# **COCO Documentation**

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This document serves as documentation for the COCO (COmparing Continuous Optimizers) software.

This software was mainly developed in the TAO team.

More information and the full documentation can be found at http://coco.gforge.inria.fr/

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### INTRODUCTION

**COMparing Continuous Optimisers** (**COCO**) is a tool for benchmarking algorithms for black-box optimisation. COCO facilitates systematic experimentation in the field of continuous optimization.

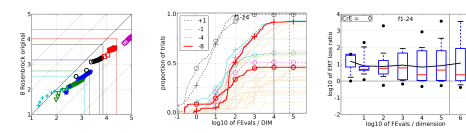


Figure 1.1: The pictures show (a) a scatterplot of run lengths (log10-scale) comparing two algorithms in different dimensions on a single function, (b) the run length distribution of two algorithms for different target difficulties  $\{10, 10^{-1}, 10^{-4}, 10^{-8}\}$  on a set of 24 functions and (c) box-whisker plots of the loss ratios of the expected run length compared to best (shortest) observed expected run length in BBOB-2009.

**COCO** has been used in several Black-Box Optimization Benchmarking (BBOB) workshops at the GECCO conference, the first in 2009 (BBOB-2009). Overall around a 100 articles and algorithms have been benchmarked.

#### COCO provides:

- 1. an **experimental framework** for testing the algorithms,
- 2. post-processing facilities for generating publication quality figures and tables,
- 3. LaTeX templates of articles which present the figures and tables in a single document.

The practitioner in continuous optimization who wants to benchmark one or many algorithms has to download COCO, plug the algorithm(s) into the provided experimental template and use the post-processing tools.

The COCO software is composed of two parts:

- 1. an interface available in different programming languages which allows to run and log experiments on multiple test functions; testbeds of functions (noisy and noiseless) are provided
- 2. a Python tool for generating figures and tables that can be used in the LaTeX templates.

The typical user of COCO renders the interface of the considered algorithm compatible with COCOs interface of the objective/fitness function, runs the provided main script (most presumably several times with increasing allowed maxfunevals until the waiting time becomes infeasible), invokes the postprocessing on the generated data and latexs the provided template.

An extensive documentation of the **test functions** can be found here.

### INSTALLATION

To install COCO, provided that the minimal requirements are met (see Requirements below), you only need to:

- 1. download the *first* archive bbobexpXX.XX.tar.gz from the COCO download page.
- 2. extract the archive

To check that COCO is running correctly, please proceed to What is the purpose of COCO?.

# 2.1 Requirements

## 2.1.1 Running Experiments

The interface for running experiments is provided in C/C++, Matlab/Gnu Octave, Java, R and Python. An algorithm to be tested with COCO has to be adapted to work with either of these versions.

Our code has been tested with:

- C/C++ gcc 4.2.1 and higher, the code provided is ANSI C,
- Matlab R2008a and higher
- Java 1.6, the code uses the Java Native Interface to call a library created from the C/C++ version of the code
- Python 2.6, 2.7
- R 2.14.2

# 2.1.2 Python and Post-Processing

The requirements below are for the post-processing tool of COCO that runs in Python.

Note: All packages are easily available for 32-bit Python 2.X.

Using 64-bit Python 2.X is possible but not straightforward. Numpy and Matplotlib do not work with Python 3.X.

#### Needed are

- python 2.6.x or 2.7.x (**not python 3.x**)
- numpy 1.2.1 or later (the version must fit to the python version)
- matplotlib 1.0.1 or later (the version must fit to the python version)

All three are part of the of the

· Enthought python distribution and

• Sage (then sage instead of python must be used).

Documents and articles that aggregate tables and figures are generated using LaTeX

#### How do I install on Windows?

For installing Python under Windows, either install Enthought or go to the Python download page and download the Python Windows Installer python-2.7.X.msi (the default Windows is for Win 32 architecture). This file requires the Microsoft Installer, which is a part of Windows XP and later releases. Numpy and Matplotlib can be installed with the standard .exe files which are respectively:

- numpy-X-win32-superpack-python2.X.exe and,
- matplotlib-X.win32-py2.X.exe.

#### How do I install on Linux?

In most Linux distributions, Python is already part of the installation. If not, use your favorite package manager to install Python (package name: python). The same needs to be done for Numpy (python-numpy) and Matplotlib (python-matplotlib) and their dependencies or install Sage or Enthought.

#### How do I install on Mac OS X?

Mac OS X comes with Python pre-installed but if your version of Python is 2.5 or earlier, you might need to upgrade.

You then need to download and install Numpy and Matplotlib:

- numpy-X-py2.X-python.org-macosx10.X.dmg
- matplotlib-X-python.org-32bit-py2.X-macosx10.X.dmg

or install Sage or Enthought.

# 2.2 Interactive Python

In order to fully enjoy a Python shell, we enable tab-completion on Unix systems like

```
>>> import rlcompleter
>>> import readline
>>> readline.parse_and_bind("tab: complete")
```

or we use IPython. In order to have a matlab-like environment with hundreds of additional commands directly available (including the plot command) use

```
>>> from pylab import *
or
>>> import pylab as pl
which provides pl.plot etc.
See also Pylab.
```

**CHAPTER** 

**THREE** 

# WHAT IS THE PURPOSE OF COCO?

Quantifying and comparing performance of optimization algorithms is one important aspect of research in search and optimization. However, this task turns out to be tedious and difficult to realize even in the single-objective case — at least if one is willing to accomplish it in a scientifically decent and rigorous way.

COCO provides tools for most of this tedious task:

- 1. choice and implementation of a well-motivated single-objective benchmark function testbed,
- 2. design of an experimental set-up,
- 3. generation of data output for
- 4. post-processing and presentation of the results in graphs and tables.

**CHAPTER** 

**FOUR** 

### FIRST TIME USING COCO

The following describes the workflow with COCO:

- Running Experiments
- Post-Processing
- Write/Compile an Article

# 4.1 Running Experiments

For more details, see Running Experiments with COCO.

argv = sys.argv[1:] # shortcut for input arguments

The code for running experiments with COCO is available in C/C++, Matlab/GNU Octave, Java, R and Python.

exampleexperiment and exampletiming are provided in each language. These files demonstrate how the code is supposed to be run: the first for doing an experiment over a set of functions, the second for measuring the time complexity of an optimizer.

The content of exampleexperiment.py shows how the experiment is invoked (see *Running Experiments with COCO* for an example in Matlab/Octave). The last 20-or-so lines define the experimentation loop over all dimension, all functions, and all instances:

```
#!/usr/bin/env python
# -*- coding: utf-8 -*-
"""Runs an entire experiment for benchmarking PURE_RANDOM_SEARCH on a testbed.

CAPITALIZATION indicates code adaptations to be made.
This script as well as files bbobbenchmarks.py and fgeneric.py need to be in the current working directory.

Under unix-like systems:
    nohup nice python exampleexperiment.py [data_path [dimensions [functions [instances]]]] > output

"""
import sys # in case we want to control what to run via command line args
import time
import numpy as np
import fgeneric
import bbobbenchmarks
```

```
datapath = 'PUT_MY_BBOB_DATA_PATH' if len(argv) < 1 else argv[0]</pre>
dimensions = (2, 3, 5, 10, 20, 40) if len(argv) < 2 else eval(argv[1])
function_ids = bbobbenchmarks.nfreeIDs if len(argv) < 3 else eval(argv[2])</pre>
# function_ids = bbobbenchmarks.noisyIDs if len(argv) < 3 else eval(argv[2])
instances = range(1, 6) + range(31, 41) if len(argv) < 4 else eval(argv[3])
opts = dict(algid='PUT ALGORITHM NAME',
            comments='PUT MORE DETAILED INFORMATION, PARAMETER SETTINGS ETC')
maxfunevals = '10 * dim' # 10*dim is a short test-experiment taking a few minutes
                         # INCREMENT maxfunevals SUCCESSIVELY to larger value(s)
minfunevals = 'dim + 2' # PUT MINIMAL sensible number of EVALUATIONS before to restart
maxrestarts = 10000
                        # SET to zero if algorithm is entirely deterministic
def run_optimizer(fun, dim, maxfunevals, ftarget=-np.Inf):
    """start the optimizer, allowing for some preparation.
    This implementation is an empty template to be filled
    # prepare
    x_start = 8. * np.random.rand(dim) - 4
    # call, REPLACE with optimizer to be tested
    PURE_RANDOM_SEARCH(fun, x_start, maxfunevals, ftarget)
def PURE_RANDOM_SEARCH(fun, x, maxfunevals, ftarget):
    """samples new points uniformly randomly in [-5,5]^dim and evaluates
    them on fun until maxfunevals or ftarget is reached, or until
    1e8 * dim function evaluations are conducted.
    dim = len(x)
   maxfunevals = min(1e8 * dim, maxfunevals)
   popsize = min(maxfunevals, 200)
   fbest = np.inf
    for iter in range(0, int(np.ceil(maxfunevals / popsize))):
        xpop = 10. * np.random.rand(popsize, dim) - 5.
        fvalues = fun(xpop)
        idx = np.argsort(fvalues)
        if fbest > fvalues[idx[0]]:
            fbest = fvalues[idx[0]]
            xbest = xpop[idx[0]]
        if fbest < ftarget: # task achieved</pre>
            break
    return xbest
t0 = time.time()
np.random.seed(int(t0))
f = fgeneric.LoggingFunction(datapath, **opts)
for dim in dimensions: # small dimensions first, for CPU reasons
    for fun_id in function_ids:
        for iinstance in instances:
            f.setfun(*bbobbenchmarks.instantiate(fun_id, iinstance=iinstance))
```

```
# independent restarts until maxfunevals or ftarget is reached
        for restarts in xrange(maxrestarts + 1):
            if restarts > 0:
                f.restart('independent restart') # additional info
            run_optimizer(f.evalfun, dim, eval(maxfunevals) - f.evaluations,
                          f.ftarget)
            if (f.fbest < f.ftarget</pre>
                or f.evaluations + eval(minfunevals) > eval(maxfunevals)):
                break
        f.finalizerun()
        print(' f%d in %d-D, instance %d: FEs=%d with %d restarts, '
              'fbest-ftarget=%.4e, elapsed time [h]: %.2f'
              % (fun_id, dim, iinstance, f.evaluations, restarts,
                 f.fbest - f.ftarget, (time.time()-t0)/60./60.)
                 date and time: %s' % (time.asctime())
print '---- dimension %d-D done ----' % dim
```

To run the script in exampleexperiment.py (it takes only about a minute), execute the following command from a command line of your operating system:

```
$ python path_to_postproc_code/exampleexperiment.py

f1 in 2-D, instance 1: FEs=20 with 0 restarts, fbest-ftarget=1.6051e+00, elapsed time [h]: 0.00
...
    f24 in 40-D, instance 15: FEs=400 with 0 restarts, fbest-ftarget=1.1969e+03, elapsed time [h]: 0
    date and time: ...
---- dimension 40-D done ----
```

This generates experimental data in folder PUT\_MY\_BBOB\_DATA\_PATH.

To run your own experiments, you will need to:

- 1. modify the method run\_optimizer to call your optimizer.
- 2. customize the experiment information: put the datapath, say 'advanced-random-search' instead of 'PUT\_MY\_BBOB\_DATA\_PATH', put the optimizer name instead of 'PUT ALGORITHM NAME', add a description instead of 'PUT MORE DETAILED INFORMATION, PARAMETER SETTINGS ETC'
- 3. after a first test run, successively increase the maximum number of function evaluations maxfunevals, subject to the available CPU resources

# 4.2 Post-Processing

For more details, see *Post-Processing with COCO*.

To post-process the data in folder advanced-random-search, execute the python post-processing script from a command line of your operating system:

```
$ python path_to_bbob_pproc/bbob_pproc/rungeneric.py advanced-random-search

BBOB Post-processing: will generate output data in folder ppdata
    this might take several minutes.

BBOB Post-processing: will generate post-processing data in folder ppdata/advanced-random-search
```

```
this might take several minutes.
Loading best algorithm data from BBOB-2009... done.
Scaling figures done.
TeX tables (draft) done. To get final version tables, please use the -f option ECDF graphs done.
ERT loss ratio figures and tables done.
Output data written to folder ppdata/advanced-random-search.
```

The bbob\_pproc folder is located in the python folder of the COCO distribution. This command generates output figures and tables in the folder ppdata/advanced-random-search.

# 4.3 Write/Compile an Article

LaTeX files are provided as template articles. These files include the figures and tables generated by the post-processing during LaTeX compilation.

The template corresponding to the post-processed figures and tables for the data of *one* optimizer on the *noiseless* testbed is template1generic.tex.

We assume that the output folder of the post-processing, by default ppdata, and the template file are in the current working directory. The path to the figures and tables is set to the default but can be edited in the LaTeX file:

```
\newcommand{\bbobdatapath} {ppdata/}
```

Also the name of the algorithm as it appears in figure captions can be edited (by default it is implied from the original data folder).

```
\renewcommand{\algname} {Advanced Random Search}
```

Then, compile the file using LaTeX, for example:

```
$ pdflatex templatelgeneric
This is pdfTeX...
entering extended mode
(./templatelgeneric.tex
...
Output written on templatelgeneric.pdf (...).
Transcript written on templatelgeneric.log.
```

This generates document templatelgeneric.pdf in the current working directory. On some systems, the combination latex and dvipdf might produce much better results.

### **BLACK-BOX OPTIMIZATION BENCHMARKING PROCEDURE**

COCO has been used in several workshops during the GECCO conference since 2009 (BBOB-2009).

For these workshops, a testbed of 24 noiseless functions and another of 30 noisy functions are provided. Descriptions of the functions can be found at http://coco.gforge.inria.fr/doku.php?id=downloads

This section describes the setup of the experimental procedures and their rationales, giving the guidelines to produce an article for a GECCO-BBOB workshop using COCO.

- Symbols, Constants, and Parameters
  - Rationale for the Choice of Ntrial = 15
  - Rationale for the Choice of f<sub>target</sub>
- Benchmarking Experiment
  - Input to the Algorithm and Initialization
  - Termination Criteria and Restarts
- Time Complexity Experiment
- Parameter Setting and Tuning of Algorithms
- Performance Measurement
  - Fixed-Cost versus Fixed-Target Scenario
  - Expected Running Time
  - Bootstrapping
  - Empirical Cumulative Distribution Functions

# 5.1 Symbols, Constants, and Parameters

For the workshops, some constants were set:

D = 2; 3; 5; 10; 20; 40 is search space dimensionalities used for all functions.

Ntrial = 15 is the number of trials for each single setup, i.e. each function and dimensionality. The performance is evaluated over all trials.

 $\Delta f = 10^{-8}$  precision to reach, that is, a difference to the smallest possible function value  $f_{\rm opt}$ .

 $f_{\mathrm{target}} = f_{\mathrm{opt}} + \Delta f$  is target function value to reach for different  $\Delta f$  values. The final, smallest considered target function value is  $f_{\mathrm{target}} = f_{\mathrm{opt}} + 10^{-8}$ , but also larger values for  $f_{\mathrm{target}}$  are evaluated.

The Ntrial runs are conducted on different instances of the functions.

#### 5.1.1 Rationale for the Choice of Ntrial = 15

All functions can be instantiated in different "versions" (with different location of the global optimum and different optimal function value). Overall Ntrial runs are conducted on different instantiations (in the 2009 setup each instance was repeated three times). The parameter Ntrial, the overall number of trials on each function/dimension pair, determines the minimal measurable success rate and influences the overall necessary CPU time. Compared to a standard setup for testing stochastic search procedures, we have chosen a small value, Ntrial = 15. Consequently, within the same CPU-time budget, single trials can be longer and conduct more function evaluations (until  $f_{\rm target}$  is reached). If an algorithm terminates before  $f_{\rm target}$  is reached, longer trials can simply be achieved by independent multistarts. Because these multistarts are conducted within each trial, more sophisticated restart strategies are feasible. Within-trial multistarts never impair the used performance measures and are encouraged. Finally, 15 trials are sufficient to make relevant performance differences statistically significant.  $^1$ 

## 5.1.2 Rationale for the Choice of f<sub>target</sub>

The initial search domain and the target function value are an essential part of the benchmark function definition. Different target function values might lead to different characteristics of the problem to be solved, besides that larger target values are invariably less difficult to reach. Functions might be easy to solve up to a function value of 1 and become intricate for smaller target values. We take records for a larger number of predefined target values, defined relative to the known optimal function value  $f_{\rm opt}$  and in principle unbounded from above. The chosen value for the final (smallest)  $f_{\rm target}$  is somewhat arbitrary. Reasonable values can change by simple modifications in the function definition. In order to safely prevent numerical precision problems, the final target is  $f_{\rm target} = f_{\rm opt} + 10^{-8}$ .

# 5.2 Benchmarking Experiment

The real-parameter search algorithm under consideration is run on a testbed of benchmark functions to be minimized (the implementation of the functions is provided in C/C++, Java, MATLAB/Octave and Python). On each function and for each dimensionality Ntrial trials are carried out (see also *Rationale for the Choice of Ntrial* = 15). Different function *instances* can be used.

The exampleexperiment. \* code template is provided to run this experiment. For BBOB-2012, the instances are 1 to 5 and 21 to 30.

# 5.2.1 Input to the Algorithm and Initialization

An algorithm can use the following input:

- 1. the search space dimensionality D
- 2. the search domain; all functions of BBOB are defined everywhere in  $\mathbb{R}^D$  and have their global optimum in  $[-5,5]^D$ . Most BBOB functions have their global optimum in the range  $[-4,4]^D$  which can be a reasonable setting for initial solutions.
- 3. indication of the testbed under consideration, i.e. different algorithms and/or parameter settings can be used for the noise-free and the noisy testbed.
- 4. the final target precision delta-value  $\Delta f=10^{-8}$  (see above), in order to implement effective termination and restart mechanisms (which should also prevent early termination)
- 5. the target function value  $f_{\text{target}}$ , however provided *only* for conclusive (final) termination of trials, in order to reduce the overall CPU requirements. The target function value must not be used as algorithm input otherwise (not even to trigger within-trial restarts).

<sup>&</sup>lt;sup>1</sup> If the number of trials is chosen *much* larger, small and therefore irrelevant performance differences become statistically significant.

Based on these input parameters, the parameter setting and initialization of the algorithm is entirely left to the user. As a consequence, the setting shall be identical for all benchmark functions of one testbed (the function identifier or any known characteristics of the function are, for natural reasons, not allowed as input to the algorithm, see also Section *Parameter Setting and Tuning of Algorithms*).

#### 5.2.2 Termination Criteria and Restarts

Algorithms with any budget of function evaluations, small or large, are considered in the analysis of the results. Exploiting a larger number of function evaluations increases the chance to achieve better function values or even to solve the function up to the final  $f_{\rm target}$ . In any case, a trial can be conclusively terminated if  $f_{\rm target}$  is reached. Otherwise, the choice of termination is a relevant part of the algorithm: the termination of unsuccessful trials affects the performance. To exploit a large number of function evaluations effectively, we suggest considering a multistart procedure, which relies on an interim termination of the algorithm.

Independent restarts do not change the main performance measure, *expected running time* (ERT, see Appendix *Expected Running Time*) to hit a given target. Independent restarts mainly improve the reliability and "visibility" of the measured value. For example, using a fast algorithm with a small success probability, say 5% (or 1%), chances are that not a single of 15 trials is successful. With 10 (or 90) independent restarts, the success probability will increase to 40% and the performance will become visible. At least four to five (here out of 15) successful trials are desirable to accomplish a stable performance measurement. This reasoning remains valid for any target function value (different values are considered in the evaluation).

Restarts either from a previous solution, or with a different parameter setup, for example with different (increasing) population sizes, might be considered, as it has been applied quite successful [Auger:2005a] [Harik:1999].

Choosing different setups mimics what might be done in practice. All restart mechanisms are finally considered as part of the algorithm under consideration.

# **5.3 Time Complexity Experiment**

In order to get a rough measurement of the time complexity of the algorithm, the overall CPU time is measured when running the algorithm on  $f_8$  (Rosenbrock function) of the BBOB testbed for at least a few tens of seconds (and at least a few iterations). The chosen setup should reflect a "realistic average scenario". If another termination criterion is reached, the algorithm is restarted (like for a new trial). The *CPU-time per function evaluation* is reported for each dimension. The time complexity experiment is conducted in the same dimensions as the benchmarking experiment. The chosen setup, coding language, compiler and computational architecture for conducting these experiments are described.

The exampletiming.\* code template is provided to run this experiment. For CPU-inexpensive algorithms the timing might mainly reflect the time spent in function fgeneric.

# 5.4 Parameter Setting and Tuning of Algorithms

The algorithm and the used parameter setting for the algorithm should be described thoroughly. Any tuning of parameters to the testbed should be described and the approximate number of tested parameter settings should be given.

On all functions the very same parameter setting must be used (which might well depend on the dimensionality, see Section *Input to the Algorithm and Initialization*). That means, *a priori* use of function-dependent parameter settings is prohibited (since 2012). In other words, the function ID or any function characteristics (like separability, multimodality, ...) cannot be considered as input parameter to the algorithm. Instead, we encourage benchmarking different

<sup>&</sup>lt;sup>2</sup> The easiest functions of BBOB can be solved in less than 10D function evaluations, while on the most difficult functions a budget of more than  $1000D^2$  function evaluations to reach the final  $f_{\text{target}} = f_{\text{opt}} + 10^{-8}$  is expected.

parameter settings as "different algorithms" on the entire testbed. In order to combine different parameter settings, one might use either multiple runs with different parameters (for example restarts, see also Section *Termination Criteria and Restarts*), or use (other) probing techniques for identifying function-wise the appropriate parameters online. The underlying assumption in this experimental setup is that also in practice we do not know in advance whether the algorithm will face  $f_1$  or  $f_2$ , a unimodal or a multimodal function... therefore we cannot adjust algorithm parameters a priori  $^3$ .

## 5.5 Performance Measurement

We advocate performance measures that are:

- quantitative, ideally with a ratio scale (opposed to interval or ordinal scale) <sup>4</sup> and with a wide variation (i.e., for example, with values ranging not only between 0.98 and 1.0)
- well-interpretable, in particular by having a meaning and semantics attached to the numbers
- relevant with respect to the "real world"
- as simple as possible

For these reasons we measure "running times" to reach a target function value, denoted as fixed-target scenario in the following.

### 5.5.1 Fixed-Cost versus Fixed-Target Scenario

Two different approaches for collecting data and making measurements from experiments are schematically depicted in Figure *Horizontal vs Vertical View*.

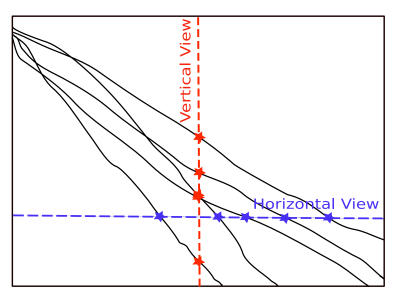


Figure 5.1: Horizontal vs Vertical View

Illustration of fixed-cost view (vertical cuts) and fixed-target view (horizontal cuts). Black lines depict the best function value plotted versus number of function evaluations.

<sup>&</sup>lt;sup>3</sup> In contrast to most other function properties, the property of having noise can usually be verified easily. Therefore, for noisy functions a *second* testbed has been defined. The two testbeds can be approached *a priori* with different parameter settings or different algorithms.

<sup>&</sup>lt;sup>4</sup> Wikipedia gives a reasonable introduction to scale types.

**Fixed-cost scenario (vertical cuts)** Fixing a number of function evaluations (this corresponds to fixing a cost) and measuring the function values reached for this given number of function evaluations. Fixing search costs can be pictured as drawing a vertical line on the convergence graphs (see Figure Horizontal vs Vertical View where the line is depicted in red).

Fixed-target scenario (horizontal cuts) Fixing a target function value and measuring the number of function evaluations needed to reach this target function value. Fixing a target can be pictured as drawing a horizontal line in the convergence graphs (Figure Horizontal vs Vertical View where the line is depicted in blue).

It is often argued that the fixed-cost approach is close to what is needed for real word applications where the total number of function evaluations is limited. On the other hand, also a minimum target requirement needs to be achieved in real world applications, for example, getting (noticeably) better than the currently available best solution or than a competitor.

For benchmarking algorithms we prefer the fixed-target scenario over the fixed-cost scenario since it gives quantitative and interpretable data: the fixed-target scenario (horizontal cut) measures a time needed to reach a target function value and allows therefore conclusions of the type: Algorithm A is two/ten/hundred times faster than Algorithm B in solving this problem (i.e. reaching the given target function value). The fixed-cost scenario (vertical cut) does not give quantitatively interpretable data: there is no interpretable meaning to the fact that Algorithm A reaches a function value that is two/ten/hundred times smaller than the one reached by Algorithm B, mainly because there is no a priori evidence how much more difficult it is to reach a function value that is two/ten/hundred times smaller. This, indeed, largely depends on the specific function and on the specific function value reached. Furthermore, for algorithms that are invariant under certain transformations of the function value (for example under order-preserving transformations as algorithms based on comparisons like DE, ES, PSO), fixed-target measures can be made invariant under these transformations by simply choosing different target values while fixed-cost measures require the transformation of all resulting data.

### 5.5.2 Expected Running Time

We use the expected running time (ERT, introduced in [Price:1997] as ENES and analyzed in [Auger:2005b] as success performance) as most prominent performance measure. The Expected Running Time is defined as the expected number of function evaluations to reach a target function value for the first time. For a non-zero success rate  $p_s$ , the ERT computes to:

$$ERT(f_{target}) = RT_{S} + \frac{1 - p_{s}}{p_{s}} RT_{US}$$

$$p_{s}RT_{S} + (1 - p_{s})RT_{US}$$
(5.1)

$$= \frac{p_{\rm s}RT_{\rm S} + (1 - p_{\rm s})RT_{\rm US}}{p_{\rm s}}$$

$$= \frac{\#FEs(f_{\rm best} \ge f_{\rm target})}{\#succ}$$
(5.2)

$$= \frac{\#\text{FEs}(f_{\text{best}} \ge f_{\text{target}})}{\#\text{succ}}$$
 (5.3)

where the running times  $RT_S$  and  $RT_{US}$  denote the average number of function evaluations for successful and unsuccessful trials, respectively (zero for none respective trial), and  $p_s$  denotes the fraction of successful trials. Successful trials are those that reached  $f_{\rm target}$ ; evaluations after  $f_{\rm target}$  was reached are disregarded. The  $\#{\rm FEs}(f_{\rm best}({\rm FE}) \geq$  $f_{\rm target}$ ) is the number of function evaluations conducted in all trials, while the best function value was not smaller than  $f_{\text{target}}$  during the trial, i.e. the sum over all trials of:

$$\max\{\text{FE s.t. } f_{\text{best}}(\text{FE}) \ge f_{\text{target}}\}.$$

The #succ denotes the number of successful trials. ERT estimates the expected running time to reach  $f_{\text{target}}$ [Auger:2005b], as a function of  $f_{\text{target}}$ . In particular, RT<sub>S</sub> and  $p_{\text{s}}$  depend on the  $f_{\text{target}}$  value. Whenever not all trials were successful, ERT also depends (strongly) on the termination criteria of the algorithm.

### 5.5.3 Bootstrapping

The ERT computes a single measurement from a data sample set (in our case from Ntrial optimization runs). Bootstrapping [Efron:1993] can provide a dispersion measure for this aggregated measurement: here, a "single data sample" is derived from the original data by repeatedly drawing single trials with replacement until a successful trial is drawn. The running time of the single sample is computed as the sum of function evaluations in the drawn trials (for the last trial up to where the target function value is reached) [Auger:2005b] [Auger:2009]. The distribution of the bootstrapped running times is, besides its displacement, a good approximation of the true distribution. We provide some percentiles of the bootstrapped distribution.

### 5.5.4 Empirical Cumulative Distribution Functions

We exploit the "horizontal and vertical" viewpoints introduced in the last Section *Fixed-Cost versus Fixed-Target Scenario*. In Figure *ECDF* we plot the ECDF (Empirical Cumulative Distribution Function) <sup>5</sup> of the intersection point values (stars in Figure *Horizontal vs Vertical View*) for 450 trials.

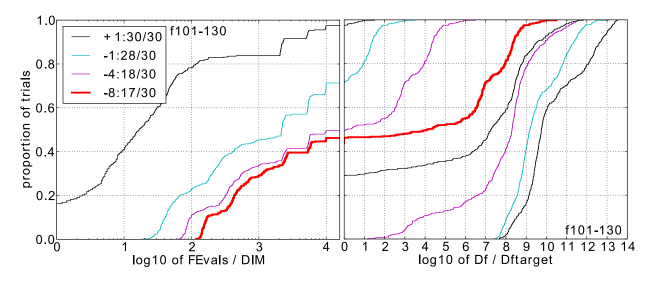


Figure 5.2: ECDF

Illustration of empirical (cumulative) distribution functions (ECDF) of running length (left) and precision (right) arising respectively from the fixed-target and the fixed-cost scenarios in Figure *Horizontal vs Vertical View*. In each graph the data of 450 trials are shown. Left subplot: ECDF of the running time (number of function evaluations), divided by search space dimension D, to fall below  $f_{\rm opt} + \Delta f$  with  $\Delta f = 10^k$ , where k = 1, -1, -4, -8 is the first value in the legend. Data for algorithms submitted for BBOB 2009 and  $\Delta f = 10^{-8}$  are represented in the background in light brown. Right subplot: ECDF of the best achieved precision  $\Delta f$  divided by  $10^k$  (thick red and upper left lines in continuation of the left subplot), and best achieved precision divided by  $10^{-8}$  for running times of D, 100 D, 1000 D... function evaluations (from the rightmost line to the left cycling through black-cyan-magenta-black).

A cutting line in Figure Horizontal vs Vertical View corresponds to a "data" line in Figure ECDF, where 450 (30 x 15) convergence graphs are evaluated. For example, the thick red graph in Figure ECDF shows on the left the distribution of the running length (number of function evaluations) [Hoos:1998] for reaching precision  $\Delta f = 10^{-8}$  (horizontal cut). The graph continues on the right as a vertical cut for the maximum number of function evaluations, showing the distribution of the best achieved  $\Delta f$  values, divided by  $10^{-8}$ . Run length distributions are shown for different target precisions  $\Delta f$  on the left (by moving the horizontal cutting line up- or downwards). Precision distributions are shown

<sup>&</sup>lt;sup>5</sup> The empirical (cumulative) distribution function  $F: \mathbb{R} \to [0,1]$  is defined for a given set of real-valued data S, such that F(x) equals the fraction of elements in S which are smaller than x. The function F is monotonous and a lossless representation of the (unordered) set S.

for different fixed number of function evaluations on the right. Graphs never cross each other. The y-value at the transition between left and right subplot corresponds to the success probability. In the example, just under 50% for precision  $10^{-8}$  (thick red) and just above 70% for precision  $10^{-1}$  (cyan).

### **RUNNING EXPERIMENTS WITH COCO**

COCO provides an interface for running experiments. This interface fgeneric has been ported in different languages:

• in Python, Matlab/GNU Octave, C/C++, R and Java,

# 6.1 exampleexperiment and exampletiming

In each language, two example scripts are provided. Below are the example scripts in Matlab/GNU Octave:

• exampleexperiment runs an experiment on one testbed,

```
% runs an entire experiment for benchmarking MY_OPTIMIZER
  % on the noise-free testbed. fgeneric.m and benchmarks.m
  % must be in the path of Matlab/Octave
  % CAPITALIZATION indicates code adaptations to be made
  addpath ('PUT_PATH_TO_BBOB/matlab'); % should point to fqeneric.m etc.
  datapath = 'PUT_MY_BBOB_DATA_PATH'; % different folder for each experiment
  % opt.inputFormat = 'row';
  opt.algName = 'PUT ALGORITHM NAME';
  opt.comments = 'PUT MORE DETAILED INFORMATION, PARAMETER SETTINGS ETC';
10
  maxfunevals = '10 * dim'; % 10*dim is a short test-experiment taking a few minutes
11
                             % INCREMENT maxfunevals successively to larger value(s)
12
  minfunevals = 'dim + 2'; % PUT MINIMAL SENSIBLE NUMBER OF EVALUATIONS for a restart
13
  maxrestarts = 1e4;
                           % SET to zero for an entirely deterministic algorithm
  dimensions = [2, 3, 5, 10, 20, 40]; % small dimensions first, for CPU reasons
  functions = benchmarks('FunctionIndices'); % or benchmarksnoisy(...)
```

```
instances = [1:5, 31:40]; % 15 function instances
19
   more off; % in octave pagination is on by default
20
21
22
   t0 = clock;
   rand('state', sum(100 * t0));
23
24
   for dim = dimensions
25
     for ifun = functions
26
       for iinstance = instances
27
         fgeneric ('initialize', ifun, iinstance, datapath, opt);
28
29
          % independent restarts until maxfunevals or ftarget is reached
30
          for restarts = 0:maxrestarts
31
            if restarts > 0 % write additional restarted info
32
              fgeneric('restart', 'independent restart')
33
            end
34
35
           MY_OPTIMIZER('fgeneric', dim, fgeneric('ftarget'), ...
                         eval(maxfunevals) - fgeneric('evaluations'));
36
            if fgeneric('fbest') < fgeneric('ftarget') || ...</pre>
37
               fgeneric('evaluations') + eval(minfunevals) > eval(maxfunevals)
38
              break:
39
            end
40
          end
41
42
          disp(sprintf([' f%d in %d-D, instance %d: FEs=%d with %d restarts,' ...
43
                         ' fbest-ftarget=%.4e, elapsed time [h]: %.2f'], ...
44
                       ifun, dim, iinstance, ...
45
                       fgeneric('evaluations'), ...
46
47
                       restarts, ...
                       fgeneric('fbest') - fgeneric('ftarget'), ...
48
                       etime(clock, t0)/60/60));
49
50
         fgeneric('finalize');
51
       end
52
                     date and time: ' num2str(clock, ' %.0f')]);
53
       disp(['
     disp(sprintf('---- dimension %d-D done ----', dim));
   end
      • exampletiming runs the CPU-timing experiment.
   % runs the timing experiment for MY_OPTIMIZER. fgeneric.m
   % and benchmarks.m must be in the path of MATLAB/Octave
   addpath('PUT_PATH_TO_BBOB/matlab'); % should point to fgeneric.m etc.
  more off; % in octave pagination is on by default
  timings = [];
   runs = [];
  dims = [];
10
   for dim = [2,3,5,10,20,40]
11
     nbrun = 0;
12
     ftarget = fgeneric('initialize', 8, 1, 'tmp');
13
     while toc < 30 % at least 30 seconds</pre>
15
       MY_OPTIMIZER(@fgeneric, dim, ftarget, 1e5); % adjust maxfunevals
16
       nbrun = nbrun + 1;
17
```

```
end % while
18
     timings(end+1) = toc / fgeneric('evaluations');
19
     dims(end+1) = dim;
                           % not really needed
20
     runs(end+1) = nbrun; % not really needed
21
22
     fgeneric('finalize');
     disp([['Dimensions:' sprintf(' %11d', dims)]; ...
23
           [' runs:' sprintf(' %11d ', runs)]; ...
24
           [' times [s]:' sprintf(' %11.1e ', timings)]]);
25
   end
```

#### 6.1.1 Matlab/GNU Octave

The above example scripts run a complete experiment. The entire interface for running experiments is defined via the function fgeneric.m. fgeneric is *initialized* with a function number, instance number and output data path. fgeneric can then be called to evaluate the test function. The end of a run or trial is signaled by calling fgeneric with the 'finalize' keyword.

#### 6.1.2 C/C++

The interface for running experiments relies on functions such as fgeneric\_initialize, fgeneric\_finalize, fgeneric\_ftarget. The evaluation function is fgeneric\_evaluate for a single vector or fgeneric\_evaluate\_vector for an array of vectors as input.

A specific folder structure is needed for running an experiment. This folder structure can be obtained by un-tarring the archive createfolders.tar.gz and renaming the output folder or alternatively by executing the Python script createfolders.py before executing any experiment program. Make sure createfolders.py is in your current working directory and from the command-line simply execute:

```
$ python createfolders.py FOLDERNAME
FOLDERNAME was created.
```

The code provided can be compiled in C or C++.

#### 6.1.3 R

#### **Quick Installation**

Run the following commands in your R session:

You should now be able to load the package by running

```
library("bbob")
```

If all went well, you can skip down to the next section. Otherwise consulte the detailed instructions which follow.

#### **Detailed Installation**

Before you start, install the required dependencies. At the R prompt enter

```
install.packages(c("BBmisc", "stringr"))
```

This should download and install the two packages. Now download the current BBOB R source package and save it somewhere on your hard drive. The most up-to-date version is always available here, but there should also be a stable version available on the COCO website. Now it's time to install the package. In R, run the following command:

```
install.packages("/path/to/bbob_current.tar.gz", repos=NULL)
```

Note that you will have to adjust the path and possibly the filename to match the location where you stored the downloaded package on your hard drive. On Windows you will also need to have the Rtools installed for this to work since the package contains C code. If you have any problems with this step, do not hesitate to contact me for assistance.

After completing the above steps, you should be able to load the package in R

```
library("bbob")
```

The help page for the *bbo\_benchmark* function should get you started if you do not want to continue reading this introduction.

```
?bbo_benchmark
```

#### Simple experiment

If you already have an optimizer ready to go in R and all you want to do is produce a BBOB dataset for post-processing, then this section will walk you through the required steps. We will use the high-level interface *bbo\_benchmark* in this example. If you want to parallelize your optimization runs, need to perform complex initializations or just want full control, skip down to the next section for a brief tour of the low-level interface.

In this example we will use the L-BFGS-B optimizer included in base R. You will need to adapt parts of the code to suit your optimizer. The first step is to wrap your optimization algorithm in a function with the following signature

```
function(par, fun, lower, upper, max_eval)
```

where *par* will be a numeric vector with the starting point for the optimization, *fun* is the function to be minimized, *lower* and *upper* are the bounds of the box constraints and *max\_eval* is the number of function evaluations left in the allocated budget. What these five parameters mean is best conveyed by an example. Here we wrap the *optim* function included in base R

If your algorithm does not have the notion of an initial parameter setting, you can safely ignore the *par* parameter in your implementation. You might also notice, that we do not strictly adhere to the *max\_eval* limit because the number of iterations is generally not equal to the number of function evaluations for L-BFGS-B. This is OK. *max\_eval* is only a hint to the optimizer how much effort it should put into optimizing the function.

Should your algorithm perform restarts internally it is possible to log these using the *bbob\_log\_restart* function. The function takes exactly one argument, a string describing the reason for the restart.

We are now ready to run our experiment. This is done by calling the bbo\_benchmark function

```
bbo_benchmark(my_optimizer, "l-bfgs-b", "optim_l-bfgs-b")
```

which will perform the BBOB experiments (caution, may take many hours). The first argument passed to bbo\_benchmark is our optimization wrapper from the previous step. Make sure that it has the correct function signature! Next we supply the so called *algorithm id*. This is a short descriptive name for our optimization algorithm. Ideally it should include the package name and version which contains the algorithm. So for for genoud from the rgenoud package, we might use rgenoud::genoud (5.7-3) as the algorithm id. The last required argument is the name of the base directory where the result files will be stored. Again, it is a good idea to include the algorithm name in the directory name. If no other arguments are given, this runs a complete BBO benchmark on the noiseless test functions. This includes all instances (1-5 and 21-30), all dimensions (2, 3, 5, 10, 20, 40). If you do not want to include runs in 40 dimensions or want to use different instances you can change the defaults using the dimensions and instances arguments to bbo\_benchmark. For details, see the manual page for bbo\_benchmark.

If no other budget is specified, bbo\_benchmark will perform random independent restarts of your algorithm until the desired target precision (1e-8) is reached or the default budget of 10000000 function evaluations is exhausted. If you want to reduce the budget, you can by specifiying it as the budget argument to bbo\_benchmark.

To run the required timing experiment, execute the following code:

```
bbo_timing(my_optimizer)
```

It will return a data frame with the relevant timing information.

#### Low-level interface

Will follow soon.

### 6.1.4 Python

25

The interface for running an experiment is figeneric which is used within example experiment.py:

# function\_ids = bbobbenchmarks.noisyIDs if len(argv) < 3 else eval(argv[2])</pre> instances = range(1, 6) + range(31, 41) if len(argv) < 4 else eval(argv[3])

```
#!/usr/bin/env python
   \# -*- coding: utf-8 -*-
2
   """Runs an entire experiment for benchmarking PURE_RANDOM_SEARCH on a testbed.
   CAPITALIZATION indicates code adaptations to be made.
6
   This script as well as files bbobbenchmarks.py and fgeneric.py need to be
   in the current working directory.
   Under unix-like systems:
10
       nohup nice python exampleexperiment.py [data_path [dimensions [functions [instances]]]] > output
11
12
13
   import sys # in case we want to control what to run via command line args
14
   import time
15
   import numpy as np
16
   import fgeneric
17
   import bbobbenchmarks
18
19
   argv = sys.argv[1:] # shortcut for input arguments
20
21
   datapath = 'PUT_MY_BBOB_DATA_PATH' if len(argv) < 1 else argv[0]</pre>
22
23
   dimensions = (2, 3, 5, 10, 20, 40) if len(argv) < 2 else eval(argv[1])
24
   function_ids = bbobbenchmarks.nfreeIDs if len(argv) < 3 else eval(argv[2])</pre>
```

```
28
   opts = dict(algid='PUT ALGORITHM NAME',
29
                comments='PUT MORE DETAILED INFORMATION, PARAMETER SETTINGS ETC')
   maxfunevals = '10 * dim' # 10*dim is a short test-experiment taking a few minutes
                              # INCREMENT maxfunevals SUCCESSIVELY to larger value(s)
32
   minfunevals = 'dim + 2'
                              # PUT MINIMAL sensible number of EVALUATIONS before to restart
33
   maxrestarts = 10000
                              # SET to zero if algorithm is entirely deterministic
34
35
36
37
   def run_optimizer(fun, dim, maxfunevals, ftarget=-np.Inf):
38
        """start the optimizer, allowing for some preparation.
       This implementation is an empty template to be filled
39
40
        .....
41
       # prepare
42
43
       x_start = 8. * np.random.rand(dim) - 4
44
        # call, REPLACE with optimizer to be tested
45
       PURE RANDOM SEARCH(fun, x start, maxfunevals, ftarget)
46
47
   def PURE_RANDOM_SEARCH(fun, x, maxfunevals, ftarget):
48
        """samples new points uniformly randomly in [-5,5]^dim and evaluates
49
       them on fun until maxfunevals or ftarget is reached, or until
50
       1e8 * dim function evaluations are conducted.
51
52
       .....
53
       dim = len(x)
54
       maxfunevals = min(1e8 * dim, maxfunevals)
55
       popsize = min(maxfunevals, 200)
       fbest = np.inf
57
58
       for iter in range(0, int(np.ceil(maxfunevals / popsize))):
59
           xpop = 10. * np.random.rand(popsize, dim) - 5.
60
           fvalues = fun(xpop)
61
           idx = np.argsort(fvalues)
62
           if fbest > fvalues[idx[0]]:
                fbest = fvalues[idx[0]]
64
                xbest = xpop[idx[0]]
65
           if fbest < ftarget: # task achieved</pre>
66
                break
67
68
       return xbest.
69
70
   t0 = time.time()
71
   np.random.seed(int(t0))
72
73
   f = fgeneric.LoggingFunction(datapath, **opts)
74
   for dim in dimensions: # small dimensions first, for CPU reasons
75
       for fun_id in function_ids:
76
            for iinstance in instances:
77
                f.setfun(*bbobbenchmarks.instantiate(fun_id, iinstance=iinstance))
78
79
                # independent restarts until maxfunevals or ftarget is reached
80
                for restarts in xrange(maxrestarts + 1):
81
                    if restarts > 0:
82
                        f.restart('independent restart') # additional info
83
                    run_optimizer(f.evalfun, dim, eval(maxfunevals) - f.evaluations,
84
                                   f.ftarget)
85
```

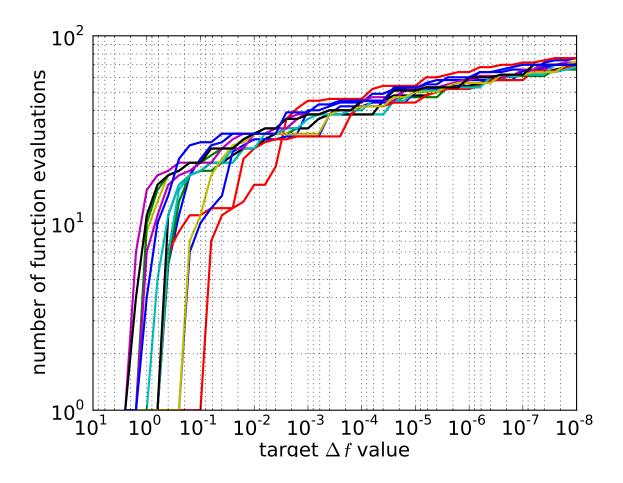
# **6.2 Testing New Functions**

We describe here how to use fgeneric to record experiments on functions that are not part of the BBOB testbeds.

**Note:** This feature is only available in Python for the moment.

Example: log experiment using the Nelder-Mead simplex algorithm (scipy.optimize.fmin) on the sphere function. The following commands from the Python Interpreter does 15 runs of the Nelder-Mead simplex algorithm on the 2-D sphere functions. The data is recorded in folder data in the current working directory.

```
>>> from pylab import *
>>> import fgeneric as fg
>>> import scipy.optimize as so
>>> f = lambda x: sum(i**2 for i in x) # function definition
>>> e = fg.LoggingFunction(datapath='data', algid='Nelder-Mead simplex',
                  comments='x0 uniformly sampled in [0, 1]^2, '
                           'default settings')
>>> for i in range(15): # 15 repetitions
       e.setfun(fun=f, fopt=0., funId='sphere', iinstance='0')
       so.fmin(e.evalfun, x0=rand(2)) # algorithm call
       e.finalizerun()
(<bound method LoggingFunction.evalfun of <fgeneric.LoggingFunction object at [...]>>, 1e-08)
Optimization terminated successfully.
         Current function value: 0.000000
         Iterations: [...]
         Function evaluations: [...]
array([...])
>>> # Display convergence graphs
>>> import bbob_pproc as bb
>>> ds = bb.load('data')
>>> ds.plot()
```



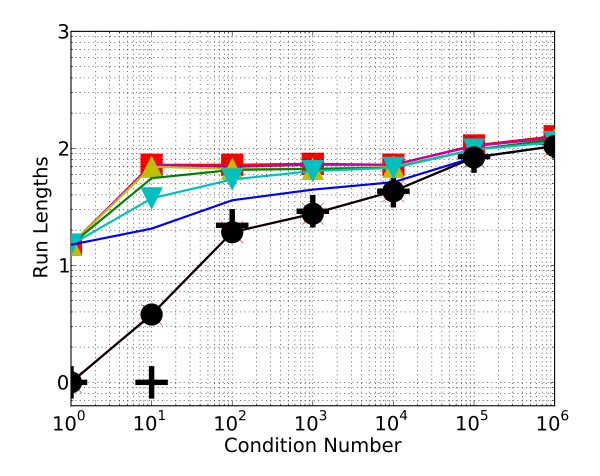
### 6.2.1 Testing Functions with Parameter

Note: This feature is only available in Python for the moment.

Example: log experiment using the BFGS algorithm (scipy.optimize.fmin) on the ellipsoid function with different condition numbers.

The following commands from the Python Interpreter does 15 runs of the BFGS algorithm on the 2-D sphere functions. The data is recorded in folder data in the current working directory and generate

```
so.fmin_bfgs(e.evalfun, x0=np.random.rand(5)) # algorithm call
            e.finalizerun()
(<bound method LoggingFunction.evalfun of <fgeneric.LoggingFunction object at [...]>>, 1e-08)
Optimization terminated successfully.
         Current function value: 0.000000
         Iterations: [...]
         Function evaluations: [...]
         Gradient evaluations: [...]
array([...])
[...]
>>> # plot data
>>> import bbob_pproc as bb
>>> import bbob_pproc.ppfigparam
>>> ds = bb.load('ellipsoid')
>>> bb.ppfigparam.plot(ds, param='condnum')
>>> bb.ppfigparam.beautify()
>>> import matplotlib.pyplot as plt
>>> plt.xlabel('Condition Number')
```



### POST-PROCESSING WITH COCO

# 7.1 Overview of the bbob\_pproc Package

We present here the content of version 10.7 of the bbob\_pproc package.

- rungeneric.py is the main interface of the package that performs different routines listed below,
- rungeneric1.py post-processes data from one single algorithm and outputs figures and tables included in the
  templates template1generic.tex, noisytemplate1generic.tex,
- rungeneric2.py post-processes data from two algorithms using modules from bbob\_pproc.comp2
   and outputs comparison figures and tables included in the template template2generic.tex,
   noisytemplate2generic.tex,
- rungenericmany.py post-processes data from 1 or more algorithms <sup>1</sup> using modules from bbob\_pproc.compall and outputs comparison figures and tables included in the template, template3generic.tex, noisytemplate3generic.tex
- genericsettings.py defines generic settings for the output figures and tables during module import,
- config.py configuration and modification of settings depending on input parameters while the module is running
- bbob\_pproc.pproc defines the classes bbob\_pproc.pproc.DataSetList and
  bbob\_pproc.pproc.DataSet which are the main data structures that we use to gather the experimental raw data,
- **bbob\_pproc.dataoutput** contain routine to output instances of bbob\_pproc.pproc.DataSet in Python-formatted data files,
- **bbob\_pproc.readalign, bbob\_pproc.toolsstats** contain routines for the post-processing of the raw experimental data,
- bbob\_pproc.pptex defines some routines for generating TeXtables,
- bbob\_pproc.ppfig defines some routines for generating figures,
- bbob\_pproc.ppfigdim, bbob\_pproc.pptable, bbob\_pproc.pprldistr, bbob\_pproc.pplogloss are used to produce figures and tables presenting the results of one algorithm,
- bbob\_pproc.compall is a sub-package which contains modules for the comparison of the performances of algorithms, routines in this package can be called using the interface of rungenericmany.py,
- **bbob\_pproc.comp2** is a sub-package which contains modules for the comparison of the performances of two algorithms, routines in this package can be called using the interface of rungeneric2.py.

<sup>&</sup>lt;sup>1</sup> for more than ten algorithm, the template shall be modified,

# 7.2 Standard Use of the bbob\_pproc Package

The main interface is in rungeneric.py and behaves differently depending on the number of folders given as input arguments (each corresponding to the data of a different algorithm).

If one folder, DATAPATH, containing all data generated by the experiments for *one* algorithm is in the current working directory, the following command executes the post-processing:

```
$ python path_to_postproc_code/bbob_pproc/rungeneric.py DATAPATH

BBOB Post-processing: will generate output data in folder ppdata
    this might take several minutes.

BBOB Post-processing: will generate post-processing data in folder ppdata/DATAPATH
    this might take several minutes.

Loading best algorithm data from BBOB-2009... done.

Scaling figures done.

TeX tables (draft) done. To get final version tables, please use the -f option

ECDF graphs done.

ERT loss ratio figures and tables done.

Output data written to folder ppdata/DATAPATH.
```

The above command creates the folder with the default name <code>ppdata/DATAPATH</code> in the current working directory, which contains the post-processed data in the form of figures and LaTeX files for the tables. This process might take a few minutes. Help on the use of the rungeneric.py script, which corresponds to the execution of method <code>bbob\_pproc.rungeneric.main</code>, can be obtained by the command <code>python path\_to\_postproc\_code/bbob\_pproc/rungeneric.py -h</code>.

To run the post-processing directly from a Python shell, the following commands need to be executed:

```
>>> import bbob_pproc as bb
>>> bb.rungeneric.main('DATAPATH'.split())

BBOB Post-processing: will generate output data in folder ppdata this might take several minutes.
...
Output data written to folder ppdata/DATAPATH.
>>>
```

This import command requires that the path to the package bbob\_pproc is in the search path of Python (e.g. the current folder or use sys.path.append to append the path).

The resulting folder ppdata/DATAPATH now contains a number of tex, eps, pdf files.

# 7.2.1 Expensive Setting

rungeneric.py accepts quite a few parameters, consider

```
$ python path_to_postproc_code/bbob_pproc/rungeneric.py --help
$ python path_to_postproc_code/bbob_pproc/rungeneric1.py --help
```

Since version 13.01, data with a smaller overall budget (number of overall evaluations less than about 1000 times dimension) are processed differently. This processing style can be enforced using the --expensive option. In this case the targets are chosen based on function values reached by the best 2009 algorithm after some fixed budgets. At this time, the option only applies to the single algorithm processing (via rungeneric.py or rungenericl.py).

### 7.2.2 Comparison of Algorithms

The sub-packages bbob\_pproc.compall and bbob\_pproc.comp2 provide facilities for the generation of tables and figures comparing the performances of algorithms.

The post-processing works with data folders as input argument, with each folder corresponding to the data of an algorithm. Supposing you have the folders ALG1, ALG2 and ALG3 containing the data of algorithms ALG1, ALG2 and ALG3, you will need to execute from the command line:

```
$ python path_to_postproc_code/bbob_pproc/rungeneric.py ALG1 ALG2 ALG3
BBOB Post-processing: will generate output data in folder ppdata
    this might take several minutes.
...
Output data written to folder ppdata.
```

This assumes that ALG1, ALG2 and ALG3 are valid directories and containing either pickle files (which contain Python-formatted data) or the raw experiment data, in particular files with info, dat and tdat extensions. Running the aforementioned command will generate the folder ppdata containing comparison figures and tables.

Outputs appropriate to the comparison of only two algorithms can be obtained by executing the following from the command line:

```
$ python path_to_postproc_code/bbob_pproc/rungeneric.py ALGO ALG1
BBOB Post-processing: will generate output data in folder ppdata
this might take several minutes.
...
Output data written to folder ppdata.
```

This assumes the folders ALGO and ALGO are in the current working directory. Running the aforementioned command will generate the folder ppdata containing the comparison figures.

**Note:** Instead of using the rungeneric.py interface, the user can directly use the sub-routines rungeneric1.py, rungeneric2.py or rungenericmany.py to generate some post-processing output. These sub-routines are to be used in the same way as rungeneric.py.

## 7.2.3 Use of the LaTeX Templates

Several LaTeX files are provided as template articles. These files compile the figures and tables generated by the post-processing in a single document. The templates have different layouts. The files with *noisy* in their names aggregate results on the noisy testbed from BBOB while the others compile results on the noiseless testbed.

The purpose of \*template1\* templates is to present the results of one algorithm on a BBOB testbed (obtained with bbob\_pproc.rungeneric1). The templates \*template2\* and \*template3\* respectively for the comparison of two (bbob\_pproc.rungeneric2) and two or more algorithms (bbob\_pproc.rungenericmany). Similarly, names with 'cmp' refer to LaTeX templates comparing two algorithms and templates with 'many' in their names can be used to compare two or more algorithms.

The usual way of using these LaTeX templates is to:

- 1. copy the necessary template and associated style files (if necessary) in the current working directory, i.e. in the same location as the output folder of the post-processing ppdata.
- 2. (optional) edit the template, in particular the algorithm name. The default data folder name is defined as \newcommand{\bbobdatapath}{ppdata/}. In ppdata the file bbob\_pproc\_commands.tex defines relevant LaTeX commands, in particular the folder in ppdata where the latest post-processed data are found.

3. compile with LaTeX.

**Note:** bbob\_pproc.rungeneric will generate *comparison* figures and tables at the base of the output folder and individual results of each algorithm for which data were provided in subfolders. The \*template1\* templates can be used by pointing to any of these subfolders with \newcommand{\algfolder}{desired-subfolder}.

The templates are compiled using LaTeX, for instance:

```
$ pdflatex template1generic

This is pdfTeX...
entering extended mode
(./template1generic.tex
...
Output written on template1generic.dvi (...).
Transcript written on template1generic.log.
```

This generates document template1generic.pdf in the current working directory.

# 7.3 Post-Processing from the Python Interpreter

- Pylab
- Pre-requisites
- · Start of the Session

The package bbob\_pproc can also be used interactively. A Matlab-like interaction is available when using the Python interpreter in Pylab mode (see below).

### 7.3.1 Pylab

Installing matplotlib also installs pylab, providing a Matlab-style interface to the functionalities in numpy and matplotlib. To start pylab mode from the Python interpreter, type:

```
>>> from pylab import *
>>>
```

which

- imports the matplotlib.pyplot and numpy packages, and
- switches the interactive mode of matplotlib on.

**Note:** We recommend using iPython, an environment enhancing the Python interpreter for interactive use. The pylab mode can then be started from the command line by typing:

```
$ ipython -pylab
Python 2...
Type "copyright", "credits" or "license" for more information.

IPython 0.X -- An enhanced Interactive Python.
? -> Introduction and overview of IPython's features.
```

```
%quickref -> Quick reference.
help -> Python's own help system.
object? -> Details about 'object'. ?object also works, ?? prints more.

Welcome to pylab, a matplotlib-based Python environment.
For more information, type 'help(pylab)'.

In [1]:
```

Some differences with the Python interpreter are:

- the prompt is different: In [1]: versus >>>
- system commands are directly available, like 1s, cd, ...
- the excellent tab completion is directly available
- run myfile.py runs a python script, just as python myfile.py from the OS-shell command line, help plot displays help, and many other so-called magic commands are available
- pasting code into the shell usually works (%doctest\_mode allows to paste code preceded with a >>> prompt).
- typing debug after an error occurs jumps into the execution code into the state before the error was raised, where all variables can be inspected and commands executed like in an interactive python shell.

## 7.3.2 Pre-requisites

For this session, we assume that:

- the folder ~/code contains the folder bbob.v10.74/python/bbob\_pproc extracted from one of the archives. The latter folder contains the Python module bbob\_pproc.
- the following files are downloaded and unarchived in the current working directory:
  - http://coco.lri.fr/BBOB2009/rawdata/BIPOP-CMA-ES\_hansen\_noiseless.tar.gz
  - http://coco.lri.fr/BBOB2009/pythondata/BIPOP-CMA-ES.tar.gz
  - http://coco.lri.fr/BBOB2009/pythondata/NEWUOA.tar.gz

The following describes a step-by-step interactive session with bbob\_pproc.

#### 7.3.3 Start of the Session

First, we start the Python interpreter here by typing python from the command line:

```
$ python
Python 2....
...
Type "help", "copyright", "credits" or "license" for more information.
>>>
```

On Unix systems, enabling the tab-completion in the python shell makes life much more convenient:

```
>>> import rlcompleter
>>> import readline # only available on Unix systems
>>> readline.parse_and_bind("tab: complete")
>>>
```

**Note:** One might like to write these commands into a startup.py file and type

```
>>> exec open('startup.py')
>>>
```

or even better put them into the file pointed to by the PYTHONSTARTUP system environment variable, e.g. ~/.pythonrc.

An alternative is to use ipython or IDLE.

In order to have a matlab-like environment we can "start pylab mode":

```
>>> from pylab import *
>>>
```

An alternative is to use ipython -pylab as noted above.

The bbob\_pproc package is then loaded into memory:

```
>>> import sys
>>> sys.path.append('~/code/bbob.v10.74/python') # where bbob_proc can be found
>>> import bbob_pproc as bb
>>> help(bb)
Help on package bbob_pproc:
[...]
```

will list the bbob\_pproc documentation.

Commands in bbob\_pproc can now be accessed by prepending 'bb.' to command names.

Typing help (bb.cococommands) provides some help on the use of bbob\_pproc in the Python interpreter.

Data is loaded into memory with the load function:

```
>>> ds = bb.load('BIPOP-CMA-ES_hansen_noiseless/bbobexp_f2.info')
Processing BIPOP-CMA-ES_hansen_noiseless/bbobexp_f2.info.
    [...]
Processing ['BIPOP-CMA-ES_hansen_noiseless/data_f2/bbobexp_f2_DIM40.tdat']: 15/15 trials found.
>>> ds
[DataSet(cmaes V3.30.beta on f2 2-D), ..., DataSet(cmaes V3.30.beta on f2 40-D)]
>>> type(ds)
<class 'bbob_pproc.pproc.DataSetList'>
>>> len(ds)
324
>>>
```

The variable ds stores an instance of class pproc.DataSetList represented between square brackets. This instance in ds is a list of pproc.DataSet instances.

```
>>> help(ds)
Help on DataSetList in module bbob_pproc.pproc object:

class DataSetList(__builtin__.list)
    | List of instances of DataSet with some useful slicing functions.
    |
    | Will merge data of DataSet instances that are identical (according to function __eq__ of DataSet).
    |
    | Method resolution order:
    | DataSetList
    | __builtin__.list
```

```
__builtin__.object
Methods defined here:
__init__(self, args=[], verbose=True)
    Instantiate self from a list of inputs.
    Keyword arguments:
    args -- list of strings being either info file names, folder containing
           info files or pickled data files.
    verbose -- controls verbosity.
    Exception:
    Warning -- Unexpected user input.
    pickle.UnpicklingError
append(self, o)
    Redefines the append method to check for unicity.
dictByAlg(self)
    Returns a dictionary of DataSetList instances by algorithm.
    The resulting dict uses algId and comment as keys and the
    corresponding slices of DataSetList as values.
dictByDim(self)
    Returns a dictionary of DataSetList instances by dimensions.
    Returns a dictionary with dimension as keys and the
    corresponding slices of DataSetList as values.
dictByFunc(self)
    Returns a dictionary of DataSetList instances by functions.
    Returns a dictionary with the function id as keys and the
    corresponding slices of DataSetList as values.
dictByFuncGroup(self)
    Returns a dictionary of DataSetList instances by function groups.
    Returns a dictionary with function group names as keys and the
    corresponding slices of DataSetList as values.
dictByNoise(self)
    Returns a dictionary splitting noisy and non-noisy entries.
extend(self, o)
    Extend a DataSetList with elements.
    This method is implemented to prevent problems since append was
    superseded. This method could be the origin of efficiency issue.
info(self, opt='all')
    Display some information onscreen.
    Keyword arguments:
    opt -- changes size of output, can be 'all' (default), 'short'
```

The central data structure in COCO is the DataSet that holds all trials on a single function on a given dimension.

```
>>> help(ds[0])
Help on instance of DataSet in module bbob_pproc.pproc:
class DataSet
| Unit element for the BBOB post-processing.
   One unit element corresponds to data with given algId and comment, a
| funcId, and dimension.
| Class attributes:
       funcId -- function Id (integer)
       dim -- dimension (integer)
       indexFiles -- associated index files (list of strings)
       dataFiles -- associated data files (list of strings)
       comment -- comment for the setting (string)
       targetFuncValue -- target function value (float)
       algId -- algorithm name (string)
       evals -- data aligned by function values (array)
       funvals -- data aligned by function evaluations (array)
       maxevals -- maximum number of function evaluations (array)
      finalfunvals -- final function values (array)
      readmaxevals -- maximum number of function evaluations read from
                       index file (array)
       readfinalFminusFtarget -- final function values - ftarget read
                                 from index file (array)
       pickleFile -- associated pickle file name (string)
       target -- target function values attained (array)
       ert -- ert for reaching the target values in target (array)
       instancenumbers -- list of numbers corresponding to the instances of the
                  test function considered (list of int)
       isFinalized -- list of bool for if runs were properly finalized
   evals and funvals are arrays of data collected from N data sets.
 | Both have the same format: zero-th column is the value on which the
 | data of a row is aligned, the N subsequent columns are either the
 | numbers of function evaluations for evals or function values for
```

```
| funvals.
  Methods defined here:
  __eq__(self, other)
      Compare indexEntry instances.
  __init__(self, header, comment, data, indexfile, verbose=True)
      Instantiate a DataSet.
      The first three input argument corresponds to three consecutive
      lines of an index file (info extension).
      Keyword argument:
      header -- string presenting the information of the experiment
      comment -- more information on the experiment
      data -- information on the runs of the experiment
      indexfile -- string for the file name from where the information come
      verbose -- controls verbosity
  __ne__(self, other)
  __repr__(self)
  computeERTfromEvals(self)
      Sets the attributes ert and target from the attribute evals.
  createDictInstance(self)
      Returns a dictionary of the instances.
      The key is the instance id, the value is a list of index.
  detERT(self, targets)
      Determine the expected running time to reach target values.
      Keyword arguments:
      targets -- list of target function values of interest
      Output:
      list of expected running times corresponding to the targets
  detEvals(self, targets)
      Determine the number of evaluations to reach target values.
      Keyword arguments:
      targets -- list of target function values of interest
      Output:
      list of arrays each corresponding to one value in targets
  generateRLData(self, targets)
      Determine the running lengths for reaching the target values.
      Keyword arguments:
      targets -- list of target function values of interest
      Output:
      dict of arrays, one array has for first element a target
```

```
function value smaller or equal to the element of inputtargets
       considered and has for other consecutive elements the
 corresponding number of function evaluations.
| info(self)
       Return some text info to display onscreen.
| mMaxEvals(self)
       Returns the maximum number of function evaluations.
  nbRuns(self)
       Returns the number of runs.
   pickle(self, outputdir=None, verbose=True)
Save DataSet instance to a pickle file.
       Saves the instance of DataSet to a pickle file. If not specified
       by argument outputdir, the location of the pickle is given by
       the location of the first index file associated to the DataSet.
       This method will overwrite existing files.
  splitByTrials(self, whichdata=None)
       Splits the post-processed data arrays by trials.
Returns a two-element list of dictionaries of arrays, the key of
the dictionary being the instance id, the value being a smaller
post-processed data array corresponding to the instance id.
>>>
```

The load function also works on pickle files or folders. For instance:

```
>>> ds = bb.load('BIPOP-CMA-ES/ppdata_f002_20.pickle')
Unpickled BIPOP-CMA-ES/ppdata_f002_20.pickle.
>>> ds = bb.load('BIPOP-CMA-ES_hansen_noiseless')
Searching in BIPOP-CMA-ES_hansen_noiseless ...
[...]
Processing ['BIPOP-CMA-ES_hansen_noiseless/data_f9/bbobexp_f9_DIM40.tdat']: 15/15 trials found.
>>>
```

To use wildcards for loading only part of the files in a folder, the glob package included in Python standard library can be used:

```
>>> import glob
>>> ds = bb.load(glob.glob('BIPOP-CMA-ES/ppdata_f002_*.pickle'))
>>>
```

Some information on loaded data can be obtained with the info method:

```
>>> ds = bb.load('BIPOP-CMA-ES/ppdata_f002_20.pickle')
>>> ds.info() # display information on DataSetList ds
1 data set(s)
Algorithm(s): cmaes V3.30.beta
Dimension(s): 20
Function(s): 2
Max evals: 20690
    min
                   10
                          med
                                          max
1.0e+01 | 10875 11049 13483 16152
1.0e+00 | 13329 14061 15155 17421
                                        16826
                                        17675
```

```
1.0e-01 |
           15392
                     15581
                              16481
                                        18382
                                                 18463
1.0e-03 |
                                                 19083
           16706
                     17282
                              18162
                                        19081
1.0e-05 |
            17854
                     17855
                              18939
                                        19629
                                                 19831
                     18749
                              19757
1.0e-08 |
            18726
                                        20599
                                                 20678
```

The actual data in a pproc. DataSet instance can be displayed on-screen:

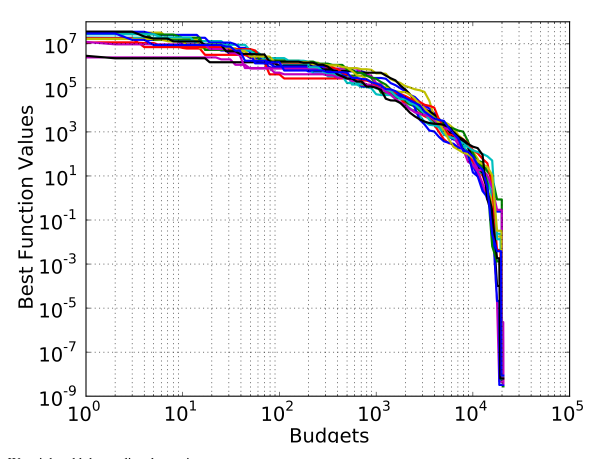
```
>>> d = ds[0] # store the first element of ds in d for convenience
>>> d.funcId # in IPython d.<TAB> displays all available attributes and methods
>>> d.instancenumbers # instance numbers of the available trials
[1, 1, 1, 2, 2, 2, 3, 3, 3, 4, 4, 4, 5, 5, 5]
>> type(d.funvals) # numpy array type is fundamental
<type 'numpy.ndarray'>
>>> d.funvals.shape # we find 95 rows in function value data
(95, 16)
>>> d.funvals[[0,1,-1], :] # let's see the first two and last row
array([[ 1.00000000e+00, 1.91046048e+07, 3.40744851e+07,
         1.16378912e+07, 1.88965407e+07,
                                          2.41571279e+06,
         3.22847770e+07, 2.85601634e+06, 3.51908185e+07,
         2.82405771e+07,
                         1.79074770e+07, 2.71439178e+07,
         1.22112898e+07,
                          1.65690802e+07,
                                           3.57969327e+07,
         2.94479644e+07],
       [ 2.00000000e+00, 1.83130834e+07, 3.40744851e+07,
                          1.88965407e+07,
                                           2.41571279e+06,
         1.16378912e+07,
         3.22847770e+07,
                          2.20298276e+06,
                                            3.51908185e+07,
         2.82405771e+07,
                          1.79074770e+07,
                                            2.71439178e+07,
         9.46258565e+06,
                           1.65690802e+07,
                                            3.57969327e+07,
         2.94479644e+07],
       [ 2.06900000e+04, 4.79855089e-09, 2.78740231e-09,
         5.28442001e-09, 5.17276533e-09, 4.54151916e-09,
         5.04810771e-09, 5.50441826e-09, 3.25114513e-09,
         7.20403648e-09,
                         6.95291646e-09,
                                           5.91512617e-09,
         3.79441190e-09,
                         8.42007353e-09,
                                           6.33018260e-09,
         8.59924398e-0911)
>>> budgets = d.funvals[:, 0] # first column contains budgets
>>> funvals = d.funvals[:, 1:] # stores columns 1,...,15 in funvals
```

The first column (with index 0) of DataSet attribute funvals contains number of function evaluations. The remaining columns contain the best function value precision (difference to the optimal function value) in the respective trial or run.

Graphs of the evolution of the best precision over time for each single run can be displayed as follows <sup>2</sup>:

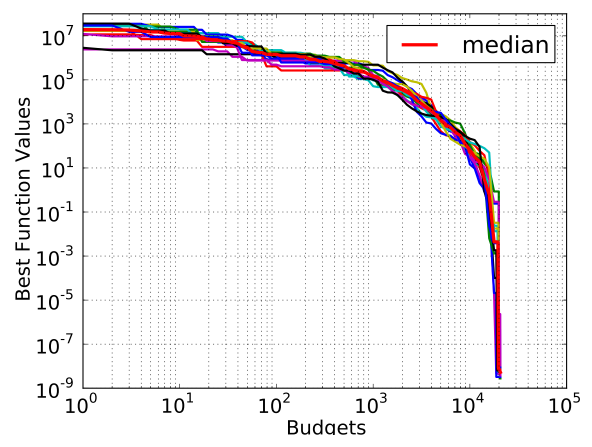
```
>>> for f in funvals.T: # T==transpose: iterate over columns of funvals
... loglog(budgets, f)
...
[<matplotlib.lines.Line2D object at 0x1[...]>]
       [...]
[<matplotlib.lines.Line2D object at 0x1[...]>]
>>> grid()
>>> xlabel('Budgets')
<matplotlib.text.Text object at 0x1[...]>
>>> ylabel('Best Function Values')
<matplotlib.text.Text object at 0x1[...]>
```

<sup>&</sup>lt;sup>2</sup> More information on plot functions from http://matplotlib.sourceforge.net/users/pyplot\_tutorial.html



### We might add the median data points.

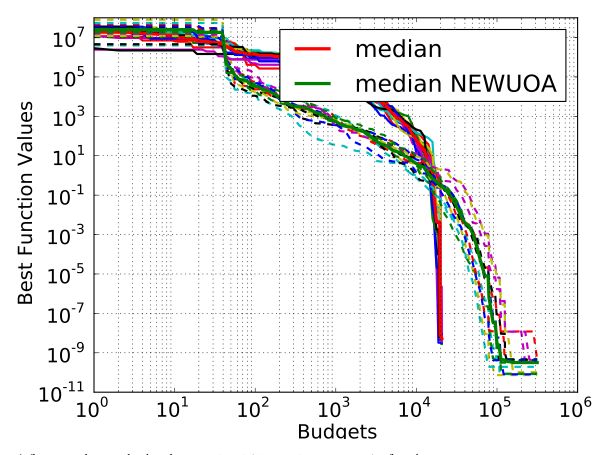
```
>>> loglog(budgets, median(funvals, axis=1), linewidth=3, color='r',
... label='median CMA-ES')
[<matplotlib.lines.Line2D object at 0x1[...]>]
>>> legend() # display legend
<matplotlib.legend.Legend object at 0x1[...]>
```



The options linewidth and color makes the line bold and red. The matplotlib.pyplot.legend function displays a legend for the figure which uses the label option.

We display another data set for comparison:

```
>>> ds1 = bb.load('NEWUOA/ppdata_f002_20.pickle')
>>> d1 = ds1[0]
>>> budgets1 = d1.funvals[:, 0]
>>> funvals1 = d1.funvals[:, 1:]
>>> for i in range(0, funvals1.shape[1]):
... loglog(budgets1, funvals1[:, i], linestyle='--')
...
[<matplotlib.lines.Line2D object at 0x1[...]>]
[...]
[<matplotlib.lines.Line2D object at 0x1[...]>]
>>> loglog(budgets1, median(funvals1, axis=1), linewidth=3,
... color='g', label='median NEWUOA')
[<matplotlib.lines.Line2D object at 0x1[...]>]
>>> legend() # updates legend
<matplotlib.legend.Legend object at 0x1[...]>
```



A figure can be saved using the matplotlib.pyplot.savefig function:

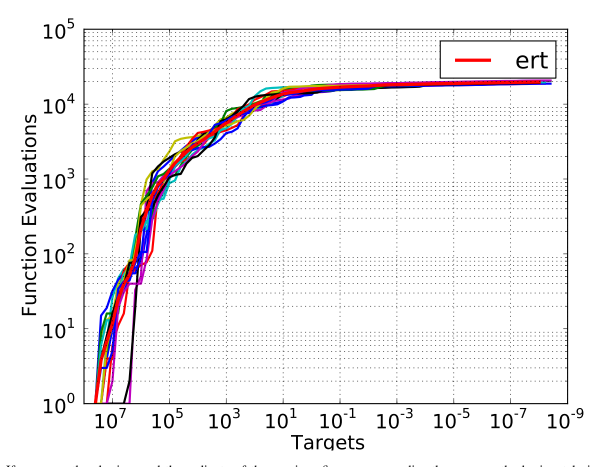
```
>>> savefig('examplefigure') # save active figure as image
```

An image file is then created. The format of the output image file depends on the extension of the file name provided to matplotlib.pyplot.savefig or the default settings of matplotlib.

In the previous figures, it is the evolution of the best function values versus time which is considered, in other words the best function values given a budget (vertical view). Instead, we can consider the horizontal view: determining the run lengths for attaining a target precision. This defines the Expected Run Time performance measure. Please refer to Experimental Set-Up Document for more details.

```
>>> targets = d.evals[:, 0]
>>> evals = d.evals[:, 1:]
>>> nruns = evals.shape[1]
>>> figure()
<matplotlib.figure.Figure object at 0x1[...]
>>> for i in range(0, nruns):
... loglog(targets, evals[:, i])
...
[<matplotlib.lines.Line2D object at 0x1[...]
[...]
[<matplotlib.lines.Line2D object at 0x1[...]>]
>>> grid()
>>> xlabel('Targets')
<matplotlib.text.Text object at 0x1[...]>
```

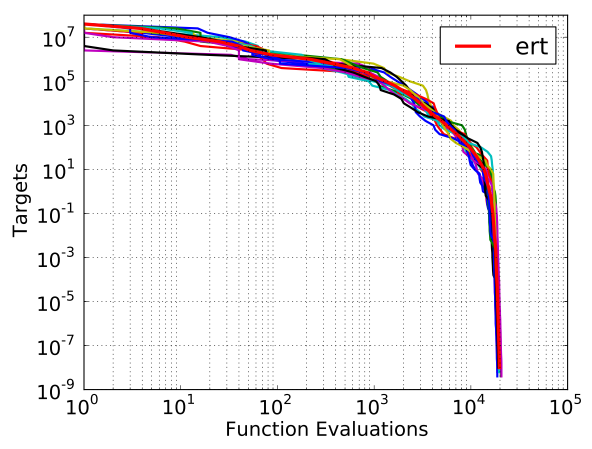
```
>>> ylabel('Function Evaluations')
<matplotlib.text.Text object at 0x1[...]>
>>> loglog(d.target[d.target>=le-8], d.ert[d.target>=le-8], lw=3,
... color='r', label='ert')
[<matplotlib.lines.Line2D object at 0x1[...]>]
>>> gca().invert_xaxis() # xaxis from the easiest to the hardest
>>> legend() # this operation updates the figure with the inverse axis.
<matplotlib.legend.Legend object at 0xa84592c>
```



If we swap the abscissa and the ordinate of the previous figure, we can directly compare the horizontal view and vertical view figures:

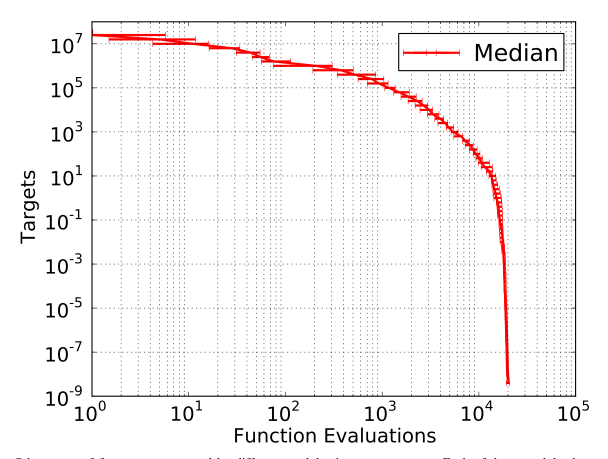
```
>>> figure()
<matplotlib.figure.Figure object at 0x1[...]>
>>> for i in range(0, nruns):
... loglog(evals[:, i], targets)
...
[<matplotlib.lines.Line2D object at 0x1[...]>]
      [...]
[<matplotlib.lines.Line2D object at 0x1[...]>]
>>> grid()
>>> xlabel('Function Evaluations')
<matplotlib.text.Text object at 0x1[...]>
>>> ylabel('Targets')
<matplotlib.text.Text object at 0x1[...]>
>>> loglog(d.ert[d.target>=1e-8], d.target[d.target>=1e-8], lw=3,
```

```
... color='r', label='ert')
[<matplotlib.lines.Line2D object at 0x1[...]>]
>>> legend()
```



The process for displaying the median function values and interquartile ranges for different targets is described thereafter:

```
>>> figure() # open a new figure
<matplotlib.figure.Figure object at 0x1[...]>
>>> from bbob_pproc.bootstrap import prctile
>>> q = array(list(prctile(i, [25, 50, 75]) for i in evals))
>>> xmed = q[:, 1]
\rightarrow \rightarrow x low = xmed - q[:, 0]
>>> xhig = q[:, 2] - xmed
>>> xerr = vstack((xlow, xhig))
>>> errorbar(xmed, targets, xerr=xerr, color='r', label='CMA-ES')
(<matplotlib.lines.Line2D object at 0x1[...]>, [...], [<matplotlib.collections.LineCollection object object]
>>> grid()
>>> xscale('log')
>>> yscale('log')
>>> xlabel('Function Evaluations')
<matplotlib.text.Text object at 0x1[...]>
>>> ylabel('Targets')
<matplotlib.text.Text object at 0x1[...]>
```



Other types of figures are generated by different modules in bbob\_pproc. Each of these modules has a plot function and a beautify function. The plot function is used for generating a type of graph. The beautify function accommodates the general presentation of the figure to its content.

The modules for generating figures in this manner are bbob\_pproc.pprldistr, bbob\_pproc.compall.ppperfprof, bbob\_pproc.ppfigdim.

The bbob\_pproc.pprldistr module generates Empirical Cumulative Distribution Function (ECDF) graphs of the number of evaluations for attaining target precisions and the function values for a given budget:

```
NAME
    bbob_pproc.pprldistr - Creates run length distribution figures.

FILE
    [...]/python/bbob_pproc/pprldistr.py

FUNCTIONS
    beautify()
        Format the figure of the run length distribution.

comp(dsList0, dsList1, valuesOfInterest, isStoringXMax=False, outputdir='', info='default', verber Generate figures of empirical cumulative distribution functions.
        Dashed lines will correspond to ALGO and solid lines to ALGO.
```

>>> help(bb.pprldistr)

```
Keyword arguments:
        dsList0 -- list of DataSet instances for ALG0.
        dsList1 -- list of DataSet instances for ALG1
        valuesOfInterest -- target function values to be displayed.
        isStoringXMax -- if set to True, the first call BeautifyVD sets the
          globals fmax and maxEvals and all subsequent calls will use these
          values as rightmost xlim in the generated figures.
        outputdir -- output directory (must exist)
        info --- string suffix for output file names.
    main(dsList, valuesOfInterest, isStoringXMax=False, outputdir='', info='default', verbose=True)
        Generate figures of empirical cumulative distribution functions.
        Keyword arguments:
        dsList -- list of DataSet instances to process.
        valuesOfInterest -- target function values to be displayed.
        isStoringXMax -- if set to True, the first call BeautifyVD sets the
          globals fmax and maxEvals and all subsequent calls will use these
          values as rightmost xlim in the generated figures.
        outputdir -- output directory (must exist)
        info --- string suffix for output file names.
        Outputs:
        Image files of the empirical cumulative distribution functions.
    plot(dsList, valuesOfInterest=(10.0, 0.100000000000001, 0.0001, 1e-08), kwargs={})
        Plot ECDF of final function values and evaluations.
DATA
    __all__ = ['beautify', 'comp', 'main', 'plot']
>>> ds = bb.load(glob.glob('BIPOP-CMA-ES/ppdata_f0*_20.pickle'))
>>> figure()
<matplotlib.figure.Figure object at 0x1[...]>
>>> pprldistr.plot(ds)
>>> pprldistr.beautify() # resize the window to view whole figure
         1.0
                         f1-24
        0.8
       proportion of trials
        0.4
```

The bbob\_pproc.compall.ppperfprof module generates ECDF graphs of the bootstrap distribution of the Expected Running Time for target precisions:

```
>>> help(bb.compall.ppperfprof)
Help on module bbob_pproc.compall.ppperfprof in bbob_pproc.compall:
```

log10 of FEvals / DIM

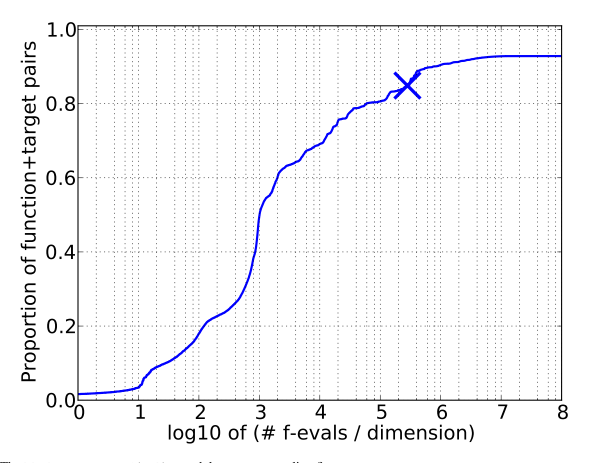
log10 of Df / Dftarget

f1-24

0.2

0.05

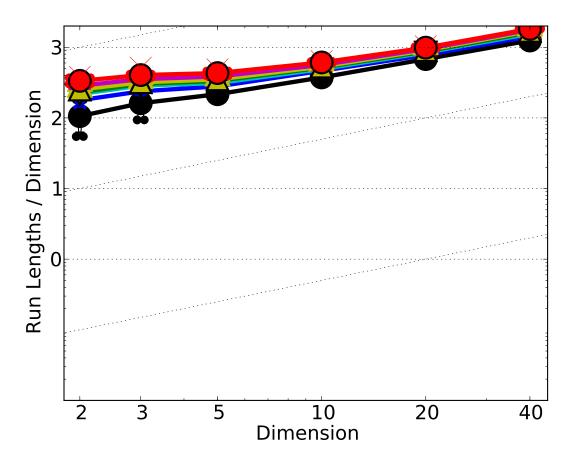
```
NAME
    bbob_pproc.compall.ppperfprof - Generates figure of the bootstrap distribution of ERT.
FILE
    [...]/python/bbob_pproc/compall/ppperfprof.py
DESCRIPTION
    The main method in this module generates figures of Empirical
    Cumulative Distribution Functions of the bootstrap distribution of
    the Expected Running Time (ERT) divided by the dimension for many
    algorithms.
FUNCTIONS
   beautify()
        Customize figure presentation.
    main(dictAlg, targets, order=None, plotArgs={}, outputdir='', info='default', verbose=True)
        Generates a figure showing the performance of algorithms.
        From a dictionary of DataSetList sorted by algorithms, generates the
        cumulative distribution function of the bootstrap distribution of
        ERT for algorithms on multiple functions for multiple targets
        altogether.
        Keyword arguments:
        dictAlg -- dictionary of dataSetList instances containing all data
        to be represented in the figure
        targets -- list of target function values
        order -- sorted list of keys to dictAlq for plotting order
    plot(dsList, targets=(1e-08, [...]), rhleg=False, kwargs={})
        Generates a plot showing the performance of an algorithm.
        Keyword arguments:
        dsList -- a DataSetList instance
        targets -- list of target function values
        kwargs
DATA
    __all__ = ['beautify', 'main', 'plot']
>>> ds = bb.load(glob.glob('BIPOP-CMA-ES/ppdata_f0*_20.pickle'))
>>> figure()
<matplotlib.figure.Figure object at 0x1[...]>
>>> ppperfprof.plot(ds)
[<matplotlib.lines.Line2D object at 0x1[...]>, <matplotlib.lines.Line2D object at 0x1[...]>]
>>> ppperfprof.beautify()
```



The bbob\_pproc.ppfigdim module generates scaling figures.

```
>>> from bbob_pproc import ppfigdim
>>> help(ppfigdim)
Help on module bbob_pproc.ppfigdim in bbob_pproc:
NAME
   bbob_pproc.ppfigdim - Generate performance scaling figures.
FILE
    [...]/python/bbob_pproc/ppfigdim.py
FUNCTIONS
   beautify()
        Customize figure presentation.
   main(dsList, _valuesOfInterest, outputdir, verbose=True)
        From a DataSetList, returns a convergence and ERT/dim figure vs dim.
    plot(dsList, _valuesOfInterest=(10, 1, [...], 1e-08))
        From a DataSetList, plot a figure of ERT/dim vs dim.
DATA
     _all__ = ['beautify', 'plot', 'main']
>>> ds = bb.load(glob.glob('BIPOP-CMA-ES/ppdata_f002_*.pickle'))
```

```
Unpickled BIPOP-CMA-ES/ppdata_f002_02.pickle.
[...]
Unpickled BIPOP-CMA-ES/ppdata_f002_40.pickle.
>>> figure()
>>> ppfigdim.plot(ds)
>>> ppfigdim.beautify()
```



The post-processing tool bbob\_pproc generates image files and LaTeX tables from the raw experimental data obtained with COCO.

Its main usage is through the command line interface. To generate custom figures, bbob\_pproc can be used alternatively from the Python interpreter.

**CHAPTER** 

# **EIGHT**

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