Supplemental Material for 'Hidden Lexicase Selection Parameters: Varying Population Size and Test Case Redundancy with Diagnostic Metrics'

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	plot pane pane lege axis axis lege	<pre>ne <- theme(c.title = element_text(face = "bold", size = 16, hjust=0.5] el.border = element_blank(), el.grid.minor = element_blank(), end.title=element_text(size=16), end.text=element_text(size=16), s.title = element_text(size=16), s.text = element_text(size=16), end.position="bottom", el.background = element_rect(fill = "#f1f2f5",</pre>	
)			

Chapter 1

Introduction

This is not intended as a stand-alone document, but as a companion to our manuscript.

1.1 About our supplemental material

As you may have noticed (unless you're reading a pdf version of this), our supplemental material is hosted using GitHub pages. We compiled our data analyses and supplemental documentation into this nifty web-accessible book using bookdown.

The code used for this supplemental material can be found in this GitHub repository.

Our supplemental material includes the following:

- Exploitation rate results (Section 2)
- Contradictory rates results w/ 100 standard test cases (Section 3)
- Contradictory rates results w/ 50 redundant test cases (Section 4)
- Contradictory rates results w/ 100 redundant test cases (Section 5)
- Contradictory rates results w/ 200 redundant test cases (Section 6)
- Contradictory rates results w/ 300 redundant test cases (Section 7)

1.2 Contributing authors

- Jose Guadalupe Hernandez
- Anil Kumar Saini
- Jason H. Moore

Chapter 2

Exploitation rate results

Here we report the **performance** and evaluation a **satisfactory solution** was found on the exploitation rate diagnostic. 50 replicates were conducted for each population size explored. Performance is defined at the average trait performance, where we collect the best performing solution in each generation over time and the best performing solution evolved. A satisfactory solution is defined as a solution that has a phenotype with all traits greater than or equal to 99.0.

2.1 Analysis setup

```
library(cowplot)
library(dplyr)
library(PupillometryR)

# over time data
over_time <- read.csv("../Paper_Data/Exploitation/ot.csv", header = TRUE, stringsAsFactors = FALS
over_time$pop_size <- factor(over_time$pop_size, levels = NAMES)

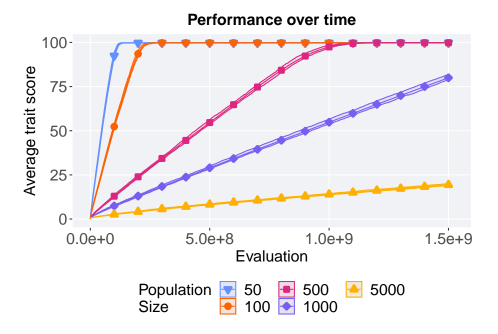
# best performance data
best <- read.csv('.../Paper_Data/Exploitation/best.csv', header = TRUE, stringsAsFactors = FALSE)
best$pop_size <- factor(best$pop_size, levels = NAMES)

# get the data
ssf <- read.csv('.../Paper_Data/Exploitation/ssf.csv', header = TRUE, stringsAsFactors = FALSE)
ssf$pop_size <- factor(ssf$pop_size, levels = NAMES)
ssf <- read.csv('.../Paper_Data/Exploitation/ssf.csv', header = TRUE, stringsAsFactors = FALSE)
ssf$pop_size <- factor(ssf$pop_size, levels = NAMES)
ssf <- filter(ssf, evaluation <= 1.5*10^9)</pre>
```

2.2 Performance over time

Performance of the best solution in the population at each generation over time.

```
# aggregate
lines = over_time %>%
  group_by(pop_size, eval) %>%
 dplyr::summarise(
   min = min(performance),
   mean = mean(performance),
   max = max(performance)
lines$pop_size <- factor(lines$pop_size, levels = NAMES)</pre>
ggplot(lines, aes(x=eval, y=mean, group = pop_size, fill = pop_size, color = pop_size,
 geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(linewidth = 0.5) +
  geom_point(data = filter(lines, eval %% 1000000000 == 0 & eval != 0), size = 1.0, str.
  scale y continuous(
   name="Average trait score",
   limits=c(0, 100),
   breaks=seq(0,100, 25),
   labels=c("0", "25", "50", "75", "100")
 ) +
  scale_x_continuous(
   name="Evaluation",
   labels = c('0.0e+0', '5.0e+8', '1.0e+9', '1.5e+9'),
   limits = c(0,1520000000)
  ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Performance over time')+
 p_theme +
  guides (
    shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```

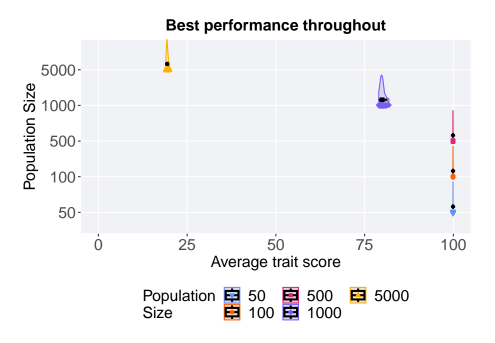


2.3 Best performance evolved

Performance of the best solution found throughout the entire evolutionary run.

```
ggplot(best, aes(x = pop_size, y = performance, color = pop_size, fill = pop_size, shape = pop_size
  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 = 0.02, height = 0.0001), size = 1.5, alpha = 1.0)
  scale_y_continuous(
   name="Average trait score",
   limits=c(0, 100),
   breaks=seq(0,100, 25),
   labels=c("0", "25", "50", "75", "100")
  ) +
  scale_x_discrete(
   name="Population Size"
  )+
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best performance throughout')+
  p_theme+ coord_flip() +
  guides(
```

```
shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
)
```



2.3.1 Summary statistics

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

```
## # A tibble: 5 x 8
## pop_size count na_cnt min median mean max IQR
## <fct> <int> <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
```

```
## 1 50
                50
                       0 99.9
                                 99.9 99.9 99.9 0.0149
## 2 100
                50
                          99.9
                                 99.9
                                       99.9 99.9 0.0166
## 3 500
                50
                       0 99.9
                                 99.9 99.9 99.9 0.0235
## 4 1000
                50
                       0 79.0
                                 79.9 80.0 81.8 0.779
## 5 5000
                50
                       0 19.0
                                 19.4 19.4 20.0 0.299
```

2.3.2 Kruskal-Wallis test

```
##
## Kruskal-Wallis rank sum test
##
## data: performance by pop_size
## Kruskal-Wallis chi-squared = 198.29, df = 4,
## p-value < 2.2e-16</pre>
```

2.3.3 Pairwise wilcoxon test

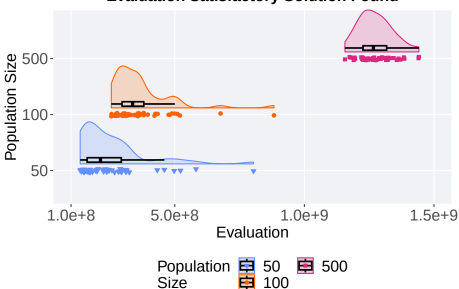
```
pairwise.wilcox.test(x = best$performance, g = best$pop_size, p.adjust.method = "bonferroni",
                     paired = FALSE, conf.int = FALSE, alternative = '1')
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: best$performance and best$pop_size
##
                                1000
##
        50
                100
                        500
## 100 1.00000 -
## 500 0.00078 0.00059 -
## 1000 < 2e-16 < 2e-16 < 2e-16 -
## 5000 < 2e-16 < 2e-16 < 2e-16 < 2e-16
##
## P value adjustment method: bonferroni
```

2.4 Evaluation satisfactory solution if found

Evaluation a satisfactory solution is found for each population size.

```
ggplot(ssf, aes(x = pop_size, y = evaluation, color = pop_size, fill = pop_size, shape
  geom_flat_violin(position = position_nudge(x = 0.12, y = 0), scale = 'width', alpha =
  geom_boxplot(color = 'black', width = .08, outlier.shape = NA, alpha = 0.0, size = 0
  geom_point(position = position_jitter(width = 0.03, height = 0.000001), size = 1.5,
  scale_y_continuous(
   name = 'Evaluation',
   labels = c('1.0e+8', '5.0e+8', '1.0e+9', '1.5e+9'),
   limits = c(100000000, 1500000000)
   ) +
 scale_x_discrete(
   name="Population Size",
 )+
  scale_shape_manual(values=SHAPE, )+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
 ggtitle(bquote('Evaluation Satisfactory Solution Found'))+
 p_theme + coord_flip() +
 guides(
    shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
   color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
   fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```

Evaluation Satisfactory Solution Found



2.4.1 Summary statistics

```
ssf %>%
 group_by(pop_size) %>%
 dplyr::summarise(
   count = n(),
   na_cnt = sum(is.na(evaluation)),
   min = min(evaluation, na.rm = TRUE),
   median = median(evaluation, na.rm = TRUE),
   mean = mean(evaluation, na.rm = TRUE),
   max = max(evaluation, na.rm = TRUE),
   IQR = IQR(evaluation, na.rm = TRUE)
## # A tibble: 3 x 8
   pop_size count na_cnt
                            min median
                                       mean
                                                max
             <int> <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
   <fct>
## 1 50
              50 0 1.35e8 2.13e8 2.52e8 8.04e8 1.32e8
## 2 100
              50
                       0 2.55e8 3.38e8 3.65e8 8.83e8 8.37e7
## 3 500
              50
                      0 1.16e9 1.27e9 1.28e9 1.44e9 9.10e7
```

2.4.2 Kruskal-Wallis test

```
##
##
Kruskal-Wallis rank sum test
##
## data: evaluation by pop_size
##
Kruskal-Wallis chi-squared = 113.38, df = 2,
## p-value < 2.2e-16</pre>
```

2.4.3 Pairwise wilcoxon test

```
##
## data: ssf$evaluation and ssf$pop_size
##
## 50 100
## 100 3.1e-08 -
## 500 < 2e-16 < 2e-16
##
## P value adjustment method: bonferroni</pre>
```

Chapter 3

Contradictory objectives 100 results

Here we report the activation gene coverage and satisfactory trait coverage was found on the contradictory objectives diagnostic. 50 replicates were conducted for each population size explored. Activation gene coverage is calculated by finding all the unique activation genes found within a given population. Satisfactory trait coverage is calculated by finding all the unique satisfactory traits found within a given population.

3.1 Analysis setup

```
library(ggplot2)
library(cowplot)
library(dplyr)
library(PupillometryR)

# over time data
over_time <- read.csv("../Paper_Data/Contradictory-100/ot.csv", header = TRUE, stringsAsFactors = over_time$pop_size <- factor(over_time$pop_size, levels = NAMES)

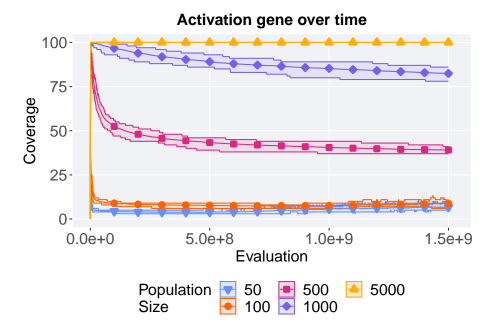
# best performance data
best <- read.csv('../Paper_Data/Contradictory-100/best.csv', header = TRUE, stringsAsFactors = FAbest$pop_size <- factor(best$pop_size, levels = NAMES)</pre>
```

3.2 Activation gene coverage

3.2.1 Coverage over time

Performance of the best solution in the population at each generation over time.

```
# aggregate
lines = over_time %>%
  group_by(pop_size, eval) %>%
 dplyr::summarise(
   min = min(activation_coverage),
   mean = mean(activation_coverage),
   max = max(activation_coverage)
lines$pop_size <- factor(lines$pop_size, levels = NAMES)</pre>
ggplot(lines, aes(x=eval, y=mean, group = pop_size, fill = pop_size, color = pop_size,
 geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(linewidth = 0.5) +
  geom_point(data = filter(lines, eval %% 1000000000 == 0 & eval != 0), size = 1.0, str.
  scale y continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks=seq(0,100, 25),
   labels=c("0", "25", "50", "75", "100")
  ) +
  scale_x_continuous(
   name="Evaluation",
   labels = c('0.0e+0', '5.0e+8', '1.0e+9', '1.5e+9'),
   limits = c(0,1520000000)
 ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene over time')+
 p_theme +
  guides (
    shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```



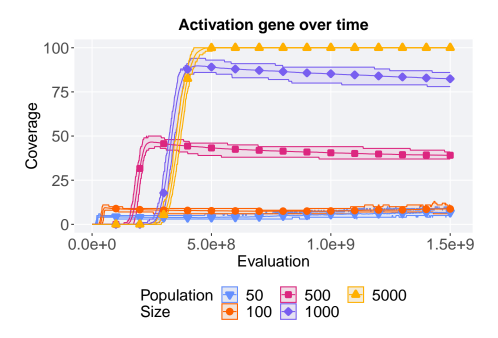
3.3 Satisfactory trait coverage

3.3.1 Coverage over time

Satisfactory trait coverage over time.

```
# aggregate
lines = over_time %>%
  group_by(pop_size, eval) %>%
  dplyr::summarise(
    min = min(satisfactory_coverage),
    mean = mean(satisfactory_coverage),
    max = max(satisfactory_coverage)
  )
lines$pop_size <- factor(lines$pop_size, levels = NAMES)</pre>
ggplot(lines, aes(x=eval, y=mean, group = pop_size, fill = pop_size, color = pop_size, shape = pop_size
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(linewidth = 0.5) +
  geom_point(data = filter(lines, eval %% 1000000000 == 0 & eval != 0), size = 1.0, stroke = 2.0,
  scale_y_continuous(
    name="Coverage",
    limits=c(0, 100),
```

```
breaks=seq(0,100, 25),
  labels=c("0", "25", "50", "75", "100")
) +
scale_x_continuous(
  name="Evaluation",
  labels = c('0.0e+0', '5.0e+8', '1.0e+9', '1.5e+9'),
  limits = c(0,1520000000)
scale_shape_manual(values=SHAPE)+
scale_colour_manual(values = cb_palette) +
scale_fill_manual(values = cb_palette) +
ggtitle('Activation gene over time')+
p_theme +
guides(
  shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
  color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
  fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```

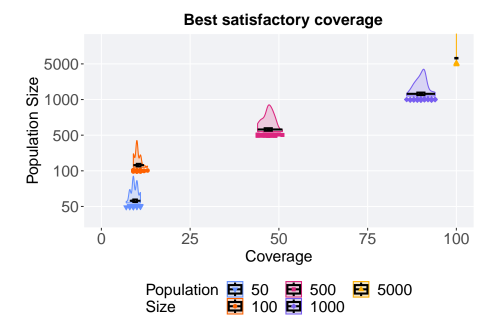


3.3.2 Best satisfactory trait coverage found throughout run

Satisfactory trait coverage of the best population found throughout an evolutionary run.

```
ggplot(best, aes(x = pop_size, y = coverage, color = pop_size, fill = pop_size, shape = pop_size)
  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 = 0.02, height = 0.0001), size = 1.5, alpha = 1.0)
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks = seq(0, 100, 25),
   labels=c("0", "25", "50", "75", "100"),
  ) +
  scale_x_discrete(
   name="Population Size"
  )+
  scale shape manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle(' Best satisfactory coverage')+
  p_theme + coord_flip() +
  guides(
    shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
   fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```

```
## Warning: Removed 29 rows containing missing values
## ('geom_point()').
```



3.3.2.1 Summary statistics

```
best %>%
  group_by(pop_size) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(coverage)),
    min = min(coverage, na.rm = TRUE),
    median = median(coverage, na.rm = TRUE),
    mean = mean(coverage, na.rm = TRUE),
    max = max(coverage, na.rm = TRUE),
    IQR = IQR(coverage, na.rm = TRUE)
)
```

```
## # A tibble: 5 x 8
    pop_size count na_cnt
                                                        IQR
                             min median
                                          mean
                                                 max
    <fct>
              <int> <int> <int>
                                  <dbl>
                                         <dbl> <int> <dbl>
## 1 50
                 50
                               7
                                          9.36
                         0
                                      9
                                                   11
## 2 100
                 50
                         0
                               9
                                     10 10.1
                                                   13
                                                          1
## 3 500
                 50
                         0
                              44
                                     47
                                         47.0
                                                  51
                                                          2
## 4 1000
                 50
                              86
                                     90
                                        90.0
                                                  94
                                                          2
## 5 5000
                             100
                 50
                                    100 100
                                                 100
```

3.3.2.2 Kruskal-Wallis test

```
##
##
Kruskal-Wallis rank sum test
##
## data: coverage by pop_size
## Kruskal-Wallis chi-squared = 232.33, df = 4,
## p-value < 2.2e-16</pre>
```

3.3.2.3 Pairwise wilcoxon test

```
pairwise.wilcox.test(x = best$coverage, g = best$pop_size, p.adjust.method = "bonferroni",
                    paired = FALSE, conf.int = FALSE, alternative = 'g')
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: best$coverage and best$pop_size
##
##
       50
            100
                     500
                            1000
## 100 0.0019 -
## 500 <2e-16 <2e-16 -
## 1000 <2e-16 <2e-16 <2e-16 -
## 5000 <2e-16 <2e-16 <2e-16
##
## P value adjustment method: bonferroni
```

Chapter 4

Contradictory objectives 150 results

Here we report the activation gene coverage and satisfactory trait coverage was found on the contradictory objectives diagnostic. 50 replicates were conducted for each population size explored. Activation gene coverage is calculated by finding all the unique activation genes found within a given population. Satisfactory trait coverage is calculated by finding all the unique satisfactory traits found within a given population.

4.1 Analysis setup

```
library(ggplot2)
library(cowplot)
library(dplyr)
library(PupillometryR)

# over time data
over_time <- read.csv("../Paper_Data/Contradictory-150/ot.csv", header = TRUE, stringsAsFactors = over_time$pop_size <- factor(over_time$pop_size, levels = NAMES)

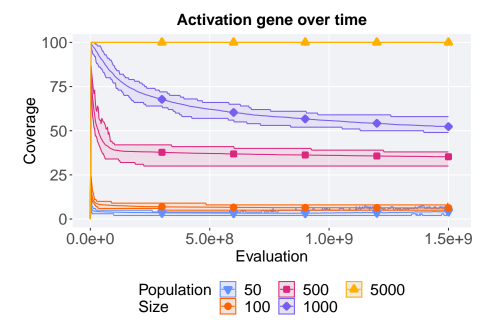
# best performance data
best <- read.csv('../Paper_Data/Contradictory-150/best.csv', header = TRUE, stringsAsFactors = FAbest$pop_size <- factor(best$pop_size, levels = NAMES)</pre>
```

4.2 Activation gene coverage

4.2.1 Coverage over time

Performance of the best solution in the population at each generation over time.

```
# aggregate
lines = over_time %>%
  group_by(pop_size, eval) %>%
 dplyr::summarise(
   min = min(activation_coverage),
   mean = mean(activation_coverage),
   max = max(activation_coverage)
lines$pop_size <- factor(lines$pop_size, levels = NAMES)</pre>
ggplot(lines, aes(x=eval, y=mean, group = pop_size, fill = pop_size, color = pop_size,
 geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(linewidth = 0.5) +
  geom_point(data = filter(lines, eval %% 1000000000 == 0 & eval != 0), size = 1.0, str.
  scale y continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks=seq(0,100, 25),
   labels=c("0", "25", "50", "75", "100")
  ) +
  scale_x_continuous(
   name="Evaluation",
   labels = c('0.0e+0', '5.0e+8', '1.0e+9', '1.5e+9'),
   limits = c(0,1520000000)
 ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene over time')+
 p_theme +
  guides (
    shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```



4.3 Satisfactory trait coverage

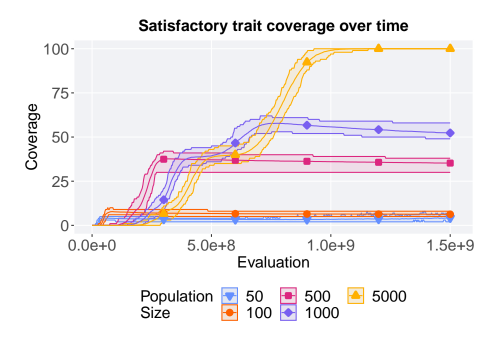
4.3.1 Coverage over time

Satisfactory trait coverage over time.

```
# aggregate
lines = over_time %>%
group_by(pop_size, eval) %>%
dplyr::summarise(
    min = min(satisfactory_coverage),
    mean = mean(satisfactory_coverage),
    max = max(satisfactory_coverage)
)
lines$pop_size <- factor(lines$pop_size, levels = NAMES)

ggplot(lines, aes(x=eval, y=mean, group = pop_size, fill = pop_size, color = pop_size, shape = pop_size, shape
```

```
breaks=seq(0,100, 25),
  labels=c("0", "25", "50", "75", "100")
) +
scale_x_continuous(
  name="Evaluation",
  labels = c('0.0e+0', '5.0e+8', '1.0e+9', '1.5e+9'),
  limits = c(0,1520000000)
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 +
guides(
  shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
  color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
  fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```

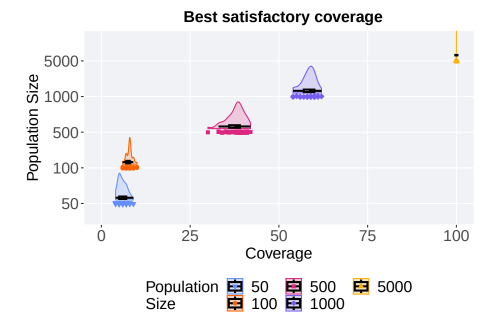


4.3.2 Best satisfactory trait coverage found throughout run

Satisfactory trait coverage of the best population found throughout an evolutionary run.

```
ggplot(best, aes(x = pop_size, y = coverage, color = pop_size, fill = pop_size, shape = pop_size)
  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 = 0.02, height = 0.0001), size = 1.5, alpha = 1.0)
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks = seq(0, 100, 25),
   labels=c("0", "25", "50", "75", "100"),
  ) +
  scale_x_discrete(
   name="Population Size"
  )+
  scale shape manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle(' Best satisfactory coverage')+
  p_theme + coord_flip() +
  guides(
    shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
   fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```

```
## Warning: Removed 29 rows containing missing values
## ('geom_point()').
```



4.3.2.1 Summary statistics

```
best %>%
group_by(pop_size) %>%
dplyr::summarise(
   count = n(),
   na_cnt = sum(is.na(coverage)),
   min = min(coverage, na.rm = TRUE),
   median = median(coverage, na.rm = TRUE),
   mean = mean(coverage, na.rm = TRUE),
   max = max(coverage, na.rm = TRUE),
   IQR = IQR(coverage, na.rm = TRUE)
)
```

```
## # A tibble: 5 x 8
     pop_size count na_cnt
                                                        IQR
                             min median
                                           mean
                                                  max
     <fct>
              <int> <int> <int>
                                   <dbl>
                                          <dbl> <int> <dbl>
## 1 50
                 50
                                           5.86
                         0
                               4
                                       6
                                                    9
                                                          2
## 2 100
                                           7.88
                 50
                         0
                               6
                                       8
                                                   10
                                                          1
## 3 500
                 50
                         0
                              30
                                      38 37.8
                                                   42
                                                          3
## 4 1000
                 50
                              54
                                      59
                                        58.4
                                                   62
                                                          3
## 5 5000
                         0
                 50
                             100
                                     100 100
                                                  100
                                                          0
```

4.3.2.2 Kruskal-Wallis test

```
##
## Kruskal-Wallis rank sum test
##
## data: coverage by pop_size
## Kruskal-Wallis chi-squared = 237.37, df = 4,
## p-value < 2.2e-16</pre>
```

4.3.2.3 Pairwise wilcoxon test

```
pairwise.wilcox.test(x = best$coverage, g = best$pop_size, p.adjust.method = "bonferroni",
                    paired = FALSE, conf.int = FALSE, alternative = 'g')
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: best$coverage and best$pop_size
##
       50
              100
                       500
                               1000
## 100 1.2e-11 -
## 500 < 2e-16 < 2e-16 -
## 1000 < 2e-16 < 2e-16 < 2e-16 -
## 5000 < 2e-16 < 2e-16 < 2e-16 < 2e-16
##
## P value adjustment method: bonferroni
```

Chapter 5

Contradictory objectives 200 results

Here we report the activation gene coverage and satisfactory trait coverage was found on the contradictory objectives diagnostic. 50 replicates were conducted for each population size explored. Activation gene coverage is calculated by finding all the unique activation genes found within a given population. Satisfactory trait coverage is calculated by finding all the unique satisfactory traits found within a given population.

5.1 Analysis setup

```
library(ggplot2)
library(cowplot)
library(dplyr)
library(PupillometryR)

# over time data
over_time <- read.csv("../Paper_Data/Contradictory-200/ot.csv", header = TRUE, stringsAsFactors = over_time$pop_size <- factor(over_time$pop_size, levels = NAMES)

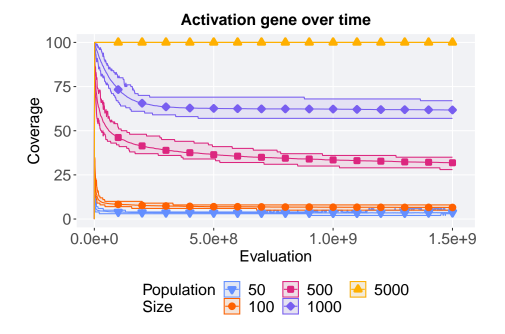
# best performance data
best <- read.csv('../Paper_Data/Contradictory-200/best.csv', header = TRUE, stringsAsFactors = FABES)
best$pop_size <- factor(best$pop_size, levels = NAMES)</pre>
```

5.2 Activation gene coverage

5.2.1 Coverage over time

Performance of the best solution in the population at each generation over time.

```
# aggregate
lines = over_time %>%
  group_by(pop_size, eval) %>%
 dplyr::summarise(
   min = min(activation_coverage),
   mean = mean(activation_coverage),
   max = max(activation_coverage)
lines$pop_size <- factor(lines$pop_size, levels = NAMES)</pre>
ggplot(lines, aes(x=eval, y=mean, group = pop_size, fill = pop_size, color = pop_size,
 geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(linewidth = 0.5) +
  geom_point(data = filter(lines, eval %% 1000000000 == 0 & eval != 0), size = 1.0, str.
  scale y continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks=seq(0,100, 25),
   labels=c("0", "25", "50", "75", "100")
  ) +
  scale_x_continuous(
   name="Evaluation",
   labels = c('0.0e+0', '5.0e+8', '1.0e+9', '1.5e+9'),
   limits = c(0,1520000000)
 ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene over time')+
 p_theme +
  guides (
    shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```



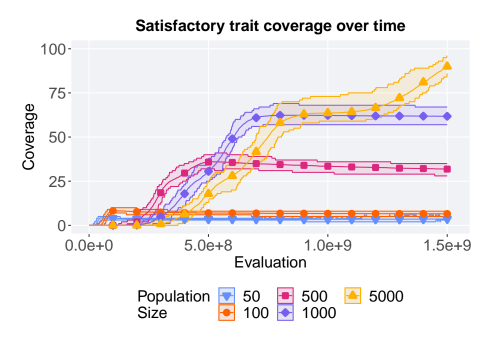
5.3 Satisfactory trait coverage

5.3.1 Coverage over time

Satisfactory trait coverage over time.

```
# aggregate
lines = over_time %>%
  group_by(pop_size, eval) %>%
  dplyr::summarise(
    min = min(satisfactory_coverage),
    mean = mean(satisfactory_coverage),
    max = max(satisfactory_coverage)
  )
lines$pop_size <- factor(lines$pop_size, levels = NAMES)</pre>
ggplot(lines, aes(x=eval, y=mean, group = pop_size, fill = pop_size, color = pop_size, shape = pop_size
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(linewidth = 0.5) +
  geom_point(data = filter(lines, eval %% 1000000000 == 0 & eval != 0), size = 1.0, stroke = 2.0,
  scale_y_continuous(
    name="Coverage",
    limits=c(0, 100),
```

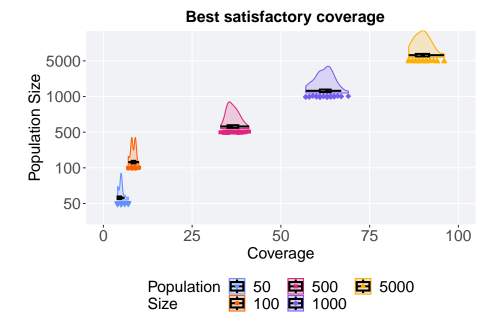
```
breaks=seq(0,100, 25),
  labels=c("0", "25", "50", "75", "100")
) +
scale_x_continuous(
  name="Evaluation",
  labels = c('0.0e+0', '5.0e+8', '1.0e+9', '1.5e+9'),
  limits = c(0,1520000000)
scale_shape_manual(values=SHAPE)+
scale_colour_manual(values = cb_palette) +
scale fill manual(values = cb palette) +
ggtitle('Satisfactory trait coverage over time')+
p_theme +
guides(
  shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
  color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
  fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```



5.3.2 Best satisfactory trait coverage found throughout run

Satisfactory trait coverage of the best population found throughout an evolutionary run.

```
ggplot(best, aes(x = pop_size, y = coverage, color = pop_size, fill = pop_size, shape = pop_size)
  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 = 0.02, height = 0.0001), size = 1.5, alpha = 1.0)
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks=seq(0,100, 25),
   labels=c("0", "25", "50", "75", "100"),
  ) +
  scale_x_discrete(
   name="Population Size"
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle(' Best satisfactory coverage')+
 p_theme + coord_flip() +
  guides(
   shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
   color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
   fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```



5.3.2.1 Summary statistics

```
best %>%
  group_by(pop_size) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(coverage)),
    min = min(coverage, na.rm = TRUE),
    median = median(coverage, na.rm = TRUE),
    mean = mean(coverage, na.rm = TRUE),
    max = max(coverage, na.rm = TRUE),
    IQR = IQR(coverage, na.rm = TRUE)
)
```

```
## # A tibble: 5 x 8
    pop_size count na_cnt
                            min median mean
                                                     IQR
    <fct>
             <int> <int> <int>
                                 <dbl> <dbl> <int> <dbl>
## 1 50
                50
                                     5 4.94
                        0
                              4
                                                7
## 2 100
                              7
                                     8 8.38
                50
                        0
                                                10 1
## 3 500
                50
                        0
                             33
                                    36 36.5
                                                41 3
## 4 1000
                50
                             57
                                    63 62.3
                                                69 3
                                                96 3.75
## 5 5000
                50
                             86
                                    90 90.0
```

5.3.2.2 Kruskal-Wallis test

```
##
## Kruskal-Wallis rank sum test
##
## data: coverage by pop_size
## Kruskal-Wallis chi-squared = 239.68, df = 4,
## p-value < 2.2e-16</pre>
```

5.3.2.3 Pairwise wilcoxon test

```
pairwise.wilcox.test(x = best$coverage, g = best$pop_size, p.adjust.method = "bonferroni",
                    paired = FALSE, conf.int = FALSE, alternative = 'g')
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: best$coverage and best$pop_size
##
##
       50
            100
                     500
                            1000
## 100 <2e-16 -
## 500 <2e-16 <2e-16 -
## 1000 <2e-16 <2e-16 <2e-16 -
## 5000 <2e-16 <2e-16 <2e-16
##
## P value adjustment method: bonferroni
```

Chapter 6

Contradictory objectives 300 results

Here we report the activation gene coverage and satisfactory trait coverage was found on the contradictory objectives diagnostic. 50 replicates were conducted for each population size explored. Activation gene coverage is calculated by finding all the unique activation genes found within a given population. Satisfactory trait coverage is calculated by finding all the unique satisfactory traits found within a given population.

6.1 Analysis setup

```
library(ggplot2)
library(cowplot)
library(dplyr)
library(PupillometryR)

# over time data
over_time <- read.csv("../Paper_Data/Contradictory-300/ot.csv", header = TRUE, stringsAsFactors = over_time$pop_size <- factor(over_time$pop_size, levels = NAMES)

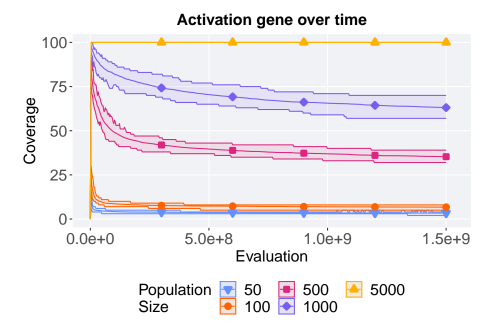
# best performance data
best <- read.csv('../Paper_Data/Contradictory-300/best.csv', header = TRUE, stringsAsFactors = FAbest$pop_size <- factor(best$pop_size, levels = NAMES)</pre>
```

6.2 Activation gene coverage

6.2.1 Coverage over time

Performance of the best solution in the population at each generation over time.

```
# aggregate
lines = over_time %>%
  group_by(pop_size, eval) %>%
 dplyr::summarise(
   min = min(activation_coverage),
   mean = mean(activation_coverage),
   max = max(activation_coverage)
lines$pop_size <- factor(lines$pop_size, levels = NAMES)</pre>
ggplot(lines, aes(x=eval, y=mean, group = pop_size, fill = pop_size, color = pop_size,
 geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(linewidth = 0.5) +
  geom_point(data = filter(lines, eval %% 1000000000 == 0 & eval != 0), size = 1.0, str.
  scale y continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks=seq(0,100, 25),
   labels=c("0", "25", "50", "75", "100")
  ) +
  scale_x_continuous(
   name="Evaluation",
   labels = c('0.0e+0', '5.0e+8', '1.0e+9', '1.5e+9'),
   limits = c(0,1530000000)
 ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene over time')+
 p_theme +
  guides (
    shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```



6.3 Satisfactory trait coverage

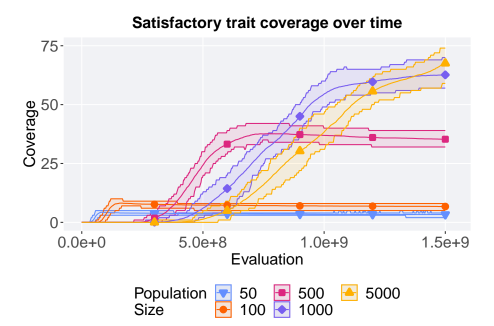
6.3.1 Coverage over time

Satisfactory trait coverage over time.

```
# aggregate
lines = over_time %>%
group_by(pop_size, eval) %>%
dplyr::summarise(
    min = min(satisfactory_coverage),
    mean = mean(satisfactory_coverage),
    max = max(satisfactory_coverage)
)
lines$pop_size <- factor(lines$pop_size, levels = NAMES)

ggplot(lines, aes(x=eval, y=mean, group = pop_size, fill = pop_size, color = pop_size, shape = pop_size, shape
```

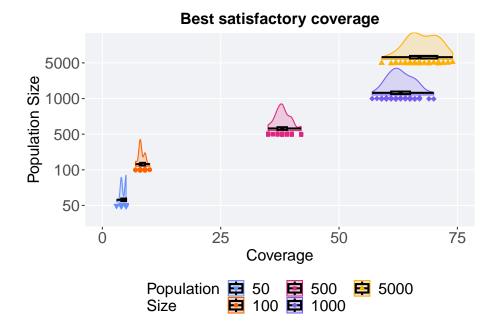
```
breaks=seq(0,75, 25),
  labels=c("0", "25", "50", "75")
) +
scale_x_continuous(
  name="Evaluation",
  labels = c('0.0e+0', '5.0e+8', '1.0e+9', '1.5e+9'),
  limits = c(0,1530000000)
scale_shape_manual(values=SHAPE)+
scale_colour_manual(values = cb_palette) +
scale_fill_manual(values = cb_palette) +
ggtitle('Satisfactory trait coverage over time')+
p_theme +
guides(
  shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
  color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
  fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```



6.3.2 Best satisfactory trait coverage found throughout run

Satisfactory trait coverage of the best population found throughout an evolutionary run.

```
ggplot(best, aes(x = pop_size, y = coverage, color = pop_size, fill = pop_size, shape = pop_size)
  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 = 0.02, height = 0.0001), size = 1.5, alpha = 1.0)
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 75),
   breaks=seq(0,75, 25),
   labels=c("0", "25", "50", "75")
  ) +
  scale_x_discrete(
   name="Population Size"
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Best satisfactory coverage')+
  p_theme + coord_flip() +
  guides(
   shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
   fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```



6.3.2.1 Summary statistics

```
best %>%
  group_by(pop_size) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(coverage)),
    min = min(coverage, na.rm = TRUE),
    median = median(coverage, na.rm = TRUE),
    mean = mean(coverage, na.rm = TRUE),
    max = max(coverage, na.rm = TRUE),
    IQR = IQR(coverage, na.rm = TRUE)
)
```

```
## # A tibble: 5 x 8
    pop_size count na_cnt
                           min median mean
                                                    IQR
    <fct>
             <int> <int> <int>
                                 <dbl> <dbl> <int> <dbl>
## 1 50
                50
                                  4.5 4.46
                        0
                             3
## 2 100
                             7
                                       8.3
                50
                        0
                                   8
                                               10 1
## 3 500
                50
                        0
                             35
                                 38
                                      38.1
                                               42 2
                                               70 4
## 4 1000
                50
                             57
                                  63
                                      63.0
                                               74 5.75
## 5 5000
                        0
                                      67.5
                50
                             59
                                 67
```

6.3.2.2 Kruskal-Wallis test

```
##
## Kruskal-Wallis rank sum test
##
## data: coverage by pop_size
## Kruskal-Wallis chi-squared = 233.56, df = 4,
## p-value < 2.2e-16</pre>
```

6.3.2.3 Pairwise wilcoxon test

```
pairwise.wilcox.test(x = best$coverage, g = best$pop_size, p.adjust.method = "bonferroni",
                    paired = FALSE, conf.int = FALSE, alternative = 'g')
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
## data: best$coverage and best$pop_size
##
##
       50
              100
                       500
                               1000
## 100 < 2e-16 -
## 500 < 2e-16 < 2e-16 -
## 1000 < 2e-16 < 2e-16 < 2e-16 -
## 5000 < 2e-16 < 2e-16 < 2e-16 2.3e-08
##
## P value adjustment method: bonferroni
```

Chapter 7

Contradictory objectives 500 results

Here we report the activation gene coverage and satisfactory trait coverage was found on the contradictory objectives diagnostic. 50 replicates were conducted for each population size explored. Activation gene coverage is calculated by finding all the unique activation genes found within a given population. Satisfactory trait coverage is calculated by finding all the unique satisfactory traits found within a given population.

7.1 Analysis setup

```
library(ggplot2)
library(cowplot)
library(dplyr)
library(PupillometryR)

# over time data
over_time <- read.csv("../Paper_Data/Contradictory-500/ot.csv", header = TRUE, stringsAsFactors = over_time$pop_size <- factor(over_time$pop_size, levels = NAMES)

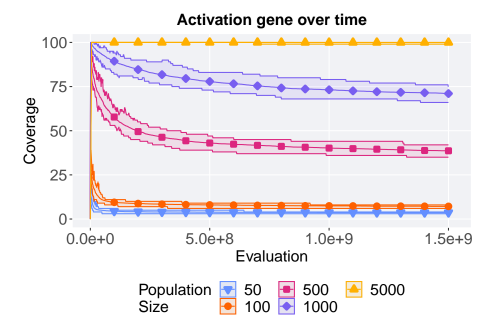
# best performance data
best <- read.csv('../Paper_Data/Contradictory-500/best.csv', header = TRUE, stringsAsFactors = FAbest$pop_size <- factor(best$pop_size, levels = NAMES)</pre>
```

7.2 Activation gene coverage

7.2.1 Coverage over time

Performance of the best solution in the population at each generation over time.

```
# aggregate
lines = over_time %>%
  group_by(pop_size, eval) %>%
 dplyr::summarise(
   min = min(activation_coverage),
   mean = mean(activation_coverage),
   max = max(activation_coverage)
lines$pop_size <- factor(lines$pop_size, levels = NAMES)</pre>
ggplot(lines, aes(x=eval, y=mean, group = pop_size, fill = pop_size, color = pop_size,
 geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(linewidth = 0.5) +
  geom_point(data = filter(lines, eval %% 1000000000 == 0 & eval != 0), size = 1.0, str.
  scale y continuous(
   name="Coverage",
   limits=c(0, 100),
   breaks=seq(0,100, 25),
   labels=c("0", "25", "50", "75", "100")
  ) +
  scale_x_continuous(
   name="Evaluation",
   labels = c('0.0e+0', '5.0e+8', '1.0e+9', '1.5e+9'),
   limits = c(0,1520000000)
 ) +
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette) +
  scale_fill_manual(values = cb_palette) +
  ggtitle('Activation gene over time')+
 p_theme +
  guides (
    shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
    fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```



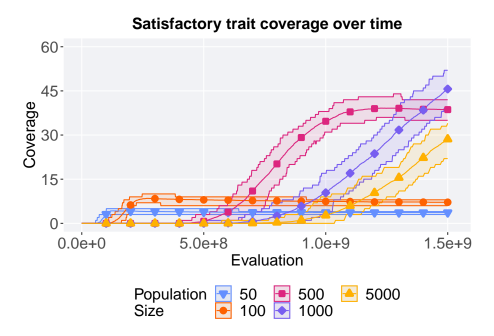
7.3 Satisfactory trait coverage

7.3.1 Coverage over time

Satisfactory trait coverage over time.

```
# aggregate
lines = over_time %>%
  group_by(pop_size, eval) %>%
  dplyr::summarise(
    min = min(satisfactory_coverage),
    mean = mean(satisfactory_coverage),
    max = max(satisfactory_coverage)
  )
lines$pop_size <- factor(lines$pop_size, levels = NAMES)</pre>
ggplot(lines, aes(x=eval, y=mean, group = pop_size, fill = pop_size, color = pop_size, shape = pop_size
  geom_ribbon(aes(ymin = min, ymax = max), alpha = 0.1) +
  geom_line(linewidth = 0.5) +
  geom_point(data = filter(lines, eval %% 1000000000 == 0 & eval != 0), size = 1.0, stroke = 2.0,
  scale_y_continuous(
    name="Coverage",
    limits=c(0, 60),
```

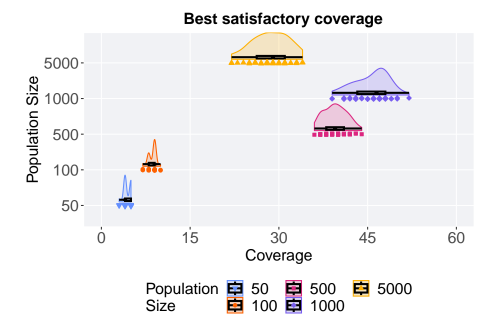
```
breaks=seq(0,60, 15),
  labels=c("0", "15", "30", "45", "60")
) +
scale_x_continuous(
  name="Evaluation",
  labels = c('0.0e+0', '5.0e+8', '1.0e+9', '1.5e+9'),
  limits = c(0,1520000000)
scale_shape_manual(values=SHAPE)+
scale_colour_manual(values = cb_palette) +
scale fill manual(values = cb palette) +
ggtitle('Satisfactory trait coverage over time')+
p_theme +
guides(
  shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
  color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
  fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```



7.3.2 Best satisfactory trait coverage found throughout run

Satisfactory trait coverage of the best population found throughout an evolutionary run.

```
ggplot(best, aes(x = pop_size, y = coverage, color = pop_size, fill = pop_size, shape = pop_size)
  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 = 0.02, height = 0.0001), size = 1.5, alpha = 1.0)
  scale_y_continuous(
   name="Coverage",
   limits=c(0, 60),
   breaks = seq(0,60, 15),
   labels=c("0", "15", "30", "45", "60")
  ) +
  scale_x_discrete(
   name="Population Size"
  scale_shape_manual(values=SHAPE)+
  scale_colour_manual(values = cb_palette, ) +
  scale_fill_manual(values = cb_palette) +
  ggtitle(' Best satisfactory coverage')+
 p_theme + coord_flip() +
  guides(
   shape=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
   color=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize'),
   fill=guide_legend(nrow=2, title.position = "left", title = 'Population\nSize')
```



7.3.2.1 Summary statistics

```
best %>%
  group_by(pop_size) %>%
  dplyr::summarise(
    count = n(),
    na_cnt = sum(is.na(coverage)),
    min = min(coverage, na.rm = TRUE),
    median = median(coverage, na.rm = TRUE),
    mean = mean(coverage, na.rm = TRUE),
    max = max(coverage, na.rm = TRUE),
    IQR = IQR(coverage, na.rm = TRUE)
)
```

```
## # A tibble: 5 x 8
    pop_size count na_cnt
                            min median mean
                                                     IQR
                                               max
    <fct>
             <int> <int> <int>
                                 <dbl> <dbl> <int> <dbl>
## 1 50
                50
                                   4
                                        4.36
                        0
                              3
                                                 5
## 2 100
                              7
                                   9
                                        8.62
                50
                        0
                                                10 1
## 3 500
                50
                        0
                             36
                                  39.5 39.6
                                                44 3
## 4 1000
                50
                        0
                             39
                                  46.5 45.7
                                                52 4.75
## 5 5000
                        0
                50
                             22
                                  29
                                       28.6
                                                34 4.75
```

7.3.2.2 Kruskal-Wallis test

```
##
## Kruskal-Wallis rank sum test
##
## data: coverage by pop_size
## Kruskal-Wallis chi-squared = 237.63, df = 4,
## p-value < 2.2e-16</pre>
```

7.3.2.3 Pairwise wilcoxon test

```
best$pop_size \leftarrow factor(best$pop_size, levels = c(1000,500,5000,100,50))
pairwise.wilcox.test(x = best$coverage, g = best$pop_size, p.adjust.method = "bonferroni",
                     paired = FALSE, conf.int = FALSE, alternative = '1')
##
## Pairwise comparisons using Wilcoxon rank sum test with continuity correction
## data: best$coverage and best$pop_size
##
       1000
               500
                      5000
                                100
## 500 5.7e-14 -
## 5000 < 2e-16 < 2e-16 -
## 100 < 2e-16 < 2e-16 < 2e-16 -
## 50 < 2e-16 < 2e-16 < 2e-16 < 2e-16
## P value adjustment method: bonferroni
```