illustrates evidence of statistical differences.

```
kruskal.test(val ~ K, data = performance)
##
##
   Kruskal-Wallis rank sum test
##
## data: val by K
## Kruskal-Wallis chi-squared = 91.122, df = 5, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = performance$val, g = performance$K, p.adjust.method = "bonferroni",
                     paired = FALSE, conf.int = FALSE, alternative = '1')
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: performance$val and performance$K
##
##
             2
                             8
                                     15
      1
## 2 1.0000 -
## 4 1.0000 1.0000
## 8 1.0000 1.0000 0.0084
## 15 0.3071 1.7e-09 5.2e-12 3.2e-06 -
## 30 1.0000 1.0000 0.1583 1.0000 1.0000
##
## P value adjustment method: bonferroni
```

## 7.4 Contradictory objectives results

Here we present the results for activation gene coverage and satisfactory trait coverage found by each selection scheme parameter on the contradictory objectives diagnostic. 50 replicates are conducted for each scheme parameters explored.

#### 7.4.1 Activation gene coverage over time

Activation gene coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations. Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'con') %>%
group_by(K, gen) %>%
dplyr::summarise(
  min = min(uni_str_pos),
  mean = mean(uni_str_pos),
  max = max(uni_str_pos)
)
```

```
## `summarise()` has grouped output by 'K'. You can override using the `.groups`
## argument.
ggplot(lines, aes(x=gen, y=mean, group = K, fill = K, color = K, shape = K)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen %% 2000 == 0 & gen != 0), size = 1.5, stroke = 2
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks=seq(0,100, 20),
   labels=c("0", "20", "40", "60", "80", "100")
  ) +
  scale_x_continuous(
    name="Generations".
   limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene coverage over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
    shape=guide_legend(nrow=1, title.position = "bottom"),
    color=guide_legend(nrow=1, title.position = "bottom"),
   fill=guide_legend(nrow=1, title.position = "bottom")
```



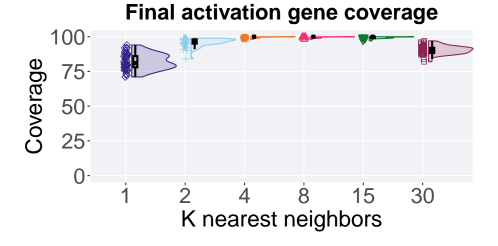
#### 7.4.2 Final activation gene coverage

Activation gene coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'con') %>%
  ggplot(., aes(x = K, y = uni_str_pos, color = K, fill = K, shape = K)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = 0.2, width
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1.0, position
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha = 1.0) +
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100)
  scale_x_discrete(
   name="K nearest neighbors"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final activation gene coverage')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```

## Warning: Removed 56 rows containing missing values (`geom\_point()`).





#### 7.4.2.1 Stats

Summary statistics for the generation a satisfactory solution is found.

```
act_coverage = filter(over_time_df, gen == 50000 & acro == 'con')
act_coverage$K = factor(act_coverage$K, levels = c('2.5','1.2','0.6','0.3','0.1','5','0.4','0.2','0.6','0.3','0.1','5','0.4','0.2','0.6','0.3','0.1','5','0.4','0.2','0.6','0.3','0.1','5','0.4','0.2','0.6','0.3','0.1','5','0.4','0.2','0.6','0.3','0.1','5','0.4','0.2','0.6','0.3','0.1','5','0.4','0.2','0.6','0.3','0.1','5','0.4','0.2','0.6','0.3','0.1','5','0.4','0.2','0.6','0.3','0.1','5','0.4','0.2','0.6','0.3','0.1','5','0.4','0.2','0.6','0.3','0.1','5','0.4','0.2','0.6','0.3','0.1','5','0.4','0.2','0.6','0.3','0.1','5','0.4','0.6','0.3','0.1','5','0.4','0.6','0.3','0.1','5','0.4','0.6','0.3','0.1','5','0.4','0.6','0.3','0.1','5','0.4','0.6','0.3','0.1','5','0.4','0.6','0.3','0.1','5','0.4','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0.1','0
```

## # A tibble: 1 x 8

```
##
                                                      IQR
           count na_cnt
                           min median mean
                                               max
##
     <fct> <int>
                  <int> <int>
                                <dbl> <dbl> <int> <dbl>
## 1 <NA>
             300
                            71
                                       94.6
                                               100
                       0
                                    98
```

Kruskal-Wallis test illustrates evidence of statistical differences.

```
\# kruskal.test(uni\_str\_pos \sim K, data = act\_coverage)
```

Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.

```
 \begin{tabular}{ll} \# \ pairwise.wilcox.test(x = act\_coverage\$uni\_str\_pos, \ g = act\_coverage\$K, \ p.adjust.method = "bonform paired = FALSE, \ conf.int = FALSE, \ alternative = 'l') \\ \end{tabular}
```

#### 7.4.3 Satisfactory trait coverage over time

Satisfactory trait coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations. Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'con') %>%
group_by(K, gen) %>%
dplyr::summarise(
  min = min(pop_uni_obj),
  mean = mean(pop_uni_obj),
  max = max(pop_uni_obj)
)
```

## `summarise()` has grouped output by 'K'. You can override using the `.groups`
## argument.

```
ggplot(lines, aes(x=gen, y=mean, group = K, fill = K, color = K, shape = K)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom_point(data = filter(lines, gen \( \frac{\psi}{k} \) 2000 == 0 \( \& \) gen != 0), size = 1.5, stroke = 2.0, alpha =
  scale_y_continuous(
    name="Coverage",
    limits=c(0, 100)
  scale_x_continuous(
    name="Generations",
    limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
    labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale colour manual(values = cb palette) +
  scale_fill_manual(values = cb_palette) +
```

```
ggtitle('Satisfactory trait coverage over time')+
p_theme + theme(legend.title=element_blank()) +
guides(
  shape=guide_legend(nrow=1, title.position = "bottom"),
  color=guide_legend(nrow=1, title.position = "bottom"),
  fill=guide_legend(nrow=1, title.position = "bottom")
```

# Satisfactory trait coverage over time



#### Final satisfactory trait coverage

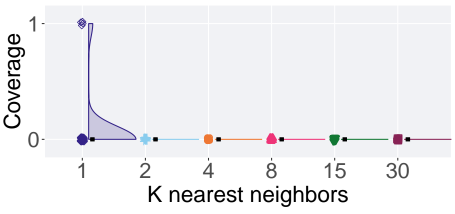
Satisfactory trait coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'con') %>%
  ggplot(., aes(x = K, y = pop_uni_obj, color = K, fill = K, shape = K)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = '
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha
  scale_y_continuous(
   name="Coverage",
   limits=c(-0.1, 1.1),
   breaks = c(0,1)
  scale_x_discrete(
    name="K nearest neighbors"
```

```
scale_shape_manual(values=SHAPE)+
scale_colour_manual(values = cb_palette, ) +
scale_fill_manual(values = cb_palette) +
ggtitle('Final satisfactory trait coverage')+
p_theme + theme(legend.title=element_blank())

plot_grid(
   plot +
        theme(legend.position="none"),
   legend,
   nrow=2,
   rel_heights = c(3,1)
)
```

# Final satisfactory trait coverage





#### 7.4.4.1 Stats

Summary statistics for the generation a satisfactory solution is found.

```
sat_coverage = filter(over_time_df, gen == 50000 & acro == 'con')
sat_coverage$K = factor(sat_coverage$K, levels = NS_LIST)
sat_coverage %>%
  group_by(K) %>%
  dplyr::summarise(
   count = n(),
   na_cnt = sum(is.na(pop_uni_obj)),
```

```
min = min(pop_uni_obj, na.rm = TRUE),
    median = median(pop_uni_obj, na.rm = TRUE),
    mean = mean(pop_uni_obj, na.rm = TRUE),
    max = max(pop_uni_obj, na.rm = TRUE),
    IQR = IQR(pop_uni_obj, na.rm = TRUE)
)
## # A tibble: 6 x 8
##
    K
           count na_cnt
                          min median mean
                                                    IQR
                                             max
    <fct> <int> <int> <dbl> <dbl> <int> <dbl>
## 1 1
             50
                      0
                            0
                                   0.08
                                               1
## 2 2
              50
                      0
                            0
                                   0 0
## 3 4
              50
                      0
                            0
                                   0 0
                                               0
                                                     0
## 4 8
              50
                      0
                            0
                                   0 0
                                               0
                                                     0
## 5 15
              50
                      0
                            0
                                   0 0
                                               0
                                                     0
## 6 30
              50
                                   0 0
                                               0
                                                      0
Kruskal–Wallis test illustrates evidence of statistical differences.
kruskal.test(pop_uni_obj ~ K, data = sat_coverage)
##
##
   Kruskal-Wallis rank sum test
##
## data: pop_uni_obj by K
## Kruskal-Wallis chi-squared = 20.203, df = 5, p-value = 0.001145
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = sat_coverage$pop_uni_obj, g = sat_coverage$K, p.adjust.method
                     paired = FALSE, conf.int = FALSE, alternative = 't')
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: sat_coverage$pop_uni_obj and sat_coverage$K
##
           2 4 8 15
##
      1
## 2 0.22 - - - -
## 4 0.22 - - - -
## 8 0.22 - - - -
## 15 0.22 - - - -
## 30 0.22 - - - -
```

## P value adjustment method: bonferroni

## 7.5 Multi-path exploration results

Here we present the results for **best performances** and **activation gene coverage** found by each selection scheme parameter on the multi-path exploration diagnostic. 50 replicates are conducted for each scheme parameter explored.

#### 7.5.1 Activation gene coverage over time

Activation gene coverage in a population over time. Data points on the graph is the average activation gene coverage across 50 replicates every 2000 generations. Shading comes from the best and worse coverage across 50 replicates.

```
lines = filter(over_time_df, acro == 'mpe') %>%
group_by(K, gen) %>%
dplyr::summarise(
  min = min(uni_str_pos),
  mean = mean(uni_str_pos),
  max = max(uni_str_pos)
)
```

## `summarise()` has grouped output by 'K'. You can override using the `.groups`
## argument.

```
ggplot(lines, aes(x=gen, y=mean, group = K, fill = K, color = K, shape = K)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(size = 0.5) +
  geom point(data = filter(lines, gen \( \frac{1}{2} \) 2000 == 0 \( \frac{1}{2} \) gen != 0), size = 1.5, stroke = 2.0, alpha =
  scale_y_continuous(
    name="Coverage",
    limits=c(0, 100),
    breaks=seq(0,100, 20),
    labels=c("0", "20", "40", "60", "80", "100")
  ) +
  scale_x_continuous(
    name="Generations",
    limits=c(0, 50000),
    breaks=c(0, 10000, 20000, 30000, 40000, 50000),
    labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale shape manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene coverage over time')+
  p theme + theme(legend.title=element blank()) +
  guides(
    shape=guide_legend(nrow=1, title.position = "bottom"),
```

```
color=guide_legend(nrow=1, title.position = "bottom"),
fill=guide_legend(nrow=1, title.position = "bottom")
)
```

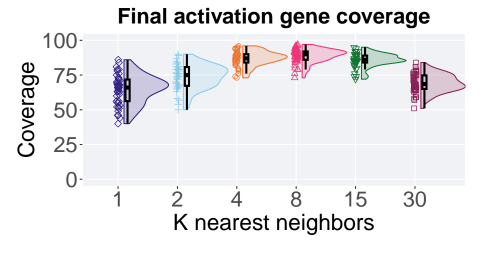


### 7.5.2 Final activation gene coverage

Activation gene coverage found in the final population at 50,000 generations.

```
plot = filter(over_time_df, gen == 50000 & acro == 'mpe') %>%
  ggplot(., aes(x = K, y = uni_str_pos, color = K, fill = K, shape = K)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = '
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha
  scale_y_continuous(
    name="Coverage",
    limits=c(0, 100)
  ) +
  scale_x_discrete(
    name="K nearest neighbors"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Final activation gene coverage')+
  p_theme + theme(legend.title=element_blank())
```

```
plot_grid(
  plot +
    theme(legend.position="none"),
  legend,
  nrow=2,
  rel_heights = c(3,1)
)
```





#### 7.5.2.1 Stats

Summary statistics for activation gene coverage found in the final population at 50,000 generations.

```
act_coverage = filter(over_time_df, gen == 50000 & acro == 'mpe')
act_coverage$K = factor(act_coverage$K, levels = c('15','8','4','2','30','1'))
act_coverage %>%
group_by(K) %>%
dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(uni_str_pos)),
    min = min(uni_str_pos, na.rm = TRUE),
    median = median(uni_str_pos, na.rm = TRUE),
    mean = mean(uni_str_pos, na.rm = TRUE),
    max = max(uni_str_pos, na.rm = TRUE),
    IQR = IQR(uni_str_pos, na.rm = TRUE)
```

```
## # A tibble: 6 x 8
##
     K
           count na_cnt
                                                     IQR
                           min median mean
                                               max
                                <dbl> <dbl> <int> <dbl>
##
     <fct> <int> <int> <int>
## 1 15
              50
                       0
                            72
                                  86
                                        85.9
                                                95
                                                    5
## 2 8
              50
                       0
                            73
                                 90
                                        88.8
                                                97
                                                    6
## 3 4
              50
                       0
                            73
                                 87
                                        87.0
                                                96 6
## 4 2
              50
                       0
                            50
                                 75
                                        73.8
                                                90 13.5
## 5 30
                       0
              50
                            51
                                 68.5
                                       69.1
                                                84 9.75
## 6 1
              50
                       0
                            40
                                  66
                                        64.7
                                                86 15.5
Kruskal-Wallis test illustrates evidence of statistical differences.
kruskal.test(uni_str_pos ~ K, data = act_coverage)
##
##
    Kruskal-Wallis rank sum test
##
## data: uni_str_pos by K
## Kruskal-Wallis chi-squared = 199.94, df = 5, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = act_coverage$uni_str_pos, g = act_coverage$K, p.adjust.method
                      paired = FALSE, conf.int = FALSE, alternative = '1')
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: act_coverage$uni_str_pos and act_coverage$K
##
##
                       4
                               2
                                        30
      15
              8
## 8 1.00000 -
     1.00000 0.38518 -
## 2 5.7e-11 2.8e-13 1.7e-11 -
## 30 2.2e-15 3.4e-16 1.2e-15 0.01313 -
## 1 2.5e-15 4.3e-16 1.4e-15 0.00016 0.32658
##
## P value adjustment method: bonferroni
```

### 7.5.3 Performance over time

Best performance in a population over time. Data points on the graph is the average performance across 50 replicates every 2000 generations. Shading comes from the best and worse performance across 50 replicates.

```
lines = filter(over_time_df, acro == 'mpe') %>%
group_by(K, gen) %>%
dplyr::summarise(
  min = min(pop_fit_max) / DIMENSIONALITY,
  mean = mean(pop_fit_max) / DIMENSIONALITY,
  max = max(pop_fit_max) / DIMENSIONALITY
)
```

```
## `summarise()` has grouped output by 'K'. You can override using the `.groups`
## argument.
ggplot(lines, aes(x=gen, y=mean, group = K, fill = K, color = K, shape = K)) +
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom line(size = 0.5) +
  geom_point(data = filter(lines, gen 1000 == 0 & gen != 0), size = 1.5, stroke = 2.0, alpha =
  scale_y_continuous(
   name="Average trait score"
  ) +
  scale_x_continuous(
   name="Generations".
   limits=c(0, 50000),
   breaks=c(0, 10000, 20000, 30000, 40000, 50000),
   labels=c("0e+4", "1e+4", "2e+4", "3e+4", "4e+4", "5e+4")
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Performance over time')+
 p_theme + theme(legend.title=element_blank()) +
  guides(
   shape=guide_legend(nrow=1, title.position = "bottom"),
    color=guide_legend(nrow=1, title.position = "bottom"),
   fill=guide_legend(nrow=1, title.position = "bottom")
```

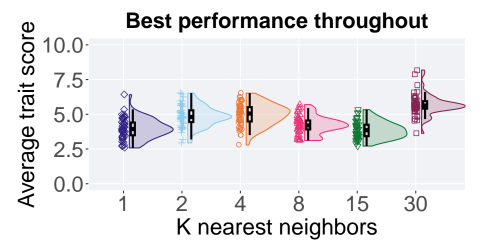


## 7.5.4 Best performance throughout

Best performance reached throughout 50,000 generations in a population.

```
plot = filter(best df, var == 'pop fit max' & acro == 'mpe') %>%
  ggplot(., aes(x = K, y = val / DIMENSIONALITY, color = K, fill = K, shape = K)) +
  geom_flat_violin(position = position_nudge(x = .1, y = 0), scale = 'width', alpha = '
  geom_boxplot(color = 'black', width = .07, outlier.shape = NA, alpha = 0.0, size = 1
  geom_point(position = position_jitter(width = 0.03, height = 0.02), size = 2.0, alpha
  scale_y_continuous(
    name="Average trait score",
    limits=c(0, 10)
  ) +
  scale_x_discrete(
    name="K nearest neighbors"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
  p_theme + theme(legend.title=element_blank())
plot_grid(
  plot +
    theme(legend.position="none"),
```

```
legend,
nrow=2,
rel_heights = c(3,1)
)
```



◆ 1 ★ 2 ◆ 4 ▲ 8 ▼ 15 ■ 30

#### 7.5.4.1 Stats

Summary statistics for the best performance.

```
performance = filter(best_df, var == 'pop_fit_max' & acro == 'mpe')
performance$K = factor(performance$K, levels = c('30','4','2','8','1','15'))

performance %>%
    group_by(K) %>%
    dplyr::summarise(
        count = n(),
        na_cnt = sum(is.na(val)),
        min = min(val / DIMENSIONALITY, na.rm = TRUE),
        median = median(val / DIMENSIONALITY, na.rm = TRUE),
        mean = mean(val / DIMENSIONALITY, na.rm = TRUE),
        max = max(val / DIMENSIONALITY, na.rm = TRUE)
        TQR = TQR(val / DIMENSIONALITY, na.rm = TRUE)
)
```

## # A tibble: 6 x 8
## K count na\_cnt min median mean max IQR

##

```
##
    <fct> <int> <int> <dbl>
                             <dbl> <dbl> <dbl> <dbl> <
## 1 30
             50
                     0 3.63
                              5.63 5.68 8.18 0.541
## 2 4
             50
                     0 2.79
                               5.06 4.97 6.52 1.03
## 3 2
             50
                     0 2.94
                               4.80 4.86
                                          6.53 0.846
## 4 8
                               4.20 4.24
             50
                     0 3.12
                                          5.70 0.638
## 5 1
             50
                     0 2.58
                               3.94 3.98
                                          6.41 0.916
## 6 15
             50
                     0 2.71
                               3.86 3.88 5.33 0.826
```

Kruskal–Wallis test illustrates evidence of statistical differences.

## 1 6.4e-14 3.0e-07 1.0e-06 0.41150 -

## P value adjustment method: bonferroni

## 15 2.1e-15 1.0e-08 2.9e-08 0.05326 1.00000

```
kruskal.test(val ~ K, data = performance)
##
## Kruskal-Wallis rank sum test
##
## data: val by K
## Kruskal-Wallis chi-squared = 134.32, df = 5, p-value < 2.2e-16
Results for post-hoc Wilcoxon rank-sum test with a Bonferroni correction.
pairwise.wilcox.test(x = performance$val, g = performance$K, p.adjust.method = "bonfer.")
                     paired = FALSE, conf.int = FALSE, alternative = '1')
##
##
   Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: performance$val and performance$K
##
##
      30
                      2
                                       1
## 4 0.00015 -
## 2 3.7e-06 1.00000 -
## 8 2.7e-13 5.7e-05 0.00031 -
```