Signatr Artifact

We also provide pdf and html versions of this README. If reading locally and not on github, we advise to use the html version.

The artifact contains the signatr tool, and the pipelines to create an R value database and to fuzz R functions with the database to find type signatures. The pipeline to create a value database is in pipeline-dbgen. The fuzzing pipeline will generate the inputs for the sle.Rmd R markdown notebook. That notebook can then be rendered to get all the results (tables, figures) we use in the paper.

To use the artifact:

- Install the docker image (see Install the docker image). Installing locally is possible but involved. Following the steps described in the docker-image/Dockerfile should help if this is the hard path you are choosing!
- 2. Experiment with the tool on a small example: see Experimenting the tool
- 3. Reproduce the analysis pipeline: see The analysis pipeline

The tool is packaged as an R library. It is hosted at https://github.com/PRL-PRG/signatr.

The artifact is also provided directly on github: https://github.com/PRL-PRG/sle22-signatr-artifact

You can get it by entering the following commands in a shell:

\$ git clone git@github.com:PRL-PRG/sle22-signatr-artifact.git

Install the docker image

Go in the artifact's folder:

\$ cd sle22-signatr-artifact

To install the docker image, you can:

- pull the docker image with docker pull prlprg/sle22-signatr, or
- build the docker image (it takes time!):

\$ cd docker-image

\$ make

After installing the docker image, **make sure** to run all the following commands in a shell inside the docker image (for Linux, macOS) from the artifact directory.

To start the docker image, go back to the root directory of the artifact (sle22-signatr-artifact/) and enter in a shell:

./enter

which should give you a bash shell prompt, like (modulo the hostname):

```
r@eaf63037fd02:/work$
```

It automatically mounts the content of the folder from which you run the command into the /work directory in the container.

Experimenting with the tool

Run the R interpreter *inside the docker image*. It will start the patched R interpreter. The tool *does not run* in the standard R interpreter.

In the following listings, \$ indicates the shell and > denotes the R REPL.

```
$ R
R version 4.0.2 (2020-06-22) -- "Taking Off again"
...
> library(signatr)
```

All following commands and instructions should be run in the docker container.

Database

To generate a database of values, we need some code to run. One way is to extract it from an existing R package, for example stringr, which provides regexes:

```
> extract_package_code("stringr", output_dir = "demo")
...
7 examples/str_detect.Rd.R examples
```

This will extract all the runnable snippets from the package documentation and tests into the given directory. For example:

```
> cat(readLines("demo/examples/str_detect.Rd.R", n = 15), sep = "\n")
...
fruit <- c("apple", "banana", "pear", "pinapple")
str_detect(fruit, "a")
str_detect(fruit, "^a")
...</pre>
```

Next, we trace the file by running it (in the patched R interpreter) and recording all the calls, using the trace_filefunction:

The database generation is also automated in the pipeline-dbgen directory in the artifact, and handles there tracing on multiple files and merging the results. See Generate the database for more details.

Fuzzing

Once the database is ready, we can start fuzzing the str_detect function of the stringr package:

```
> R <- quick_fuzz("stringr", "str_detect", "demo.sxpdb", budget = 100, action = "infer")
started a new runner:PROCESS 'R', running, pid 4157
fuzzing stringr:::str_detect [=====] 100/100 (100%) 39s
stopped runner:PROCESS 'R', running, pid 4157</pre>
```

The infer action will infer types for each call argument and return value using the type annotation language described in Designing types for R, empirically. It returns an R data frame with the inferred call signature in the result column:

If you are repeating these steps, it is possible that your results will be different since fuzzing is non-deterministic.

The listing shows two calls: a failed one (non-zero status) with an error message, and a successful one with an inferred signature. The args_idx column contains the indices of the values of the arguments in the database: the actual argument values can be obtained by looking up the args_idx in the database:

```
> library(sxpdb)
> db <- open_db("demo.sxpdb")
> get_val_idx(db, 0) # value at index 0
[1] "a"
> close(db)
```

One advantage of using R is that we can use R's many data analysis functions. For example, we can look at the resulting signatures:

```
3 (list<integer>, character[], list<integer>) => logical[] 1
4 NA 97
```

This shows that in 3 cases, the fuzzer managed to generate a call that was successful, and so the signatures of those calls.

The analysis pipeline

The following tutorial demonstrates how to run the analysis pipeline to reproduce the results of the paper. It consists of a series of steps that at the end generates the input for the analysis.

In this write up, we will run it on a small subset of the original packages (cf. data/packages.txt). The reason is that the size of the data require is fairly large. For example, just the value database is over 287GB and its generation take over half a day (on a 72 core Intel Xeon 6140 2.30GHz server). Also one would have to download and install all the packages and their dependencies which again takes space and time. If you are however interested and have the computational resource, we will be happy to share the data, please contact the AEC chair.

Note: - You will be running code downloaded from a public repository. Despite that CRAN is a curated repository, it should be done with caution. Run it inside the container.

• Most steps takes a few minutes at most, long running ones are indicated with an estimate.

Steps

The following is essentially what is in the Figure 1 and Figure 2 in the paper, packaged in scripts for simpler use using GNU parallels for parallel execution. All steps should be run inside a docker container. As a reminder, to enter the container, run:

./enter.sh

Anytime you want to kill a task, it is good to exit the container and enter it again so all the child processes are properly killed.

0. get the sample sxpdb database

For the experiment we need a value database (sxpdb database) that will be used for the fuzzing. You can either build one yourself, or download one we have prepared using the same steps.

To get the prebuilt, one do the following:

cd data

wget -0 cran_db.tar.xz https://owncloud.cesnet.cz/index.php/s/aHprMbas4haELVf/download
tar xvJf cran_db.tar.xz

The extracted database has about 10GB.

Building it yourself The database generation uses targets to orchestrate the pipeline.

The database for the SLE paper is obtained by tracing 400 packages from data/packages-typer-400.txt. The packages to be traced have to be specified in data/packages.txt, which contains a new-line separated list of packages to include in the corpus.

To start tracing, after opening an R session and specifying an adequate number of parallel workers:

```
cp data/packages-typer-400.txt data/packages.txt
cd pipeline-dbgen
R -e 'targets::tar_make_future(workers = 64)'
```

The extracted code of the packages will be located in output/extracted-code. The resulting database will be generated as output/sxpdb/cran_db. You should move it to data to follow the next steps. Depending on your machine, the generation of the database for the 400 packages can take from a few hours to a few days.

We provide other variants of packages.txt. For instance, packages-4.txt includes 2 huge and common R packages, dplyr and ggplot2.

1. create a corpus

The corpus consists of the following:

- R package sources in data/sources
- installed R packages data/library
- extracted code from R packages data/extracted-code
- corpus metadata file data/corpus.csv

This is bootstrapped using the data/packages.txt file.

To create a corpus, run the following:

```
./create-corpus.R
```

Depending on the number of packages (and their transitive dependencies), it might take a while. For the sample of 5 packages (small corpus, though of the very popular packages), it might be $\sim\!20$ minutes.

It could happen that some dependencies won't install.

The result should be something like:

```
data/extracted-code <--- extracted code from R packages
data/library <--- installed R packages
data/sources <--- R package sources
data/corpus.csv <--- corpus metadata
```

2. fuzz the installed functions

Next, we will run the fuzzer using the values from the sample database:

```
./run-fuzz.sh
```

By default this will sample 100 functions from the corpus.csv and fuzz each 100 times. Both can adjusted by setting the FUNS and BUDGET environment variables. Using all the functions (e.g. FUNS=\$(wc -l data/corpus.csv) and 5000 runs (e.g. BUDGET=5000), the experiment might take about a day. That is why we recommend to scale it down so it runs within 30 minutes. By default, it will run 16 jobs in parallel. The can be changed using the JOBS environment variable.

The result will be:

```
data/fuzz <--- directory with the fuzzer output data/run-fuzz.csv <--- metadata about the run, duration, exitcodes, ...
```

You could view the intermediate results using the qcat.sh utility. For example:

```
./qcat.R 'data/fuzz/dplyr::arg_name'
```

shall show results for a function arg_name from dplyr package:

```
# A tibble: 100 × 9
   args_idx error
                          exit status dispatch
                                                  result ts
                                                               fun_n... 1 rdb_p... 2
   st>
             <chr>
                         <int> <int> <list>
                                                   <int> <drt> <chr>
                                                                      <chr>
 1 <int [2]> "Error in ...
                                                        NA 0.08... dplyr:... ../rdb...
                             NA
                                      1 <named list>
 2 <int [2] > "Error in ...
                              NA
                                      1 <named list>
                                                        NA 0.11... dplyr:... ../rdb...
 3 <int [2]> "Error in ...
                             NA
                                      1 <named list>
                                                        NA 0.14... dplyr:..../rdb...
                                      1 <named list>
                                                        NA 0.15... dplyr:..../rdb...
 4 <int [2] > "Error in ...
                             NA
                                                        NA 0.09... dplyr:... ../rdb...
 5 <int [2]> "Error in ...
                                      1 <named list>
                             NA
 6 <int [2] > "Error in ...
                              NA
                                                        NA 0.53... dplyr:..../rdb...
                                      1 <named list>
 7 <int [2]> "Error in ...
                                                       NA 0.11... dplyr:..../rdb...
                             NA
                                      1 <named list>
 8 <int [2]> NA
                            NA
                                    0 <named list>
                                                       30 0.09... dplyr:... ../rdb...
 9 <int [2] > NA
                                    0 <named list>
                                                       31 0.09... dplyr:... ../rdb...
                            NA
10 <int [2]> NA
                            NA
                                    0 <named list>
                                                       32 0.09... dplyr:... ../rdb...
```

It indicates 7 failed calls and 3 good ones. Please note that due to random sampling your results will likely be different.

To simplify, we merge all the individual trace files into one using:

```
./fix-fuzz-rdb.R
./merge-traces.R
```

which should create data/all-traces.qs file with properly adjusted paths.

3. type the results

To type the traces, run the following:

```
./run-type.sh
```

By default, it will run 16 jobs in parallel. The can be changed using the JOBS environment variable.

The result will be:

```
data/types <--- directory with the type output
data/run-type.csv <--- metadata about the run, duration, exitcodes, ...</pre>
```

We can again peek the results:

A tibble: 40 × 3

```
./qcat.R 'data/types/dplyr::arg_name'
```

which should show types inferred from the fuzzed calls:

```
fun name
                     id signature
  <chr>
                  <int> <chr>
                      8 (list<list<class<unit, unit_v2> | double | integer> | ...
1 dplyr::arg_name
                      9 (class<gList>, list<class<factor> | double | integer>)...
2 dplyr::arg_name
                     10 (pairlist, list<character | double[]>) => class<glue, ...
3 dplyr::arg_name
                     13 (list<list<class<matrix> | double[] | integer | intege...
4 dplyr::arg_name
5 dplyr::arg_name
                     14 (character[], list<character | logical>) => class<glue...
                     15 (list<class<unit, unit_v2>>, list<list<class<expectati...
6 dplyr::arg_name
```

- 7 dplyr::arg_name 17 (list<class<call>>, double[]) => class<glue, character>
 8 dplyr::arg_name 24 (list<class<margin, simpleUnit, unit, unit_v2> | class...
 9 dplyr::arg_name 28 (class<matrix>, list<class<expectation_success, expect...

4. fuzz coverage

Computing the function source code coverage from the fuzzed calls is done by running the following:

```
./{\tt run-coverage.sh}
```

This will use the traced data to recreate the calls while using the covr tool to record code coverage. By default, it will run 16 jobs in parallel. The can be changed using the JOBS environment variable.

The result will be:

```
data/coverage <--- directory with the coverage output data/run-coverage.csv <--- metadata about the run, duration, exitcodes, ...
```

5. baseline

To have a comparison, we need to need to get the baseline data. Instead of fuzzing, we will simply run the extracted code from the packages. There are three steps:

- 1. run the extracted code to get the traces
 - ./run-baseline.sh
 - ./traces-baseline.R

This might be a bit longer running - about 15 minutes.

- 2. type the traces
 - ./run-type-baseline.sh
- 3. compute the coverage from these traces
 - ./run-coverage-baseline.sh

This might be a bit longer running - about 15 minutes.

By default, all will run 16 jobs in parallel. The can be changed using the JOBS environment variable.

The results will be in

```
data/baseline <--- baseline traces
data/baseline-types <--- baseline types
data/baseline-coverage <--- baseline coverage
data/run-*-baseline.csv <--- metadata about the runs, duration, exitcodes, ...
```

6. create a report

Finally, to render the results, run:

```
R --slave --quiet -e 'rmarkdown::render("sle.Rmd")'
```

This should create a file sle.html which you can open in a browser (navigate to the directory where you run the ./enter.sh). It also creates two more files: - experiment-uf.tex the data for the paper - uf-call-signatures.pdf the figure 4 in the paper

Note:

• Regarding the coverage, most likely a small number of fuzzed calls won't find a new paths, so in the report you will see 0 - as to better coverage.