The Multi-Run Simulation Environment SimEnv

User Guide for Version 1.13

by M. Flechsig, U. Böhm, T. Nocke & C. Rachimow



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The Multi-Run Simulation Environment

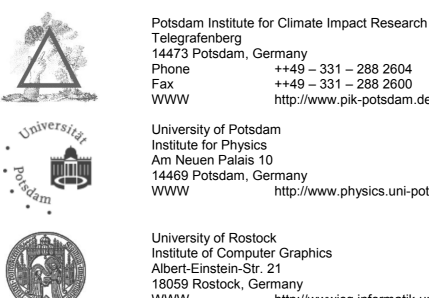
SimEnv

User Guide for Version 1.13 (03-Aug-2004)

by

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Executive Summary

SimEnv is a multi-run simulation environment that focuses on model evaluation and usage mainly for quality assurance matters and scenario analyses using sampling techniques. Interfacing models to the simulation environment is supported for a number of programming languages by minimal source code modifications and in general at the shell script level. Pre-defined experiment types are the backbone of SimEnv, enabling experimenting with numerical parameter, initial value, or driving forces adjustments of the model. The resulting multi-run experiment can be performed sequentially or in parallel. Interactive experiment postprocessing makes use of built-in operator definitions, optionally supplemented by user-defined operators and applies operator chains on model output and reference data. Result output functions generated during postprocessing can be evaluated within SimEnv with advanced visualization techniques.

Simulation is one of the cornerstones for research in Global Change. The aim of the SimEnv project is to develop a toolbox oriented simulation environment that enables the modeller to handle model related quality assurance matters (Saltelli et al., 2000, Saltelli et al., 2004) and scenario analyses. Both research foci require complex simulation experiments for model inspection, validation and control design without changing the model in general.

SimEnv (Flechsig et.al, 2004) aims at model evaluation by performing simulation runs with a model in a coordinated manner and running the model several times. Co-ordination is achieved by pre-defined experiment types representing multi-run simulations.

According to the strategy of a selected experiment type for a set of so-called targets t which represent drivers, parameters, boundary and initial values of the model **M** a sample is generated before simulation and the targets t are re-adjusted numerically before each single simulation run during the experiment. Each experiment results in a sequence of model outputs over the single runs for selected state variables z dependent on the target adjustments of the model M. Model outputs can be processed and evaluated across the run ensemble specifically after simulation.

The following experiment types form the base of the SimEnv multi-run facility:

Behavioural analysis

Inspection of the model's behaviour in a space spanned from targets t with discrete numerical adjustments and a flexible inspection strategy for the whole space.

For model verification, numerical validation, deterministic error analysis, deterministic control design, scenario analysis and spatial patch model applications.

- Monte Carlo analysis Perturbations of targets t according to probability density functions. Determination of moments, confidence intervals and heuristic probability density functions for z in the course of post-processing. For error analysis, uncertainty analysis, verification and validation of deterministic models.
- Local sensitivity analysis Determination of model (state variable's z) local sensitivity to targets t. Is performed by finite difference derivative approximations from M.
 - For numerical validation purposes, model analysis, sub-model sensitivity.
- Optimization

Iterative determination of optimal targets t for a cost functions derived from z by a simulated annealing methods.

For model validation (system - model comparison), control design, decision making.

SimEnv makes use of modern IT concepts. Model preparation for interfacing them to SimEnv is based on minimal source code manipulations by implementing function calls into Fortran-, C/C++-, Python- or GAMSmodel source code for target adjustments and model output. Additionally, an interface at shell script level is available.

In experiment preparation an experiment type is selected and equipped numerically. Experiment performance supports local, remote, and parallel architectures.



Experiment-specific model output post-processing enables navigation in the experiment - model output space and interactive filtering of model output and reference data by application of built-in and user-defined post-processing operator chains.

Result evaluation is dominated by application of pre-formed visualization modules.

SimEnv model output as well as model output post-processing offer data interfaces for NetCDF, IEEE compliant binary and ASCII format for a more detailed post-processing outside SimEnv.

SimEnv key features:

- Support of key working techniques in experimenting with models: SimEnv enables model evaluation, uncertainty and scenario analyses in a structured, methodologically sound and pre-formed manner applying sampling techniques.
- Run ensembles instead of single model runs: Model evaluation by multi-run simulation experiments
- Availability of pre-defined multi-run simulation experiment types: To perform an experiment only the targets (parameters, drivers, initial values, ...) to experiment with and a strategy how to sample the target space have to be specified.
- Simple model interface to the simulation environment: There are model interface functions mainly to re-adjust an experiment target and to output model results for later post-processing. Model interfacing and finally communication between the model and SimEnv can be done at the model language level by incorporating coupling functions into model source code (C/C++, Fortran and Python) or can be done at the shell script level within shell-scripts. Additionally, there is a special interface for GAMS models.
- Support of distributed models: Independently on the kind distributed models are coupled they can be interfaced to SimEnv.
- Parallelization of the experiment: This is a prerequisite for a lot of simulation tasks.
- Operator-based experiment post-processing: Chains of built-in and user-defined operators enable interactive experiment post-processing based on experiment model output and reference data including general purpose and experiment-specific operators.
- Graphical experiment evaluation: For post-processed model output
- Support of standard data formats: Output from the model as well from the post-processor can be stored in NetCDF or IEEE compliant binary format.

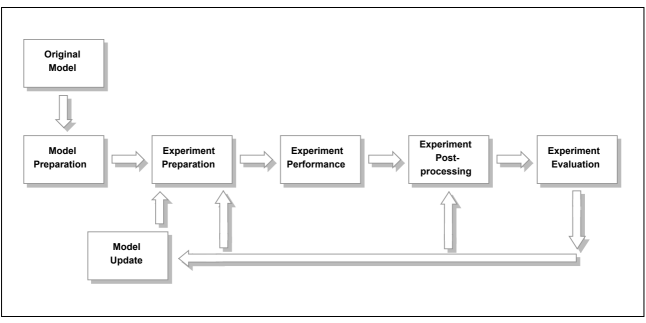


Fig. 0.1

SimEnv system design

-2-

1 About this Document

In this chapter document conventions are explained. Within the whole document one reference example model is used to explain application of SimEnv. Examples are always located in grey boxes.

1.1 Document Conventions

Character / string	Meaning	
< >	angle brackets enclose a placeholder for a string	
{ }	praces enclose an optional element	
[]	square brackets enclose a list of choices, separated by a vertical bar	
· '	quotation marks enclose a keyword or sub-keyword for user-defined files	
<nil></nil>	stands for the empty string (nothing)	
monospace	indicates SimEnv example code	

Tab. 1.1Document conventions

Tab. 1.2 summarizes the main placeholders used in this document.

Placeholder	Description	
<direct></direct>	path to a file directory	
<file_name></file_name>	name of a data file	
<gams_model></gams_model>	name of a GAMS model	
<model></model>	model name to start a SimEnv service with	
<nil></nil>	the empty string	
<res></res>	integer post-processor output file number 1, 2,, 99	
<res_char></res_char>	character post-processor output file number 01, 02,, 99	
<run></run>	integer single run number 0, 1, within an experiment	
<run_char></run_char>	character single run number 000000, 000001, within an experiment	
<sep></sep>	sequence of white spaces as item separators in user-defined files	
<string></string>	any string	
<target_def_val></target_def_val>	default value of a target according to <model>.edf</model>	
<target_name></target_name>	name of a target to experiment with	
<value_list></value_list>	list of values in explicit or implicit notation according to Tab. 11.6	

Tab. 1.2

Main placeholders in this document



1.2 Example Layout

All examples in this document refer to a hypothetical global simulation **model world**. It is to describe dynamics of atmosphere and biosphere at the global scale over 200 years. Lateral (latitudinal and longitudinal) model resolution differs for different model implementations (see below), temporal resolution is at decadal time steps. Additionally, atmosphere is structured vertically into levels.

The model world is assumed to map lateral and vertical (level) fluxes and demands that's why for computing state variables for the whole globe.

The model world is a generic model. Model implementation in several programming languages results in models world_<lng> where <lng> is an identifier for the programming language (and the lateral model resolution).

In the model pixel_f state variables are calculated for one grid cell (one single latitude - longitude constellation) without consideration of lateral fluxes.

Model state variable	Description	Defined on	Data type
atmo	aggregated atmospheric state	lat x lon x level x time	float
bios	aggregated biospheric state at land masses (defined between 83°N and 60°S latitude at land masses, i. e., without Antarctic)	lat x lon x time	float
atmo_g (not for model pixel_f)	aggregated global state derived from atmo for level 1	time	int
bios_g (not for model pixel_f)	aggregated global state derived from bios	-	int

Dynamics of all model variables depend on model parameters p1, p2, p3 and p4.

With this SimEnv release the following model implementations are distributed:

	Model	Resolution		
Model	interface example for	lateral: lat x lon	vertical: number of levels	temporal: number of time steps
world_f	Fortran	4 x 4	4: 1, 7, 11, 16	20
world_c	С	4 x 4	4: 1, 7, 11, 16	20
world_cpp	C++	4 x 4	4: 1, 7, 11, 16	20
world_py	Python	4 x 4	4: 1, 7, 11, 16	20
world_sh	script level	4 x 4	4: 1, 7, 11, 16	20
pixel_f	Fortran	without, implicitly by experiment as 4 x 4	4: 1, 7, 11, 16	20
world_f_1x1	Fortran	1 x 1	16: 1 - 16	20

The only example that does not refer to the above model type is that for the GAMS model interface to SimEnv (see chapter 5.5 at page 29).

Examples are generally placed in grey-shaded boxes. Examples that are available from the corresponding examples directory of the SimEnv release are marked as such in the lower right corner of an example box.

Example 1.1 General example layout in the User Guide

2 Getting Started

In this chapter a quick start tour is described. Without going into details the user can get an impression how to apply SimEnv and which user files are essential to use the simulation environment.

- SimEnv is implemented under AIX at IBM's RS6000 to run in a Korn-shell ksh.
- Set the operating system environment variable **SE_HOME** to /usr/local/simenv/bin, export it and include this setting in your file \$HOME/.profile.
- Set the operating system environment variable **PYTHONPATH** according to your needs, extend it by \$SE_HOME, export it, and include this setting in your file \$HOME/.profile. For more information on PYTHONPATH see Tab. 10.11 at page 106.
- Change to a working directory you have full access rights.
- Start **\$SE_HOME/simenv.hlp** to acquire basic information on how to use SimEnv.
- Select an implementation language <Ing> you want to check SimEnv with the model from Example 1.1 at page 4 and

<ing></ing> = f	for Fortran
C	for C
срр	for C++
ру	for Python
sh	for shell script level

For a GAMS model example check chapter 5.7 at page 32.

• Start the shell script

\$SE_HOME/simenv.cpy world_<lng>

to copy model world_<lng> model and experiment related files to the working directory. Copy the file world.edf_c to world_<lng>.edf

Check

U	HEUK		101
٠	The SimEnv configuration file	world_ <ing>.cfg</ing>	a general configuration
٠	The model output description file	world_ <ing>.mdf</ing>	available model variables
٠	The model	world_ <ing>.<ing></ing></ing>	implementation of the model
٠	The model shell script	world_ <ing>.run</ing>	wrapping the model executable
٠	The experiment description file	world_ <ing>.edf</ing>	experiment definition
٠	The post-processing input file	world.post_c	post-processor result sequence
٠	The macro description file	world_ <ing>.mac</ing>	macros for the post-processor
٠	The operator description file	world_ <ing>.opr</ing>	description of user-defined operators
٠	The user-defined operators	usr_opr_ <opr>.f</opr>	code of user-defined operator <opr></opr>
~	test a secondate Oise Essential has		

for

Start a complete SimEnv session by

\$SE_HOME/simenv.cpl world_<lng> -1 world.post_c

- SimEnv files will be checked
- The experiment will be prepared

• The experiment will be performed machine (select the login machine on request)

- Model output post-processing will be started for this experiment
 - With the post-processing input file world_post_c and following
 - Interactively: Enter any result and finish post-processing by entering a single <return>
- Visualization of post-processed results will be started
- Model or result output files will be dumped
- or
 - Start \$SE_HOME/simenv.chk world_<Ing>
- to check model and experiment files.
- Start **\$SE_HOME/simenv.run** world_<Ing> to prepare and perform a simulation experiment.

-5-

(*)

- Start **\$SE_HOME/simenv.res** world_<Ing> {[new | append | replace]} {<run>} to post-process the last simulation experiment over the whole run ensemble or for run number <run> and to create a new / append to / replace the result file <model>.res<res_char>.[nc | ieee | ascii] with the highest two-digit number <res_char>. <res_char> (can range from 01 to 99).
- Start **\$SE_HOME/simenv.vis** world_<Ing> {[latest | <res_char>]} (*) to visualize output from the latest post-processing output file world_<Ing>.res<res_char>.nc or that with number <res_char> with the highest two-digit number <res_char>. <res_char> can range from 01 to 99.
- Start **\$SE_HOME/simenv.dmp** world_<lng> | more to dump a SimEnv model or post-processor output file.
- Check in the working directory the model interface, native model terminal output, and experiment performance log-files

world_<lng>.mlog world_<lng>.nlog world_<lng>.elog.

- Start \$SE_HOME/simenv.cln world_<lng> to wrap up a simulation experiment.
- Get the usage of any SimEnv service by entering the appropriate command without arguments.
- To run other simulation experiments and/or output in other data formats modify
 - world_<lng>.cfg
 - world_<lng>.edf
 - world_<lng>.mdf
 - world_<Ing>.<Ing> and/or
 - world_<lng>.run
- To experiment with other models replace world_<lng> by <model> as a placeholder for the name of any other model.

(*): To get access rights for the visualization server check the SimEnv service \$SE_HOME/simenv.key <user_name> in chapter 10.2 at page 97.

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3 Version 1.13

This chapter summarizes differences between the current and the previous SimEnv release, limitations, and bugs and their workarounds.

3.1 What is New?

Туре	Check / see	At page	Description
new	chapter 5.1	21	 Experiment performance: unused targets / undefined model output Targets that are not get to the model (by simenv_get_*) as well as total or partial unwritten model output to SimEnc data structures (by simenv_put_*) is protocolled to <model>.mlog as a warning</model>
update	Tab. 10.5	101	 Experiment performance: additional log-file <model>.nlog</model> Model native output to terminal is no longer collected in the model interface performance log-file <model>.mlog but redirected to the native model output log-file <model>.nlog.</model></model> <model>.mlog and <model>.nlog are now organized in such a way that all information for one single run is lumped together. This makes it possible to understand single run performance by check-ing <model>.nlog.</model></model></model>
new	5.7.3	37	 Experiment performance: GAMS models An additional native single run related log-file <model>.nlog is stored to the sub-directory run<run_char> where a single run is performed.</run_char></model>
			Bug fixes

Tab. 3.1SimEnv changes in version 1.13

Upgrade type	Upgrade action	
mandatory	Re-link all models	

Tab. 3.2

User actions to upgrade to version 1.13



3.2 Limitations and Their Workarounds

Where / Limitation / Workaround	Description
Where	Overall
Limitation	Current SimEnv technical limitations as specified in Tab. 15.4 at page 129
Workaround	None
Where	Overall
Limitation	Only accessible under Unix / AIX
Workaround	None
Where	Overall but visual result evaluation
Limitation	Without graphical user interface
Workaround	None
Where	Experiment performance: Experiment type optimization
Limitation	Can not be performed in parallel
Workaround	Perform experiment in sequential mode
Where	Experiment post-processing: Optional specification / automated identification of result description and result unit
Limitation	Not stored to NetCDF result output
Workaround	Specify IEEE or ASCII result output instead
Where	Experiment post-processing: User-defined operators
Limitation	No C-interface to write user-defined operators
Workaround	Wrap user-defined operators written in C into Fortran before interfacing them to SimEnv
Tab. 3.3	Known limitations and their workarounds

3.3 Known Bugs / Problems and Their Workarounds

Where Bug / Problem Workaround	Description	
Where	Experiment performance: Model output to NetCDF	
Problem	Check on undefined model output results in additional CPU-time consumption. Example: to check 8 Mill of real*8 values takes per single run additionally 80 sec for single nc-file model output and 200 sec for common nc-file output.	
Workaround	Specify in <model>.cfg for sub-keyword 'message_level' value = 'error'</model>	
Where	Experiment performance: Optimization / model output to a common NetCDF file for the whole experiment	
Bug	Write error	
Workaround	Specify IEEE model output or single NetCDF file output in <model>.cfg</model>	
Where	Experiment performance: NetCDF model output of distributed models (distributed = yes in <model>.cfg)</model>	
Bug	May not store all model output	
Workaround	Specify IEEE model output in <model>.cfg</model>	
Where	Experiment performance: Experiment restart / model output to a common NetCDF file for the whole experiment	
Bug	Read error in experiment post-processing	
Workaround	Specify IEEE model output or single NetCDF file output in <model>.cfg</model>	
Where	Experiment post-processing: Behavioural analysis / result output to NetCDF	
Bug	When applying operator behav non-monotonic target adjustments are transferred to NetCDF output in a wrong manner.	
Workaround	Specify only monotonic target adjustments in <model>.edf</model>	
Where	Experiment status: (SimEnv service simenv.sts)	
Bug	Does not work properly for experiment re-starts.	
Workaround	Check the file <model>.edf manually for experiment progress</model>	
Tab 34	Known bugs/ problems and their workarounds	

Tab. 3.4Known bugs/ problems and their workarounds



-10- SimEny

4 Experiment Types

SimEnv supplies a set of pre-defined multi-run experiment types. Each experiment type addresses a special experiment class for performing a simulation model several times in a co-ordinated manner. In this chapter an overview on the available experiment types is given from the viewpoint of system's theory.

4.1 General Approach

M:

SimEnv supplies a set of pre-defined multi-run experiment types, where each type addresses a special multirun experiment class for performing a simulation model or any algorithm with an input - output transition behaviour.

In the following, the general SimEnv approach will be described for time dynamic simulation models, because this class forms the majority of SimEnv applications. All information can be transformed easily to any other algorithm.

Based on systems' theory, each time dynamic model *M* can be formulated - without limitation of generality - for the time dependent, time discrete, and state deterministic case as

 $Z(t) = ST \left(Z(t - \Delta t), \dots, Z(t - k^* \Delta t), P, X(t), Z_0, B \right)$

with

	., .	
ST Z P		state transition description state variables' vector parameter vector
X		input (driving forces) vector
Zo		initial value vector
В		boundary value vector
t		time
Δt		time increment
k		time delay

The output vector Y is a function of the state vector Z, parameters P, drivers X, and initial values Z_0 :

$$Y(t) = OU (Z(t), P, X(t), Z_0).$$

Model behaviour Z is determined for fixed k and Δt by state transition description ST, parameters P, driving forces X, initial values Z_0 , and boundary values B. Manipulating and exploring model behaviour in any sense means changing these four model components. While state transition description ST reflects mainly model structure and is quite complex to change, each component of the driving forces vector X normally is a time-dependent vector.

Introduction of additional technical parameters P_{tech} can reduce the complexity of handling a model with respect to the five model components, described above: Changes in state transition description *ST* can be predetermined in the model by assigning values of a technical parameter p_{tech} to alternative sub-model versions, which are switched on or off by these values. Additionally, each component of the driving forces vector *X* can be combined with technical parameters in different ways:

- By selecting special driving forces dependent on the technical value
- By manipulating the driving forces with the parameter value
- (e.g., as an additive or multiplicative adjustment)
- By parametrizing the shape of a driving force

When this has been done, the model behaviour finally depends only on the parameters P, the initial values Z_0 , and the boundary values B. From the methodical point of view there is no difference between parameters, initial values and boundary values, because all are considered as constant during one model run. That



is why in SimEnv all the four model components parameters, drivers, initial values and boundary values are lumped together and the term **target**¹ stands as a placeholder for them. All targets form the target set T:

and

$$\mathsf{T} = \{ P, X, Z_0, B \}$$

Z = ST(T).

In the following,

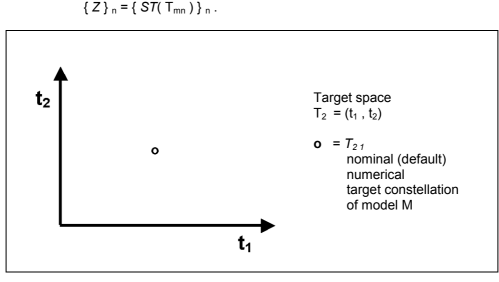
$$T_m = (t_1, ..., t_m)$$
 m > 0

stands for a subset of the target set T that spans up an m-dimensional sub-space of T by selected model targets (t_1 ,..., t_m) from T and

 $T_{mn} = \begin{pmatrix} t_{11} & \dots & t_{1m} \\ \dots & \dots \\ t_{n1} & \dots & t_{nm} \end{pmatrix} \qquad m > 0, \ n > 1$

stands for a numerical sample for T_m of size n and finally for m*n values representing in any sense T_m . In the set of all T_{mi} (i > 1) one extraordinary sample T_{m1} exists that matches the nominal (default) numerical target constellation for the model *M*.

If $\{ \}_n$ denotes the dynamics of the model *M* over a sample of size n then it holds:





SimEnv supports different sampling stategies and performance of multi-run experiments where m targets are readjusted numerically for each of n single simulation runs. Central goal is to study depencency of the model dynamics on target adjustments. For simulation purposes in SimEnv experimentation with the model *M* over T_{mn} is based on the assumption that dynamics of *M* for each representative from the sample is indepent from all other representatives, which is fulfilled in general. This results in the possibility to form a run ensemble for performing the model *M* with n single model runs from the sample T_{mn} .

SimEnv experiment types differ in the way T_m is sampled to get T_{mn} . There are deterministic and non-deterministic sampling strategies that offer a broad range of techniques for

- Experimentation with models
- Post-processing model output results

Interpreting results with respect to uncertainty and sensitivity matters of models.

The experiment types are described in detail in the following.

¹ The term target was selected as an analogue to experimentation with real systems: Often a target is under investigation to study the change in the real system when the state of the target is changed by the experimentor.

4.2 Behavioural Analysis

Behavioural analysis uses a deterministic strategy to sample T_m . It is the inspection of the model in the target space T_m where inspection points are set in a regular and well structured manner.

Behavioural analysis can be interpreted and used in different ways:

- For scenario analysis:
- to show how model behaviour changes with changes of target values
- For numerical validation purposes: to determine target values in such a way that the output vector matches with measurement results of the real system
- For deterministic error analysis:
- to analyse how the model error is dependent on target errors
- For a simulation-based control design:
- to determine target values in such a way that a goal function becomes an extreme

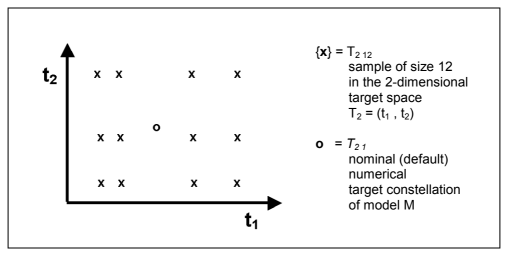


Fig. 4.2 Sample for a behavioural analysis

SimEnv behavioural analysis sampling strategy is a generalization of the one-dimensional case for T_1 , where the model behaviour is scanned in dependence on deterministic adjustments of one target t_1 . The general case for T_m demands a strategy for scanning m-dimensional spaces in a flexible manner. Based on the predecessors of SimEnv (Wenzel *et al.*, 1990, Wenzel *et al.*, 1995, Flechsig, 1998) subspaces of the m-dimensional target space can be scanned on the subspace diagonal (parallel in a one-dimensional hyper-space) or completely for all dimensions (combinatorially on a grid) and both techniques can be combined. Besides this regular scanning method an irregular technique is possible.

The resulting number of single simulation runs for the experiment depends on the number of target samples per dimension of the scanned target space and from the selected scanning method. An experiment is described by the names of the involved targets, their numerical adjustments and their combination (scanning method). Model output post-processing resolves the scanning method again and outputs results as projections on multi-dimensional target subspaces.

Fig. 4.3 describes the regular scanning technique by an example. In the left scheme (a) the two-dimensional target space $T_2 = (p_1, p_2)$ is scanned combinatorially, resulting in 4*4 = 16 model runs, while the middle scheme (b) represents a parallel scanning of these two targets at the diagonal by 1+1+1+1 = 4 model runs. The scheme (c) at the right side shows a complex scanning strategy of the 3-dimensional target space $T_3 = (p_1, p_2, p_3)$ with $(1+1+1+1)^*3 = 12$ model runs. Each filled dot • in Fig. 4.3 correspond to an cross **x** in Fig. 4.2 and represents a sample point in the target space and finally a single model run of the experiment.



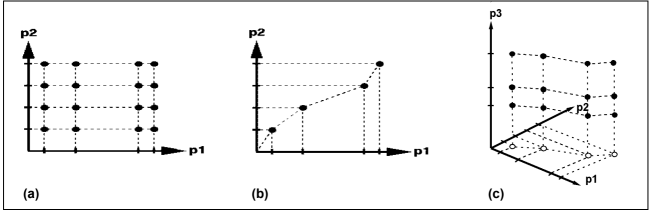


Fig. 4.3 Beha

Behavioural analysis: Scanning multi-dimensional target spaces

4.3 Monte Carlo Analysis

Monte Carlo analysis uses a non-deterministic strategy to sample T_{mn} . A Monte Carlo experiment in SimEnv is a perturbation analysis with pre-single run target perturbations.

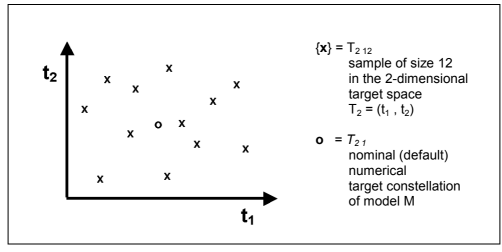


Fig. 4.4

with

Sample for a Monte Carlo analysis

Theoretically, with a Monte Carlo analysis moments of a state variable z can be computed as

$$M^{(k)}{z} = \int_{m}^{\dots} z(T_m)^k * pdf(T_m) dT_m$$
$$T_m$$
k-th moment of the

M ^(K) {z}	k-th moment of the state variable z with respect to the
	probability density function pdf
z(T _m)	state variable z as a function of T_m
pdf(T _m)	probability density function of T _m

By interpreting the probability density function $pdf(T_m)$ as the error distribution in the target space T_m it is possible to study error propagation in the model. On the other hand Monte Carlo analysis can be interpreted as a stochastic error analysis, if there are measurements of the real system for z.

For a numerical experiment in SimEnv it is assumed that the probability density function $pdf(T_m)$ can be decomposed into independent probability density functions pdf_i for all targets t_i of T_m :

$$pdf(T_m) = \prod_{i=1}^{m} pdf_i(t_i)$$

and the m-dimensional integral is approximated by a sequence of n single simulation runs of the model where the numerical target values t_{ij} of t_i ($1 \le i \le m$, $1 \le j \le n$) are sampled according to the probability density function pdf_i.

On the basis of these assumptions, the statistical measures in Tab. 4.1 can be computed during performance of a post-processing session from a Monte Carlo analysis with n simulation runs resulting in n realizations z_1 ,..., z_n of the model's state variables z, z1 and z2:

Statistical measure	Definition (*)	
minimum	$min(z) = min(z_i)$	
maximum	$max(z) = max(z_i)$	
sum	sum(z) = Σz_i	
arithmetic mean	$avg(z) = \sum z_i / n$	
variance	var(z) = $\sum (z_i - avg(z))^2 / (n - 1)$	
skewness	skw(z) = $\Sigma (z_i - avg(z))^3 / n^* (\Sigma (z_i - avg(z))^2 / (n - avg(z))^2)$	1)) ^{3/2}
kurtosis	krt(z) = $(\sum (z_i - avg(z))^4 / n * (\sum (z_i - avg(z))^2 / (n + avg(z))^2))$	– 1)) ²) - 3
range	rng(z) = max(z) - min(z)	
geometric mean	$avgg(z) = (\Pi z_i)^{1/n}$	
harmonic mean	$agvh(z) = n / \Sigma(1 / z_i)$	
weighted mean	avgw(z) = $\sum z_i * w_i / \sum w_i$ w : weight	
correlation	cor(z1,z2) = $\sum (z1_i - avg(z1)) * (z2_i - avg(z2)) /$	
	$\frac{\sqrt{\Sigma} (z1_i - avg(z1))^2 * \Sigma (z2_i - avg(z2))^2}{\sum (z1_i - avg(z1))^2 + \sum (z1_i - avg(z2))^2}$	<u></u>
covariance	$cov(z1,z2) = \Sigma (z1_i - avg(z1)) * (z2_i - avg(z2)) / (n - 1)$)
linear regression coefficient	reg(z1,z2) = $(\sum (z1_i - avg(z1)) * (z2_i - avg(z2))) / (\sum (z1_i - avg(z1))^2)$	
median	med(z) = middle value from increasingly ordered { z _i } mean of the two middle values from { z _i }	(n = odd) (n = even)
quantile	<pre>qnt^(p)(z) = that value from increasingly ordered { z_i } which corresponds to a cumulative frequence qnt^(0.5)(z) = med(z)</pre>	· · · · · · · · · · · · · · · · · · ·
confidence interval boundaries	$cnf^{(\alpha)}(z) = avg(z) \pm t_{\alpha,n-1} \sqrt{var(z) / n}$ α : level of error $t_{\alpha,n}$: significance boundaries of Student	t distribution
heuristic probability density function	$\label{eq:class} \begin{array}{ll} \text{hgr}^{(\text{class})}(z) & = & \text{number of } z_i \text{ with } \text{class}_{\text{min}} \leq & z_i < & \text{class}_{\text{max}} \\ & & \text{class}_{\text{min}}, \text{class}_{\text{max}} : \text{boundaries of equidi} \end{array}$	stant classes

Tab. 4.1Statistical measures

n

n

(*): indices for sums Σ , products Π and extremes run from 1 to n: $\sum_{i=1}^{n} \prod_{i=1,...,n}^{n} \max_{i=1,...,n}$

-15-

Tab. 4.2 summarizes these probability density functions (Bohr, 1998) that are pre-defined in SimEnv for targets to be perturbed. Additionally, SimEnv offers to import random number samples in the course of experiment preparation.

Distribution	Short- cut	Probability density function pdf		Distr	ibution parameters
uniform	U(a,b)	$pdf(x) = \frac{1}{b-a}$	$\text{if } x \in [a,b]$	a b	lower boundary upper boundary > a
		pdf(x) = 0	otherwise	it is:	mean = (a+b) / 2 standard deviation = $\sqrt{(b-a)^2 / 12}$
normal	$N(\mu, \sigma^2)$	$pdf(x) = \frac{1}{\sigma\sqrt{2\pi}} exp\left(-\frac{(x-\mu)^2}{2\sigma^2}\right)$		$\mu \sigma$	mean standard deviation > 0
lognormal	$L(\mu, \sigma^2)$	$pdf(x) = \frac{1}{x\sigma\sqrt{2\pi}} exp\left(-\frac{(\ln x - \mu)^2}{2\sigma^2}\right)$	if x > 0	μ σ	> 0
		pdf(x) = 0	otherwise	it is:	$\ln(x) \sim N(\mu, \sigma^2)$
exponential	Ε(μ)	$pdf(x) = \frac{1}{\mu} exp\left(-\frac{x}{\mu}\right)$	if x > 0	μ	mean > 0
		pdf(x) = 0	otherwise	it is:	standard deviation = μ

Tab. 4.2Probability density functions

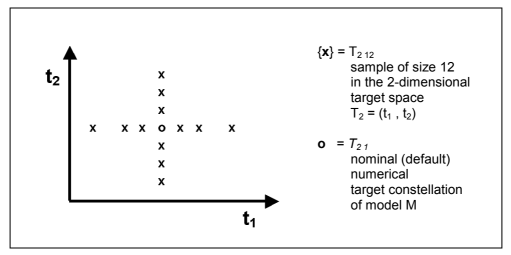
The number of runs to be performed during a Monte Carlo analysis has to be specified. An experiment is described by the targets involved in the analysis, their distribution and the appropriate distribution parameters.

4.4 Local Sensitivity Analysis

Local sensitivity analysis uses a deterministic sampling stategy in ε -neighbourhoods of the numerical default constellation T_{m1} of the model M. For each target t_i from the nominal target constallation T_{m1} and each ε_j from the ε -neighbourhoods (ε_1 ,..., ε_k) two members (t_1 ,..., t_{i-1} , $t_i \pm \varepsilon_j$, t_{i+1} ,..., t_m) of the resulting sample are generated. The sample size n is given by 2*m*k. Running the model at this sampling set serves to determine sensitivity functions.

In classical systems' theory, model sensitivity of a model state variable z with respect to a target t is the partial derivative of z after t $\delta z/\delta t$. In the numerical simulation of complex systems a finite sensitivity function is preferred, because it can be obtained without model enlargements or re-formulations. It is a linear approximations of the classical model sensitivity measure (Wierzbicki, 1984).

Local sensitivity measures as well as measures which reflect model output linearity and/or symmetry nearby T_{m1} can be used for localizing modification-relevant model parts as well as control-sensitive targets in control problems. On the other hand, identification of robust parts of a model or even complete robust models makes it possible to run a model under internal or external disturbances. Sensitivity analysis in SimEnv post-processing is based on finite sensitivity, linearity, and symmetry measures, which are defined as in Tab. 4.3.





Sample for a sensitivity analysis

Local	Defir	Definition		
measure	Absolute measure	Relative measure		
sensitivity measure	sens_abs(z, $\pm\epsilon$) = $\frac{z(t \pm \epsilon) - z(t)}{\pm \epsilon}$	sens_rel(z, $\pm\epsilon$) = sens_abs(z, $\pm\epsilon$) $\frac{t}{z(t)}$		
linearity measure	$\lim_{\varepsilon \to 0} abs(z,\varepsilon) = \frac{(z(t+\varepsilon) - z(t)) + (z(t-\varepsilon) - z(t))}{\varepsilon}$	$lin_rel(z,\epsilon) = lin_abs(z,\epsilon) \frac{t}{z(t)}$		
symmetry measure	sym_abs(z, ϵ) = $\frac{z(t + \epsilon) - z(t - \epsilon)}{\epsilon}$	sym_rel(z, ϵ) = sym_abs(z, ϵ) $\frac{t}{z(t)}$		

Tab. 4.3Local sensitivity, linearity, and symmetry measures
for a selected target t from T_{m1} and a selected ε from $(\varepsilon_1, ..., \varepsilon_n)$

Accordingly, local measures of the model with respect to a target are always expressed as a measure of a model's state variable z, usually at a selected time step within a surrounding neighborhood ϵ of a target value t. That is why the conclusions drawn from a local sensitivity analysis are only valid locally at T_{m1} with respect to the whole target space T_m . Additionally, local measures only describe the influence of one target t_i from the whole vector T_m on the model's dynamics.

As stated above, the sensitivity measures reflect the classical sensitivity functions in a neighborhood of T_{m1} . The larger the absolute value of the measure the higher is the influence of an incremental change of the target t on the model output z. The linearity measures map the linear behaviour of z nearby T_{m1} . If the linear measure is zero z shows a linear behaviour with respect to t. The symmetry measures map the symmetric behaviour of the z nearby T_{m1} . If the symmetry measure is zero z shows a symmetric behaviour with respect to t. The larger the absolute values of the latter two measures the higher is the nonlinear / non-symmetric behaviour of z with respect to t.

The absolute measures are best suited to compare the influence of different targets $\{t\}$ on the same state variable z while due to their normalization factor the relative measures enable comparison of the influence of one target t on different state variables $\{z\}$.

From the local measures of table Tab. 4.3 additional measures can be derived on demand, e.g., $abs(sym_abs(z, \epsilon))$.

A local sensitivity experiment is described by the names of the targets t to be involved and the increments ε . The number of runs for the experiment results from the number of targets and increments: two runs per target for each increment plus one run with the default values of the targets. Local sensitivity functions are calculated during model output post-processing.

4.5 Optimization

The optimization experiment in SimEnv uses a stochastic strategy to sample T_m . It is the only experiment type where the sample is generated during experiment performance and not at experiment preparation. The general approach of optimization is to find the global minimum of a cost function (synonym: objective function)

$$F(Z) = F(ST(T_m))$$

that depends on model's state variables Z and consequently on the experiment targets $T_m = (t_1, ..., t_m)$:

minimize	F(t ₁ ,, t _m)	
subject to	$t_{i \min} \le t_i \le t_{i \max}$	for i = 1 ,, m

Often, F represents a distance measure in a specific metric between selected model state variables and reference data (measurement values of the real system or simulation results from an other model). Consequently, optimization can be used for model validation and control design to find optimal values of model targets in such a way that model state variables are close to reference data. In SimEnv the cost function is specified in result post-processing as a result formed from model output (and reference data) where an operator chain is applied on (check chapters 6.5 and 8). The value of the cost function is calculated directly after the current single run has been performed.

SimEnv uses a gradient free optimization approach that is called "**Simulated Annealing**" and is a generalization of a Monte Carlo method for examining the state equations of n-body systems. The concept is based on the manner in which metals recrystalize in the process of annealing. In an annealing process a melt, initially at high temperature Temp and disordered, is slowly cooled so that the system at any time is approximately in thermodynamic equilibrium. As cooling proceeds, the system becomes more ordered and approaches a "frozen" ground state at Temp = 0. Hence the process can be thought of as an adiabatic approach to the lowest energy state E. If the initial temperature of the system is too low or cooling is done insufficiently slowly the system may become quenched forming defects or freezing out in metastable states (i.e. trapped in a local minimum energy state).

The annealing scheme is that an initial state of a thermodynamic system is chosen at energy E and temperature Temp, holding Temp constant the initial configuration is perturbed and the change in energy dE is computed. If the change in energy is negative or zero the new configuration is accepted. If the change in energy is positive it is accepted with a probability given by

$$p = exp(-dE/(k_B *Temp))$$

where k_B denotes the Boltzmann constant. This process is then repeated sufficient times to give good sampling statistics for the current temperature, and then the temperature is decremented and the entire process repeated until a frozen state is achieved at Temp = 0.

By analogy the generalization of this Monte Carlo approach to optimization problems is straight forward:

- The current state of the thermodynamic system is analogous to the current solution to the optimization problem
- The energy equation for the thermodynamic system is analogous to the objective function F, and
- The ground state at Temp = 0 is analogous to the global minimum of F.

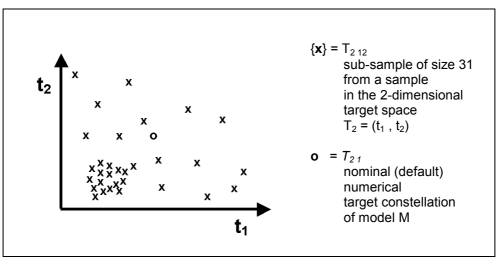
The major difficulty (art) in implementation of a simulated annealing algorithm is that there is no obvious analogy for the temperature Temp with respect to a free parameter in the optimization problem. Furthermore, avoidance of entrainment in local minima (quenching) is dependent on the "annealing schedule", that is, the choice of initial temperature, how many iterations are performed at each temperature, and how much the temperature is decremented at each step as cooling proceeds (after Gray *et al.*, 1997). Ideally, when local optimization methods are trapped in a poor local minimum, simulated annealing can 'climb' out.

The algorithm applied in SimEnv is a very fast simulated re-annealing method, named Adaptive Simulated Annealing ASA (Ingber 2004, Ingber 1989 and Ingber 1996). For the above stated probability p the term k_B * Temp is chosen as

$$k_B * Temp = Temp_0 * exp(-c*k^{1/m})$$

where k is the annealing time.

The ASA schedule is much faster than Boltzmann annealing, where $k_B * \text{Temp} = \text{Temp}_0 / \ln(k)$ and faster than fast Cauchy annealing, where $k_B * \text{Temp} = \text{Temp}_0 / k$. With the ASA method the global minimum of a nonlinear non-convex cost function F over an m-dimensional bounded target space T_m is determined.





Part of a sample for an optimization experiment, generated during the experiment



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5 Model Interface

To use any model within SimEnv it has to be coupled to the simulation environment. SimEnv offers easy coupling techniques at programming language and shell script level. While at language level SimEnv function calls have to be implemented into model source code to adjust experiment targets, i. e. model parameters, initial values or boundary values of the current single run out of the run ensemble numerically and to output simulation results, at the shell script level communication between the simulation environment and the model can be based on operating system information exchange methods. To plug the model into the simulation environment the variables of the model to be output during experiment performance and to be post-processed during model output processing have to be declared in the model output description file <model>.mdf. Additionally, the model itself has to be wrapped into a shell script <model>.run.

Model interfacing is related to transferring adjusted numerical values of model targets under investigation from the simulation environment to the model and to transferring model variables under investigation from the model to the simulation environment for later post-processing. Coupling is supported at the programming language level for C/C++, Fortran, Python, and GAMS programming languages, the model is implemented in and at shell script level.

5.1 General Approach

SimEnv model interface has supply a link between the simulation environment and the model and has to address two aspects:

For each single run from the run ensemble

- All numerical adjustments of experiment targets as defined in the experiment description file <model>.edf (check chapter 6.1) have to be associated to the corresponding model entities (parameters, initial or boundary values, drivers) and these entities have to be modified numerically in the model according to the specified adjustments.
- All model output variables as defined in the model output description file <model>.mdf (check chapter 5.3) have to be associated to the corresponding model entities (in general, model state variables) and these entities have to be output to SimEnv data structures during the performance of the model.

Realisation of this general approach is based on minimal source code manipulation of the model. SimEnv supplies model language related libraries with a set of six simple functions to interface the model to the simulation environment. Generally speaking,

- Every experiment target and
- Every model output variable

demand one additional SimEnv function call in the model source code. According to Tab. 5.1 model interface functions are generic.

Function name	Description
simenv_ini_ <ing></ing>	open model coupling interface
simenv_get_ <lng></lng>	associate a model source code entity with an experiment tar- get from <model>.edf and get the target adjustment</model>
simenv_get_run_ <ing></ing>	get the current single run number of the run ensemble
simenv_put_ <ing></ing>	associate a model source code entity with a model output variable from <model>.mdf and output it to SimEnv data structures</model>
simenv_slice_ <ing></ing>	enable slicing, i.e., a repetitively partial output of model output variables.
simenv_end_ <ing></ing>	close model coupling interface

Tab. 5.1

Generic SimEnv interface functions (for <Ing> check Tab. 5.2) The function simenv_slice_<Ing> announces output of a slice of the data of a defined model variable. This is good for models with multi-dimensional variables where at least one dimension is omitted in the state variable declaration in the model the source code because the dynamics for this dimension is calculated in place (e.g., time). The assigned variable then has a lower dimensionality than the corresponding variable in the model output description file. Nevertheless, the slice-function ensures that model output over the omitted dimension can be handled in model output post-processing in common.

The alignment of the contents of the SimEnv description files and the used SimEnv interface functions in the model source code is dominated by the description files: These files determine the experiment and the model source code is expected to be well adapted. Nevertheless, this approach is implemented in a flexible manner:

- Function calls in the source code where an experiment target from <model>.edf and/or a model varibale from <model>.mdf is not associated with are handled during the model performance in such a way that the targets are unadjusted and/or the model variable is not output. This enables adaption of the model source code for a number of potential experiment targets and model outputs where only a subset of these targets is under consideration in special experiments and/or requested for model output.
- Vice versa, model entities that are requested by the corresponding experiment and/or model description file for target adjustments and/or model output and where the corresponding SimEnv functions in the model source code are missing are identified as such.

A regular matching between the model description files and the used SimEnv interface functions in the model source code as well as the above exceptions are reported to the interface log-file <model>.mlog (check Tab. 10.5).

Native model output does not influence performance of the model in SimEnv and there is no necessity to disable this output for SimEnv. The user only has to ensure that for a parallel performance of an experiment the output of different single runs does not conflict with each other. Normally, this can be ensured by performing each single run in a special related sub-directory (check Example 15.5). Native user model output to terminal is redirected to the file <model>.nlog.

For running an interfaced model outside SimEnv there are dummy SimEnv libraries to link / run the model with. They ensure the same model dynamics as before interfacing the model to SimEnv (check chapter 5.9).

Currently, there are SimEnv interfaces for Fortran, C/C++, Python and GAMS models. Additionally, there is an interface implementation at shell script level. Mixed language models as well as distributed models (check 5.8) can be run with SimEnv.

<ing></ing>	for model source code		
С	C/C++		
f	Fortran		
ру	Python		
sh	Shell script level		

Tab. 5.2

Language suffices for SimEnv interface functions (for the GAMS interface check chapter 5.7)

5.2 Grid and Coordinate Assignments to Variables

To each variable

- Dimensionality dim(variable)
- Extents ext(variable,i) with i=1,...,dim(v)
- Coordinates coord(variable,i)

with i=1,...,dim(variable) with i=1,...,dim(variable)

are assigned to. The dimensionality is the number of dimensions, an extent is related to each dimension and represents the number of elements in that dimension. Extents are always greater than 1. To each dimension a coordinate is assigned to. Coordinates have a name and from all coordinate values the coordinate is defined for a subset is assigned to the extent of the dimension of the variable. Variables of dimensionality 0 do not have a coordinate assignment.

A variable of dimensionality n corresponds with an n-dimensional array, a variable of dimensionality 0 is a scalar.

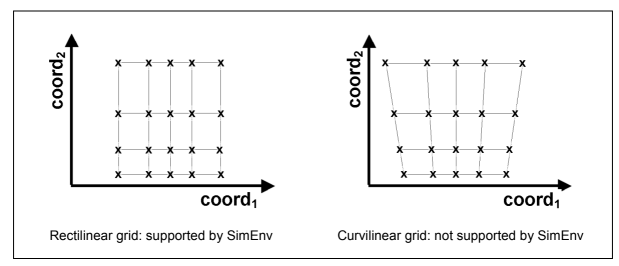


Fig. 5.1Grid types

Additionally, coordinate axes are defined. Each coordinate axis a strictly monotonic sequence of coordinate values, a description and a unit is assigned to. For reasons of simplification in model output post-processing coordinate axes are assumed as curvilinear.

Each dimension of a variable with a dimensionality > 0 a complete coordinate axis or a part of a coordinate axis is assigned to. Consequently, each variable with a dimensionality > 0 is defined on a coordinate system formed from the assigned coordinates. For reasons of simplification in result evaluation with visualization techniques coordinate systems are assumed as rectilinear (orthogonal with variable distances between adjacent coordinate values). The model variable values then exist on the grid, spanned up from the coordinate values of the coordinate axes (see Fig. 5.1).

Since coordinate axes can be assigned to model variable dimensions in a flexible manner, model variables can exist on the same coordinate system or completely or partially disjoint coordinate systems.



5.3 Model Output Description File <model>.mdf

In the model output description file <model>.mdf the model variables are declared that are to be output by a SimEnv model coupling interface function in the model (code) and are to be post-processed after experiment performance. Additionally, coordinate axes are defined and flexibly assigned to model variables. Consequently, a model variable always is defined on a coordinate system, formed from the assigned coordinates to the variable.

keyword	name	sub- keyword	Line type	Max. line nmb.	value	Explanation
mdf	<nil></nil>	descr	0	any	<string></string>	model description
coordinate	<coordinate_< td=""><td>descr</td><td>0</td><td>1</td><td><string></string></td><td>coordinate axis description</td></coordinate_<>	descr	0	1	<string></string>	coordinate axis description
	name>	unit	0	1	<string></string>	coordinate axis unit
	(<co_name>)</co_name>	values	m	1	<value_list></value_list>	strictly monotonic sequence of coordinate values <co_vals> (for syntax see Tab. 11.6)</co_vals>
variable	<variable_< td=""><td>descr</td><td>0</td><td>1</td><td><string></string></td><td>variable description</td></variable_<>	descr	0	1	<string></string>	variable description
	name>	unit	0	1	<string></string>	variable unit
		type	m	1	see Tab. 5.4	variable type in the simulation model
		coords	c1	1	<co_name<sub>1> , , <co_name<sub>n></co_name<sub></co_name<sub>	assigns a coordinate axis by its name to each dimension of the variable. Determines in this way implicitly the dimen- sionality n of the variable.
		coord_extents	c2	1	<co_val<sub>11>: <co_val<sub>12> , , <co_val<sub>n1>: <co_val<sub>n2></co_val<sub></co_val<sub></co_val<sub></co_val<sub>	assigns start and end coordi- nate value from each coordi- nate axis to the variable. If missing all coordinate values will be used from all assigned coordinates.
		var_extents	c1	1	<vi_ext<sub>11>: <vi_ext<sub>12> , , <vi_ext<sub>n1>: <vi_ext<sub>n2></vi_ext<sub></vi_ext<sub></vi_ext<sub></vi_ext<sub>	assigns start and end index for each dimension to the variable. Indices can be used to address the variable during post-processing.

Tab. 5.3 Elements of a model output description file <model>.mdf

Each model variable has a name, a dimensionality and assigned extents, a data type, a description and a unit. The name should correspond with the name of the variable in the simulation model code. Association between these two names is achieved by the SimEnv coupling function simenv_put_* (see below).

<model>.mdf is an ASCII file that holds this information. It follows the coding rules in chapter 11.1 at page 107 with the keywords, names, sub-keywords, and values as in Tab. 5.3.

To Tab. 5.3 the following additional rules and explanations apply:

- For the description of line type check Tab. 11.4 at page 109.
- Coordinate and variable names must differ from target names in experiment description (see chapter 6.1) and from built-in and user-defined operator names for model output post-processing (see chapter 8.5.2).
- Assignment of coordinate axes to variable dimensions and consequently of a grid to a variables is only
 valid for model output post-processing. Normally, the simulation model itself will also exploit the same
 grid structure. Nevertheless, the grid structures of the model are defined autonomously in the model in a

explicit or implicit manner and do only correspond with the grid structure in the model output description file symbolically.

- Model variables with dimensionality 0 are not assigned to a coordinate axis.
- The values of a coordinate have to be ordered in a strictly monotonic sequence. They may be nonequidistant and may be ordered in a decreasing sequence.
- With the sub-keyword 'coord_extents' only a portion of coordinate values of a coordinate axis can be assigned to a dimension of a variable. This portion is addressed by its begin and end value <co_val_{i1}> and/or <co_val_{i2}>. The number of coordinates values of the portion has to be greater than 1.
 <co_val_{i1}> < <co_val_{i2}> for strictly increasing values of coordinates

<co_val_{i1}> < <co_val_{i2}> for strictly decreasing values of coordinates

With the sub-keyword 'var_extents' portions of variables are made addressable during SimEnv post-processing. In the same way multi-dimensional variables are equipped with indices in the simulation model they also have an index description in the model output description file for purposes of model output post-processing. It is advisable, that these two descriptions coincide. The index range is described by a start and an end index <vi_ext_{i1}> and/or <vi_ext_{i2}>.

The index set is a strictly increasing, equidistant set of integer values with an index increment of 1, $\langle v_i | ext_{i1} \rangle \langle \langle v_i | ext_{i2} \rangle$,

<vi_ext_{i1}> \leq 0 is possible.

- Coordinate values and index values are assigned in a one-to-one manner.
- For multi-dimensional variables that do not exist on an assigned grid completely or partially, simply assign formal coordinate axes to.
- Specify at least one model output variable in <model>.mdf.

SimEnv data type		Desc	ription	Restriction	
byte	or	int*1	1 byte	integer	not for Python models
short	or	int*2	2 bytes	integer	not for Python models
int	or	int*4	4 bytes	integer	
float	or	real*4	4 bytes	real	
double	or	real*8	8 bytes	real	not for Python models

Tab. 5.4SimEnv data types

For the following example of a model output description file and the assigned grid for model variable bios check Example 1.1 at page 4:

	descr descr descr descr descr	World with a resolution of 4° lat x 4° lon x 4 levels x 20 time steps Data centred per lat-lon cell This file is valid for all models world_[f c cpp py sh]
lat	descr unit values	geographic latitude deg equidist_end 88(-4)-88
lon	descr unit values	geographic longitude deg equidist_end -178(4)178
level		atmospheric vertical level level no list 1,7,11,16
	lat lat lon lon lon level level	descr descr descr descr descr descr descr descr descr descr descr lat unit lat values lon descr lon unit lon values level descr unit

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coordinate	time time time	descr unit values	time in decades 10 years equidist_nmb 1(1)20
variable	atmo atmo atmo atmo atmo		aggregated atmospheric state without float lat , lon , level , time 1:45 , 1:90 , 1:4 , 1:20
variable variable	bios bios bios bios bios bios	coords coord_extents	aggregated biospheric state g/m ² float lat , lon , time 84.:-56. , -178.:178. , 1:20 1:36 , 1:90 , 1:20
variable variable variable variable	atmo_g atmo_g		int time 1:20 int
			Example-file: world_[f c cpp py sh].mdf

Example 5.1 Model output description file <model>.mdf

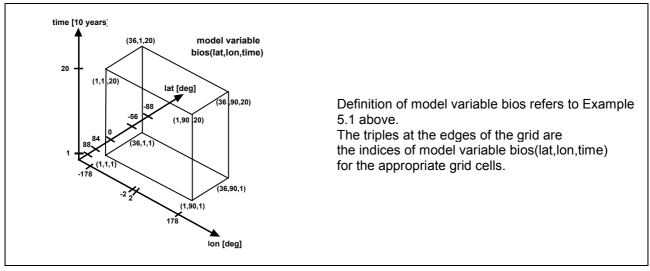


Fig. 5.2 Model variable definition: Grid assignment

5.4 Model Interface for Fortran and C/C++ Models

Tab. 5.5 describes the interface functions that can be used in user models written in Fortran or C/C++ to adjust experiment targets for the current single run of the run ensemble and to output model results from the current single run. In this table the input and output data types are documented for functions used in Fortran. For C/C++ the corresponding data types are valid.

All functions have a 4-byte integer function value (integer*4 and/or int). Implementation of the functions for C/C++ is based on a call by reference for the function arguments.



Function name	Function description	Arguments / function value	Argument / function value description
simenv_ ini_[f c] ()	initialize model coupling interface Apply always before the first call of the other SimEnv functions in the model	integer*4 simenv_ini (function value)	return code = 0 ok = 2 I/O error for model output file = 3 error memory allocation = 4 I/O error for <model>.edf_bin = 5 I/O error for <model>.mdf_bin = 6 I/O error for <model>.edf_adj = 7 wrong single run number</model></model></model>
simenv_ get_[f c] (target_name,	gets the numerical modification (adjustment) for the target (pa-	character*(*) target_name (input) real*4	name of the target in <model>.edf nominal / default (non-adjusted) target value.</model>
target_val_def, target_val_adj)	rameter / initial value / boundary value) to be ex- perimented with in	target_val_def (input) real*4 target_val_adj	If target_name is not defined in <model>.edf then target_val_adj is set to target_val_def adjusted target value</model>
	the current single run	(output) integer*4 simenv_get (function value)	return code = 0 ok = 1 target_name undefined: target_val_adj := target_val_def = 3 for Monte Carlo analysis: warning w.r.t. distribution paramter adjustment (check Tab. 6.6 at page 45)
simenv_ gets run number of get_run_[f c] current run as an (integer value and a run_int, character string)		character*6 run_char (output) integer*4	current run number with leading zeros
		run_int (output) integer*4 simenv_get_run	return code = 0 ok
simenv_ put_[f c] (outputs model results to native SimEnv output	(function value) character*(*) var_name (input)	name of the variable in <model>.mdf to be output</model>
var_name, field)	file(s)	dimension field(), type according to <model>.mdf (input)</model>	data of variable var_name to be stored as simula- tion results
		integer*4 simenv_put (function value)	return code = 0 ok = 1 var_name undefined = 2 I/O error model output file
simenv_ slice_[f c] (announces to out- put at next simenv_put call	character*(*) var_name (input)	name of the variable in <model>.mdf to be sliced</model>
var_name, idim, ifrom, ito	only a slice of vari- able var_name. This announce- ment becomes	integer*4 idim (input)	dimension to be sliced slice to start at position ifrom
)	inactive after per- formance of the appropriate simenv_put	integer*4 ifrom (input)	

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Function name	Function description	Arguments / function value	Argument / function value description
		integer*4 ito (input) integer*4	slice to end at position ito return code
		simenv_slice (function value)	 = 0 ok = 1 var_name undefined = 3 inconsistency between variable and idim, ifrom, ito = 4 slice storage exceeded = 5 warning: slice overwritten
simenv_ end_[f c] ()	close model cou- pling interface Apply always after the last call of the other SimEnv functions in the model	integer*4 simenv_end (function value)	return code = 0 ok = 2 I/O error for model output file

Tab.	5.5	Model coupler functions at language level
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- Make sure consistency of type and dimension declarations between the model variables in model source code and the corresponding variable declarations in the model output description file <model>.mdf.
- Model variables that are not output completely or partially within the user model are handled in resultpost-processing as their corresponding nodata-values (see Tab. 10.10).
- Application of simenv_slice_* for NetCDF model output may result in a higher consumption of computing time for each single run of the experiment compared with NetCDF model output without simenv_slice_*. For this case, keep in mind the trade-off between the demand for computing time and the demand for main memory.

User models implemented in C/C++ or Fortran have to be linked with the following libraries to couple them to the simulation environment

- \$SE_HOME/libsimenv.a
- /usr/local/lib/libnetcdf.a

In

- Example 15.1 at page 131 the model world_f.f
- Example 15.2 at page 133 the model world_c.c
- Example 15.3 at page 135 the model world_cpp.cpp

are explained.

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5.5 Model Interface for Python Models

Due to the special features of Python the coupling interface to SimEnv differs from that for Fortran and C/C++ in chapter 5.4. Additionally, Python supports only some data types (check Tab. 5.4). Tab. 5.6 summarizes the coupling functions to use for a Python model.

Function name	Function description	Arguments / function value	Argument / function value description
simenv_ ini_py ()	initialize model coupling interface Apply always before the first call of the other SimEnv functions in the model	string ini_py (function value)	return code of the spawn function for a SimEnv executable
simenv_ get_py (target_name, target_def_val))	gets the numerical modification (adjustment) for the target (pa- rameter / initial value / boundary value) to be ex- perimented with in the current single	string target_name (input) float target_val_def float get_py	name of the target in <model>.edf nominal / default (non-adjusted) target value. If target_name is not defined in <model>.edf then target_val_adj is set to target>_val_def adjusted target value target_val_adj</model></model>
simenv_ get_run_py (run gets run number of current run as a character string	(function value) string get_run_py (function value)	current run number as string of the length 6 with leading zeros. If an error occurred then run_char = ''
y simenv_ put_py (var_name, field)	outputs model results to native SimEnv output file(s)	string var_name (input) declaration of field() according to <model>.mdf (input) put_py</model>	name of the variable in <model>.mdf to be output data of variable var_name to be stored as simula- tion results. Maximum length of field is limited to 12.000 characters.</model>
simenv_ slice_py (var_name, idim, ifrom, ito	Currently not available for Python models	(function value)	
) simenv_ end_py ()	close model cou- pling interface		



Model coupler functions for Python models

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- Python coupling functions are declared in the file \$SE_HOME/simenv.py. To use these functions in a Python model import it by
 - from simenv import *
 - and refer to it for example by simenv_get_py.
- Errors that occur during performance of one of the above functions are directly reported to <model>.nlog.

In Example 15.4 at page 136 the model world_py.py is described in detail.

5.5.1 Standard User-Defined Files for Python Models

<model>.ini

<model>.ini (see chapter 7.1 at page 51) is for Python models a mandatory script and has to have the same contents for all Python models:

```
$SE_HOME/py_model_ini
rc_py_model_ini = $?
# additional user-model specific commands can be implemented up from here
if test $rc_py_model_ini = 0
then
...
fi
exit $rc_py_model_ini
```

For an experiment restart with a Python model (check chapter 7.3 at page 53) <model>.ini has to be performed again. To force this specify in <model>.cfg (check chapter 10.1 at page 95) for the sub-keyword 'restart_ini' the value yes.

5.6 Model Interface at Shell Script Level

For models that do not allow to implement the model coupling interface at programming language level (e.g., because source code is not available) SimEnv supplies a coupling interface at shell script level: the shell script <model>.run (see chapter 7.1 at page 51) is used to wrap the model and optionally to have at disposal corresponding functionality of the SimEnv coupling functions of Tab. 5.5).

Command name	Command description	Arguments	Argument description
. \$SE_HOME/ simenv_ ini_sh	initialize current single run Apply always as the first command in <model>.run and <model>.rst</model></model>	SE_RUN (output)	operating system environment variable SE_RUN is set to the current run number of the simulation experiment
target_name= '' target_def_val=	gets a numerical modification (adjustment) for the target (pa-	script variable target_name (input) script variable	name of the target in <model>.edf nominal / default (non-adjusted) target value.</model>
 . \$SE_HOME/ simenv_	rameter / initial value / boundary	target_def_val (input)	If target_name is not defined in <model>.edf then target_val_adj is set to target>_val_def</model>
get_sh	value) to be ex- perimented with in the current single run	script variable target_name (output)	shell script variable with the same name as the value of target_name. Script variable value is the adjusted target value target_val_adj.
. \$SE_HOME/ simenv	gets run number of current run as an	run_char (output)	shell script variable with the current run number with leading zeros
get_run_sh	integer and a character script variable	run_int (output)	shell script variable (type integer) with the current run number
. \$SE_HOME/ simenv_ put_sh	Not available at shell script level		Write your own simenv_put_sh at the language level using the SimEnv coupling functions from Tab. 5.5 or Tab. 5.6
. \$SE_HOME/ simenv_ slice_sh	Not available at shell script level		
. \$SE_HOME/ simenv_ end_sh	wrap up current single run		
	Apply always as the last command in <model>.run and <model>.rst</model></model>		

Tab. 5.7

Model coupler functions at shell script level

• For the model interface at the shell script level, i.e., within the shell script <model>.run the adjusted experiment targets for the current single run from the whole run ensemble can be made available within <model>.run to forward them by any means the modeller is responsible for to the model under investigation.

One common way to forward experiment targets to the model is to place current numerical target values as arguments to the model at model command line. Another way could be to read the targets from a special file in a special file format.

- Directly before performing simenv_get_sh make sure that the shell script variables target_name and target_def_val have be specified. At the end of each simenv_get_sh these variables are set again to empty strings.
- After running . \$SE_HOME/simenv_get_sh an experiment target <target_name> from the experiment description file <model>.edf is available in <model>.run as a shell script variable <target_name> and the adjusted value of the target is available as \$<target_name>.
- After running the model model output has to be identified and potentially transformed within <model>.run for SimEnv output. To do this simply write your own simenv_put_sh as a transformation program that reads in all the native model output and outputs it to SimEnv by applying the coupling functions simenv_*_* from the SimEnv model coupler at language level.



• Tab. 10.8 lists the built-in (pre-defined) shell script variables that are used in \$SE_HOME/simenv_*_sh and finally in <model>.run.

In Example 15.5 at page 137 the model shell script world_sh.run is described in detail.

. \$SE HOME/simenv ini sh # get adjusted value for the a target p def, defined in the edf-file target name='p def' target def val=2. . \$SE HOME/simenv get sh # now shell script variable p def is available is according to edf-file # value of shell script variable p def # get adjusted value for a target p undef, not defined in edf-file target name='p undef' target def val=-999. . \$SE HOME/simenv get sh # now shell script variable p undef is available # value of shell script variable p undef is -999. # \$SE HOME/simenv end sh Example file: world sh.run

Example 5.2 Addressing target names and values for the model interface at shell script level

5.7 Model Interface for GAMS Models

SimEnv allows to couple GAMS models to the experiment shell. A GAMS model for SimEnv can consist of a GAMS main model and GAMS sub-models.

Therefore, two additional include-statements have to be inserted into these GAMS model source code files where experiment targets are to be adjusted or model variables are to be output. GAMS model source code files to be interfaced to SimEnv are one GAMS main model and a number of GAMS sub-model that are called directly from the main model. All these GAMS model source code files have to be located in the current working directory. Additional GAMS sub-programs (included files) are not affected bei SimEnv, but you should keep in mind that the GAMS code within SimEnv will be executed in a subdirectory of the current working directory (see below) and so the include statements have to be changed, if the files are addressed in a relative manner (see below).

 The include files are <GAMS_model>_simenv_get.inc
 <GAMS_model>_simenv_put.inc

where <GAMS_model> is the name of a GAMS model file without extension .gms under consideration.

- During experiment preparation the file <GAMS_model>_simenv_put.inc and during experiment performance files <GAMS_model>_simenv_get.inc are generated automatically to forward GAMS model output to SimEnv data structures and to adjust investigated experiment targets, respectively. These include files correspond with the simenv_put and simenv_get functions at the language level (see
- chapter 5.4).
 The GAMS include statement \$include <GAMS_model>_simenv_get.inc has to be placed in the GAMS model file at such a position where all the GAMS variables are declared. Directly before the include

statement the target default values have to be assigned to target variables, that are introduced addition-

ally in the model. Directly after the include statement the target variables with the adjusted target values have to be assigned to the model variables.

- The GAMS include statement \$include <GAMS_model>_simenv_put.inc has to be placed in the GAMS model file at such a position where all the variables from the model output description file can be output by GAMS put-statements.
- In the course of experiment preparation the GAMS model and all sub-models that are specified in <model>.gdf (see below) are transformed automatically. Each GAMS model single run from the run ensemble is performed in a separate sub-directory of the current working directory. Transformed GAMS models and sub-models are copied to this sub-directory and are performed from there. Keep this in mind if you specify in any GAMS model include statements with relative paths.

In Example 15.6 at page 139 the model gams_model.gms is described in detail.

Additionally, the following settings are valid:

- An ASCII GAMS description file **<model>.gdf** (see below) has to be supplied to specify the GAMS submodels and assigned targets and model variables in detail.
- Maximum dimensionality of any model output variable declared in <model>.mdf is 4 for GAMS models.

Note the following information:

• To output the GAMS model status to SimEnv a

PARAMETER modstat

has to be declared and the statement

modstat = <model_name>.modelstat

has to be incorporated in the GAMS model above the \$include <GAMS_model>_simenv_put.inc line. The variable modstat has to be stated in the model output description file <model>.mdf and the GAMS description file <model>.gdf.

• GAMS information, normally output to the screen, is output to the native model log file <model>.nlog.

5.7.1 Standard User-Defined Files for GAMS Models

<model>.ini

<model>.ini (see chapter 7.1 at page 51) is for GAMS models a mandatory script and has to have the contents for all GAMS models:

```
$SE_HOME/gams_model_ini
rc_gams_model_ini = $?
# additional user-model specific commands can be implemented up from here
if test $rc_gams_model_ini = 0
then
...
fi
exit $rc gams model ini
```

For an experiment restart with a GAMS model (check chapter 7.3 at page 53) <model>.ini has to be performed again. To force this specify in <model>.cfg (check chapter 10.1 at page 95) for the sub-keyword 'restart_ini' the value yes.

<model>.run

<model>.run (see chapter 7.1 at page 51) has for each GAMS model the same contents:

. \$SE_HOME/simenv_ini_sh \$SE_HOME/gams_model_run . \$SE_HOME/simenv_end_sh

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<model>.end

<model>.end (see chapter 7.1 at page 51) is for GAMS models a mandatory script and has to have the contents for all GAMS models:

\$SE_HOME/gams_model_end

additional user-model specific commands can follow

Python script language is used to prepare, run and to end a GAMS model.

<model>.edf

Corresponding experiment targets in the experiment description file <model>.edf (see chapter 6.1 at page 39) and in the GAMS model source code must have same names. In the GAMS model code the targets specified in the experiment description file have to be of type PARAMETER and have be defined before the include statement \$include simenv_get.inc.

<model>.mdf

Corresponding variables in the model output description file and in the GAMS model source code must have same names. The variable type has to be always float in the model output description file. In GAMS model code the model variables declared in the model output description file can be of the numeric types VARIABLES or PARAMETER. Currently, maximal dimensionality of GAMS model output is restricted to 4.

With respect to Example 15.6 the model output description file could look like						
coordinate	plant plant plant	descr unit values	canning plants plant number equidist_end 1(1)2			
coordinate	market market market	descr unit values	canning markets market number equidist_end 1(1)3			
variable variable variable variable variable	a a a a	descr unit type coords var_extents	plant capacity cases float plant 1:2			
variable variable variable variable variable	X X X X X	descr unit type coords var_extents	<pre>shipment quantities cases float plant , market 1:2 , 1:3</pre>			
variable variable variable variable variable	z z modstat modstat	descr unit type descr type	total transportation costs 10^3 US\$ float model status float			
		21 ⁻	Example file: gams_model.mdf			

Example 5.3 Model output description file for a GAMS model



5.7.2 GAMS Description File <model>.gdf

The ASCII GAMS description file <model>.gdf is intented to create a block of lines for each GAMS submodel with a simenv_get.inc file and/or a simenv_put.inc file. The block holds the specific characteristics of GAMS model input and output needed by SimEnv to generate GAMS put-statements. All model variables from the model output description file and all targets from the target description file have to be used in this file again.

<model>.gdf is an ASCII file that follows the coding rules in chapter 11.1 at page 107 with the keywords, names, sub-keywords, and values as in Tab. 5.3.

keyword	name	sub- keyword	Line type	Max. line nmb.	value	Explanation
gdf	<nil></nil>	descr	0	any	<string></string>	GAMS coupling description
		keep_runs	0	1	<value_list></value_list>	value list of run numbers where single GAMS model runs are to be stored by keeping their corresponding sub-directories (for syntax see Tab. 11.6)
		time_limit	0	1	<positive_ integer></positive_ 	CPU limit in seconds for each GAMS model single run
		options	0	1	<string></string>	string of options, GAMS main model is started with from command line
model	<model_< td=""><td>descr</td><td>0</td><td>1</td><td><string></string></td><td>(sub-)model description</td></model_<>	descr	0	1	<string></string>	(sub-)model description
	name>	type	m	1	[main sub]	identifies GAMS main or sub- model
	(without extension .gms)	get	m	exactly number of targets	<target_name></target_name>	get resulting adjustment for <target_name> to this model</target_name>
		put	m	exactly number of model vari- ables	<pre>(<var_name> {.<suffix_set>} {(<index_set>)}) {<format>}</format></index_set></suffix_set></var_name></pre>	put values of SimEnv model output variable <var_name> from this model to SimEnv output. GAMS variable <var_name> has the specified suffix and index sets and is interfaced from GAMS to SimEnv ac- cording to <format></format></var_name></var_name>

Tab. 5.8

Elements of a GAMS description file <model>.gdf

To Tab. 5.8 the following additional rules and explanations apply:

- For the description of line type check Tab. 11.4 at page 109.
- Each target and each model variable as declared in <model>.edf and <model>.mdf respectively has to be used in the value-field of <model>.gdf exactly one time.
- To each GAMS model <model_name> an arbitrary number of targets and model variables can be assigned to by the corresponding sub-keyword 'get' and/or 'put'.
 To each sub-model (type = sub) at least one 'get' or one 'put' sub-keyword must be assigned to. The main model (type = main) and he configured without any sub-keyword 'get' and 'sut'. This is useful when
- main model (type = main) can be configured without any sub-keyword 'get' and 'put'. This is useful when the main model simply calls sub-models.
- Each model <model_name> in <model>.gdf with at least one sub-keyword 'get' has to have an \$include <model_name>_simenv_get.inc statement in the corresponding GAMS model file <model_name>.gms



- Each model <model_name> in <model>.gdf with at least one sub-keyword 'put' has to have an \$include <model_name>_simenv_put.inc statement in the corresponding GAMS model file <model_name>.gms
- There has to be exactly one main GAMS model, identified by the sub-keyword 'type' value main. All other models have to be of sub-keyword type value sub.
- The value-field for the sub-keyword 'put' is adapted to GAMS syntax to output GAMS model variables. Afterwards this output is used to generate the appropriate SimEnv output.
 <index_set> is mandatory for variables with a dimensionality > 0. Otherwise, specification of <index set> is forbidden. Indices as used in the GAMS model are separated from each other by comma.
- The sub-keyword 'time_limit' enables limitation of each GAMS model single run in the run ensemble to a
 maximum CPU-time consumption. If this threshold is reached the single run is aborted and the following
 single run started. In general, SimEnv nodata values will be assigned to the results of the aborted single
 runs. The sub-keyword 'time_limit' can be necessary since each GAMS model single run itself is an optimization procedure which could result in an unfeasible CPU time consumption. If the sub-keyword is
 not used in the gdf-file CPU-time cosumption per single run is unlimited.

gdf gdf		descr descr	GAMS model output description for the examples in the SimEnv
gdf		descr	User Guide
gdf		keep_runs	list 0,1
model	gams_model	descr	this is the only GAMS model to u
model	gams_model	type	main
model	gams_model	get	dem_ny
model	gams_model	get	dem_ch
model	gams_model	put	x.l(i,j):10:5
model	gams_model	put	a(i):10:5
model	gams_model	put	z.l
model	gams_model	put	modstat



GAMS description file <model>.gdf

If the model gams_model from the above Example 5.5 would be coupled with two additional GAMS sub-models sub_m1 and sub_m2 where both sub-models interact with SimEnv the GAMS description file could look like

model	gams_model	type	main
model	gams_model	put	modstat
model	sub_m1	type	sub
model	sub_m1	get	dem_ny
model	sub_m1	put	x.l(i,j):10:5
model	sub_m1	put	a(i):10:5
model	sub_m2	type	sub
model	sub_m2	get	dem_ch
Model	sub_m2	put	z.l



model	gams_model	type	main
model	sub_m1	type	sub
model	sub_m1	get	dem_ny
model	sub_m1	put	x.l(i,j):10:5
model	sub_m1	put	a(i):10:5
model	sub_m2	type	sub
model	sub_m2	get	dem_ch
model	sub_m2	put	z.l
model	sub_m2	put	modstat

Example 5.5 GAMS description file for coupled GAMS models

5.7.3 Files Created during GAMS Model Performance

Additionally to the files listed in Tab. 10.5, during the performance of a GAMS model the files <gams_model>_[pre | main | post].inc are created temporarily in the current working directory by <model>.ini and are deleted after the whole experiment where <gams_model> is a placeholder for the model of type main and all models of type sub in the gdf-file.

During experiment performance of a GAMS model each single run from the experiment is performed individually in a directory run<run_char> of the current working directory. Each directory is generated automatically before performing the corresponding single run and removed after perfomance of this single run. With the sub-keyword 'keep_runs' the user can force to keep sub-directories for later check of the transformed model code and its performance.

Unlike the other interface implementations GAMS main model terminal output for each single run is redirected to the file <model>.nlog in the directory run<run_char>. The modeler is responsible for re-direction of the terminal output from sub-models and from solvers. It is recommended to call all GAMS sub-models with the GAMS option string

ll=0 lo=2 lf=gams model.nlog dp=0

(see Example 15.6) which is also applied for the main model. With the options sub-keyword 'options' additional options can be specified for the main model.

5.8 Distributed Models

SimEnv supports performance of distributed models. A distributed model in SimEnv consists from a web or a chain of stand-alone sub-models, i.e., the model dynamics are computed by performing a set of stand-alone sub-models that normally exchange information.

Each of these sub-models can use SimEnv functionality, i.e., simenv_get_*, simenv_get_run_*, simenv_put_*, or simenv_slice_*. In each sub-model with SimEnv functionality simenv_ini_* and simenv_end_* calls have to be incorporated in. Sub-models can be implemented in different programming languages. Additionally, the corresponding SimEnv model coupling functionality at shell script level (simenv_*_sh modules) can be applied. As usual, the overall model is wrapped into a shell script <model>.run (see chapter 7).

The model description file <model>.mdf collects all the model output variables from all sub-models and the experiment description file <model>.edf collects all the targets from all sub-models.

In the model configuration file <model>.cfg the sub-keyword 'distributed' of the keyword 'model' announces a distributed model to the system. Its value has to be set to "yes" if



- More than one sub-model uses SimEnv functionality by the simenv_*_*-functions and
- Sub-models may get and send data from and/or to SimEnv data files in parallel. A distributed model where the sub-models are performed sequentially one by one can run with distributed = no.

Sub-models can reside on different machines. The only prerequisite is that the current working directory and the model output directory can be mapped to each of these machines.

5.9 Using Interfaced Models Outside SimEnv

To run a model interfaced to SimEnv outside the simulation environment in its native mode as before code adaptation the following simple changes have to be applied to the model:

- For Fortran and C/C++ models:
 - Link the model with the object library

\$SE_HOME/libsimenvdummy.a

instead of

\$SE_HOME/libsimenv.a.

For this library

- SimEnv function values (return codes) are zero
- simenv_get_* forwards target_val_def to target_val_adj
- simenv_get_run_* returns integer run number 0 and character run string ' ' (six blanks).
- For Python models:

Replace in the model source code from simenv import *

by

from simenvdummy import *

For this module

- SimEnv function values (return codes) are zero
- simenv_get_py forwards target_val_def to target_val_adj
- simenv_get_run_py returns run 000000.
- For GAMS models:

Handle in the model source code the include statements \$include <GAMS_model>_simenv_get.inc

and

\$include <GAMS_model>_simenv_put.inc
as comment.

6 Experiment Preparation

Experiment preparation is the first step in experiment performance of a model coupled to the environment. In an experiment description file <model>.edf all information to the selected experiment type and its numerical equipment is gathered in a structured way.

6.1 Experiment Description File <model>.edf

<model>.edf is an ASCII file that follows the coding rules in chapter 11.1 at page 107 with the keywords, names, sub-keywords, and value as in Tab. 6.1.

keyword	name	sub- keyword	Line type	Max. line nmb.	value	Explanation
edf	<nil></nil>	descr	0	any	<string></string>	experiment description
		type	m	1	[behaviour monte-carlo sensitivity optimization]	experiment type
target	<target_< td=""><td>descr</td><td>0</td><td>1</td><td><string></string></td><td>target description</td></target_<>	descr	0	1	<string></string>	target description
	name>	unit	0	1	<string></string>	target unit
		type	m	1	see Tab. 6.2	adjustment type
		default	m	1	<real_value></real_value>	target default value <target_def_val></target_def_val>
		adjusts	c3	1	<experiment- specific></experiment- 	experiment-specific informa- tion
specific	<nil></nil>	<experiment- specific></experiment- 	m	<ex- peri- ment- spe- cific></ex- 	<experiment- specific></experiment- 	experiment-specific informa- tion

 Tab. 6.1
 Elements of an experiment description file <model>.edf

To Tab. 6.1 the following additional rules and explanations apply:

- For the description of line type check Tab. 11.4 at page 109.
- Target names must differ from model variables and coordinate names in the model output description file (see chapter 5.1) and from built-in and user-defined operator names for model output post-processing (see chapter 8.5.2).
- A target name is the symbolic parameter / driver / initial value / boundary value name, corresponding to targets of the investigated model. Correspondence is achieved by applying the SimEnv model coupling function simenv_get_* in the model.
- The default value as specified in <model>.edf and not the default value from the model code is used to derive the adjusted value.
- For adjustment type multiply default <real_value> = 0. is forbidden.
- All experiment-specific information is explained in the appropriate chapters.
- Specify at least one experiment target.
- When preparing an experiment an experiment input file <model>.edf_adj is generated with the values to be finally used for the resulting adjustments. These values are applied to the default values of the targets according to the specified adjustment type (see Tab. 6.2 below) before finally influencing the dynamics of the model. The sequence of elements (columns) of each record of <model>.edf_adj corre-



sponds with the sequence of targets in the target name space (see chapter 11.1 at page 107), the sequence of records corresponds with the sequence of single model runs of the experiment. For each experiment a single model run with run number 0 is generated automatically as the nominal run of the model without adjustments. This run does not have an assigned record in <model>.edf_adj.

Adjustment type	Meaning
set	value setting: Use the adjustment to the target default value within the SimEnv function simenv_get_* as the final adjusted value. Not available for local sensitivity analysis
add	addition: Add the declared adjustment to the target default value within the SimEnv function simenv get * to get the final adjusted value to use.
multiply	multiplication: Multiply the declared adjustment with the target default value within the SimEnv function simenv_get_* to get the final adjustment to use. Differing implementation for local sensitivity analysis (check chapter 6.4.1).

Tab. 6.2 Adjustment types in experiment preparation

6.2 **Behavioural Analysis**

The experiment-specific information for experiment description files in Tab. 6.1 at page 39 is defined for behavioural analysis as follows:

keyword	name	sub- keyword	Line type	Max. line nmb.	value	Explanation
target	<target_ name></target_ 	adjusts	а	1	<value_list></value_list>	value list of target value ad- justments <adj_val> to apply. (for syntax see Tab. 11.6)</adj_val>
specific	<nil></nil>	comb	m	1 or any	[default <combination> file {<direct>/} <file_name>]</file_name></direct></combination>	information how to scan the spanned target space

Tab. 6.3 Experiment-specific elements of an edf-file for behavioural analysis

To Tab. 6.3 the following additional rules and explanations apply:

- For the description of line type check Tab. 11.4 at page 109. •
- For sub-keyword 'comb' the following rule holds: • value = [default | <combination>] for available sub-keyword 'adjusts'
 - for unavailable sub-keyword 'adjusts'
- value = [file {<direct>/}<file_name>] for unavailab
 Values of a value list have to be unique for available sub-keyword 'adjusts' and each target Assigned values from file {<direct>/}<file_name> can be multiple defined for each target.

6.2.1 Adjustments

Adjustment type	Set	Add	Multiply
adjusted target value =	<adj_val></adj_val>	<target_def_val> + <adj_val></adj_val></target_def_val>	<target_def_val> * <adj_val></adj_val></target_def_val>

6.2.2 The Combination

- The combination <combination> defines the way in which the space spanned by the experiment targets will be inspected by SimEnv: This is done by applying operators "" and "," to all stated experiment targets.
 - **The operator** "*" combines adjustments of different targets and so their resulting values combinatorially ("for all mesh points in a grid").
 - Compare with experiment description file (a) from Example 6.1 below.
 - The operator "," combines adjustments of different targets and so their resulting values parallel ("on the diagonal").

For the operator "," the targets must have the same number of adjustments.

Compare with experiment description file (b) from Example 6.1 below.

- The operator "," has a higher priority than the operator "*". Parentheses are not allowed: For example, p1 * p2, p3 * p4 always combines p2 and p3 in parallel and this combinatorially with p1 and p4. A parallel combination of p1 * p2 with p3 * p4 by (p1 * p2), (p3 * p4) is not possible. Compare with experiment description file (c) from Example 6.1 below.
- In <combination> each target has to be used exactly once.
- By the default combination **default** all experiment targets are combined combinatorially.
 - comb default of the experiment description file (a) from Example 6.1 below is equivalent to comb p1 * p2.
- Specification of file is only allowed for unused sub-keywords 'adjusts' all over the edf-file.
 - The adjustments are read from the adjustment data file {<direct>/}<file_name>.
 - All targets are assumed to be combined in parallel. Each record of the data file represents one simulation run. The sequence of the adjustments (sequence of columns) in each record corresponds with the sequence of the targets in the target name space (see chapter 11.1 at page 107).
 - Syntax rules for value lists at page 107 hold.
 - Identical adjustments for a target are allowed.
 - During model output post-processing restricted capabilities for the operator behav apply for this experiment layout.
 - Compare with experiment description file (d) from Example 6.1 below. Combination is implicitly as comb p1, p2. Experiment description files (b) and (d) in Example 6.1 below describe the same experiment.

6.2.3 Example

	The first three experiment description files (a) to (c) represent an experiment description according to Fig. 4.3 (a) to (c) at page 14.						
					Results in values		
(a)	edf		descr	Experiment description for	the examples		
	edf		descr	in the SimEnv User Guide (H	Fig. 4.3 (a))		
	edf		type	behaviour			
	target	p1	descr	parameter pl			
	target	p1	unit	without			
	target	p1	type	add			
	target	p1	default	1.			
	target	p1	adjusts	list 1, 3, 7, 8	2,4,8,9 for p1		



(b)	target target target target specific edf edf	p2 p2 p2 p2 p2	descr unit type default adjusts comb descr type	<pre>parameter p2 without multiply 2. list 1, 2, 3, 4 default Fig. 4.3 (b) behaviour</pre>	2,4,6,8 for p2
	target target target target target target specific	p1 p1 p2 p2 p2	type default adjusts type default adjusts comb	<pre>multiply 1. list 1, 3, 7, 8 multiply 2. equidist_end 1(0.5)2.5 pl,p2</pre>	1,3,7,8 for p1 2,3,4,5 for p2
(c)	edf edf target target target target target target target target target specific	p1 p1 p2 p2 p3 p3 p3	descr type type default adjusts type default adjusts type default adjusts comb	<pre>Fig. 4.3 (c) behaviour set 1. list 1, 3, 7, 8 set 2. equidist_end 1(1)4 multiply 3. list 1.1, 1.5, 2.4 p2,p1*p3</pre>	1,3,7,8 for p1 1,2,3,4 for p2 3.3,4.5,7.2 for p3
(d)	edf edf target target target specific	p1 p1 p2 p2	descr type type default type default comb	Fig. 4.3 (b) file world behaviour file world multiply 1 0 1. 3 1 add 7 2 2. 8 3 file world.dat_d 7 2	(1,2),(3,3),(7,4),(8,5) for (p1,p2)
				Example files: w	vorld.edf_a to world.edf_d

Example 6.1

Experiment description file <model>.edf for behavioural analysis

6.2.4 Experiment Performance

- Firstly, a model run 000000 with the default values of the experiment targets is performed.
- According to the sub-keyword 'comb' the appropriate runs are generated.
- The sequence of the runs corresponds with the sequence of the adjustments in the ASCII file <model>.edf_adj (check chapter 6.1 at page 39 for more information).

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6.3 Monte Carlo Analysis

The experiment-specific information for experiment description files in Tab. 6.1 at page 39 is defined for Monte Carlo analysis as follows:

keyword	name	sub- keyword	Line type	Max. line nmb.	value	Explanation
target	<target_ name></target_ 	adjusts	m	1	[<distribution> file {<direct>/} <file_name>]</file_name></direct></distribution>	distribution and distribution parameters to be applied for the target or import of an external sample <distr_val> from <file_name></file_name></distr_val>
		sample	c4	1	[random latin hypercube]	sampling strategy: random or latin hypercube sampling LHS
specific	<nil></nil>	runs	m	1	<nr_of_runs></nr_of_runs>	number of runs > 10 to be performed for the experiment

Tab. 6.4	Experiment-specific elements of an edf-file for Monte Carlo analysis
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To Tab. 6.4 the following additional rules and explanations apply:

- For the description of line type check Tab. 11.4 at page 109.
- <distribution> = <distr_shortcut> (<distr_param_1> { , <distr_param_2> }) (check Tab. 6.5)
- For implicitly specified distributions according to Tab. 6.5 adjustments are applied to the specified distribution parameters of the distributions. Afterwards, a sample <distr_val> is generated from the distribution with the adjusted distribution parameters. Adjustment types add and multiply are not applied to the distribution parameter <distr_param> = standard deviation. Instead, the specified standard deviation from the experiment description file is used (adjustment type set is applied).
- For explicitly specified samples of any distribution by the ASCII file <file_name> adjustments are applied directly to the sample values <distr_val> from the file. For syntax rules for files check chapter 11.1. Each record of the ASCII file can hold only one sample value. Sample size has to be identical to <nr_of_runs> from the keyword 'specific'.
- In random sampling, there is no assurance that sampling points will cover all regions of the selected distribution. With Latin hypercube sampling LHS this shortcoming is reduced: The sampling range of the target is divided into <nr_of_runs> intervals of equal probability according to the selected distribution and from each interval exactly one sampling point is drawn. For more information on LHS check Fig. 6.1 below and see Imam & Helton (1998) and Helton & Davis (2000).
- The number of runs <nr_of_runs> must be greater than 10.



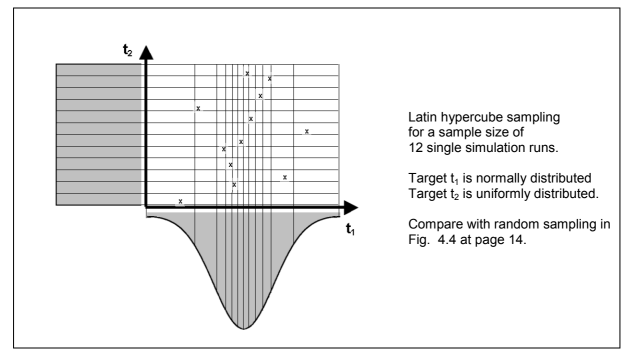


Fig. 6.1 Monte Carlo analysis: Latin hypercube sampling

6.3.1 Adjustments

Adjustment type	Set	Add	Multiply	
for distribution:	<distr_param></distr_param>	<target_def_val> +</target_def_val>	<target_def_val> *</target_def_val>	
adjusted distr_param =		<distr_param></distr_param>	<distr_param></distr_param>	
		not for standard deviation instead, adjustment type value "set" is applied		
for file:	<distr_val></distr_val>	<target_def_val> +</target_def_val>	<target_def_val> *</target_def_val>	
adjusted target_value =		<distr_val></distr_val>	<distr_val></distr_val>	

6.3.2 Distribution Functions and their Parameters

Distribution function	distr_ shortcut	distr_param_1	distr_param_2	Restriction
uniform	U	lower boundary	upper boundary	lower boundary < upper boundary
normal	Ν	mean value	variance	variance > 0
lognormal	L	mean value of a normally distributed target	variance of a nor- mally distributed target	variance > 0
exponential	E	mean value		mean value > 0

Tab. 6.5Probability density functions and their parameters

For more information on the distribution functions see chapter 4.3 and Tab. 4.2.

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6.3.3 Example

```
Experiment description for the examples
(e) edf
                    descr
    edf
                    descr
                                 in the SimEnv User Guide
    edf
                                Monte-Carlo
                    type
              p2
                    descr
                                parameter pl
    target
   target p2 unit without
target p2 type multiply
target p2 default 2.
target p2 sample latin hypercube
                                                       p2 is a realization of a uniform distrib.
    target p2 adjusts
                              distr U(0.5,1.5)
                                                       between 0.5*2 and 1.5*2
                                add
    target pl type
    target p1 default
                                1.
    target p1 sample
                               random
                                                       p1 is a realization of a normal distribution
    target p1 adjusts
                                distr N(0,0.4)
                                                       with mean = 1+0 and variance = 0.4
              pЗ
                                 add
    target
                   type
              p3 default
    target
                                 3.
                                                       realization of p3 is read from file
    target
              p3 adjusts
                                 file world.dat e
                                                       world.dat e and afterwards 3 is added
    specific
                    runs
                                 250
                                                                     Example file: world.edf e
```

Example 6.2 Experiment description file <model>.edf for Monte Carlo analysis

6.3.4 Experiment Performance

- Firstly, a model run 000000 with the default values of the experiment targets is performed which represents the deterministic case.
- The sequence of the runs corresponds with the sequence of the adjustments in the ASCII file <model>.edf_adj. <model>.edf_adj is generated from random numbers of the appropriate distributions U(0,1), N(0,1), L(0,1), and/or E(1). For more information on <model>.edf_adj check chapter 6.1 at page 39.
- If the resulting distribution parameters do not fulfill the restrictions in Tab. 6.5 the following adaptations are applied and corresponding warnings are output to the model interface log-file <model>.mlog.

Distribution	Condition	Adaptation
U	lower boundary > upper boundary	boundaries are interchanged
U	lower boundary = upper boundary	lower boundary := lower boundary - 0.5 upper boundary := upper boundary + 0.5
E	mean < 0	mean := -mean
E	mean = 0	mean := abs(model default value) for model default value ≠ 0 1 else

Tab. 6.6

Probability density functions: Distribution parameters - conditions and adaptation



6.4 Local Sensitivity Analysis

The experiment-specific information for experiment description files in Tab. 6.1 at page 39 is defined for local sensitivity analysis as follows:

keyword	name	sub- keyword	Line type	Max. line nmb.	value	Explanation
target	<target_ name></target_ 	adjusts	f	0		sub-keyword is forbidden for this experiment type
specific	<nil></nil>	incrs	m	1	<value_list></value_list>	increments <incr_val> > 0. for all targets. <incr_val> in <value_list> has to be ordered in a strictly monotonic in- creasing manner. (for syntax see Tab. 11.6)</value_list></incr_val></incr_val>

 Tab. 6.7
 Experiment-specific elements of an edf-file for local sensitivity analysis

To Tab. 6.7 the following additional rules and explanations apply:

- For the description of line type check Tab. 11.4 at page 109.
- Values from the value list must be positive and unique.
- **Note** that computation of adjusted values for adjustment type multiply in local sensitivity analysis differs from all other experiment types (see chapter 6.4.1 below.

6.4.1 Adjustments

Adjustment type	Set	Add	Multiply
adjusted target value =	undefined for this	<target_def_val> ±</target_def_val>	<target_def_val> *</target_def_val>
	experiment type	<incr_val></incr_val>	(1 ± <incr_val>)</incr_val>

As an example, the absolute sensitivity function (see Tab. 4.3 at page 17) is then as follows:

for adjustment type Add sens_abs(<target_def_val>,±<incr_val>) = <u>z(<target_def_val> ± <incr_val>) - z(<target_def_val)</u> <u>± <incr_val></u>

for adjustment type Multiply

6.4.2 Example

```
(f) edf
                            Experiment description for the examples
                  descr
   edf
                  descr
                            in the SimEnv User Guide
   edf
                            sensitivity
                  type
          pl
   target
                  descr
                            parameter pl
          p1
   target
                  unit
                            without
          p1
   target
                            add
                  type
   target
           p1
                  default
                            1.
   target p2 type
                            multiply
   target
           p2
                 default
                            2.
           pЗ
   target
                type
                            multiply
           pЗ
   target
                  default
                            3.
                            list 0.001,0.01,0.05,0.1
   specific
                  incrs
                                                        Example file: world.edf
```

Example 6.3 Experiment description file <model>.edf for local sensitivity analysis

6.4.3 Experiment Performance

- Each experiment target will be adjusted by the same increments as those stated in the incrs info-field
- For finite sensitivity functions several runs have to be performed:
 - A nominal run with the default values of the experiment targets (run number 000000)
 - Per target and per increment two runs with the default values of all targets except that one under consideration, where the adjustment is applied according to the above adjustment rules
 - Accordingly, the number of resulting runs is 2 * number_of_targets * number_of_increments + 1
- Results of each model run are stored and sensitivity functions are applied during model output postprocessing.

The following sensitivity functions can be performed:

Linear, squared, absolute, relative as well as a symmetry test.

- The sequence of the simulation runs is determined in the following manner:
- nominal run

Іоор	over increment sequence loop over experiment targets end loop	
end loop		
Іоор	over negative increment sequence loop over experiment targets end loop	

end loop



6.5 Optimization

The experiment-specific information for experiment description files in Tab. 6.1 at page 39 is defined for local sensitivity analysis as follows:

keyword	name	sub- keyword	Line type	Max. line nmb.	value	Explanation
target	<target_ name></target_ 	adjusts	m	1	<lower_bound>: <upper_bound></upper_bound></lower_bound>	real valued lower bound and upper bound to define the target range where the cost function is to be minimized on. <lower_bound> ≤ <target_def_val> ≤ <upper_bound></upper_bound></target_def_val></lower_bound>
specific	<nil></nil>	cost_fct	m	1	<result></result>	cost function to minimize. 0-dimensional result formed according to the rules of the SimEnv post-processor. Do not apply multi-run operators.

 Tab.
 6.8
 Experiment-specific elements of an edf-file for an optimization experiment

To Tab. 6.8 the following additional rules and explanations apply:

• For the description of line type check Tab. 11.4 at page 109.

6.5.1 Adjustments

Adjustment type	Set	Add	Multiply
adjusted target value =	<adj_val></adj_val>	<target_def_val> + <adj_val></adj_val></target_def_val>	<target_def_val> * <adj_val></adj_val></target_def_val>

6.5.2 Example

(g)	edf edf edf		descr descr type	Experiment description for the examples in the SimEnv User Guide optimization
	target target target target target	p1 p1 p1 p1 p1	descr unit type default adjusts	<pre>parameter p1 without set 112:12 minimize cost function for p1c <-12, 12></pre>
	target target target target target target	p2 p2 p3 p3 p3	type default adjusts type default adjusts	set 2. 1:10 set 3. -12:12

```
target
            p4
                   type
                               set
                   default
            p4
target
                               4.
            p4
                   adjusts
                               1:10
target
specific
                   cost fct
                               -sum(bios)
                                             maximize bios over land masses
                                                                 Example file: world.edf g
```

Example 6.4

Experiment description file <model>.edf for an optimization experiment

6.5.3 Experiment Performance

- This is the only experiment type where the adjustments for the targets of the single runs are not determined before the experiment but in the course of the experiment by the optimization algorithm. Consequently, the file <model>.edf_adj is not created during experiment prepration but is written during experiment performance.
- In parallel to the file model>.edf_adj an ASCII file **<model>.edf_cf** is written during experiment performance with the value of the cost function for each of the single runs.
- The status of an optimization experiment can be acquired by the SimEnv service simenv.sts. For more information check Tab. 10.3.
- The optimization algorithm itself is controlled by additional technical parameters and options that are normally fixed by SimEnv. To modify them copy from \$SE_HOME the ASCII file **simenv.oopt** to the directory you want to start an optimization experiment and edit this file. During the experiment the edited file is used instead of the file with the default constellation in \$SE_HOME. The description of the options and parameters can be found in Ingber (2004).
- Optimization experiments can not restarted by the SimEnv service simenv.rst.
- The values for the sub-keywords 'begin_run' and 'end_run' in the configuration file <model>.cfg are ignored for an optimization experiment. The experiment always starts with run number 0 and ends if one of the criteria in the file simenv.oopt (see above) is fullfilled.
- The optimization return code, the optimal targets, the corresponding value of the cost function and the number of the corresponding single run as the results of the optimization experiment are documented at the end of the model interface log-file <model>.mlog.
- A protocol from the optimization procedure is delivered by SimEnv in the ASCII file <model>.olog.
- The initial seed for the optimization technique is fixed. That's why the algorithm results for the same optimization problem always in the same sampling sequence in the target space.

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7 Experiment Performance

After experiment preparation experiment performance is the second step in running a model coupled to SimEnv. Each multi-run experiment can be performed sequentially or in parallel. Besides a new-start of an experiment a restart after an experiment interrupt or only for an experiment slice can be handled by SimEnv.

7.1 General Approach

SimEnv enables performance of an experiment sequentially on the login-machine and in parallel and/or sequential mode in a job class controlled by the LoadLeveler.

In parallel mode the single runs of the run ensemble are distributed to all allocated nodes with their assigned processors. One communication processor is responsible for experiment management. Parallel experiment performance is controlled by the parallel operating environment POE and the LoadLeveler.

Experiments may be performed partially only for a run slice out of the run ensemble. Experiment slices are controlled by the general configuration file <model>.cfg.

For successive performance of run slices and/or after abnomal experiment interrupt experiments can be restartet. The experiment log-file <model>.elog is analyzed to identify these single runs out of the run ensemble that have to be performed the first time and/or anew and the corresponding output data structure is appended to the output data that already exists for this experiment.

For all experiment settings the user model has to be wrapped in a shell script <model>.run.

7.2 Experiment Start

• The user can define an experiment preparation shell script **<model>.ini** that is performed additionally after standard experiment preparation when starting a new experiment. For experiment restart <model>.ini is performed only on request (see chapter 7.3 below).

In <model>.ini additional settings / checks can be performed. For return codes unless zero from <model>.ini the experiment will not be started.). Make sure that <model>.ini has execute permission by UNIX command chmod u+x <model>.ini. For Python and GAMS models <model>.ini is a mandatory script with pre-defined contents. Check chapters 5.5.1 and 5.7.1 for more information.

- The model to be applied within the SimEnv experiment has to be wrapped in the shell script <model>.run. <model>.run is performed for each single run within the run ensemble.
 - Make sure that
 - . \$SE_HOME/simenv_ini_sh is the first command and
 - . \$SE_HOME/simenv_end_sh is the last command
 - in <model>.run (see Tab. 5.7 at page 31 and Example 7.2 below).
 - Ensure by UNIX command chmod u+x <model>.run that <model>.run has execute permission.
 - To cancel the whole experiment after the performance of the current single run due to any condition of this run make sure a file **<model>.\$run_char.err** exists as an indicator to stop. You can create this file in the model or in <model>.run. For the latter
 - Perform . \$SE_HOME/simenv_get_run_sh to get the current run number <run_int> and <run_char> (see Tab. 5.7 at page 31 and Example 7.2 below).
 - Touch the file <model>.\$run_char.err.
 - For GAMS models <model>.run has a pre-defined structure. Check chapter 5.7.1 for more information.
- The model variables to be output during experiment performance are declared in the model output description file <model>.mdf



- The type and the targets of the experiment to be performed are declared in the experiment description file <model>.edf
- Mapping between experiment targets and targets in the model source code is achieved by application of the generic SimEnv function simenv_get_* in the model code or at shell script level.
- Output of model variables declared in <model>.mdf into SimEnv structures is achieved by the application of the generic SimEnv function simenv_put (and simenv_slice) in the model source code.
- Model output from run number <run> is stored in the file <model>.out<run_char>.[nc | ieee] if the sum over all model output variables of a single run is less than the appropriate value specified in <model>cfg. Otherwise, model output from the complete experiment is stored in <model>.outall.[nc | ieee].
- For each experiment type a run number 0 with the default values of all experiment targets will be performed additionally to the runs declared in the experiment description file <model>.edf.
- During experiment performance a model interface log-file <model>.mlog is written where adjustments of
 experiment target values and possibly workarounds for wrong re-adjustments (only for experiment type
 Monte Carlo analysis, see Tab. 6.6) are stored. All model output to the terminal is re-directed within
 SimEnv to the model interface log-file <model>.mlog.
- During experiment performance an experiment log-file <model>.elog is written with the minutes of the experiment.
- Do not start and/or submit another experiment from a working directory where an experiment is still running.
- After the experiment has been finished the model-specific output from the experiment can be wrapped up with the optional shell script **<model>.end**.
- After the experiment has been finished an e-mail is send on demand (check chapter 10.1) to the address as specified in <model>.cfg.
- For more information check Fig. 7.1.

For the shell script world_*.ini the following contents could be defined:

```
# coarse 0.5° x 0.5° land-sea mask from file land_sea_mask.05x05
# in the current directory
# to a 4° x 4° resoluted land-sea-mask in file land_sea_mask.coarsed
# in the current directory to use for all single runs
land_sea_mask 4 4
rc_land_sea_mask=$?
# exit from world_*.ini with return code != 0
# as an indicator not to start the experiment
exit $rc_land_sea_mask
```

Example files: world_[f | c | cpp | py | sh].ini

Example 7.1 Shell script <model>.ini for user-model specific experiment preparation

For the shell script world_f.run the following contents could be defined:
always perform at begin:
. \$SE_HOME/simenv_ini_sh
run the model:
world_f
assuming a model return code != 0 as an indicator to stop
the whole experiment for any reason.
Touch the file below in the current working directory \$SE_WD
as an indicator to SimEnv for this.

```
if test $? -ne 0
then
    . $SE_HOME/simenv_get_run_sh
    touch $SE_WD/world_f.$run_char.err
fi
# always perform at end:
. $SE_HOME/simenv_end_sh
```

Example file: world_f.run

Example 7.2 Shell script <model>.run to wrap the user model

For the shell script world_f.end the following contents could be defined:

```
# remove the file of the coarsed land-sea mask
rm -f land_sea_mask.coarsed
```

Example file: world_[f | c | cpp | py | sh].end

Example 7.3 Shell script <model>.end for user-model specific experiment wrap-up

7.3 Experiment Restart

When an experiment was interrupted / has failed due to any reason or in the case of partial experiment performance (see chapter 7.4 below) it can be restarted:

- Simply restart the experiment by simenv.rst without changing any of the SimEnv files describing the experiment and/or the model. The only exception may be the values for the sub-keywords of the experiment-keyword in the general model configuration file <model>.cfg.
- simenv.rst has the same usage as simenv.run
- Restart can be launched on an other machine / in an other job class than that of the interrupted experiment.
- Dependent on the experiment log-file <model>.elog, written in the interrupted experiment a single model run from the complete run ensemble in the restart experiment will be
 - Performed if this run has neither a start nor a finish information in the elog-file
 - Not performed if this run has a start as well as a finish information in the elog file
 - Performed anew if the run has a start information but no finish information in the elog-file. For this case a model restart shell script <model>.rst can be provided by the user optionally to prepare restart of this single model run (e.g., by deleting non-SimEnv temporary or output files). Make sure that <model>.rst has execute permission by UNIX command chmod u+x <model>.rst.
 Make sure that
 - . \$SE_HOME/simenv_ini_sh is the first command and
 - . \$SE_HOME/simenv_end_sh is the last command

in <model>.rst (see Tab. 5.7 at page 31 and Example 7.4 below).

After running \$SE_HOME/simenv_get_run_sh the shell script variables run_int and run_char are available in <model>.rst (see above).

- Experiment restart works without standard SimEnv experiment preparation. Instead, experiment preparation files and other information from the interrupted experiment will be used.
- For a restart, the optional experiment preparation shell script <model>.ini will be performed only on demand. This request is specified in the configuration file <model>.cfg with the sub-keyword 'restart_ini'.
 For Python and GAMS models interfaced to SimEnv restart_ini has to be set to yes (check chapters 5.5.1 and 5.7.1)



- <model>.cfg will be checked anew for experiment restart. Do not change for a restart any of the information related to the keyword 'model' in <model>.cfg.
- Minutes of the restarted experiment will be appended to the <model>.mlog and <model>. elog files, respectively from the interrupted experiment.
- Restart can be applied to an experiment several times successively.
- Experiment restart can be performed also as partial experiments, independently on the partial status of the original model
- Experiment re-start is not possible for the experiment type optimization.

For the model world_sh (check Example 15.5 at page 137) the following contents could be defined for the restart script world_sh.rst:

```
# always perform at begin
. $SE_HOME/simenv_ini_sh
# get run number
. $SE_HOME/simenv_get_run_sh
# remove all files from the temporary directory and the directory itself
if test -d run$run_char
then
    cd run$run_char
    rm -f *
    cd ..
    rmdir run$run_char
fi
# always perform at end
. $SE_HOME/simenv_end_sh
Example file: world_sh.rst
```

Example 7.4 Shell script <model>.rst to prepare model performance during experiment restart

7.4 Experiment Partial Performance

- SimEnv enables to perform an experiment partially by performing only a run slice out of the whole run ensemble.
- Therefor assign appropriate run numbers to the corresponding sub-keywords 'experiment' in <model>.cfg.
- Make sure that begin run number and end run number represent run number from the experiment (including run number 0) and that begin run number ≤ end run number.
- A partial experiment performance is also possible for an experiment restart.
- For more information check Fig. 7.1.

7.5 Job Control by POE and LoadL

- For parellel experiment performance controlled by the parallel operating environment POE and the LoadLeveler make sure that the environment variable SE_HOME is set in your .profile-file correctly.
- On a login node to a parallel machine there is an additional SimEnv dialogue whether the experiment is to be submitted by POE and the LoadLeveler to a parallel or sequential job class of this parallel machine or is to be performed locally at the login node.
- Default job control files are supplied by SimEnv to ensure communication with POE and LoadLeveler. These job control files may be copied to the current working directory, can be modified and will then be used instead of the default job control files to start an experiment at a parallel or sequential job class. If necessary, copy \$SE_HOME/simenv.jcf_par and/or \$SE_HOME/simenv.jcf_seq to the current working directory SimEnv is started from, modify the file(s) according to the needs of the experiment you want to perform and / or the machine you want to use and start afterwards simenv.run (or simenv.rst). If available in the current working directory, these modified job control files are used instead of the original files in \$SE_HOME.

simenv.jcf_seq submits a job to a sequential batch class, simenv.jcf_par to a parallel batch class.

• Default job control files enable automatic restart of the experiment by the LoadLeveler after an interrupt of the job in a parallel or sequential job class caused by POE, the LoadLeveler or the operating system. The user does not need to restart the experiment manually after such an event.



Experiment-Related User Scripts and Files 7.6

Script / file	Explanation	Used for (*)	Exist status
	Scripts (**)		
<model>.run</model>	 model shell script to wrap the model executable .\$SE_HOME/simenv_ini_sh has to be the first command in <model>.run</model> .\$SE_HOME/simenv_end_sh has to be the last command in <model>.run</model> Model coupler functions at shell script level can be applied in <model>.run</model> Pre-defined contents for GAMS models (check chapter 5.6) 	S R	mandatory
<model>.rst</model>	<pre>model shell script to prepare single model run restart for such single runs that were started by not finished during the previ- ous experiment start / restart . \$SE_HOME/simenv_ini_sh has to be the first command in <model>.rst . \$SE_HOME/simenv_get_run_sh can be applied in <model>.rst (check chapter 5.6)</model></model></pre>	R	optional
<model>.ini</model>	 model shell script to prepare simulation experiment additionally to standard SimEnv preparation Experiment will be not performed if return code from this script is unequal zero. For experiment re-start <model>.ini will be performed only on request.</model> 	S (R)	optional, for Python and GAMS models mandatory
<model>.end</model>	model shell script to clean up simulation experiment from non- SimEnv files	SR	optional
	Files		
<model>. <run_char>.err</run_char></model>	touch this file in the model, in <model>.run and/or <model>.rst as an indicator to stop the complete experiment after single run <run_char> has been finished</run_char></model></model>	A	optional
simenv.jcf_par	user-specific job control file to submit an experiment to a par- allel class by the LoadLeveler Copy from \$SE_HOME on demand	L	optional
simenv.jcf_seq	user-specific job control file to submit an experiment to a se- quential class by the LoadLeveler. Copy from \$SE_HOME on demand	L	optional
simenv.oopt	user-specific control and option file for experiment type Opti- mization Copy from \$SE_HOME on demand	0	optional

(*): script applied for

Restart of an experiment by \$SE_HOME/simenv.rst <model> R:

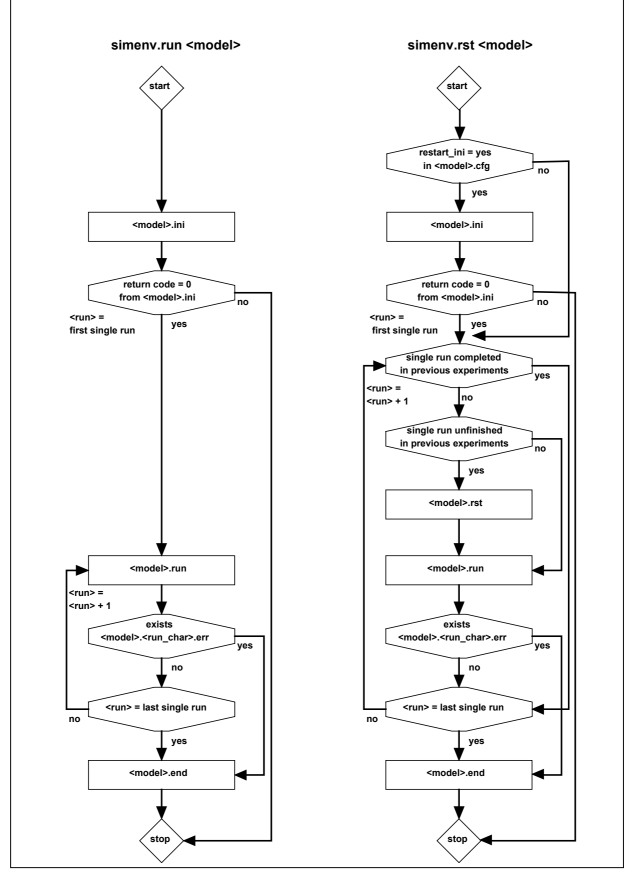
S: Start of an experiment by \$SE_HOME/simenv.run <model> file applied for

- All experiment perform. at the login machine or by LoadLeveler submission А:
- L: LoadLeveler experiment submission
- О: **O**ptimization experiment performance

(**): make sure by Unix command chmod u+x <model>.???

that the shell script <model>.??? has execute permission

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Flowcharts for performing simenv.run and simenv.rst First and last single run always refer to the corresponding settings in <model>.cfg



7.7 Saving Experiments

To save experiments for later use, e.g., by SimEnv post-processing, make sure to store from the experiment the following files:

- <mdel>.out[all | <run_char>].[nc | ieee]
- <model>.cfg
- model>.mdf
- <model>.edf
- <model>.edf_adj (for optimization)
- <model>.edf_cf (for optimization)
- <model>.elog (optional)
- <model>.mlog (optional)
- <model>.nlog (optional)
- <model>.olog (optional, for optimization)

from the model output directory from the current working directory

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8 Experiment Post-Processing

Goal of post-processing is to navigate within the model / experiment output space by deriving interactively output functions / data that are to be visualized in experiment evaluation afterwards. Therefor SimEnv supplies operators that can be applied to model output and reference data. There are built-in basic and advanced operators and built-in experiment-specific operators. The user can define its own private operators and easily couple them to the post-processor. Operator chains and recursions are possible. Macros can be defined as abbreviations for operator chains.

8.1 General Approach

8.1.1 Post-Processor Results

In SimEnv post-processing post-processor results (synonym: output functions) are derived from model output of the experiment and from reference data. A post-processor result is specified by a post-processor expression, optionally prefixed by a result description and a result unit string:

<result> = { { <result_description> } { [<result_unit>] } = } <result_expression>

<result></result>	by the string "Enter a result" the user is requested to enter a result. Input lines with a character # as the first non-white spaced character are treated as comments. The post-processor session is finished by entering <ret> or a sequence of white spaces instead of a result. For case sensitivity of <result> check Tab. 10.9 at page 105.</result></ret>
<result_description> <result_unit></result_unit></result_description>	must not contain an apostrophe character '. characters "[" and "]" belong to the syntax and are not a part of the document convention as defined in Tab. 1.1
<result_expression></result_expression>	is a chain of SimEnv operators applied to applied to model output variables and/or reference data. Can be continued on a new input line (continue expression:) if the current input line ends on one of the operators "+", "-", "*", "/", or "**" or on the operand separator "," in operators. White spaces are filtered out from the result expression string, also from character arguments.

<result_description> or <result_unit> are used to describe the result in the corresponding result output file (see chapter 12). For the case one of these entities is not specified SimEnv analyses the result expression: For a result expression formed without any operator or only from one operator and using exactly one model output variable and/or one experiment target <result_description> and/or <result_unit> is copied from the corresponding information for the sub-keyword 'descr' in <model>.mdf (for a model output variable as an operand of this operator) and/or from <model>.edf (for an experiment target as an operand of this operator). The only operator used in this expression must not transform the contents of the operand in general (must be invariant with respect to description and unit). For all other cases <result_description> is set to the string res_<xy> and <result_unit> is undefined.

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Having a model variable definition as in Example 5.1 at page 26 then in model output post-processing

abs(atmo)+3	applies operator abs to atmo and adds 3 (multi-operator result expression) <result_description> = 'res_<xy>' <result_unit> undefined</result_unit></xy></result_description>
<pre>Energy [MWh] = abs(atmo)+3</pre>	as above, but: <result_description> = 'Energy' <result_unit> = 'MWh'</result_unit></result_description>
[MWh] = abs(atmo)+3	as above, but: <result_description> = 'res_<xy>' <result_unit> = 'MWh'</result_unit></xy></result_description>
sign(atmo)	applies operator sign to atmo (operator sign is not invariant w.r.t. the contents of its operand) <result_description> = 'res<xy>'</xy></result_description>
abs(atmo)	<result_unit> undefined applies operator abs to atmo (operator abs is invariant w.r.t. the contents of its operand) <result_description> = 'aggregated atmospheric state' (according to <model>.mdf) <result_unit> = 'without' (according to <model>.mdf)</model></result_unit></model></result_description></result_unit>
Energy = abs(atmo)	applies operator abs to atmo <result_description> = 'Energy' (according to <model>.mdf) <result_unit> = 'without' (according to <model>.mdf)</model></result_unit></model></result_description>

Example 8.1 Addressing results in model output post-processing

8.1.2 Operands

Operands in result expressions can be

- Model output variables (see below)
- Example: atmoExperiment targets
- Example: p1
- Constants in integer or float notation Example: 10.
- Character strings Example: `tie_avg'
- Operator results Example: abs(atmo)
- Macros (see chapter 8.7) Example: equ_100yrs_m

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As for model variables (see chapter 5.1) also to each operand (with the exception of character string operands)

 Dimensionality Extents
 Coordinates
 dim(operand) ext(operand,i)
 dim(operand) ext(operand,i)
 with i=1,...,dim(operand) with i=1,...,dim(operand)

are assigned to. The dimensionality is the number of dimensions, an extent is related to each dimension and represents the number of elements in that dimension. Extents are always greater than 1. To each dimension a coordinate is assigned to. Coordinates have a name and from all coordinate values the coordinate is defined for a subset is assigned to the extent of the dimension of the operand. Coordinate specification for operands follows that for model output variables. For more information see chapter 5.1.

- Operators transform dimensionality, dimensions, and coordinates of the their non-character operator arguments into unique dimensionality, dimensions and coordinates of the operator result (see chapter 8.1.4).
- Consequently, the output of an operator and finally a post-processor result as a sequence of operators applied to operands also has unique dimensionality, extents and coordinates.
- Experiment targets and constants always have a dimensionality of 0.
- Operands of dimensionality 0 and character string operands do not have a coordinate assignment.

8.1.3 Model Output Variables

- A variable of dimensionality n corresponds with a n-dimensional array and is defined at an n-dimensional grid, spanned up from the coordinate values of the assigned coordinates The complete data field of a model output variable or parts of it can be addressed in model output post-processing (see below). Dimensionality, dimensions and coordinate description of this data field is derived from the model variable description in <model>.mdf.
- Model output variables are specified in the ASCII model output description file <model>.mdf (see Tab. 5.3 at page 24) by their
 - Name
 - Dimensionality
 - Extents
 - Coordinate assignment to each dimension
 - Data type (see Tab. 5.4 at page 25).
 - Use the service simenv.chk to check variables description in model output description file <model>.mdf
- Addressing of model output data fields or parts of it is done in model output post-processing by corresponding model output variables names.

• For variables with a dimensionality greater than 0 it is possible to address only a part of the whole variable field by

- Specifying for a dimension an index range by

 i = <index_value₁> { : <index_value₂> }
 index_value₁> ≤ <index_value₂>
 <index_value₂> = <index_value₁> if <index_value₂> is missing.
 i = stands for index addressing
- Specifying for a dimension a coordinate range by c = <coordinate_value₁> { : <coordinate_value₂> } <coordinate_value₁> ≤ <coordinate_value₂> for strictly increasing coordinate values <coordinate_value₁> ≥ <coordinate_value₂> for strictly decreasing coordinate values <coordinate_value₁> = <coordinate_value₂> if <coordinate_value₂> is missing c= stands for coordinate addressing
- Index and coordinate ranges are separated from each other by a comma, the sequence of ranges for all dimensions is enclosed in brackets and is appended after the variable name.
- For one variable c= and i= can be used in mixed mode for different dimensions.
 - * denotes the complete range of a dimension. c= * is identical to i= * is identical to *
- In the general SimEnv configuration file <model>.cfg (see chapter 10.1 at page 95) a global default for index and/or coordinate addressing is established for the whole post-processing session. This global default can be overwritten locally by using c= and/or i=.



Having a model variable definition as in Example 5.1 at page 26 then in model output post-processing result expressions can be

atmo	and
atmo(*,*,*,*)	and
atmo(c=*,*,i=*,*)	and
atmo(c=88:-88,c=-178:178,c=1:16,	
atmo(i=1:45,i=1:90,i=1:4,i=1:20)	and
atmo(i=1:45,c=-178:178,*,*)	and
atmo(1:45,1:90,1:4,1:20)	and (with address_default = index in model.cfg)
atmo(1:45,c=-178:178,1:4,1:20)	and (with address_default = index in model.cfg)
	all address all 45*90*4*20 values and
	the following holds true for this addressed variable:
	Dimensionality = 4
	Coordinates = lat , lon , level , time
	Extents = 45, 90, 4, 20
atmo(*,*,*,c=11:20)	addresses all values of last 10 decades Dimensionality = 4
	Coordinates = lat , lon , level , time
	Extents = 45 , 90, 4, 10
atmo(*,*,c=1,c=1)	addresses all values of the first decade for level 1
	Dimensionality = 2
	Coordinates = lat , lon
	Extents = 45, 90
atmo(c=0,*,1,i=20)	addresses all values of level 1 for the last decade at
	equator
	Dimensionality = 1
	Coordinates = Ion
	Extents = 90
atmo(i=23,*,1,i=20)	addresses all values of level 1for the last decade at
	equator Dimensionality = 1
	Coordinates = Ion
	Extents = 90
atmo(c=0, c=2, c=1, c=20)	addresses the value for the last decade at
	$(lat,lon,level,time) = (0^{\circ}, 2^{\circ}, 1, 20)$
	Dimensionality = 0
	Coordinates = (without)
	Extents = (without)
atmo(c=0,c=1:9,c=1,c=20)	addresses the values for the last decade at
	$(lat, lon, level, time) = (0^{\circ}, 2^{\circ}, 1, 20) and (0^{\circ}, 6^{\circ}, 1, 20)$
	Dimensionality = 1
	Coordinates = lon
	Extents = 2
atmo(c=0, c=1, c=1, c=20)	error in addressing: c=1 for lon does not exist
	Example file: world.post_bas
	, , <u>.</u>

Example 8.2 Addressing model output variables in model output post-processing

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8.1.4 Operators

• Operators transform dimensionality, dimensions, and coordinates of the their non-character operator arguments into unique dimensionality, dimensions and coordinates of the operator result (check chapter 8.1.2).

There are

- Single-argument operators that replicate dimensionality, dimensions and coordinates from the only
 argument to the operator result
 Example: sin(atmo)
- Multi-argument operators that demand a certain relation between dimensionalities, dimensions and coordinates of their arguments

Example: mod(atmo(c=84:-56,*,c=1,*),bios)

- Operators that increase the dimensionality of the operator result and assign new coordinates to the additional dimensions (check Tab. 10.7) or form new coordinates from resulting target adjustments Example: ens(atmo)
- SimEnv post-processing operators may have two special types of arguments:
 - Character arguments: Only character strings enclosed in ' ' are valid as arguments. Some built-in operators (e.g., count) have a pre-defined set of valid character argument strings (e.g., for operator count strings all, def, and undef). Some built-in operators allow an empty string (e.g., behav) Example: run (`0', atmo)
 - Integer or float constant arguments: Only constants in appropriate format are valid as arguments. Model variables of dimensionality 0 or general operands with dimensionality 0 are invalid. Example: classify(20, atmo)
 - If defined, character and constant arguments are always the first arguments of an operator. If both argument types are defined for an operator then the character arguments are followed by the constant arguments.
 - Example: hgr_l('1000', 20, atmo)
- Operators are generic with respect to the data types of their operands: Each non-character and nonconstant argument can be used with operands of all defined data types (see chapter 5.1). Internally, arguments of any type are converted to a float representation. This may lead to undefined double arguments in float representation.
- Results of SimEnv post-processing operators are always of the type float.
- SimEnv post-processing follows the standard approach for description of operators for basic as well as advanced built-in or user-defined operators.

Advanced built-in or user-defined operators

- Have a unique name and a number of operands
- The sequence of operands is enclosed in parentheses directly after the operator name
- Operands are separated from each other by a comma.
- Recursions of the same operator (also for user-defined operators) are possible.
- Example: log10 (min_n(3, min_n(log10 (atmo(*,*,1,c=20)), 400), 10*bios_g)) Elemental operators use the common form of notation:

Example: atmo_g + 345



8.1.5 Operator Classification, Flexible Coordinate Checking

Tab. 8.1 lists for all built-in operators a classification of argument restrictions and result description that are used in the following for the explanation of built-in operators.

Argument restriction(s) / result description		Argument restriction(s)	Result description (check chapter 8.1.2 for syntax)
(1)		dimensionality, extents and coordinates of the only non-character / non-constant argument <u>arg</u> can be arbitrary	same dimensionality, extents and coordinates as the only non-character / non-constant argument: dim(res) = dim(arg) ext(res,j) = ext(arg,j) for all j coord(res,j) = coord(arg,j) for all j
(2)	(2.1)	all non-character / non-constant arguments <u>arg</u> with same dimensionality, extents and coordi- nates (*)	same dimensionality, extents and coordinates as all the non-character / non-constant arguments: dim(res) = dim(arg) ext(res,j) = ext(arg,j) for all j coord(res,j) = coord(arg,j) for all j
(2) = (2.1) or (2.2)	(2.2)	some non-character / non-constant arguments arg with same non-zero dimensionality, extents and coordinates (*), all the other non-character arguments with dimensionality zero	same dimensionality, extents and coordinates as all the non-character / non-constant arguments with non- zero dimensionality: dim(res) = dim(arg) ext(res,j) = ext(arg,j) for all j coord(res,j) = coord(arg,j) for all j the zero-dimensional argument is applied to each element of the non- zero dimensional argument
(;	3)	dimensionality, extents and coordinates of the only non-character / non-constant argument can be arbitrary	dimensionality 0: dim(res) = 0
(4)	(4.1)	all non-character / non-constant arguments with same dimensionality, extents and coordinates (*)	dimensionality 0: dim(res) = 0
(4.1) or (4.2) (4.2)		some non-character / non-constant arguments with same non-zero dimensionality, extents and coordinates (*), all the other non-character / non-constant arguments with dimensionality zero	dimensionality 0: dim(res) = 0 the zero-dimensional argument is applied to each element of the non- zero dimensional argument
(5)		dimensionality, extents and coordinates of the first non-character / non-constant argument arg can be arbitrary, all the other following argu- ments have to have dimensionalities, extents and coordinates (*) of this argument or have to have dimensionality 0	same dimensionality, extents and coordinates as the first non-character / non-constant argument: dim(res) = dim(arg) ext(res,j) = ext(arg,j) for all j coord(res,j) = coord(arg,j) for all j
(6)		without arguments	dimensionality 0: dim(res) = 0

Tab. 8.1

Classified argument restriction(s) / result description

(*): for the different levels of checking a coordinate description see above

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The requirement for a lot of operators to have same coordinates for same dimensions may restrict application of post-processing especially for hypothesis checking heavily. To enable a broader flexibility with respect to this situation a general solution is provided by SimEnv post-processing: With the sub-keyword 'coord_check' in the general configuration file <model>.cfg three different modi can be assigned globally to the SimEnv complete post-processing session:

coord_check = strong

To ensure for two arguments with same dimensionalities and extents to have same coordinates it is necessary that

- Assigned coordinate values for corresponding dimensions are unique
- Assigned coordinate names for corresponding dimensions are unique
- coord_check = strong is the default
- coord_check = weak

To ensure for two arguments with same dimensionalities and extents to have same coordinates it is necessary that

- Assigned coordinate values for corresponding dimensions are unique
- Assigned coordinate names may differ.

Coordinate description of the appropriate operator result is delivered from its first non-character / non-constant operand.

• coord_check = without

To ensure for two arguments with same dimensionalities and extents to have same coordinates

• Neither coordinate names nor coordinate values for corresponding dimensions are checked Coordinate description of the appropriate operator result is delivered from its first non-character / non-constant operand.

Check Example 8.3 for examples.

Result expression	CC	coordinat	k =
	strong	weak	without
<pre>bios(*,*,*) + atmo(c=84:-56,*,c=1,*) (same coordinate names, same coordinate values)</pre>	yes	yes	yes
atmo_g(*) + hgr('bin_no', 20, 0., 0., atmo) (differing coordinate names, same coordinate values)	no	yes	yes
<pre>atmo_g(c=6:16) + atmo_g(c=8:18) (same coordinate names, differing coordinate values)</pre>	no	no	yes
<pre>atmo_g(c=20) + atmo(c=0,c=2,c=1,c=1) (two operands with dimensionality 0)</pre>	yes	yes	yes

coordinate information is determined by the first summand for coord check = [weak | without].

Example 8.3 Checking rules for coordinates



8.2 Built-In Generic Standard Aggregation / Moment Operators

The generic operators in Tab. 8.2 can be applied during model output post-processing to derive aggregations and moments from operands in different ways by appending suffixes (_n, _l, _e, without suffix) to the generic operator name or by incorporating them into the filter argument for experiment-specific operators of bahavioural analysis:

<u>Generic</u> aggregation and moment operator	Meaning
max	maximum of values
min	minimum of values
sum	sum of values
avg	arithmetic mean of values
var	variance of values
avgg	geometric mean of values
avgh	harmonic mean of values
avgw	weighted mean of values
hgr	histogram of values
count	number of values
maxprop	maximal, suffix related property of values
minprop	minimal, suffix related property of values

 Tab.
 8.2
 Built-in generic standard aggregation / moment operators

For more information check chapters 8.3.3 and 8.4.1.

8.3 Built-In Elemental, Basic, and Advanced Operators

8.3.1 Elemental Operators

Name	Meaning	Argument restriction(s) / result description (see Tab. 8.1)	Argument value restriction	Precedence
(left parenthesis	-		first
)	right parenthesis	-		first
arg1 ** arg2	exponentiation	(2)	arg1 > 0	second
arg1 * arg2	multiplication	(2)		third
arg1 / arg2	division	(2)	arg2 ≠ 0	third
arg1 + arg2	addition (dyadic +)	(2)		fourth
arg1 – arg2	subtraction (dyadic -)	(2)		fourth
+ arg	identity (monadic +)	(1)		fourth
– arg	negation (monadic -)	(1)		fourth

Tab. 8.3Built-in elemental operators

- n-dimensional matrix algebra of built-in elemental operators is performed element by element
 Example: atmo(*, *, 1, *) * bios(*, *, *) = "atmo(i,j,1,k) * bios(i,j,k)" for all addressed (i,j,k)
- If an argument value restriction is not fulfilled for an operand element the corresponding element of the operator result is undefined.
- For examples check chapter 8.3.5.

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8.3.2 Basic and Trigonometric Operators

Name	Meaning	Argument restriction(s) / result description (see Tab. 8.1)	Argument value restriction	Example	
	Basic operators				
abs(arg)	absolute value	(1)		abs(-3) = 3.	
dim(arg1,arg2)	positive difference	(2)		$\dim(10,5) = 5.$ $\dim(5,10) = 0.$	
exp(arg)	exponential function	(1)		$\exp(1.) = 2.7183$	
int(arg)	truncation value	(1)		int(7.6) = 7. int(-7.6) = -7	
log(arg)	natural logarithm	(1)	arg > 0	log(2.7183) = 1.	
log10(arg)	decade logarithm	(1)	arg > 0	log10(10) = 1.	
mod(arg1,arg2)	remainder	(2)	arg2 ≠ 0	mod(10, 4) = 2.	
nint(arg)	round value	(1)		nint(7.6) = 8.	
sign(arg)	sign of value	(1)		sign(-3) = -1. sign(0) = 0.	
sqrt(arg)	square root	(1)	arg ≥ 0	sqrt(4) = 2.	
	Trigo	onometric operators			
sin(arg)	sine	(1)		sin(0) = 0.	
cos(arg)	cosine	(1)		$\cos(0) = 1.$	
tan(arg)	tangent	(1)	arg ≠ π/2±n*π		
cot(arg)	cotangent	(1)	arg ≠ ±n*π	$\cot(1.5708) = 0.$	
asin(arg)	arc sine	(1)	abs(arg) ≤ 1	asin(0) = 0.	
acos(arg)	arc cosine	(1)	abs(arg) ≤ 1	acos(1) = 0.	
atan(arg)	arc tangent	(1)		atan(0) = 0.	
acot(arg)	arc cotangent	(1)		acot(0) = 1.5708	
sinh(arg)	hyperbolic sine	(1)		$\sinh(0) = 0.$	
cosh(arg)	hyperbolic cosine	(1)		$\cosh(0) = 1.$	
tanh(arg)	hyperbolic tangent	(1)		tanh(0) = 0.	
coth(arg)	hyperbolic cotangent	(1)	arg ≠ 0	coth(3.1416) = 1.	

Tab. 8.4Built-in basic and trigonometric operators

The following explanations hold for the operators in Tab. 8.4:

- All operators are applied to each element of the argument(s). These operators deal with an unfulfilled argument value restriction for an operand element in a way that the corresponding element of the operator result will be undefined.
- For examples check chapter 8.3.5.



8.3.3 Standard Aggregation / Moment Operators

The generic standard aggregation / moment operators in Tab. 8.2 can be applied during model output postprocessing to derive aggregations and moments from operands in different ways by appending suffixes to the generic operator name:

• Appending no suffix:

Aggregate the only non-character argument(s)

Result is a scalar (an operator result of dimensionality zero) for all but operators hgr, minprop and maxprop.

For operator hgr dimensionality of the result is 1, the extent is the specified number of bins for the histogram and the coordinate assigned has the name bin. Coordinate values are equidistant with 1 as the first value and an increment of 1.

For operators minprop and maxprop dimensionality of the result is 1. For argument dimensionality greater / equal 1 extent of the result is equal to the argument dimensionality. Assigned coordinate name is index. Coordinate values are equidistant with 1 as the first value and an increment of 1. For argument dimensionality 0 result dimensionality is 0.

- Appending suffix _n (for n arguments)
 Aggregate an arbitrary number of arguments with argument restriction(s) / result description according to
 (2) in Tab. 8.1 at page 64 element by element
 Currently, only operators min_n and max_n are implemented.
 Result has same dimensionality, extents and coordinates as the arguments
- Appending **suffix** _I (for loop)

Aggregate the only non-character argument(s) separately for selected dimensions. Dimensions to select are described by an additional loop character argument (corresponds with the group by-clause of the standard query language SQL of relational database management systems).

Result has a lower dimensionality as the only non-character argument according to the loop character argument.

For operator hgr_l, dimensionality is increased additionally by one, the additional extent is the specified number of bins for the histogram and the additional coordinate assigned to has the name bin. Coordinate values are equidistant with 1 as the first value and an increment of 1.

For operators minprop_I and maxprop_I dimensionality is modified in the same manner like for operators minprop and maxprop, respectively.

• For **examples** check chapter 8.3.5.

Aggregation and moment operator		Argument restriction(s) / result description (see Tab. 8.1)			
max(arg)					
min(arg)					
sum(arg)					
avg(arg)		(3)			
var(arg)					
avgg(arg)					
avgh(arg)					
avgw(arg1,arg2)	arg2	(4.1) = weight			
hgr(arg1,arg2,arg3,	dim(res)	= dim(arg4)+1			
arg4,arg5)	ext(res,dim(res))				
	for arg1 = 'bin_no				
	coord(res,dim(res))= name = bin_no			
		values = equidist_end 1(1) number of bins			
	for arg1 = 'bin_mi				
	coord(res,dim(res	coord(res,dim(res))= name = bin_mid			
		values = equidist_end 1 st bin mid (bin width) number of bins			
	arg1 see above				
	character argument				
	argz	arg2 = number of bins: 4 ≤ arg2 ≤ number_of_values or = 0: automatic determination:			
		number of bins = max(4,number of values/10)			
		integer constant argument			
	arg3	left bin bound for bin number 1			
		real constant argument			
	arg4	right bin bound for bin number arg2			
		real constant argument			
	arg3 = arg4 = 0.:	determine bounds by min(arg5) and max(arg5)			
		min(arg5) = max(arg5): all result values are undefined			
count(arg1,arg2)		(3)			
	arg1	= character argument			
		= [all def undef]			
maxprop(arg)	dim(res)	= 1 for dim(arg) > 1			
	ext(res,1)	= dim(arg) = 0 else			
minprop(arg)	dim(res)				
minprop(arg)	time according to	of that element of arg where the extreme is reached the first the processing sequence of the argument field arg by the odel (see Glossary).			

 Tab.
 8.5
 Built-in standard aggregation / moment operators without suffix

Aggregation and moment operator	Argument restriction(s) / result description (see Tab. 8.1)
max_n(arg1,,argn)	(Λ)
min_n(arg1,,argn)	(4)
maxprop_n(arg1,,argn)	(4)
minprop_n(arg1,,argn)	returns per result element the argument position (1 n) where the extreme is reached the first time. Processing sequence starts with arg1.

 Tab.
 8.6
 Built-in standard aggregation / moment operators with suffix _n



Aggregation and moment operator		nt restriction(s) / It description
min_l(arg1,arg2) max_l(arg1,arg2) sum_l(arg1,arg2) avg_l(arg1,arg2) var_l(arg1,arg2) avgg_l(arg1,arg2) avgg_l(arg1,arg2) avgh_l(arg1,arg2) avgw_l(arg1,arg2)	dim(non-character argument(s)) > 1 ext(non-character argument(s)) = arbitrary dim(res), ext(res,i) according to arg1 and the non-character argu- ment(s) arg1 = loop character argument	dim(arg2) = dim(arg3) ext(arg2,i) = ext(arg3,i) arg3 = weight
hgr_l(arg1,arg2,arg3, arg4,arg5,arg6) count_l(arg1,arg2,arg3)		dim(res) = 1 + dim(res) of all other operators ext(res,dim(res)) = number of bins for arg2 = 'bin_no' (bin number): coord(res,dim(res)) = name = bin_no values = equidist_end 1(1) number of bins for arg2 = 'bin_mid' (bin mid): coord(res,dim(res)) = name = bin_mid values = equidist_end 1 st bin mid (bin width) number of bins arg2 see above character argument arg3 number of bins 4 ≤ arg3 ≤ number_of_values or 0: automatic determination = max(4,number_of_values/10) integer constant argument arg4 left bin bound for bin number 1 real constant argument arg5 right bin bound for bin number 1 real constant argument arg4 = arg5 = 0.: determine bounds by min(arg6) and max(arg6) min(arg6) = max(arg6): all result values are undefined arg2 = [all def undef] character argument
minprop_l(arg1,arg2)	as above, but: dim(res) is increased by 1 w.r.t.	returns the indices of those elements of arg2 where the extreme is reached the first
maxprop_l(arg1,arg2)	above. ext(res,dim(res)) = dim(arg1) coord(res,dim(res)): name = index values = equidist_end 1(1)"n"	time according to arg1 and to a Fortran-like processing sequence / storage model (see Glossary) of the argument field arg2.

Tab. 8.7 Built-in standard aggregation / moment operators with suffix _I

The loop character argument arg1 is characterised as follows:

- The length of the string is equal to the dimensionality of the non-character argument
- The string consists of 0 and 1
- 0 at position n means: aggregate over the corresponding dimension n of the argument

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- 1 at position n means: do not aggregate over the corresponding dimension n of the argument
- Loop character arguments completely formed of 0 or 1 are forbidden

For the operator hgr_l bins are determined on the base of the minimum and maximum value of the total argument arg2.

8.3.4 Advanced Operators

Name	Meaning	Argument restriction(s) / result description (see Tab. 8.1)	Argument value restriction	Example
classify(arg1, arg2,arg3,arg4)	classify arg4 into arg1 classes; potentially re- strict classification to interval (arg2 , arg3).	(1) dim(arg4) > 0 arg1 = number of clas 2 ≤ arg1 ≤ num of arg = 0: automatic number of clas max(2,number integer constar arg2 = minimum boun in class # 1 real constant a arg3 = maximum boun in class # arg1 real constant a arg2 = 0. and arg3 = 0 automatic boun	aber_of_values g2 determination: ses = _of_values/10) at argument d for values rgument and for values rgument .:	classify((10,0.,0.,atmo)
clip(arg1,arg2)	clip arg2 according to arg1	dim(arg2) > 0 dim(res), ext(res,i) de and arg2 arg1 = clip range character argu	pend on arg1	clip('0,*,1,10', atmo)
cumul(arg1,arg2)	cumulates arg2 accord- ing to arg1	(1) dim(arg2) > 0 arg1 = cumulation ind per dimension character argu	icator	cumul('0001', atmo)
flip(arg1,arg2)	flips arg2 according to arg1	(1), but coordinates dim(arg2) > 0 arg1 = flip indicator pe character argu	is also flipped er dimension	flip('0001', atmo)
get_experiment(arg1,arg2, arg3,arg4)	include an other experi- ment	 (1) arg1 = experiment dirucharacter argui arg2 = model experimicharacter argui arg3 = file to transformicoordinates character argui arg4 = result from this 	ment ented with ment n result ment	<pre>get_experiment('mod_res', 'mod', 'mod.trf', avg(atmo)-400.)</pre>
get_table_fct(arg1,arg2)	table function with linear interpolation of table arg1 applied to arg2	(1) arg1 = file name character argu		<pre>get_table_fct (`table.usr', atmo)</pre>

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Name	Meaning	Argument restriction(s) / result description (see Tab. 8.1)	Argument value restriction	Example
if(arg1,arg2, arg3,arg4)	conditional if-construct	(5) arg1 = comparison op character argur arg2 = comparator arg3, arg4 = new assig	ment	if('<',atmo,400, atmo)
mask(arg1, arg2,arg3)	masks values of arg2 (set them undefined) by comparing arg2 and arg3 using operator arg1	(5) arg1 = comparison op character argur		<pre>mask(`<',atmo, 400)</pre>
matmul(arg1, arg2)	matrix multiplication	dim(arg1) = dim(arg2) = 2 ext(res,i) accord. to ma cation rules		<pre>matmul(atmo(*,*,1,1), transpose('21', atmo(*,*,1,1)))</pre>
move_avg(arg1, arg2,arg3,arg4)	moving average of arg5	(1) dim(arg4) > 0 arg1 = moving average per dimension arg2 = average type = lin: linear exp: expone moving averag arg3 = running length arg3 > 1 arg3 = 0: automatic dete = max(3, ext(ar integer constar	char. arg. ntial e char. arg. for average rmination: rg4,i)/20.	<pre>move_avg(`001', `lin',0,atmo)</pre>
nr_of_runs	number of single runs in the experiment	(6)		<pre>nr_of_runs()</pre>
rank(arg1,arg2)	assigns rank numbers to arg2 according to ranking type argument arg1	(1) dim(arg2) > 0 arg1 = ranking type [tie_plain tie_min tie_avg]		<pre>rank(`tie_avg', atmo)</pre>
run(arg1,arg2)	values of arg 2 for the selected single run number explicitly or im- plicitly coded in arg1	(1) arg1 = run number sel = 0 for default (all experim = <run_number (for Monte C 0 ≤ arg1 ≤ nu = filter argumer (for behavio same as filte operator bel check chapt</run_number 	ection t run ent types) r> Carlo analysis, imber_of_runs) nt oural analysis, er argument of hav, ter 8.4.2)	
transpose(arg1, arg2)	transpose arg2 according to sequence in arg1	dim(arg2) > 1 dim(res) = dim(arg2) ext(res,i) = ext(arg2,j) arg1 = transpose sequ character argur	ience	<pre>transpose (`3142',atmo)</pre>
undef()	undefined value	(6)	-	undef()

Tab. 8.8

Built-in advanced operators

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The following explanations hold for the operators in Tab. 8.4:

- All operators but experiment and matmul are applied to each element of the argument(s). These operators deal with an unfulfilled argument value restriction for an operand element in a way that the corresponding element of the operator result will be undefined.
- The operator classify transforms the values of an operand arg4 that has dimensionality > 0 into the class numbers 1,..., arg1 of arg1 classes. Classes are assumed to be equidistant.
 If both arguments arg2 and arg3 are zero then min(arg4) forms the lower boundary of class number 1 and max(arg4) forms the upper boundary of class number arg1. For min(arg4) = max(arg4) all result values of the operator classify are undefined.
 For arg2 ≠0 or arg3 ≠ 0 arg2 and arg3 are used as boundaries for the classification and all of those re-

For arg2 \neq 0 or arg3 \neq 0 arg2 and arg3 are used as boundaries for the classification and all of those result values are undefined where values of argument arg4 are outside the specified boundary range.

• The **operator clip** clips an operand that has dimensionality > 0. The portion to clip from the operand arg2 is described by character argument arg1. Argument arg1 uses syntax for model output variable addressing (see chapter 8.1.3 at page 61). Note, that for all dimensions of argument arg2 lower bound index is 1. This applies also to model variables where the lower bound index is unequal 1 in the model output description file. In general, extents differ between the result of the operator clip and the argument arg2. Clip reduces the dimensionality of the result with respect to the argument arg2 to clip if the portion to be clipped is limited to one value for at least one dimension.

A character argument arg1 = '*,..., *' results for operator clip in the identity of argument arg2.

- The **operator cumul** cumulates an operand that has dimensionality > 0. Cumulation is performed for all values of the argument arg2 from the first addressed index position up to the current index position. With the character argument arg1 these dimensions are identified that are to be cumulated. Character 1 at position i means cumulation across dimension i while a 0 stands for no accumulation. cumul('0...0',arg) results in the identity to arg.
- The **operator flip** enables flipping of variable fields. For a one-dimensional field (a vector) flip changes the value of the first index position with the value of the last position, the value of the second position with that of the last but one position, etc. With the character argument arg1 these dimensions are identified that are due to flip. Character 1 at position i means flipping also for dimension i while a 0 stands for no flipping at this dimension. Flipping includes adaptation of coordinates and the assigned grid. cumul('0...0',arg) results in the identity to arg.
- The operator get_experiment is to access to external SimEnv model output from the same or an other model performed with the same or another experiment type and stored in the same or in an other model output format. Model variables can differ from that used for the current model. Use for the experiment directory arg1 always that working directory the external experiment was started from. The external experiment is always post-processed completely over all single runs. Environment variables from operating system level in the specification of the directory are not allowed. Argument arg3 is the coordinate transformation file. It can be used to transform coordinates from the external result for usage in the current result of the current experiment. If no coordinate transformation file is to be used argument arg3 is empty (' '). If after potential application of an coordinate transformation file the imported result has same coordinate names as defined in the original experiment coordinate descriptions are checked against each other, otherwise coordinate descriptions are imported from the external into the original experiment. For syntax of coordinate transformation files check chapter 11.2.

Attention: Make sure no SimEnv service is running from the directory arg1 of the external experiment.

• With the **operator get_table_fct** a table function arg1 is applied to each element of the operand arg2. If necessary, table values are interpolated linearly. Outside the definition range of the table function the first and/or the last table value is used. File arg1 to hold the table function must be an ASCII file with two columns: The first column of each line is the argument value x, the second column the function value f(x). Arguments have to be ordered in a strictly increasing manner. Syntax rules for comments and separators in the table function file are the same as for user defined files (check chapter 11.3). Environment variables from operating system level in the specification of the file name arg1 are not allowed. Check the table function world.dat_tab in the examples directory of \$SE_HOME for more information.



• The **operator if** supplies a general conditional if-construct. It operates for each element of the operand arg2 in the following way:

if (condition(arg1,arg	g2)) then	
res=arg3		
else		
res=arg4		
endif		
with condition(arg1,arg2):	arg2 < 0	(arg1 = '<')
	arg2 ≤ 0	(arg1 = '<=')
	arg2 > 0	(arg1 = '>')
	arg2 ≥ 0	(arg1 = '>=')
	arg2 = 0	(arg1 = '=')
	arg2 != 0	(arg1 = '!=')
	arg2 def	(arg1 = 'def')
	arg2 undef	(arg1 = 'undef')

• The **operator mask** supplies a method to mask values. It operates for each element of the operand arg2 in the following way:

if (condition(arg1,arg2, res=undef()	arg3)) then	
else		
res=arg2		
endif		
with condition(arg1,arg2,arg3):	arg2 < arg3	(arg1 = '<')
	arg2 ≤ arg3	(arg1 = '<=')
	arg2 > arg3	(arg1 = '>')
	arg2 ≥ arg3	(arg1 = '>=')
	arg2 = arg3	(arg1 = '=')
	arg2 != arg3	(arg1 = '!=')

- The **operator matmul** performs a simple matrix multiplication for 2-dimensional arguments arg1 and arg2.
- The **operator move_avg** performs a moving average operation successively for selected dimensions of the argument arg4.

For a vector $(a_1, a_2, ..., a_{len})$ the moving average of running length rl is a vector $(ma_1, ma_2, ..., ma_{len})$ with elements

$$ma_{i} = \frac{1}{\sum_{i=max(1,i-rl+1)}^{i} \cdots \sum_{j=max(1,i-rl+1)}^{l} w_{ij} \cdot a_{j}}$$

where w_{ij} are weights. Value ma_i is averaged from the rl values a_i , a_{i-1} , ..., a_{i-rl+1} . Accordingly, the first rl-1 values ma_1 , ma_2 , ..., ma_{rl-1} are averaged from less than rl values.

For the linear moving average the weight is

$$\begin{split} w_{ij} &= 1 \qquad \text{and} \quad \sum_{j=max(1,i-rl+1)}^{l} w_{ij} = min(rl,i)\,, \\ w_{ij} &= e^{-\frac{i-j}{rl}} \,\,. \end{split}$$

for the exponential moving average the weight is

While the moving average is normally applied to time-dependent one-dimensional data vectors the operator move_avg allows processing of multi-dimensional data fields in a general and succesive manner. For example, if arg4 is the three-dimensional variable bios(1:lat,1:lon,1:time) then the linear moving average could be applied to the dimension time successively for all combinations of lat and lon. This means that (lat1 = 1,...,lat) * (lon1 = 1,...,lon) = lat*lon moving averages will be performed for the vector

(bios(lat1,lon1,1) , bios(lat1,lon1,2) , ... , bios(lat1,lon1,time)).

Afterwards this moving averaged temporary result tmp could be moving averaged for all values of lat: (lon1 = 1,...,lon) * (time1 = 1,...,time) = lon*time moving averages will be performed for the vector

(tmp(1,lon1,time1), tmp(2,lon1,time1), ..., tmp(lat,lon1,time1)).

The operator that allows for this double averaging would have the arguments

move_arg('201', 'lin', 0, bios).

The character argument arg1 supplies those dimensions that are to be involved in the moving average operation. If the n-th digit of arg1 is a digit > 0 then the moving average for dimension n of argument arg4 is performed at position number "digit" (i.e. after performing moving averages for those dimensions that correspond to digits smaller than the current one). If the n-th digit of arg1 is 0 then the moving average for the dimension n of arg4 will not be performed.

Keep in mind that the sequence of moving averages for single coordinates influences the result of the operator.

- The **operator nr_of_runs** returns the number of performed single runs of the current post-processed experiment without the run number 0 of the nominal constellation. It does not have an argument.
- The operator rank transforms all values of an operand arg2 that has dimensionality > 0 into their ranks. Small values get low ranks, large values get high ranks. The smallest rank is 1. Character argument arg1 determines how to rank ties, i.e., values of arg2 that are identical or have a maximum absolute difference of 10⁻⁶: Assume an argument arg2 with 6 values (4, 2, 4, 4, 4, 8).

ume an argument arg2 with 6 values	(4., 2., 4., 4., 4., 8.).
arg1 = 'tie_plain' returns ranks	(2, 1, 2, 2, 2, 3) same minimal rank 2; next rank is 3, does not take into account the number of identical values
arg1 = 'tie_min' returns ranks	(2,1,2,2,2,6) same minimal rank 2; next rank is 6, taking into account the number of identical values
arg1 = 'tie_avg' returns ranks	(3.5 , 1 , 3.5 , 3.5 , 3.5 , 6) same mean rank 3.5; next rank is 6, taking into account number of identical values

• The **operator run** selects a single run from the run ensemble. The operator run must not contain experiment-specific (multi-run) operators as operands, while these operators may refer to the operator run. Additionally, run must not contain itself as an argument.

Character argument arg1 can hold explicitly the run number string Monte Carlo analysis. Run number 0 corresponds with the default single run 0 and is permitted as arg1 for all experiment types. For behavioural analysis a filter of the operator behav (see chapter 8.4.2) is applied as arg1 to select a unique run number unequal zero. For this purpose, a single run can be selected by the select-operator (check Tab. 8.11) of the operator behav. For Monte Carlo analysis, single runs with a run number unequal zero are selected explicitly. Therefore, the file <model>.edf_adj holds the targets to be adjusted to the default values for the current experiment. Run number n corresponds with record number n of this file. For more information on <model>.edf_adj check chapter 6.1 at page 39. For examples see Example 8.5 and Example 8.6.

• The **operator transpose** enables to transpose an operand that has a dimensionality > 1. Sequence of extents of the transposed result is described by character argument 1: It consists of figures 1, ..., dim(arg2) where the figure sequence corresponds with the re-ordered sequence of the operator result extents.

A character argument arg1 = '123...' results for operator transpose in the identity of argument arg2.

- The **operator undef** supplies a 0-dimensional result as undefined. This operator can be used as an argument for the if-operator.
- For **examples** check chapter 8.3.5.



8.3.5 Examples

```
Having a model variable definition as in Example 5.1 at page 26 and
assuming address default=coordinate in <model>.cfg then in model output post-processing
atmo g+2*atmo g
                                     value of result 3*atmo g
                                     Dimensionality = 1
                                     Coordinates = time
                                     Extents = 20
                                     square root of atmo g
sqrt(atmo g)
                                     Dimensionality = 1
                                     Coordinates = time
                                     Extents = 20
                                     last two decades for level 1 at equator
clip('i=23,*,1,19:20',atmo)
                                     equivalent with atmo(i=23,*,1,19:20)
                                     Dimensionality = 2
                                     Coordinates = lon, time
                                     Extents = 90.2
atmo - get experiment('./other dir', 'other model', '', atmo)
                                     Difference for atmo between the current experiment and
                                     another model other model, located in directory ./other dir
                                     without application of an coordinate transformation file
                                     Dimensionality = 4
                                     Coordinates = lat , lon , level , time
                                     Extents = according to definition of atmo in other_model
get table fct('world.dat tab',atmo)
                                     Operator table fct with table world.dat tab applied to
                                     each element of atmo
                                     Dimensionality = 4
                                     Coordinates = lat, lon, level, time
                                     Extents = 45, 90, 4, 20
                                     maximum from atmo and 10 for each element of atmo
if('<',atmo-10,10,atmo)
                                     equivalent with max n(atmo, 10)
                                     Dimensionality = 4
                                     Coordinates = lat , lon , level , time
                                     Extents = 45, 90, 4, 20
avg(atmo(*,*,*,19:20))
                                     global all-level mean over the last two decades
                                     Dimensionality = 0
                                     Coordinates = (without)
                                     Extents = (without)
                                     indices of this element of atmo where the maximum of atmo
maxprop(atmo)
                                     is reached the first time
                                     Dimensionality = 1
                                     Coordinates = index
                                     Extents=4
min n(atmo(84:-56,*,1,19:20),10.)
                                     minimum per grid cell for level 1 without polar regions
                                     for the last two decades from atmo and 10
                                     Dimensionality = 3
                                     Coordinates = lat , lon , time
                                     Extents = 36, 90, 2
min l('10',atmo(20:-20,*,1,20))
                                     zonal tropical minima of atmo for the last decade and
                                     level 1
                                     Dimensionality = 1
                                     Coordinates = lat
                                     Extents = 11
```

```
minprop l('10',atmo(20:-20,*,1,20))
                                    zonal tropical indices of those elements of
                                    atmo for the last decade and level 1 where the minimum is
                                    reached the first time
                                    Dimensionality = 2
                                    Coordinates = lat, index
                                    Extents = 11, 2
hgr l('10', 'bin no',8,0.,0.,atmo(20:-20,*,1,20))
                                    zonal tropical histograms with 8 bins of atmo for the
                                    last decade and level 1. Bin bound extremes are deviated
                                    from the values of atmo
                                    Dimensionality = 2
                                    Coordinates = lat , bin_no
                                    Extents = 11, 8
avg l('100',min l('1011',atmo(20:-20,*,*,*)))
                                    temporally averaged all-level zonal tropical minima
                                    Dimensionality = 1
                                    Coordinates = lat
                                    Extents = 11
                                                                  Example file: world.post adv
```



Post-processing with advanced operators

8.4 Built-In Experiment-Specific Operators

- Experiment-specific operators are to navigate and process in the experiment space.
- Experiment specific operators must not be applied recursively.
- Addressing a variable within an experiment specific operator normally results in application of the operator on the whole run ensemble or parts of it and in aggregating across the run ensemble according to the operator.
- Addressing a variable outside an experiment specific operator results in application of the basic, advanced and/or user-defined operator on the variable for the default run number 0 of the experiment.
- If the dimensionality of an operator result is higher than that of one of its operands the additional dimensions of the result are appended to the dimensions of the operand. Examples for such operators are ens (for Monte Carlo analysis post-processing) and behav (for certain constellations of behavioural analysis post-processing).

8.4.1 Standard Aggregation / Moment Operators

Tab. 8.9 summarises multi-run standard aggregation / moment operators for behavioural analysis, Monte Carlo analysis and optimization. They work on the whole run ensemble (for Monte Carlo analysis and optimization) or parts of it (for certain constellations of behavioural analysis post-processing). They are used with suffix _e for Monte Carlo analysis and optimization and without any suffix for behavioural analysis. For a definition of these operators check Tab. 8.2 at page 66.

Aggregation and moment operator	Argument restriction(s) / result description (see Tab. 8.1)		
min(arg)			
max(arg)			
sum(arg)			
avg(arg)	(1)		
var(arg)			
avgg(arg)			
avgh(arg)			
avgw(arg1,arg2)	(2.1)		
	arg2 = weight		
hgr(arg1,arg2,arg3,	dim(res) =dim(arg2)+1		
arg4,arg5)	ext(res,dim(res)) = number of bins		
	for arg1 = 'bin_no' (bin number):		
(heuristic probability	coord(res,dim(res))= name = bin_no		
density function)	values = equidist_end 1(1) number of bins		
	for arg1 = 'bin_mid' (bin mid):		
	coord(res,dim(res))= name = bin_mid values = equidist end 1 st bin mid (bin width) number of bins		
	• = • •		
	arg2 = number of bins		
	$4 \le \arg 2 \le \text{number of June of } $		
	0: automatic determination = max(4,number of runs/10)		
	integer constant argument		
	arg3 left bin bound for bin number 1		
	real constant argument		
	arg4 right bin bound for bin number arg2		
	real constant argument		
	arg3 = arg4 = 0.: determine bounds by min(ens(arg5)) and max(ens(arg5))		
	min(ens(arg5)) = max(ens(arg5)): all result values are		
	undefined		
count(arg1,arg2)	(1)		
	arg1 = [all def undef]		
	character argument		
minprop(arg)	(1)		
maxprop(arg)	returns the run number where the extreme is reached the first time.		
	Processing sequence starts with run number 1.		

 Tab.
 8.9
 Multi-run standard aggregation / moment operators

8.4.2 Behavioural Analysis

There is only one experiment specific operator for behavioural analysis. With this operator behav

- A single run can be selected from the run ensemble
- The complete run ensemble can be addressed
- Sub-spaces from the experiment space can be addressed and
- Sub-spaces can be projected by aggregation and moment operators

dependent on the way the experiment target space was to be scanned according to the sub-keyword 'comb' in the experiment description file.

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To show the power of the operator behav the simple experiment layouts as described in Fig. 4.3 at page 14 are used as examples.

- With behav it is possible to address for any operand a single run out of the run ensemble by fixing values of experiment targets p1 and p2 (for Fig. 4.3 (a)), a value of the parallel targets p1 or p2 (for Fig. 4.3 (b)), and values of targets p3 and p1 or p2 (for Fig. 4.3 (c)). Dimensionality and extents of the operator result is the same as that of the operand.
- Without any selection in the target experiment space (p1,p2) and/or (p1,p2,p3) the dimensionality of the operator result is formed from the dimensionality of the operand enlarged by the dimensionality of the experiment space. Two additional dimensions are appended to the operand for Fig. 4.3 (a), one additional dimension for Fig. 4.3 (b), and two additional dimensions for Fig. 4.3 (c). For the latter two cases it is important which of the axis p1 and p2 is used for further processing and/or output of the operator result. The extents of the appended dimensions are determined by the number of target adjustments.
- As a third option it is possible to select only a sub-space out of the experiment space to append to the operand. For Fig. 4.3 (a) this could be the sub-space formed from the first until the third adjustment value of p1 and all adjustment values of p2 between 3 and 7. Dimensionality of the operator result increases by 2 and extents of these additional dimensions are 3 and 2 with respect to the corresponding Example 6.1 (a) in chapter 6.2.3 at page 41.
- The operator behav also enables to aggregate operands in the experiment space. In correspondence with the example in the last bullet point for Fig. 4.3 (a) the operand could be aggregated (e.g., averaged) over the first until the third adjustment value of p1 autonomously for all runs with different values of p2 and afterwards this intermediate result (that now depends only on p2) could be summed up for all adjustment values of p2 between 3 and 7. Consequently the result has the same dimensionality as the operand of behav. Sequence of performing aggregations is important.

Name	Meaning	Argument restriction(s) / result description	Argument value restriction
behav(arg1,arg2)	navigation and aggregation in the experiment space for arg2 ac- cording to arg1	arg1= selection / aggregation filter character argument according to Tab. 8.14 dim(res) = dim(arg2) + appended dimensions according to arg1	

 Tab.
 8.10
 Experiment-specific operators for behavioural analysis

Placeholder	Explanation		
<filter></filter>	$\{ \langle operator_1 \rangle \{, \langle operator_2 \rangle \dots \{, \langle operator_n \rangle \} \dots \} \}$		
<operator></operator>	[<select_operator> <aggreg_operator> <show_operator>]</show_operator></aggreg_operator></select_operator>		
<select_operator></select_operator>	<pre>sel { _<target_value_type>} (<target_name> { <target_value_range> })</target_value_range></target_name></target_value_type></pre>		
<aggreg_operator></aggreg_operator>	<pre><aggreg_type> {_<target_value_type>} (<target_name> { <target_value_range> })</target_value_range></target_name></target_value_type></aggreg_type></pre>		
<show_operator></show_operator>	show(<target name="">)</target>		
<target_name></target_name>	name of the experiment target according to the experiment description file		
<target_value_type></target_value_type>	specification how to interpret <value_range></value_range>		
	i as adjustment indices (indices always count from 1)		
	v as adjustment values		
	t as resulting target values		
<target_value_range></target_value_range>	[(<value<sub>1> { : <value<sub>2> }) (*)]</value<sub></value<sub>		
	for $ = : = $		
	(*): use all values from <target_name></target_name>		

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Placeholder	Explanation		
<aggreg_type></aggreg_type>	 an aggregation / moment operator from Tab. 8.9 at page 78. The following restrictions apply: aggregations avgw and hgr can not be used aggregation count has a differing syntax: count_<target_value_type> ([all def undef],</target_value_type> <target_name> { <target_value_range> })</target_value_range></target_name> 		

Tab. 8.11Syntax of the filter argument 1 for operator behav

The following rules hold for the operator **behav**:

• Generally, by the filter argument arg1 those runs from the run ensemble are selected and/or aggregated (here interpreted as filtered) that are used for the formation of the result.

Consequently, if no filter is specified all runs are used:

behav(` ',atmo_g)

The select operator has to be specified only if values are to be restricted by a corresponding target value range.

For the aggregation and the select operator the target value type is redundant if the value range represents the full range of values by <target_name> or <target_name>(*):

- sel(p1) = sel(p1(*)) = sel_i(p1) = sel_v(p1) = sel_t(p1) and all are redundant.
- The show-operator can be used to force a certain experiment target to be used in the result of the operator behav if this target is used in parallel with other targets. By default, the first target of a parallel target sub-space as declared in the comb-line of the experiment description file is used in the behav-result.
- Aggregation operators reduce dimensionality of the covered experiment target space in the behav-result. The sequence of aggregation operators the first argument of the operator behav influences the result: Computation starts with the first aggregation operator and ends with the last: avg (p1), min (p2) normally differs from min (p2), avg (p1)
- An unused experiment target in the selection and aggregation filter contributes with an additional dimension to arg2 to the result of the operator behav. The extent of this additional dimension corresponds with the number of adjustments to this target in the experiment description file.
 A target that is restricted by any of the select operators also contributes with an additional dimension to the result of the operator behav if the number of selected values is greater than 1. The extent of the additional dimension corresponds with the number of selected values of this target by the select operator. Consequently, an empty character string arg1 forces to output the operand arg2 over the whole target

space of the experiment.

The name of the coordinate that is assigned to an additional dimension is the name of the corresponding target. Coordinate description and coordinate unit (see 5.1 at page 21) are associated with the corresponding information for the target from the experiment description file.
 Coordinate values are formed from resulting target values. For strictly ordered target adjustments in the experiment description file.

experiment description file and finally for strictly ordered resulting target values the coordinate values are ordered accordingly in an increasing or decreasing manner. Unordered target adjustments and finally unordered target values are ordered in an increasing manner for coordinate usage. The result of the operator behav is always arranged according to ascending coordinate values for all ad-

The result of the operator behav is always arranged according to ascending coordinate values for all additional dimensions.

- Independently from the sequence of the applied aggregation-, select- and show-operators the targets
 that contribute to additional dimensions of the result of the operator behav are appended to the dimensions of the operand arg2 of behav according to the sequence they are declared in the experiment description file (and **not** to the sequence they are used in the comb-line of the experiment description file).
 From parallel changing targets that target is used in this sequence that is addressed explicitly or implicitly by the show-operator.
- For experiment targets that are changed in the experiment in parallel, that increase dimensionality of the result and where a show-operator is missing the first target from this parallel sub-space in the comb-line is used in the result.
- For experiments that use an adjustment file (<value_list = file ...) instead of adjustment definitions (<value_list = comb ...) all experiment targets are assumed to be adjusted in parallel.

Having a model variable definition as in Example 5.1 at page 26 and assuming address_default = coordinate in <model>.cfg Assume the experiment layout in Example 6.1 (c) at page 42 and the corresponding experiment description file (c) from Example 6.1 at page 41 then in result-processing last time step of bios dependent on (p2,p1) and p3 behav(' ', bios(*,*,20)) Dimensionality = 4Coordinates = lat , lon , p2 , p3 Extents = 36, 90, 4, 3 behav ('show (p1)', bios (*, *, 20)) last time step of bios dependent on (p1,p2) and p3 Dimensionality = 4Coordinates = lat , lon , p1 , p3 Extents = 36, 90, 4, 3behav('sel t(p2(4)), sel i(p3(1))', atmo(*,*,1,*)) select the single run out of the run ensemble for level 1 p2 = 4 and p3 = 3.3Dimensionality = 3 Coordinates = lat , lon , time Extents = 45, 90, 20 behav('sel i(p2(1:3)), sel v(p3(1:2))', atmo(*,*,1,20)) last time step of atmo for level 1 depend. on (p2,p1) and p3 use only runs for $p_{2} = 1, 2, 3$ and for $p_{3} = 3.3, 4.5$ Dimensionality = 4Coordinates = lat , lon , p2 , p3 Extents = 45, 90, 3, 2 behav('avg_i(p2(1:3)),sel_i(p3(2:3))',atmo(*,*,1,*)) mean of atmo for level 1 and for runs with p2 = 1, 2, 3for each value of p3 = 4.5, 7.2 Dimensionality = 4Coordinates = lat, lon, time, p3Extents = 45, 90, 20, 2 behav('min(p2), max(p3)', avg(atmo(*, *, 1, 19:20))) determine single minima of avg(atmo) for level 1 and the last two decades for each value of p2 afterwards determine from that the maximum over all p3. Dimensionality = 0Coordinates = (without) Extents = (without) behav('max(p3),min(p2)',avg(atmo(*,*,1,19:20))) Result differs normally from min(p2),max(p3) (previous result expression) behav('count(def,p3),sel i(p2=1)',bios(*,*,20))/3 determine single numbers of defined values of bios for last decade for runs with p2=1. Result consists of values 0 (for water) and 1 (for land) Dimensionality = 2Coordinates = lat , lon Extents = 36, 90behav(' ',atmo(*,*,1,20)-run('sel i(p1(1)),sel i(p3(3))', atmo(*,*,1,20))) deviation of the last time step of atmo for level 1 from the run with p1=1, p2=1, p3=3.3dependent on (p2,p1) and p3 Dimensionality = 4Coordinates = lat , lon , p2 , p3 Extents = 45, 90, 4, 3Example file: world.post c

Example 8.5 Post-processing operator behav for behavioural analysis



8.4.3 Monte Carlo Analysis

Tab. 8.12 shows experiment specific operators for Monte Carlo analysis that can be used in post-processing besides the general multi-run aggregation operators listed in Tab. 8.9 at page 78 and supplemented with a suffix _e.

Name	Meaning	Argument restriction(s) / result description (see Tab. 8.1)	Argument value restriction
cnf(arg1,arg2)	positive distance of confidence measure from mean avg_e(arg2)	(1) arg1 error probability	arg1 = [0.001 0.01 0.05 0.1] real*4 constant argu- ment
cor(arg1,arg2)	correlation coefficient between arg1 and arg2	(2.1)	
cov(arg1,arg2)	covariance between arg1 and arg2	(2.1)	
ens(arg)	whole Monte Carlo run ensemble	dim(res) = dim(arg)+1 ext(res,dim(res)) = number_of_runs coord(res,dim(res)) = name = run values = equidist_end 1(1) number_of_runs	
krt(arg)	kurtosis (4 th moment)	(1)	
med(arg)	median	(1)	
qnt(arg1,arg2)	quantile of arg2	(1) arg1 quantile value	0. ≤ arg1 ≤ 100. real*4 constant argu- ment
reg(arg1,arg2)	linear regression coefficient to forecast arg2 from arg1: arg2 = reg(arg1,arg2)*arg1 + n	(2.1)	
rng(arg)	range = max_e(arg) - min_e(arg)	(1)	
skw(arg)	skewness (3 rd moment)	(1)	
stat_full(arg1, arg2,arg3, arg4,arg5)	basic statistical measures of arg5	dim(res) = dim(arg)+1 ext(res,dim(res)) = 10 coord(res,dim(res)) = name = stat_measure values = equidist_end 1(1)10	real*4 constant argu- ments 0. ≤ arg3 < arg4 ≤ 100. quantile values real*4 constant argu- ments
stat_red(arg1, arg2,arg3)	basic statistical measures of arg3	dim(res) = dim(arg)+1 ext(res,dim(res)) = 7 coord(res,dim(res)) = name = stat_measure values = equidist_end 1(1)7	arg1, arg2 = [0.001 0.01 0.05 0.1] arg1 < arg2 error probability for confidence distance measure real*4 constant argu- ments

Tab. 8.12

Experiment-specific operators for Monte Carlo analysis (without standard aggregation / moment operators)

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The following explanations hold for the operators in Tab. 8.12:

The operators stat_full and stat_red supply basic statistical measures for their last argument. Both operators are stand-alone operators: They must not be operands of any other operator. Contrary, their last argument can be composed from other non-multi-run operators. To store the statistical measures, dimensionality of both operators is that of their last argument, appended by an additional dimension with an extent of 10 and/or 7. Appended coordinate description is pre-defined by SimEnv (check Tab. 10.7).

These ten data fields (for operator stat_full) and/or seven data fields (operator stat_red) correspond with the following statistical measures:

- 1. Deterministic run (run number 0)
- 2. Run ensemble minimum
- 3. Run ensemble maximum
- 4. Run ensemble mean
- 5. Run ensemble variance
- 6. Run ensemble positive distance of confidence measure from run ensemble mean for value arg1
- 7. Run ensemble positive distance of confidence measure from run ensemble mean for value arg2
- Only for operator stat_full:
- 8. Run ensemble median
- 9. Run ensemble quantile of quantile value arg3
- 10. Run ensemble quantile of quantile value arg4

The operator stat_red was introduced because determination of the median and quantiles consume a lot of auxiliary storage space. For the definition of the statistical measures check the corresponding single operators in Tab. 8.9 and Tab. 8.12. Both operators were designed for application of an appropriate visualization technique in result evaluation in future.

assuming address_default=coordinate	Having a model variable definition as in Example 5.1 at page 26 and assuming address_default=coordinate in <model>.cfg Assume the Monte Carlo experiment from Example 6.2 (e) at page 45 then in model output post-processing</model>		
avg_e(p1*atmo(*,*,1,19:20))	global run ensemble mean of p1*atmo for level 1 and the last two decades Dimensionality = 3 Coordinates = lat , lon , time Extents = 45 , 90 , 2		
avg(atmo(*,*,1,19:20))	global mean of atmo for level 1 and the last two decades for run number 0 Dimensionality = 0 Coordinates = (without) Extents = (without)		
ens(atmo(*,*,1,20)	run ensemble values of atmo for level 1 and the last decade Dimensionality = 3 Coordinates = lat , lon , run Extents = 45 , 90 , 250		
<pre>minprop_e(atmo(*,*,1,19:20))</pre>	run ensemble run number for level 1 and the last two decades where the minimum of atmo is reached the first time Dimensionality = 3 Coordinates = lat , lon , time Extents = 45 , 90 , 2		
var_e(atmo(*,*,1,19:20))-atm			

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```
var e(atmo(*,*,1,19:20)-run('0',atmo(*,*,1,19:20)))
                                    global run ensemble variance of the anomaly of atmo for
                                    level 1 and the last two decades.
                                    Differs normally from the previous result expression
                                    Dimensionality 4
                                    Coordinates = lat , lon , time
                                    Extents = 45, 90, 4, 20
hgr_e(`bin_no',0,0.,0.,min_l(`10',atmo(20:-20,*,1,20)))
                                    histogram with 25 bins for the zonal tropical minima
                                    for level 1 and the last decade. Bin bound extremes are
                                    derived from the values of the last argument of the operator
                                    hgr e.
                                    Dimensionality = 2
                                    Coordinates = lat, bin no
                                    Extents = 11, 25
stat full(0.01,0.05,25,75, min l('10',atmo(20:-20,*,1,20)))
                                    basic statistical measures for the zonal tropical minima
                                    of atmo for level 1 and the last decade
                                    Dimensionality = 2
                                    Coordinates = lat , stat_measure
                                    Extents = 11, 10
                                                                    Example file: world.post_e
```

Example 8.6 Post-processing operators for Monte Carlo analysis

8.4.4 Local Sensitivity Analysis

Tab. 8.13 shows the experiment specific operators for local sensitivity analysis that can be used in post-processing. For a definition of these operators check Tab. 4.3 at page 17.

Name	Meaning	Argument restriction(s) / result description	Argument value restriction
sens_abs(arg1,	absolute sensitivity measure for	arg1= selection /	
arg2)	arg2 according to arg1	aggregation filter	
sens_rel(arg1,	relative sensitivity measure for	character argument	
arg2)	arg2 according to arg1	dim(res) = dim(arg2) +	
lin_abs(arg1,	absolute linearity measure for	appended dimensions	
arg2)	arg2 according to arg1	according to arg1	
lin_rel(arg1,	relative linearity measure for arg2		
arg2)	according to arg1		
sym_abs(arg1,	absolute symmetry measure for		
arg2)	arg2 according to arg1		
sym_rel(arg1,	relative symmetry measure for		
arg2)	arg2 according to arg1		

 Tab.
 8.13
 Experiment-specific operators for local sensitivity analysis

Placeholder		Explanation			
<filter></filter>	' { <sele< td=""><td>ct_operat</td><td>or₁> {, <select_operator<sub>2> {, <select_op< td=""><td>perator₃> } } } '</td></select_op<></select_operator<sub></td></sele<>	ct_operat	or ₁ > {, <select_operator<sub>2> {, <select_op< td=""><td>perator₃> } } } '</td></select_op<></select_operator<sub>	perator ₃ > } } } '	
<select_operator></select_operator>	[selt s	eli sels]	{ _ <value_type>} (<value_range>)</value_range></value_type>		
	wi	th selt =	 select target range 		
		seli =	 select increment range 		
		sels =	 select sign range (only for sens_abs ar 	nd sens_rel)	
<value_type></value_type>	specifica	specification how to interpret <value range=""></value>			
	i	i as position indices (always count from 1) for selt and seli			
	v	v as increment values for seli		for seli	
	n	n as target n ames for selt		for selt	
		as	signs (+ or -)	for sels	
<value_range></value_range>	[(<valu< td=""><td colspan="3">[(<value<sub>1> { : <value<sub>2> }) (*)]</value<sub></value<sub></td></valu<>	[(<value<sub>1> { : <value<sub>2> }) (*)]</value<sub></value<sub>			
•		for $\langle value_2 \rangle = \langle nil \rangle$: $\langle value_2 \rangle = \langle value_1 \rangle$			
	(*):	(*): use all values from <target name=""></target>			

Tab. 8.14	Syntax of the filter argument 1 for local sensitivity operators
-----------	---

The following rules hold for the filter argument in local sensitivity operators:

• Generally, by the filter argument arg1 those runs from the run ensemble are selected (here interpreted as filtered) that are used for the formation of the result.

Consequently, if no filter is specified all runs are used:

sens_abs(' ',atmo_g)

The filter operator has to be specified only if values are to be restricted by corresponding target values, increment values and/or sign ranges.

• For the above three select operators selt, seli and sels the value type is redundant if the value range represents the full range of values by [selt | seli | sels] (*):

selt(*) = selt_n(*) = selt_i(*)

and all are redundant.

- Each select operator can be applied only once within the filter argument.
- For <value_type> = i, i.e. if a value range is specified by position indices those targets are selected for selt and/or those increments are selected for seli that correspond with the specified position indices. Position indices are assigned from index 1 to the targets and or increments according to their specification sequence in the corresponding experiment description file <model>.edf.
- If more than one target, increment value and/or sign was selected by the filter argument arg1 it contributes with an additional dimension to the result of the local sensitivity operator:
 - For targets an additional dimension target_sequ
 - For increments an additional dimension incr
 - For signs an additional dimension sign

is appended to the dimensions of the argument arg2 to form the result of the local sensitivity operator. The extent of this additional dimension corresponds with the defined and/or selected number of targets, increment values and/or signs. For a definition of the additional dimensions check Tab. 10.7.

Firstly, dimension target_sequ is appended on demand, secondly dimension incr and thirdly dimension sign.

Having a model variable definition as in Example 5.1 at page 26 and assuming address_default=coordinate in <model>.cfg Assume the experiment description file (f) from Example 6.3 at page 47 then in result-processing

sens_abs(` ',atmo_g)

absolute sensitivity measure for atmo_g for all targets, increments and signs Dimensionality = 4 Coordinates = time, target_sequ, incr, sign Extents = 20, 3, 4, 2



```
sens rel('sels n(+), selt i(1)', atmo g)
                                     relative sensitivity measure for atmo_g
                                     for target p1 and all positive increments
                                     Dimensionality = 2
                                     Coordinates = time , incr
                                     Extents = 20, 4
sens abs('seli v(0.001:0.05)',atmo g)
                                     absolute sensitivity measure for atmo g
                                     for all targets, increment values 1 to 3 and all signs
                                     Dimensionality = 4
                                     Coordinates = time , target_sequ , incr , sign
                                     Extents = 20, 3, 3, 2
lin abs('seli v(0.001:0.05)',atmo g)
                                     absolute linearity measure for atmo g
                                     for all targets and increment values 1 to 3
                                     Dimensionality = 3
                                     Coordinates = time, target sequ, incr, sign
                                     Extents = 20, 3, 3
                                                                      Example file: world.post
```

Example 8.7 Post-processing operators for local sensitivity analysis

8.4.5 Optimization

The goal of an optimization experiment is to minimize a cost function by determining the corresponding optimal point in the target space. Nevertheless, the specified model output from all single runs is stored during the experiment.

While the single run that corresponds with the optimal cost function can be post-processed in the single-run modus, the whole experiment can be post-processed as a Monte Carlo analysis. Keep in mind that the targets do not follow an pre-defined distribution.

8.5 User-Defined Operators

8.5.1 Declaration of Operator Dynamics

- User-defined operators consist of a declarative and a computational part.
 - In the declarative part consistency of the non-character operands are checked and dimensionality, extents and coordinates of the result are defined.
 - In the computational part the result of the operator in dependency of the operands is computed.
- User-defined operators are specified in the ASCII operator description file <model>.odf. This file is used to check user-defined operators syntactically during result-post-processing.
- Check usr_opr_<opr>.f and apply the assigned operator <opr> for examples of user-defined operators.
- In SimEnv the declarative and computational part of an user-defined operator <opr> is hosted in a file usr_opr_<opr>.f. The assigned executable has the name <opr>.opr and has to be located in this directory that is stated in <model>.cfg as the hosting directory opr_directory for user-defined operators.
- Use the shell script operator_f.lnk <opr> to compile and link from usr_opr_<opr>.f an executable <opr>.opr that represents the user-defined operator <opr>.
- Use the simenv.chk to check user-defined operators
- Any user-defined operator can be transformed directly without changes to a built-in operator
- The functions to declare and compute user-defined operators listed below use a named common block simenv.

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Function name	Function description	Inputs / outputs / function value	Inputs / outputs / function value description
	Functions to host	declarative and	computational part in <model>.f</model>
icheck_user_ def_operator ()	checks consis- tency of operator arguments and defines dimen- sionality and di- mensions of result	integer*4 icheck_user_ def_operator (function value)	return code = 0 ok ≠ 0 inconsistency between operands
icompute_user_ def_operator (result)	computes result of the operator in dependency on operands	real*4 result(1) (output) integer*4 icompute_user_ def_operator (function value)	result vector of the operator return code = 0 ok ≠ 0 user-defined interrupt of calculation

Tab. 8.15Operator functions: Declarative and computational part

Function name	Function description	Inputs / outputs / function value	Inputs / outputs / function value description
Functio	ns to get and put st	ructure informati	ion in declarative and computational part
iget_char_arg (iarg,	gets string and length of the string of a character	integer*4 iarg (input)	argument number
char)	argument	character*100 char (output)	string of the character argument
		integer*4 iget_char_arg (function value)	length of character argument
iget_dim_arg (iarg,	iarg4 > 0: gets dimensionality and extents of an	integer*4 iarg (input)	argument number, 0 for result
iext)	argument iarg4 = 0: gets dimensionality	integer*4 iext(9) (output)	extents iext(1) iext(iget_dim_arg) of argument / result
	and extents of the result	integer*4 iget_dim_arg (function value)	dimensionality of argument / result
iget_len_arg (iarg	iarg4 > 0: gets length of an argument	integer*4 iarg (input)	argument number, 0 for result
)	iarg4 = 0: gets length of re- sult	integer*4 iget_len_arg (function value)	length of argument / result
iget_nr_arg ()	gets number of arguments of the current operator	integer*4 iget_nr_arg (function value)	number of arguments

Function name	Function description	Inputs / outputs / function value	Inputs / outputs / function value description
(gets data type ofiargan argument)iarg4 = 0:		integer*4 iarg (input) integer*4 iget_type_arg (function value)	argument number, 0 for result type of argument / result = -1 byte = 4 float = -2 short = 8 double
iget_co_chk_ modus ()	gets level of coor- dinate check for arguments ac- cording to <model>.cfg</model>	integer*4 iget_co_chk_ modus (function value)	 = -4 int level of coordinate check for arguments = 0 without = 1 weak = 2 strong
iget_co_arg (iarg, ico_blk, ico_beg)	gets coordinate block numbers and coordinate begin numbers of an argument	integer*4 iarg (input) integer*4 ico_blk(9) (output) integer*4 ico_beg(9) (output) integer*4 iget_co_arg	argument number block number of the coordinate ico_blk(1) ico_blk(idimens) begin numbers of the coordinate ico_beg(1) ico_beg(idimens) return code = 0 ok
iget_co_val (ico_blk, ico_pos, co_val)	gets coordinate value at a position from a coordinate	(function value) integer*4 ico_blk (input) integer*4 ico_pos (input) real*4 co_val (output) integer*4 iget_co_arg (function value)	block number of the coordinate position of the value to get within all coordinate values coordinate value return code = 0 ok = 1 ico_pos out of range = 2 storage exceeded
ichk_2args (iarg1, iarg2)	checks two argu- ments on same dimensionality, extents and coor- dinates	integer*4 iarg1 (input) integer*4 iarg2 (input) integer*4 ichk_2args (function value)	argument number argument number return code = 0 ok = 1 differing dimensionalities = 2 differing extents = 3 differing coordinates according to <model>.cfg = 4 iarg1=iarg2</model>

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Function name	Function description	Inputs / outputs / function value	Inputs / outputs / function value description
iput_struct_res (inplace, idimens {, iext,	puts potential in- place-storage, dimensionality, extents, coordinate bock and begin numbers of the	integer*4 inplace (input)	potential inplace-indicator for result. result can be computed in-place with the following non-character arguments = -1 all = 0 none > 0 e.g. = 135 with args 1, 3 or 5
ico_blk, ico_beg })	Currently, only coordinates from the arguments can be assigned to the	integer*4 idimens (input) integer*4 iext(9) (input)	dimensionality of the result only for idimens > 0: extents iext(1) iext(idimens) of the result
	result. Apply only in the declarative part.	integer*4 ico_blk(9) (input) integer*4 ico_beg(9) (input) integer*4 iput dim res	only for idimens > 0: coordinate block numbers ico_blk(1) ico_blk(idimens) of the result only for idimens > 0: coordinate begin numbers in block ico_blk ico_beg(1) ico_beg(idimens) of the result return code = 0 ok
		(function value)	≠ 0 inconsistency between operands

Tab.8.16Operator functions to get and put structural information

All of these functions return -999 as an error indicator if the argument iarg is undefined.

Function name	Function description	Inputs / outputs / function value	Inputs / outputs / function value description					
Functi	Functions to get and check argument values and put results in computational part							
arg1 (index	gets value of a non-character argument with index index	integer*4 index (input)	vector index of an argument					
/ arg9 (index)		real*4 arg1 arg9 (function value)	value of an argument arguments of any type are transferred to float representation					
clip_undef (value	overflow: checks a real*8 value on an undefined	real*8 value (input)	value to be checked					
)	real*4 result underflow: sets a real*8 value to zero if appropriate	real*4 clip_undef (function value)	clipped value normally identified with a result res e.g., res(i)=clip_undef(value8)					
is_undef (value	checks whether value is undefined before processing	real*4 value (input)	argument to be checked					
)	it	integer*4 is_undef (function value)	= 0 value is defined= 1 value is undefined					

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Function name	Function description	Inputs / outputs / function value	Inputs / outputs / function value description
set_undef	sets a result to be	real*4	normally identified with a result res
(undefined	set_undef	e.g., res(i)=set_undef()
)		(function value)	

 Tab.
 8.17
 Operator function to get / check / put arguments and results

n-dimensional matrices are forwarded to user-defined operators as one-dimensional vectors, using the Fortran storage model (see Glossary).

In Example 15.7 at page 141 implementation of the user-defined operator mat_mul is described in detail.

8.5.2 Operator Definition File <model>.odf

<model>.odf is an ASCII file that follows the coding rules in chapter 11.1 at page 107 with the keywords, names, sub-keywords, and values as in Tab. 8.18. <model>.odf describes the user-defined operators.

keyword	name	sub- keyword	Line type	Max. line nmb.	values	Explanation
odf	<nil></nil>	descr	0	any	<string></string>	general operator descriptions
operator	<operator_< td=""><td>descr</td><td>0</td><td>1</td><td><string></string></td><td>operator description</td></operator_<>	descr	0	1	<string></string>	operator description
	name>	nr_args	m	1	<integer_value></integer_value>	number of arguments defined for the operator operator_name 0< <integer_value> < 10</integer_value>
		nr_charargs	m	1	<integer_value></integer_value>	from <integer_val>: number of character argu- ments defined for the operator operator_name 0 ≤ <integer_value> ≤ nr_args</integer_value></integer_val>

 Tab.
 8.18
 Elements of an operator description file <model>.odf

To Tab. 8.18 the following additional rules and explanations apply:

• For the description of line type check Tab. 11.4 at page 109.

```
odf
                                 Operator description for the
                    descr
                                 examples in the SimEnv User Guide
odf
                    descr
operator char_test descr
                                 test character arguments
operator char_test nr_args
                                 3
operator char test nr char args 2
operator corr coeff descr
                                 correlation coefficient
operator corr_coeff nr_args
                                 2
operator corr_coeff nr_char_args 0
operator div
                                 division
                    descr
operator div
                   nr args
                                 2
operator div
                    nr char args 0
```

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```
division without special cases
          simple div descr
operator
operator simple div nr args
                                     2
operator simple div nr char args
                                     0
operator mat mul
                       descr
                                     matrix multiplication
operator
         mat mul
                       nr args
                                     2
         mat mul
operator
                       nr char args
                                     0
                                         Example files: world_[f | c | cpp | py | sh ].odf
```



User-defined operator description file <model>.odf

8.5.3 Handling Undefined Results

In user-defined operators

- Check always whether an argument value val is undefined by is_undef(val) before it is processed.
- Set a result to be undefined by the function set_undef()
- Check usr_opr_mat_mul.f in Example 15.7 or usr_opr_div.f in the examples directory for a more detailed example.
- If things go so wrong that computation of the whole result expression has to be stopped alternatively it is
 possible to
 - Set all elements of the results to be undefined
 - Set icompute_user_def_operator ≠ 0 (otherwise set it always = 0)
 - In both cases application of the next operators will be suppressed and consequently computation of the result expression will be stopped
 - Check usr_opr_char_test f for a detailed example

8.6 Undefined Results

- By performing operator chains and due to possibly unwritten model output during simulation parts of the intermediate and/or final result values can be undefined within the float data representation.
- If an operand is completely undefined the computation of the result is stopped without evaluating the following operands and operators.
- For nodata value representation check Tab. 10.10.

8.7 Macros and Macro Definition File <model>.mac

- A macro in model output post-processing is an abbreviation for a result expression, consisting of an operator chain applied on operands.
- Generally, they are model related and they are defined by the user.
- Macros are identified in result post-processing expressions by the suffix _m.
- A macro is plugged into a result expression by putting it into parentheses during parsing:

Example: equ_100yrs_m*test_mac_m from Example 8.9 below is identical to

- (avg(atmo(c=20:-20, *, c=1, c=11:20)) 400) * (1+(2+3) * 4)
- Macros must not contain macros.
- Use simenv.chk to check macros. During the macro check validity of the following information is not checked:
 - Un-pre-defined character arguments of built-in operators (check Tab. 15.8)
 - Constant arguments of built-in operators (check Tab. 15.9)
 - Character arguments of user-defined operators
 - Operators with respect to dimensionality and dimensions of its operands



In SimEnv macros are defined in the file <model>.mac. <model>.mac is an ASCII file that follows the coding rules in chapter 11.1 at page 107 with the keywords, names, sub-keywords, and values as in Tab. 8.19. <model>.mac describes the user-defined macros.

keyword	name	sub- keyword	Line type	Max. line nmb.	value	Explanation
mac	<nil></nil>	descr	0	any	<string></string>	general macro descriptions
macro	<macro_< td=""><td>descr</td><td>0</td><td>1</td><td><string></string></td><td>macro description</td></macro_<>	descr	0	1	<string></string>	macro description
	name>	unit	m	1	<string></string>	unit of the value of the macro
		define	m	any	<string></string>	macro definition string macro definition can be ar- ranged at a series of define- lines in analogy to the rules for result expressions (see chapter 8.1.1).

 Tab.
 8.19
 Elements of an macro description file <model>.mac

To Tab. 8.19 the following additional rules and explanations apply:

- For the description of line type check Tab. 11.4 at page 109.
- Values for sub-keywords 'descr' and 'unit' are not evaluated during parsing a result expression.

```
Macro definitions for the
mac
                        descr
                                    examples in the SimEnv User Guide
mac
                        descr
                                    2^{\,nd} century tropical level 1 average
macro equ 100yrs
                        descr
macro equ 100yrs
                       unit
                                    without
macro equ 100yrs
                       define
                                    avg(atmo(c=20:-20,*,c=1,c=11:20))
macro tst
                        descr
                                    test macro
macro tst
                        define
                                    1+(2+3)*
macro tst
                        define
                                    4
                                          Example files: world_[f | c | cpp | py | sh ].mac
```

Example 8.9 User-defined macro definition file <model>.mac

8.8 Saving Results

The result files <model>.res<res_char>.[nc | ieee | ascii] and <model>.inf<res_char>.[ieee | ascii] contain all the model and experiment information for further processing of results.

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9 Visual Experiment Evaluation

Experiment evaluation is based on application of visualization techniques to the output data, computed during experiment post-processing and stored in NetCDF format. Currently, a preliminary version is implemented.

Analysis and evaluation of post-processed data selected and derived from large amount of relevant model output benefits from visualization techniques. Based on metadata information of the post-processed experiment type, the applied operator chain, and the dimensionalities of the post-processor output pre-formed visualization modules are evaluated by a suitability coefficient how they can map the data in an appropriate manner.

The visualization modules offer a high degree of user support and interactivity to cope with multi-dimensional data structures. They cover among others standard techniques such as isolines, isosurfaces, direct volume rendering and a 3D difference visualization techniques (for spatial and temporal data visualization). Furthermore, approaches to navigate intuitively through large multi-dimensional data sets have been applied, including details on demand, interactive filtering and animation. Using the OpenDX visualization platform techniques have been designed and implemented, suited in the context of analysis and evaluation of simulated multi-run output functions.

Currently, visual experiment evaluation is the only SimEnv service that comes with a graphical user interface. In this user interface a help-services is implemented that should be used to gather additional information on how to select post-processed results for visualization and on visualization techniques provided by SimEnv.

To get access rights to the SimEnv visualization server use the SimEnv service simenv.key one time. Check chapter 10.2 for more information.



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10 General Control, Services, User Files, and Settings

In a general configuration file <model>.cfg the user controls general settings for the simulation environment. Besides simulation performance and model output post-processing SimEnv supplies a set of auxiliary services to check status of the model, to dump model and post-processor output and files and to clean a model from output files. General settings reflect case sensitivity, nodata values and other information related to SimEnv.

10.1 General Configuration File <model>.cfg

In the ASCII file <model>.cfg general SimEnv control variables can be declared. <model>.mdf is an ASCII file that follows the coding rules in chapter 11.1 at page 107 with the keywords, names, sub-keywords, and info as in Tab. 10.1.

keyword	name	sub- keyword	Line type	Max. line nmb.	value	Explanation
cfg	<nil></nil>	descr	0	any	<string></string>	general configuration description
general	<nil></nil>	message_level	0	1	[info warning error]	specifies which message types to show during simenv.chk and in <model>.mlog</model>
model	<nil></nil>	out_directory	0	1	<direct></direct>	model output directory
		out_format	0	1	[netcdf ieee]	model output format
		out_size_ threshold	0	1	<non_negative_ integer_value></non_negative_ 	file size threshold in kBytes for lumped model output
		out_ieee_ blocksize	0	1	<positive_ integer_value></positive_ 	block size in kBytes for IEEE model output
		distributed	0	1	[no yes]	indicates a distributed model
experiment	<nil></nil>	restart_ini	0	1	[no yes]	perform <model>.ini for ex- periment re-start</model>
		begin_run	0	1	<non_negative_ integer_value></non_negative_ 	begin single run number
		end_run	0	1	[last <non_negative_ integer_value>]</non_negative_ 	end single run number
		email	0	1	<string></string>	email notification address
postproc	<nil></nil>	out_directory	0	1	<direct></direct>	post-processing output direc- tory
		out_format	0	1	[netcdf ieee ascii]	post-processing output format
		address_default	0	1	[coordinate index]	post-processing address de- fault for model variables
		coord_check	0	1	[strong weak without]	post-processing coordinate check by operators
		opr_directory	0	1	<direct></direct>	directory the post-processors looks for user-defined opera- tor executables
		visualization	0	1	[yes no]	determine whether to directly visualize an entered result during result post-processing

 Tab. 10.1
 Elements of a general configuration file <model>.cfg

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To Tab. 10.1 the following additional rules and explanations apply:

- For the description of line type check Tab. 11.4 at page 109.
- <string>, <direct>, <non_negative_integer_value> and <positive_integer_value> are placeholder for corresponding strings.
- For keyword 'general', sub-keyword 'message_level': Message output during simenv.chk and to the model interface log-file <model>.mlog is controlled by this information.
 Specify info to output errors and warnings and additional information warning to output errors and warnings
 - error to output

during simenv.chk and to <model>.mlog.

For keyword 'model', sub-keyword 'out_size_threshold':

errors

Specify here the threshold in kBytes for the sum of the size of all model output variables (according to their extents and data types) that is used to decide whether the SimEnv model output data for the whole run ensemble is stored into one file <model>.outall.[nc | ieee] or in single output files <model>.out<run_char>.[nc | ieee].

• For keyword 'model', sub-keyword 'out_ieee_blocksize': IEEE compliant model output for single files is written in single records with a length of <out_ieee_blocksize> kBytes. If <out_size_threshold> is less than this value, this value is adapted to <out_size_threshold>.

• For keyword 'model', sub-keyword 'distributed':

Ist value has to be set to "yes" for distributed models if

- More than one distributed sub-models use SimEnv functionality by the simenv_*_*-functions and
- Sub-models may get and send data from and/or to SimEnv data files in parallel. A distributed model where the sub-models are performed sequentially one by one can run with distributed = 'no'.
- For keyword 'experiment', sub-keyword ['begin_run' | 'end_run']: With the exception of an optimization experiment SimEnv enables to perform an experiment partially by performing only an experiment slice out of the whole run ensemble (see chapter 7.4 at page 54). Therefor assign appropriate run numbers to this two descriptors. Make sure that begin and end run represent run number from the experiment (including run number 0) and that begin run ≤ end run. The string 'last' always represents the last simulation run of the whole run ensemble. For an optimization experiment these two sub-keywords are ignored.
- For keyword 'experiment', sub-keyword 'email':

After performing an experiment an email is sent to the email address specified in <string>.

- For keyword 'postproc', sub-keyword 'address_default': During post-processing portions of multi-dimensional model output variables can be addressed by coordinate (c= ...) or index (i= ...) reference. A default is established here.
- For keyword 'postproc', sub-keyword 'coord_check': During post-processing feasibility of application of an operator on its operands is checked with respect to the coordinate description of the operands. Different levels of this check are possible. A default is established here.
- For keyword 'postproc', sub-keyword 'visualization':

Specifies whether to directly visualize an entered result during post-processing.

Please keep in mind to ensure consistency of control settings in <model>.cfg across different SimEnv services. As an example you have to run experimentation, post-processing and dump with the same model output file size threshold out_size_threshold for binary output in <model>.cfg.

Tab. 10.2 lists the default values in the general configuration file in the case of absence of the appropriate sub-keyword.

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keyword	sub-keyword	value-default (*)	For more information see
cfg	descr	<nil></nil>	above
general	message_level	info	above
model	out_directory	./	above
	out_format	NetCDF	chapter 12
	out_size_threshold	10	above
	out_ieee_blocksize	50	above
	distributed	no	chapter 5.8 and above
experiment	restart_ini	no	chapter 7.3
	begin_run	0	chapter 7.1 - 7.4
	end_run	last	chapter 7.1 - 7.4
	email	<nil></nil>	chapter 7.1
postproc	out_directory	./	above
	out_format	NetCDF	chapter 12
	address_default	coordinate	chapter 8.1.3 and above
	coord_check	strong	chapter 8.1.5 and above
	opr_directory	./	chapter 8.5
	visualization	yes	above

Tab. 10.2

Default values for the general configuration file (*): in the case of absence of the appropriate sub-keyword

cfg cfg	descr descr	General configuration file for the examples in the SimEnv User Guide
general	message_level	info
model	out_directory	mod_out
model	out_format	netcdf
model	out_size_threshold	100
experiment	begin run	0
experiment	end_run	last
postproc	out_directory	res_out
postproc	out format	netcdf
postproc	address_default	index
postproc	coord_check	strong
postproc	opr_directory	./
postproc	visualization	no

Example 10.1 User-defined general configuration file <model>.cfg

10.2 Main and Auxiliary Services

The following SimEnv service commands are available from the SimEnv home directory \$SE_HOME. Besides experiment performance and model output post-processing there are additional auxiliary SimEnv services to check input information consistency, to monitor the status of simulation experiments, to dump files of model and post-processor output and to wrap up the SimEnv workspace.



SimEnv service	Use to				
Main Services					
simenv.run	prepare and run an experiment				
<model></model>	(see chapter 7.1)				
simenv.rst	restart an experiment				
<model></model>	(see chapter 7.3)				
simenv.res	perform experiment result post-processing for run number <run> or for the whole run</run>				
<model></model>	ensemble (<run> = -1, default).</run>				
{ [new append replace] }	Before entering post-processing those output files <model>.res<res_char>.[nc ieee ascii] and <model>.inf<res_char>.[ieee ascii] with the highest two-digit number</res_char></model></res_char></model>				
{ <run>}</run>	<pre>char> are identified and new result files for <res+1> are written / the results are</res+1></pre>				
(and)	appended / or the result files are replaced by a new ones.				
simenv.vis	perform visual post-processor output visualization for that NetCDF post-processor				
<model></model>	output file with the highest two digit number <res char=""> (<res char=""> = latest, default)</res></res>				
{ [latest <res>] }</res>	or with the file number <res_char>. Visualization runs on a remote server.</res_char>				
	Auxiliary Services				
simenv.chk	check on model script files (<model>.run, <model>.rst, <model>.ini, <model>.end)</model></model></model></model>				
<model></model>	check <model>.cfg</model>				
	<model>.odf</model>				
	<model>.mdf</model>				
	<model>.edf</model>				
	<model>.gdf</model>				
	<model>.mac</model>				
	existing model and post-processor output files				
aimany ata	generate pre-experiment output statistics get the current status of an				
simenv.sts <model></model>					
{ <sleep> }</sleep>	• experiment that was started from a login node of a parallel machine and that is running in a parallel or sequential job class of this machine				
	Start this service from a login-node of the parallel machine				
	 optimization experiment 				
simenv.dmp	dump SimEnv model output and post-processor output files				
<model></model>	Files to dump have to match the SimEnv file name convention for model and/or post-				
	processor output and are expected to be in the directories as stated in <model>.cfg.</model>				
	Model output variables and post-processor results in IEEE and/or ASCII format with a				
	dimensionality greater than 1 are listed according to Fortran storage model for multi-				
	dimensional fields (see Glossary).				
simenv.cpl	complete sequence of SimEnv commands				
<model> { <run> }</run></model>	simenv.chk, simenv.run, simenv.res, simenv.vis, simenv.dmp simenv.res is performed with input file <file> (if available) and interactively, for both</file>				
{ <file> }</file>	optionally only for single run <run>.</run>				
simenv.cln	clean up model and post-processor output files				
<model></model>	Deletes all model output files, post-processor output files, log-files, and auxiliary files.				
simenv.cpy	copy all SimEnv example files <model>* from the examples directory of \$SE HOME</model>				
<model></model>	to the current directory.				
	Additionally, example files of user-defined operators and for models world_[f c cpp				
	py sh]* common user defined files are copied. All files are only copied if they do not				
	already exist in the current directory, this SimEnv service is started from.				
simenv.hlp <topics></topics>	acquire basic SimEnv help information for the specified topics				
simenv.key	generate a ssh2-key to get password-free access to the visualization server.				
<user_name></user_name>	Start this service at machine aix02 only one time before the first access to simenv.vis				
_	or if the ssh2-key does not work properly.				
	You will get an email when your password-free server access is possible.				
Tab 103	SimEnv services				

Tab. 10.3SimEnv services

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- With the exception of the simenv.cpy, simenv.hlp and simenv.key: Start a services only from the current working directory.
- With the exception of simenv.sts: Do not start a SimEnv service from a working directory where an other SimEnv service is still active.

Script / file (in the current working direc- tory \$SE_WD)	Explanation	Exist status	For more information see chapter
<model>.cfg</model>	ASCII user-defined general configuration file	optional	10.1
<model>.mdf</model>	ASCII user-defined model (variables) description file	mandatory	5.1
<model>.edf</model>	ASCII user-defined experiment description file	mandatory	6.1
<model>.mac</model>	ASCII user-defined macro description file	optional	8.7
<model>.odf</model>	ASCII user-defined operator description file	optional	8.5.2
<model>.gdf</model>	ASCII user-defined GAMS model output description file	for GAMS mod- els mandatory	5.7.2
<model>.run (*)</model>	model shell script to wrap the model executable	mandatory	7.6
<model>.rst (*)</model>	model shell script to prepare single model run restart	optional	7.6
<model>.ini (*)</model>	model shell script to prepare simulation experiment additionally to standard SimEnv preparation	optional, for Python and GAMS models mandatory and standardized	7.6
<model>.end (*)</model>	model shell script to clean up simulation experiment	optional, for GAMS models man- datory and standardized	7.6
<model>. <run_char>.err</run_char></model>	touch this file in the model, in <model>.run and/or <model>.rst as an indicator to stop the complete ex- periment after <model>.run has been finished for sin- gle model run <run char=""></run></model></model></model>	optional	7.6
simenv.jcf_par	user-specific job control file to submit a job by the LoadLeveler to a parallel class	optional	7.6
simenv.jcf_seq	user-specific job control file to submit a job by the LoadLeveler to a sequential class	optional	7.6
simenv.oopt	user-specific control and option file for experiment type optimization	optional	6.5.3
<opr>.opr (in the opr_directory according to <model>.cfg)</model></opr>	executable for user-defined operator <opr></opr>	optional	8.5

10.3 User Scripts and Files

Tab. 10.4

User files and scripts to perform any SimEnv service (*): make sure by the UNIX command chmod u+x <model>.??? that the shell script <model>.??? has execute permission

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File / location	Generated in	Explanation			
Permanent files					
<model>.edf_adj</model>	experiment preparation (all but optimization)	ASCII adjustment input file for the run ensemble derived from <model>.edf. Record no. n+1 corresponds to single run no. n.</model>			
	experiment performance	Column no. m of each record is the adjustment for			
\$SE_WD <model>.out<run_char></run_char></model>	(optimization) experiment performance	experiment target no. m in the edf-file model output of run number <run></run>			
.[nc ieee]	(if model output of a single run ≥ out_size_threshold in <model>.cfg)</model>	to be processed by the post-processor (for experiment performance in a parallel job class at a parallel machine files <model>.out<run_char>.[nc ieee] are created</run_char></model>			
model out_directory		temporarily)			
<model>.outall .[nc ieee]</model>	experiment performance (if model output of a single run < out_size_threshold in <model>.cfg)</model>	model output of all runs to be processed by the post-processor			
model out_directory <model>.elog</model>	experiment performance	ASCII minutes file of experiment performance			
\$SE WD	experiment performance	(simenv.run and all successive simenv.rst)			
<model>.mlog</model>	experiment performance	ASCII minutes file of m odel interface performance			
\$SE WD		(simenv.run and all successive simenv.rst)			
<model>.nlog</model>	experiment performance	ASCII minutes file of n ative model output, redi-			
\$SE_WD		rected from terminal (simenv.run and all successive simenv.rut)			
<model>.res<res_char></res_char></model>	experiment post-processing	output file of a post-processor session			
.[nc ieee ascii]					
postproc out_directory					
<model>.inf<res_char> .[ieee ascii]</res_char></model>	experiment post-processing	output structure description file of a post-processor session			
postproc out_directory					
run <run_char> \$SE_WD</run_char>	experiment performance (only for GAMS models)	sub-directory for GAMS model performance that are kept according to the sub-keyword 'keep_runs'			
<pre><model>.olog</model></pre>	experiment performance	in <model>.gdf ASCII minutes file of optimization experiment per-</model>			
	(only for experiment type	formance			
\$SE_WD	optimization)				
<model>.edf_cf</model>	experiment performance (only for experiment type	ASCII file of cost function values. Record no. n+1 corresponds to single run no. n.			
\$SE_WD	optimization)				
Temporary files (do not delete during performing the corresponding service)					
simenv.cfg	all services	structured ASCII representation of <model>.cfg</model>			
\$SE_WD					
<model>.</model>	experiment preparation,	structured binary representation			
[mdf edf odf mac] _bin	experiment post-processing	of <model>.[mdf edf odf mac]</model>			
\$SE_WD					

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File / location	Generated in	Explanation
<model>.parid</model>	experiment performance	ASCII file with the job-id.
\$SE_WD	(only for performance in a Loadl job class)	Used for performance of simenv.sts
asa_opt	experiment performance	auxiliary files for experiment type optimization
asa_out	(only for experiment type	
asa_usr_out	optimization)	
\$SE_WD		
run <run_char></run_char>	experiment performance (only for GAMS models)	sub-directory for GAMS model performance that are not kept according to the sub-keyword
sub-direct. of \$SE_WD		'keep_runs' in <model>.gdf</model>
<model>_</model>	experiment performance	auxiliary files
[pre main post].inc	(only for GAMS models)	<model> = GAMS main and all interfaced sub- models</model>
\$SE_WD		
simenv_*.tmp	different services	auxiliary files
\$SE_WD		

Tab. 10.5

Files generated during performance of SimEnv services For the current working directory \$SE_WD see Tab. 10.11.



Fig. 10.1 sketches usage of main SimEnv user scripts and files in the course of model interfacing, experiment preparation and performance, post-processing, and evaluation.

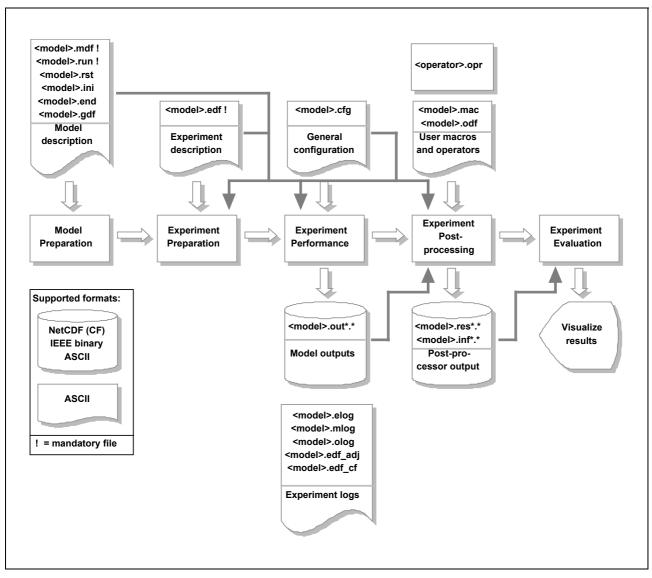


Fig. 10.1

SimEnv user scripts and files

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10.4 Built-In Names

SimEnv has a number of built-in model variable, coordinate and shell script variable names that can not be used for corresponding user-defined names.

Tab. 10.6 lists the built-in (pre-defined) model variables that are output during experiment performance to SimEnv model output structures and are available in model output post-processing without defining them in the model output description file <model>.mdf and without using the corresponding simenv_put-function calls in the model.

Built-in model variable name	Dimen- sionality	Extents	Data type	Meaning
sim_time	0		float	elapsed simulation time in seconds (rounded to 2 decimal places) per single run for <model>.run</model>

Tab. 10.6Built-in model variables

Tab. 10.7 lists the built-in (pre-defined) coordinates that are used in model output post-processing when additional dimensions are generated by an operator.

Built-in coordinate name	Generated by operator	Meaning	Definition (check Tab. 11.6)
bin_mid	hgr, hgr_e, hgr_l	bin mid value	equidist_end <xx>(<yy>) 999999 with <xx> = first bin mid <yy> = bin width</yy></xx></yy></xx>
bin_no	hgr, hgr_e, hgr_l	bin number	equidist_end 1(1)999999
incr	lin_abs, lin_rel, sens_abs, sens_rel, sym_abs, sym_rel	increment values	dependent on experiment description and operator arguments
index	maxprop, maxprop_l, minprop, minprop_l,	index number	equidist_end 1(1)999999
run	ens	run number	equidist_end 1(1)999999
sign	sens_abs, sens_rel	sign of incremental change: +ε : sign +1, -ε : sign -1	equidist_end -1(2)1
stat_measure	stat_full, stat_red	basic statistical measures	equidist_end 1(1)999999
target_sequ	lin_abs, lin_rel, sens_abs, sens_rel, sym_abs, sym_rel	target sequence: 1 st target in edf-file = 1 2 nd target in edf-file = 2 	equidist_end 1(1)999999
<target_name></target_name>	behav	target values	dependent on experiment description and operator arguments

Tab. 10.7Built-in coordinates



Tab. 10.8 lists the built-in (pre-defined) shell script variables that are used in \$SE_HOME/simenv_*_sh and finally in <model>.run.

Built-in shell script variable name	Meaning
run_int	current run number as integer
run_char	current run number as character string
target_name	target name for simenv_get_sh
target_def_val	default target value for simenv_get_sh

Tab. 10.8

Built-in shell script variables in <model>.run

10.5 Case Sensitivity

As stated in Tab. 10.9 all names used in SimEnv are case insensitive. Internally, they are mapped on a lowercase representation and this lowercase representation is used also for model and/or post-processor output files in NetCDF, IEEE and/or ASCII format.

Where?	Entity	Case sensitivity	Example
overall	model name	sensitive	simenv.chk World_f
user-defined files (see Tab. 11.1)	 keyword name exception: GAMS model file name in <model>.gdf</model> sub-keyword 	insensitive	experiment END_RUN last
	 information <value> exceptions:</value> <direct> and <file name=""></file></direct> 	insensitive	experiment end_run LAST cfg descr This is exception:
	 for <sub-keyword> = '<string>_directory' - and in <value_list></value_list></string></sub-keyword> <value> for <sub-keyword> = ['descr' 'unit'])</sub-keyword></value> 		specific comb file AbC.d
model interface	 variable and target name 	insensitive	<pre>call simenv_put_f(`ATMO',atmo) target_name=`P1' target_value=1 \$SE_HOME/simenv_get_sh</pre>
post-processing	 optional result description and unit 	sensitive	<pre>Energy [kW] = my_opr(atmo)</pre>
	 variable and target name operator name number macro name macro identifier _m 	insensitive	exp(atmo) + 3*EXP(ATMO)

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Where?	Entity	Case sensitivity	Example
	 character arguments of built-in operators with pre-defined values (check Tab. 15.8) 	insensitive	<pre>count('ALL' , atmo)</pre>
	 character arguments of built-in operators without pre-defined values 	check Tab. 15.8	<pre>get_table_fct('MyFile.dat' ,</pre>
	 character arguments of user-defined operators 	sensitive	<pre>char_test(`arg11' , `Arg21' ,</pre>

Tab. 10.9Case sensitivity of SimEnv entities

10.6 Nodata Representation

For model output with the SimEnv model coupling interface functions and for post-processor output the following data type specific nodata values are used to represent undefined (unwritten) model output or undefined post-processor output:

Data type	Nodata value
byte	127
short	32767
int	2147483648
float	3.4E+38
double	1.79D+308

Tab. 10.10

Data type related nodata values



10.7 Environment Variables

The following operating system environment variables are used by SimEnv:

Environment variable	Meaning	Explanation
SE_HOME	SimEnv home directory	has to be defined by the user Value = [/usr/local/simenv/bin any previous SimEnv version at /usr/local/simenv/version_archive] Has to be included in the file \$HOME/.profile. Recommended value is: SE_HOME= /usr/local/simenv/bin
PYTHONPATH	path to search Python and Python files	has to be defined by the user Value = machine / user dependent and has to be expanded by \$SE_HOME Has to be included in the file \$HOME/.profile. Recommended value is: PYTHONPATH= ./: /usr/local/lib/python2.1: /usr/local/lib/site-packages/Numeric: /home/rachimow/usr/local/lib/python2.1/ site-packages: \$SE_HOME
SE_RUN	run number of a single run	defined automatically in <model>.run and <model>.rst Value = <run_int></run_int></model></model>
SE_1STRUN	first single run of an experiment	defined automatically in <model>.run and <model>.rst Value = [yes no]</model></model>
SE_WD	current SimEnv working directory	defined automatically within any SimEnv service Value = <direct></direct>

Tab. 10.11Environment variables

In the file \$HOME/.profile specify first the operating system environment variable SE_HOME and then the environment variable PYTHONPATH.

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11 Structure of User-Defined Files, Coordinate Transformation Files, Value Lists

Basic information to describe general control settings of SimEnv, model output variables, the experiment itself, macros and user-defined operators as well as GAMS model specific information is stored in user-defined files. They are ASCII files and have a common structure that is described in this chapter. Additionally, coordinate transformation files are described and value lists are defined in general.

11.1 General Structure of User-Defined Files

All user-defined files listed in Tab. 11.1 have the same structure. They are ASCII-files with the following record structure:

{ <sep> } <keyword> <sep> { <name> <sep> } <sub-keyword> <sep> <value> { <sep> }

with

with	า	
•	<name></name>	 is the name of a model variable GAMS model source file experiment target coordinate user-defined operator or macro
		Declaration of <name> depends on the related keyword <keyword></keyword></name>
•	<keyword></keyword>	is a string Normally, more than one lines with differing sub-keywords belong to one "keyword-block".
•	<sub-keyword></sub-keyword>	is a string Sub-keywords are defined only in relation to the user file and the keyword under consideration.
•	<value></value>	<pre>= <substring> { <sep> <substring> } is a string with user file, keyword and sub-keyword related information.</substring></sep></substring></pre>
•	<sep></sep>	is a sequence of white spaces

Sequence of keyword and sub-keyword lines can be arbitrary. For reasons of readability it is recommended to use a block structure like in the example below. Sequence of names in the separated name spaces (name spaces of coordinates, model variables, experiment targets, user-defined operators, macros) during processing is determined by the sequence the name occur the first time in the appropriate user file. Lines consisting only from separator characters as well as lines starting with a # as the first non-separator

character are handled as comment lines. For case sensitivity of the contents of user-defined files check Tab. 10.9 at page 105.

File	Contents	See des in chapter	cription at page
<model>.cfg</model>	general configuration file	10.1	95
<model>.mdf</model>	model output description file	5.1	21
<model>.gdf</model>	GAMS description file	5.7.2	35
<model>.edf</model>	experiment description file	6.1	39
<model>.odf</model>	operator description file	8.5.2	90
<model>.mac</model>	macro description file	8.7	91
arbitrary file name	coordinate transformation file	11.2	110

Tab. 11.1

User-defined files with general structure



The following restrictions hold for user-defined files:

Element	Constraints
line length	max. 160 characters
<name></name>	max. 20 characters
	(*) first character has to be a letter
	(*) must not end on _m
	(*) must not contain elemental operators and characters . and :
	(check Tab. 8.3 at page 66)
<value></value>	for sub-keyword = 'descr' without <name>: max. 512 characters</name>
	(total sum over all lines)
	for sub-keyword = 'descr' with <name>: max. 128 characters</name>
	for sub-keyword = ' <string>_directory': max. 70 characters</string>
	must not contain operating system environment variables
	for sub-keyword = 'unit': max. 32 characters

Tab. 11.2Constraints in user-defined files
(*): with the exception for GAMS model source code file names

Tab. 11.3 lists the reserved (forbidden) names, file names and directories to files that can not be declared in user-defined files.

Element	Reserved (forbidden) names
<name></name>	built-in model variables
excepted for GAMS model	according to Tab. 10.6
source code file names	built-in coordinates
	according to Tab. 10.7
	built-in shell script variables
	according to Tab. 10.8
	built-in shell environment variables
	according to Tab. 10.11
	special keywords in <model>.edf for behavioural</model>
	analysis:
	[default file]
<direct></direct>	must not contain operating system environment vari-
	ables (\$)
	If <direct> is specified in a relative manner it relates to</direct>
	the current working directory, the / a SimEnv service
	where <direct> is referred was started from.</direct>
<file_name></file_name>	SimEnv file names
	according to Tab. 10.4 and Tab. 10.5

 Tab. 11.3
 Reserved names and file names in user-defined files

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The **line type** in the description table for a user-defined file specifies whether a keyword / sub-keyword combination can be omitted.

Abbre- viation	User-defined file	Explanation	
m	all files	m andatory	
0	all files	o ptional	
c1	<model>.mdf keyword 'variable' sub-keyword ['coords' 'var_extents']</model>	conditional 1:forbiddenfor variables with dirmandatoryfor variables with dir	
c2	<pre><model>.mdf keyword 'variable' sub-keyword 'coord_extents'</model></pre>	conditional 2: forbidden for variables with dir optional for variables with dir	
c3	<model>.edf keyword 'target' sub-keyword 'adjusts'</model>	forbidden for experiment type	= Monte Carlo analysis = local sensitivity analysis = behavioural analysis
c4	<model>.edf for Monte Carlo analysis keyword 'target' sub-keyword 'sampling'</model>	conditional 4:mandatoryfor adjusts = distrforbiddenfor adjusts = file	
а	<pre><model>.edf for behavioural analysis keyword 'target' sub-keyword 'adjusts'</model></pre>	alternatively: either mandatory for all exponent or forbidden for all expo	
f	<pre><model>.edf for local sensitivity analysis keyword 'target' sub-keyword 'adjusts'</model></pre>	forbidden	

Tab. 11.4

Line types in user-defined files

descr This is a macro description file mac for the SimEnv User Guide mac descr atmo outside polar reg., final time, level 1 macro pol atmo descr macro pol_atmo unit without macro pol atmo define atmo(c=84:-56,*,c=1,c=20) macro m1 define avg(atmo g(c=11:20)). . .

Example 11.1 Structure of a user-defined file



11.2 Coordinate Transformation File

Some operators (currently, get_experiment) enable access to external data. Most of these operators supply in general a multi-dimensional result that has to be equipped - as usual in SimEnv post-processing - with a coordinate assignment. By applying these operators it can be necessary to transform the coordinate description from the external data to fit the data to the current model and/or experiment under consideration. The following cases can be distinguished:

- A dimension of the external data does not have a coordinate assignment. A coordinate has to be assigned to this dimension.
- A coordinate description of the external data has to be modified in a way that it matches with a defined coordinate of the model / experiment under consideration.
- A coordinate description of the external data has to be incorporated with and/or without modifications into the coordinate set of the model / experiment under consideration.

Coordinate transformations for external data are supported by a coordinate transformation file that is assigned to the operator result as an argument of the operator. Coordinate transformation files follow the same syntax rules as all other user-defined files (see chapter 10.1).

keyword	name	sub- keyword	Line type	Max. line nmb.	value	Explanation
trf	<nil></nil>	descr	0	any	<string></string>	general transformation de- scription
modify	<external_< td=""><td>rename</td><td>0</td><td>1</td><td><new_name></new_name></td><td>renames external coordinate</td></external_<>	rename	0	1	<new_name></new_name>	renames external coordinate
	coordinate_ name>	position_shift	0	1	<position_shift_ val></position_shift_ 	shifts all values of the external coordinate by the specified real value <position_shift_val></position_shift_val>
		values_shift	0	1	<values_shift_ val></values_shift_ 	shifts the result values on the coordinate by integer <values_shift_val> positions</values_shift_val>
		values_add	0	1	<value_list></value_list>	defines <values_shift_val> values to add to the coordi- nate values (for syntax see Tab. 11.6)</values_shift_val>
assign	<external_ coordinate_ name></external_ 	coord	0	1	<coord_name></coord_name>	assign to the dimension with <external_coordinate_name> an internal coordinate or a coordinate defined by the keyword 'coordinate'</external_coordinate_name>
		coord_extent	0	1	<co_val<sub>1>: <co_val<sub>2></co_val<sub></co_val<sub>	assigns start and end coordi- nate value to the dimension of the result under consideration.
coordinate	<new_< td=""><td>descr</td><td>0</td><td>1</td><td><string></string></td><td>coordinate axis description</td></new_<>	descr	0	1	<string></string>	coordinate axis description
	coordinate_	unit	0	1	<string></string>	coordinate axis unit
	name>	values	0	1	<value_list></value_list>	strictly monotonic sequence of coordinate values (for syntax see Tab. 11.6)

Tab. 11.5 Elements of an coordinate transformation file

To Tab. 11.5 the following additional rules and explanations apply:

- For the description of line type check Tab. 11.4 at page 109.
- With the sub-keyword 'values_shift' result values can be shifted on the corresponding coordinate by values_shift_val coordinate values. Consequently <values_shift_val> coordinate values have to be appended at the end of the coordinate for a positive value of <values_shift_val> and/or have to be inserted

at the begin of the coordinate for a negative value of <values_shift_val>. Coordinate values that are obsolete because of this shift are removed from the coordinate definition.

For a coordinate that is defined with equidistant coordinate values the extent of the coordinate is specified automatically by simply applying the equidistant rule for this coordinate.

For a coordinate with non-equidistant coordinate values the coordinate values necessary for the coordinate extension are defined by the sub-keyword 'values_add'.

If both **position_shift** and values_shift are specified for one coordinate, firstly position shift is applied to the coordinate and then the additional coordinate values from values_shift are added the the coordinate without applying the position_shift value.

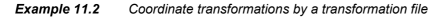
- For the sub-keyword '**coord_extent**' the same rules apply as for the sub-keyword 'coord_extents' from the model output description file <model>.mdf.
- For the keyword '**coordinate**' the same rules apply as for the keyword 'coordinate' from the model output description file <model>.mdf.
- Coordinates are incorporated additionally into the original coordinate set only for the current result.

Unlike all other user-defined files coordinate transformation files can not be checked.

assuming addres Assume the expe the correspondin Assume addition	es_default = coo eriment layout ir g experiment de ally result from a	on as in Example 5.1 at page ordinate in <model>.cfg n Example 6.1 (c) at page 42 escription file (c) from Examp another experiment with a m is defined for the following co</model>	and ble 6.1 at page 41. odel named model and there	
dimension	coordinate i	name	coordinate definition	
1	dim1			
2	dim2		equidist_end 2(2)20	
3	dim3		equidist_end 3(3)30	
4	dim4		equidist_end 4(1)43	
5	dim5		equidist_end 5(1)50	
Further, assume	the coordinate t	transformation file model.trf a descr descr	example of a coordinate	
LLI		desci	transformation file	
modify modify modify modify	dim1 dim1 dim1 dim1	rename position_shift values_shift values_add	new1 3. +2 list 1006, 1009	
modify	dim3	values_shift	-3	
assign assign	dim4 dim4	coord coord_extent	lat 88.:-68.	
assign assign	dim5 dim5	coord coord_extent	new2 50.:5.	
coordinate coordinate	new2 new2	descr values	new coordinate equidist_end 50(-1)5	



In SimEnv p	ost-processing	the result of the expression	
get_exper	iment(`mydi	r', `model', `model.trf',mod	var1+modvar2)
is a 5-dimen	sional data stru	ucture with	
dimension	coordinate name	coordinate definition	coordinate use
dimension			
dimension 1 2	name	definition	use
1	name new1	definition list 103,1003,1006,1009	use = coordinate definition
1 2	name new1 dim2	definition list 103,1003,1006,1009 equidist_end 2(2)20	use = coordinate definition = coordinate definition



11.3 Value Lists

For variables, coordinates and experiment targets value lists are supplied by the value-item in user-defined files. Value lists describe a sequence of values together with an order. The number of described values has to be greater than 1. Value lists may be restricted to strictly monotonic sequences. They follow the syntax rules in Tab. 11.6.

Value-list type		Syntax	Explana	ation
explicit	list	<value<sub>1> , , <value<sub>n></value<sub></value<sub>	explicit list of value same syntax rules record of a file with (see below)	as for one
by reference	file	{ <direct>/}<file_name></file_name></direct>	file { <direct>/}<file contains the explic</file </direct>	
implicit with end-element	equidist_end	<beg_val> (<incr_val>) <end_val></end_val></incr_val></beg_val>	description of an e of values with	quidistant list
			begin value increment end value <beg_val> ≠ <end <incr val=""> ≠ 0.</incr></end </beg_val>	<incr_val> <end_val></end_val></incr_val>
implicit with number of values	equidist_nmb	<beg_val> (<incr_val>) <nmb_vals></nmb_vals></incr_val></beg_val>	description of an e of values with begin value increment number of values <beg_val> ≠ <er <incr_val> ≠ 0. <nmb_vals> > 0, ii</nmb_vals></incr_val></er </beg_val>	<pre></pre>

Tab. 11.6Syntax rules for value lists

Syntax rules for a file {<direct>/}<file_name> with a list of values

- Has to be an ASCII file
- May be a multi-record file
- Max. record length is 1000 characters
- Values are separated from each other by white spaces or comma

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- A series of connected (running) separators is treated as a single separator
- Record end is handled as a separator
- Real values can be stated in integer, real or exponential (scientific) format
- Records formed only from white spaces or records starting with the first non-white space character # are handled as comments

```
1. list 3, 5, 7, 9, 11
                                 describes the five values 3, 5, 7, 9, 11
2. equisist end 3 (2) 11
                                 is equivalent to 1.
3. equidist nmb 3 (2) 5
                                 is equivalent to 1.
                                 is equivalent to 1. with my_values.dat = 3, , 5,
4. file my values.dat
                                                                          7
                                                                          9,11
5. equidist end 3 (2) 11.9
                                 is equivalent to 1.
  equidist end 11 (-2) 3
                                 differs from 1. - 4.:
6.
                                 values are identical, ordering sequence differs
```

Example 11.3 Examples of value lists



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12 Model and Post-Processor Output Data Structures

This chapter summarizes information on available data structures for model and post-processor output. SimEnv supports several output formats from the experiment and the post-processor. NetCDF is a selfdescribing data format and can be used for model and post-processor output. Another format specifications for both outputs is IEEE compliant binary format and ASCII for post-processor output. This chapter describes all the used data structures.

Dependent on the specification of the supported post-processor output formats in <model>.cfg model output can be stored in NetCDF format and post-processor output in NetCDF, IEEE or ASCII format.

During experiment performance model output is written either to single output files <model>.out<run_char>. [nc | ascii] per experiment single run or to a common output file <model>.outall.[nc | ieee] for all single runs from the experiment run ensemble. Output to single or a common file(s) depends on specification of the value for the sub-keyword 'out_size_threshold' in <model>.cfg. <run_char> is a six-digit placeholder for the corresponding single run number.

During model output post-processing output and structure of results is written to <model>.res<res_char>.[nc | ieee | ascii] and <model>.res<res_char>.[ieee | ascii]. <res_char> is a two-digit placeholder for the number of the result file. It ranges from 01 to 99.

For IEEE and ASCII model output and post-processor output formats, multi-dimensional data is organized in the Fortran storage model (see Glossary).

Use the SimEnv service command simenv.dmp for browsing model and result output files. See Tab. 10.3 for more information.

12.1 NetCDF Model and Post-Processor Output

The intention for supplying NetCDF format for model and post-processor output is to provide the possibility to generate self-describing, platform-independent data files with metadata that can be interpreted by subsequent visualization techniques. The conventions applied for SimEnv represent a compromise between existing standards and the metadata requirements for a flexible and expressive visualization that is adapted to the requirements of the specific data sets of concern. SimEnv follows the NetCDF Climate and Forecast (NetCDF CF) metadata convention 1.0-beta4. Currently, SimEnv supports only up to 4-dimensional NetCDF output during experiment and post-processor performance.

In principle, any NetCDF file can be viewed by the NetCDF service program ncdump <NetCDF_file>

Model output data types as declared in the model output description file <model>.mdf are transferred into NetCDF data types automatically (check the Table below). By default, post-processor output data is of type float.

SimEnv data type (see Tab. 5.4)	NetCDF data type
byte	NF_BYTE
short	NF_SHORT
int	NF_INT
float	NF_FLOAT
double	NF_DOUBLE

Tab. 12.1NetCDF data types



12.1.1 Global Attributes

The global attributes used in SimEnv from the CF standard are :institution and :convention. In addition, the following global attributes are defined for model and post-processor output:

Name	Value	Data type
:creation_time	<yyyy-mm-dd hh:mm:ss=""></yyyy-mm-dd>	char
:model_name	<model></model>	char
:model_description	model description according to <model>.mdf</model>	char
:model_description_file	{ <direct>/}<model>.mdf</model></direct>	char
:experiment_type	[behaviour monte carlo sensitivity]	char
:experiment_description	experiment description according to <model>.edf</model>	char
:experiment_description_file	{ <direct>/}<model>.edf</model></direct>	char
:number_of_runs	<number of="" runs=""></number>	int

Tab. 12.2Additional global NetCDF attributes

12.1.2 Variable Labelling and Variable Attributes

For coordinate variables, two cases of labelling are distinguished:

- If for a given predefined variable, target, model variable or post-processor result one of its coordinates spans the entire range of its general dimension, the already existing coordinate definition is used.
- Otherwise, this concerned coordinate is re-defined using the notation <variable_name>_dim_<coordinate_name>.

The following variable attributes are used according to the CF 1.0-beta4 standard:

Name	Value	Data type
<variable_name>:standard_name</variable_name>	[<coordinate_name> <predef_coordinate_name> <predef_var_name> <target_name> <variable_name> <result_name>]</result_name></variable_name></target_name></predef_var_name></predef_coordinate_name></coordinate_name>	char
<variable_name>:long_name</variable_name>	[<coordinate_description> <predef_coordinate_description> <predef_variable_description> <target_description> <variable_description> <result_applied_operator_sequence>]</result_applied_operator_sequence></variable_description></target_description></predef_variable_description></predef_coordinate_description></coordinate_description>	char
<variable_name>:missing_value</variable_name>	<variable missing="" type-depending="" value=""></variable>	type-dep.
<pre><variable_name>:axis (single coordinate variables only)</variable_name></pre>	[X Y Z T bin_no run]	char
<pre><variable_name>:unit</variable_name></pre>	[<coordinate_unit> <predef_coordinate_unit> <predef_variable_unit> <target_unit> <variable_unit> <result_unit>]</result_unit></variable_unit></target_unit></predef_variable_unit></predef_coordinate_unit></coordinate_unit>	char
<variable_name>:coordinates (multi-dimensional coordinate variables only)</variable_name>	<par1_lon> <par1_lat></par1_lat></par1_lon>	char
<variable_name>:fill_value</variable_name>	<variable fill="" type-depending="" value=""></variable>	type-dep.

 Tab. 12.3
 Variable NetCDF attributes

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- For post-processor output, the **:standard_name attribute** simply counts the number of applied operations because the result name of an arbitrary operation is not known in general. For that reason, the :long_name attribute would re-sample the :standard_name attribute and it is used instead to provide the complete description of the applied operator sequence without defining an additional attribute. If macros are included, these are resolved and elementary operations are included only.
- For the :axis attribute of a coordinate variable exist defaults.
 For each post-processor result, the first coordinate is assumed to be the "X-axis", the second and third coordinate are assumed to represent the "Y-" and "Z-axis", and the fourth dimension is time T.
 For model results, these attribute values are assigned to coordinate variables describing geographical longitude, geographical latitude, level or height and time. In case other coordinate names are used, these are simply also used for the axis attribute.
- The **:unit attribute** is actually estimated for model output only depending on the description of the corresponding sub-keywords for the keyword 'variable' in the <model>.mdf file. For post-processing output, it is only used as a placeholder and not calculated from the applied operator sequence so far.
- The :coordinates attribute serves to define coordinates depending on other ones and so to allow coordinate transformations. Actually, this attribute is not used.
- Actually, the **:fill_value attribute** is not applied to coordinate variables. It is identically to the :missing_value attribute but open for other definitions.

For visualization requirements, the following additional variable attributes have been defined for SimEnv:

Name	Value	Data type
<variable_name>:monotony (coordinate variables only)</variable_name>	[increasing decreasing none]	char
<variable_name>:coo_type</variable_name>	[1 2]	integer
<variable_name>:data_range</variable_name>	<min> <max></max></min>	char
<variable_name>:index_range_<coordinate> (coordinate variables only)</coordinate></variable_name>	<min_index> <max_index></max_index></min_index>	int
<variable_name>:simenv_data_kind</variable_name>	[predefined model variable model target model output variable postproc_result]	char
<variable_name>:var_representation</variable_name>	[positions connections] or both	char
<variable_name>:grid_shift</variable_name>	<shift_x> <shift_y></shift_y></shift_x>	real, dimension(2)
<variable_name>:north_pole</variable_name>	<lon_pole> <lat_pole></lat_pole></lon_pole>	real, dimension(2)

Tab. 12.4Variable NetCDF attributes for visualization

- The **:monotony attribute** is applied to coordinate variables only and estimated from the coordinate values as defined in the <model>.mdf file. During post-processing additional coordinates can be generated for which no monotony may be estimated. In such cases, the attribute is set to "none".
- The :coo_type attribute describes the grid representation of a given coordinate. A value of 1 indicates that all coordinate values are provided explicitly (suitable, e.g., for irregular grids). A value of 2 indicates a regular grid and a coordinate representation by its start value, increment and end value.
- The :data_range attribute provides the real range that is covered by the related variable in the recent NetCDF file.
- The **:index_range attribute** is used only in case a predefined variable, target, model variable or postprocessing result covers not the complete range of a dimension as defined for a coordinate variable. It describes that sub-space for which the concerned target, variable or result is defined.
- The :var_representation attribute is introduced to specify what operations are allowed on the data.
- The **:grid_shift attribute** is actually still a placeholder for variables that are not defined in the centre of a grid box when quasi-regular grids are used.
- The :north_pole attribute can be used if rotated grids are applied.



12.2 IEEE Compliant Binary Model Output

IEEE compliant binary model output is written in records of fixed length to <model>.out<run_char>.ieee and/or <model>.outall.ieee. Record length is determined by the sub-keyword 'out_ieee_blocksize' and in interrelation to the sub-keyword 'out_size_threshold' in <model>.cfg. For these two sub-keywords and potential modification of the value for 'out_ieee_blocksize' check Tab. 10.1. Sequence of data for each single run is as follows:

- Experiment targets as specified in <model>.edf Sequence as in <model>.edf
- Built-in (pre-defined) model output variables Sequence as in Tab. 10.6
- Model output variables Sequence as in <model>.mdf

Storage demand for each model variable / target is according to its dimensionality, extents and data type. Storage demand in bytes for each model variable / target is readjusted to the smallest number of bytes divisible by 8, where the data can be stored. Multi-dimensional data fields are organized in the Fortran storage model (see Glossary).

In <model>.outall.ieee each single run starts with a new record. Sequence of single runs corresponds with sequence of the single run numbers <run>. Consequently, data from default single run 0 is stored in the first and potentially the following records.

Target / model variable	Extents	Data type	Storage demand [Byte]	Storage demand adjusted [Byte]
p1	1	float	4	8
p2	1	float	4	8
sim time	1	float	4	8
atmo	45 x 90 x 4 x 20	float	1.296.000	1.296.000
bios	36 x 90 x 20	float	259.200	259.200
atmo_g	20	int	80	80
bios_g	1	int	4	8
				1.555.312

Example 12.1 IEEE compliant model output data structure



12.3 IEEE Compliant Binary and ASCII Post-Processor Output

For IEEE and ASCII post-processor output result information is stored in two files:

- <model>.res<res_char>.[ieee | ascii] holds the result dynamics
- <model>.inf<res_char>.[ieee | ascii] holds structure and coordinate information

The IEEE post-processor output files <model>.res<res_char>.ieee and <model>.inf<res_char>.ieee are unformatted binary files with IEEE float / int number representation, while for the ASCII post-processor version <model>.res<res_char>.ascii and <model>.inf<res_char>.ascii formatted ASCII files are used. Files for both output file formats have for each result subsequently the following structure:

Record structure of <model>.inf<res_char>.[ieee | ascii] for each result:

result number 01: record no. 1 record no. 2	max. 512 chars max. 128 chars	result expression string result description string
record no. 3	max. 32 chars	result unit string (or 1 space if unit is undefined)
record no. 4	10 int	dim ext(1) ext(dim) 0 0
record no. 4 record no. 5	max. 20 chars 10 float	coordinate name of dimension 1 coordinate values of dimension 1 in records of 10 values (last record may have less values)
 record no. xxx record no. xxx+1	max. 20 chars 10 float	coordinate name of dimension dim coordinate values of dimension dim in records of 10 values (last record may have less values)
result number 02:		

...

Record structure of <model>.res<res_char>.[ieee | ascii] for each result:

result number 01: record no. 1	10 float	in records of 10 values (last record	d may have less values).
	10 hoat	result_value(1) result_valu	
		dim	
		with length_result = $\prod ext(i)$	for dim > 0
		i=1	
		= 1	else
result number 02:			

result number 02:

The vector result_value is stored in the Fortran storage model (see Glossary). The nodata element for undefined result values is set to 3.4E38.

The Fortran code in Example 15.8 reads post-processing ASCII output files <model>.res<res_char>.ascii and <model>.inf<res_char>.ascii in their general structure. In the examples-directory of SimEnv it is accompanied by the corresponding version for IEEE result output.



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13 SimEnv Prospects

SimEnv development and improvement is user-driven. Here you can find a list of the main development pathways in future.

General

- Graphical user interface
- Linux and Windows portability
- Unique number representations for binary output of distributed models (big endians vs. small endians)

Model interface

Experiment preparation

• Experiment type stochastic analysis

Experiment performance

- Experiment performance for distributed models across networks
- Multi-file model output storage

Experiment post-processing

- Additional advanced operators (netcdf_data, regrid, coarse, sort, ...)
- Advanced uncertainty and global sensitivity analyses operators
- C-interface for user-defined operators
- Flexible assignment of data types to operator results (currently: only float)
- Shared memory access for user-defined operators to avoid data exchange by external files
- Wrapping of pure C/C++-operators in Fortran to use them as built-in operators

Experiment evaluation

 Advanced techniques for graphical representation of post-processor output, especially for multi-run operators



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14 References

Bohr, J. (1998) A summary on Probabilities http://ic.net/~jnbohr/java/CdfDemoMain.html European Commission, Joint Research Centre – IPSC (2004): SimLab 2.2 Reference Manual http://www.jrc.ce.eu.int/uasa/primer-sa.asp Flechsig, M. (1998) SPRINT-S: A Parallelization Tool for Experiments with Simulation Models. PIK-Report No. 47, Potsdam Institute for Climate Impact Research, Potsdam http://www.pik-potsdam.de/reports/pr-47/pr47.pdf Flechsig, M., Böhm, U., Nocke, T., Rachimow, C. (2004): Techniques for Quality Assurance of Models in a Multi-Run Simulation Environment. 4th International Conference on Sensitivity Analysis of Model Output SAMO 2004, March 8-11, 2004, Santa Fe, U.S.A. http://www.pik-potsdam.de/~flechsig/simenv/simenv poster samo2004.pdf Gray, P., Hart, W., Painton, L., Phillips, C., Trahan, M., Wagner, J. (1997) A Survey of Global Optimization Methods. Sandia National Laboratories, Albuquergue, U.S.A. http://www.cs.sandia.gov/opt/survey Helton, J.C., Davis, F.J. (2000): Sampling-Based Methods. In: Saltelli et.al (2000) Imam, R.L., Helton, J.C. (1998): An Investigation of Uncertainty and Sensitivity Analysis Techniques for Computer Models, Risk Anal, 8(1), 71-90 Ingber, L. (1989): Very fast simulated re-annealing. Math. Comput. Modelling, 12(8), 967 – 973 http://www.ingber.com/asa89 vfsr.pdf Ingber, L. (1996) Adaptive simulated annealing (ASA): Lessons learned. Control and Cybernetics, 25(1), 33-54 http://www.ingber.com/asa96 lessons.pdf Ingber, L. (2004) ASA-Readme. http://www.inger.com/ASA-README.pdf Saltelli, A., Chan, K., Scott, E.M. (eds.) (2000) Sensitivity Analysis. J. Wiley & Sons, Chichester Saltelli, A., Tarantola, S., Campolongo, F., Ratto, M. (2004) Sensitivity Analysis in Practice: A Guide to Assessing Scientific Models. J. Wiley & Sons, Chichester Waszkewitz, J., Lenzen, P., Gillet, N. (2001) The PINGO package: Procedural interface for Grib formatted objects. Max-Planck-Institute for Meteorology, Hamburg and http://www.mad.zmaw.de/Pingo/pingohome.html Wenzel, V., Kücken, M., Flechsig, M. (1995) MOSES - Modellierung und Simulation ökologischer Systeme.

PIK-Report No. 13, Potsdam Institute for Climate Impact Research, Potsdam

http://www.pik-potsdam.de/pik_web/publications/pik_reports/sum_pr13.htm

Wenzel, V., Matthäus, E., Flechsig, M. (1990) One Decade of SONCHES. Syst. Anal. Mod. and Sim. 7, 411-428

Wierzbicki, A.P. (1984) Models and Sensitivity of Control Systems. Studies in Automation and Control. Vol. 5. Elsevier, Amsterdam



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15 Appendices

The appendices summarize the current version implementation, list the examples for model interfaces, user-defined operators and result import interfaces, and they compile all post-processor built-in operators. Finally, a glossary of the main terms as used in this User Guide is supplied.



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15.1 Version Implementation

15.1.1 Example Models and User Files

For the following models corresponding files of Tab. 10.4 of can be copied from the corresponding examples-directory of \$SE_HOME to the user's working directory by running the SimEnv command simenv.cpy <model> from the working directory:

model	Language / source code	Explanation
world_f	Fortran world_f.f	global atmosphere - biosphere model at resolution of (lat x lon x level x time) = (45 x 90 x 4 x 20)
world_c	C world_c.c	
world_cpp	C++ world_cpp.cpp	
world_py	Python world_py.py	
world_sh	Shell script level world_sh.f world_shput.f	
world_f_1x1	Fortran world_f_1x1.f	global atmosphere - biosphere model at a resolution of (lat x lon x level x time) = (180 x 360 x 16 x 20)
pixel_f	Fortran Pixel_f.f	global atmosphere - biosphere model for one lat-lon grid cell at a resolution of (level x time) = (4 x 20)
gams_model	GAMS gams_model.gms	GAMS example model

Tab. 15.1Implemented models for current version
for model = world_* check also Example 1.1

Additionally, the following files are available in the corresponding examples directory of \$SE_HOME:

File	Explanation		
<model>.[f c cpp py gms]</model>	model source code (check also example files in chapter 15.2)		
<model></model>	model executable compiled and linked from <model>.[f c cpp]</model>		
world.edf_[a b c d e f]	experiment description files corresponding to Example 6.1, Example 6.2, and Example 6.3 to be copied to world_[f c cpp py sh].edf and/or world f 1x1.edf		
world.post_[c e f bas adv]	<pre>post-processor input file (complete experiment) for world.edf_[c e f] (simenv.res world_[f c cpp py sh] [new append replace]</pre>		
world.dat_[d e tab]	data files for world.edf_[d e] and/or world.post_adv		

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File	Explanation		
usr_opr_ <opr>.f</opr>	source code for user-defined operator <opr></opr>		
<opr>.opr</opr>	executable for user-defined operator <opr></opr>		
model_[f c cpp].lnk <model></model>	compile <model>.[f c cpp] and link to executable <model></model></model>		
usr_opr_ <opr>.f</opr>	source code file for user-defined post-processing operator <opr></opr>		
operator_f.lnk <opr> compile usr_opr_<opr>.f and link it to executable <opr>.o</opr></opr></opr>			
	defined post-processing operator <opr></opr>		
land_sea_mask[<nil> .f]</nil>	executable and source code to derive a coarsed land-sea-mask from the		
	file land_sea_mask.05x05		
land_sea_mask.05x05	global ASCII land-sea-mask file with a resolution of 0.5° lat x 0.5° lon		
read_result_file[<nil> .f]</nil>	executable and source code for the result file import interface of ASCII		
	and IEEE compliant result output		

Tab. 15.2Implemented model-related user files for current versionFor <opr>> see Tab.15.3 below

15.1.2 Example User-Defined Operators

The following user-defined operators are available from the corresponding examples directory of \$SE_HOME:

Name	Explanation / restriction	Example		
char_test('char1','char2',arg)	character test	see source code		
corr_coeff(arg1,arg2)	correlation coefficient R	<pre>corr_coeff(bios,-bios) = -1.</pre>		
div(arg1,arg2)	division as an example how the cor- responding built in basic operator works	div(-2, -4) = 0.5		
mat_mul	matrix multiplication of 2- dimen- sional operands	<pre>mat_mul(mat1,mat2)</pre>		
simple_div(arg1,arg2)	division without consideration of overflow, underflow, division by zero	$simple_div(-2,-4) = 0.5$		

Tab. 15.3Available user-defined operators

15.1.3 Linking User Models and User-Defined Operators

- User models implemented in C/C++ or Fortran have to be linked with the following libraries to couple them to the simulation environment
 - \$SE_HOME/libsimenv.a
 - /usr/local/lib/libnetcdf.a
- User-defined operators to be used in result post-processing have to be linked with the following library to couple them to the simulation environment
 - \$SE_HOME/libsimenv.a

For running interfaced models outside SimEnv check chapter 5.9.

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15.1.4 Technical Limitations

Entity		Limitation
User-defined files entities (check also ch	napter 11.1)	
max. length of a record in a user-defined file	[characters]	160
max. length of all global descriptions descr	[characters]	512
max. length of a local description descr	[characters]	128
max. length of a unit	[characters]	32
max. length of a { <direct>/}<file_name> string</file_name></direct>	[characters]	70
max. length of a record of a referred data file	[characters]	1000
Model interface and experiment preparat	tion entities	
max. length of a name	[characters]	20
max. dimensionality of a model output variable		9
max. dimensionality of a model output variable for Python mod	els	4
max. dimensionality of a model output variable for GAMS model	els	4
max. dimensionality of a model output variable stored in NetCI	DF format	4
max. number of model output variables in <model>.mdf</model>		50
max. number of coordinates in <model>.mdf</model>		30
max. number of experiment targets in <model>.edf</model>		50
max. number of slice definitions during interfacing a model		30
max. number of single model runs in an experiment		999.999 200.000
max. number of coordinate values and target adjustment values		
Post-processing entities (per res	ult)	
max. length of the optional result description	[characters]	128
max. length of the optional result unit	[characters]	32
max. number of arguments of an operator		9
max. dimensionality of a result stored in NetCDF format		4
max. number of post-processor output files		99
max. number of characters of a result		512
max. number of all operands and operators of a result		200
max. length of a constant	[characters]	20
max. number of constants		30
max. number of user-defined operators in <model>.odf</model>		45
max. number of allocatable main memory segments		10
max. allocatable main memory	[MBytes]	240

Tab. 15.4

Current SimEnv technical limitations



15.2 Examples for Model Interfaces, User-Defined Operators, and Result Import Interfaces

15.2.1 Example Implementations for Model world

According to Example 1.1 at page 4 dynamics of the model world depend on four model paramters p1, p2, p3, and p4:

Mod targe	tluctob	Internal model parameter	Unit	Meaning
p1	1.	phi_lat	π/12	latitudinal phase shift
p2	2.	omega_lat	2*π	latitudinal frequency
р3	3.	phi_lon	π/12	longitudinal phase shift
p4	4.	omega_lon	2*π	longitudinal frequency

 Tab. 15.5
 Parameters for the model world

 Mapping between model targets and internal model parameters is performed by simenv_get_*

For reasons of simplification these parameters influence state variables atmo and bios by the product of two trigonometric terms value_lat and value_lon in the following manner:

value_lat(lat)	= sin(2*п*omega_lat	* f(lat)	+ phi_lat*n/12)
value_lon(lon)	= sin(2*π*omega_lon	* f(lon)	+ phi_lon*π/12)

The function f(.) norms value_lat and value_lon by lat and/or lon in a way, that holds

Finally,

```
atmo(lat,lon,level,time) = value_lat(lat) * value_lon(lon) * (100*time+level)
bios(lat,lon,time) = value_lat(lat) * value_lon(lon) * 100*time
```

and

```
atmo_g(time) = avg_l('001',abs(atmo(lat,lon,1,time)))
bios_g = avg(abs(bios(lat,lon,time)))
```

Mean avg and avg_l are calculated in a box around (lat,lon) = $(0^{\circ}, 0^{\circ})$.

15.2.2 Fortran Model

With respect to Example 5.1 the following Fortran code **world_f.f** could be used to describe the model coupled to SimEnv. SimEnv modifications are marked in **bold**.

```
program world f
c declare SimEnv coupling functions
  integer*4
             simenv_ini_f, simenv_get_f, simenv_get_run_f
  integer*4
              simenv_slice_f, simenv_put_f, simenv_end_f
c declare atmo without dimensions level and time and bios without time
c because they are computed in place and simenv slice is used
  real*4 atmo(0:44,0:89)
              bios(0:35,0:89)
  real*4
  integer*4 atmo g(0:19)
  integer*4
             bios g
   integer*4
              run int
   character*6 run char
  istatus = simenv ini f()
c check return code for the model coupling functions at least here
  if(istatus.ne.0) call exit (1)
c only if necessary:
  istatus = simenv get run(run int,run char)
  p1 = 1.
  p2 = 2.
  p3 = 3.
  p4 = 4.
  istatus = simenv get f(`p1',p1,p1)
  istatus = simenv get f(`p2',p2,p2)
  istatus = simenv_get_f(`p3',p3,p3)
  istatus = simenv get f(`p4',p4,p4)
c compute dynamics of atmo and bios over space and time,
c of atmo g over time, all dependent on p1,p2,p3,p4
  do idecade = 0, 19
      do level= 0, 3
         istatus = simenv_slice_f(`atmo',3,level,level)
         istatus = simenv slice f('atmo',4,idecade,idecade)
         istatus = simenv put f(`atmo',atmo)
      enddo
      istatus = simenv slice f('bios',3,idecade,idecade)
      istatus = simenv put f(`bios',bios)
   enddo
   . . .
   istatus = simenv put f(`atmo g',atmo g)
c compute dynamics of bios g
   . . .
   istatus = simenv_put_f(`bios_g',bios_g)
   istatus = simenv end f()
   end
                                                         Example file: world f.f
```

Example 15.1 Model interface for Fortran models - model world_f.f

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15.2.3 C Model

With respect to Example 5.1 the following C code **world_c.c** could be used to describe the model coupled to SimEnv. SimEnv modifications are marked in **bold**.

```
#include <stdio.h>
#include <stdlib.h>
#include <string.h>
/* declare SimEnv coupling functions (compile with -I$SE HOME) */
#include <simenv.h>
/* declare atmo without dimensions level and time and bios without time*/
/* because they are computed in place and simenv slice is used */
static float atmo[45][90];
static float bios[36][90];
static int
             atmo g[20];
static int
              bios g;
main(void)
{
   float p1,p2,p3,p4;
   int run int;
   char run char[6];
  int level, idecade, istatus, idim;
  istatus = simenv ini c();
/* check return code of model coupling functions at least here */
  if(istatus != 0) return 1;
/* only if necessary: */
  istatus = simenv get run c(&run int,run char);
  p1 = 1.;
  p2 = 2.;
  p3 = 3.;
  p4 = 4.;
  istatus = simenv_get_c(`p1',&p1,&p1);
  istatus = simenv get c('p2',&p2,&p2);
  istatus = simenv_get_c(`p3',&p3,&p3);
  istatus = simenv_get_c(`p4',&p4,&p4);
/* compute dynamics of atmo and bios over space and time, */
/* of atmo g over time, all dependent on p1,p2,p3,p4 */
   for (idecade=0; idecade<=19; idecade++)</pre>
   {...
     for (level=0; level<=3; level++)</pre>
     { . . .
       idim=3;
       istatus = simenv slice c('atmo',&idim,&level,&level);
       idim=4;
       istatus = simenv slice c(`atmo',&idim,&idecade,&idecade);
       istatus = simenv_put_c('atmo',(char *) &atmo);
     ļ
     idim=3;
     istatus = simenv slice c(`bios',&idim,&idecade,&idecade);
     istatus = simenv put c('bios',(char *) &bios);
   }
   istatus = simenv put c('atmo g',(char *) &atmo g);
```

```
/* compute dynamics of bios_g */
...
istatus = simenv_put_c(`bios_g', ,(char *) &bios_g);
istatus = simenv_end_c();
return 0;
}
Example file: world_c.c
```

Example 15.2 Model interface for C models – model world_c.c



15.2.4 C++ Model

With respect to Example 5.1 the following C++ code **world_cpp.cpp** could be used to describe the model coupled to SimEnv. SimEnv modifications are marked in **bold**.

```
#include <stdio.h>
#include <stdlib.h>
/* declare SimEnv coupling functions (compile with -I$SE HOME) */
#include <simenv.h>
class World
{
/* declare atmo without dimensions level and time and bios without time*/
/* because they are computed in place and simenv slice is used */
  public: float atmo[45][90];
  public: float bios[36][90];
  public: int
                    atmo g[20];
  public: int
                    bios g;
  private: int
                    level, idecade, istatus, idim;
  public: void computeAtmo(float p1, float p2, float p3, float p4)
/* compute dynamics of atmo over space and time, */
/* and of atmo g over time, all dependent on p1,p2,p3,p4 */
   {
     for (idecade=0; idecade<=19; idecade++)</pre>
     {...
       for (level=0; level<=3; level++)</pre>
       {...
         idim=3;
         istatus = simenv slice c('atmo',&idim,&level,&level);
         idim=4;
         istatus = simenv slice c(`atmo',&idim,&idecade,&idecade);
         istatus = simenv put c('atmo', (char *) &atmo);
      }
     }
   }
public: void computeBios(float p1, float p2, float p3, float p4)
/* compute dynamics of bios over space and time, */
/* and of bios g all dependent on p1,p2,p3,p4 */
     for (idecade=0; idecade<=19; idecade++)</pre>
     {...
       idim=3:
       istatus = simenv slice c(`bios',&idim,&idecade,&idecade);
       istatus = simenv_put_c(`bios',(char *) &bios);
     }
/* compute dynamics of bios g */
     . . .
   }
}
```

```
main(void)
{
   int run int, istatus;
   char run char[6];
   istatus = simenv_ini_c();
/* check return code of model coupling functions at least here */
   if(istatus != 0) return 1;
/* only if necessary: */
  istatus = simenv_get_run_c(&run_int,run_char);
   float p1 = 1., float p2 = 2., float p3 = 3., float p4 = 4.;
   istatus = simenv_get_c(`p1',&p1,&p1);
   istatus = simenv_get_c(`p2',&p2,&p2);
   istatus = simenv_get_c(`p3',&p3,&p3);
   istatus = simenv_get_c(`p4',&p4,&p4);
  World world;
  world.computeAtmo(p1,p2,p3,p4);
   istatus = simenv_put_c(`atmo_g',(char *) &(world.atmo_g));
  world.computeBios(p1,p2,p3,p4);
  istatus = simenv_put_c('bios_g',(char *) &(world.bios_g));
  istatus = simenv_end_c();
  return 0;
}
                                                      Example file: world_cpp.cpp
```

Example 15.3 Model interface for C++ models – model world_cpp.cpp



15.2.5 Python Model

With respect to Example 5.1 the following Python code **world_py.py** could be used to describe the model coupled to SimEnv. SimEnv modifications are marked in **bold**.

```
#!/usr/local/bin/python
import string
import os
from simenv import *
from math import *
from Numeric import *
atmo=zeros([45,90,4,20], Float)
bios=zeros([36,90,20], Float)
atmo g=zeros([20], Float)
simenv_ini_py()
# only if necessary:
run_int = int(simenv_get_run_py())
p1=1.
p2=2.
p3=3.
p4=4.
p1 = float(simenv_get_py('p1',p1))
p2 = float(simenv_get_py('p2',p2))
p3 = float(simenv_get_py('p3',p3))
p4 = float(simenv_get_py('p4',p4))
# compute dynamics of atmo and bios over space and time,
# of atmo g over time, all dependent on p1,p2,p3,p4
for idecade in range(20):
• • •
  for level in range(4):
   . . .
atmo=reshape(atmo, 45*90*4*20,))
simenv_put_py(`atmo',atmo)
bios=reshape(atmo, 45*90*20,))
simenv_put_py(`bios',bios)
simenv put py(`atmo g',atmo g)
# compute dynamics of bios g
# ...
simenv put py(`bios g',bios g)
simenv end py()
                                                         Example file: world_py.py
```

Example 15.4 Model interface for Python models – model world_py.py

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15.2.6 Model Interface at Shell Script Level

Assume any experiment. Assume model executable world_sh to target values p1 to p4 as arguments from the command line.

The shell script **world_sh.run** with an interface at shell script level to run the model world_sh and to transform model output to SimEnv could look like:

```
# always perform at begin
. $SE HOME/simenv ini sh
# create temporary directory run<run char> to perform the model
# and model output transformation from native to SimEnv structure there
. $SE HOME/simenv get run sh
mkdir run$run char
cd run$run char
# get adjustments for p1 ... p4
target name='p1'
target def val=1.
. $SE HOME/simenv get sh
target name='p2'
target_def_val=2.
. $SE HOME/simenv get sh
target name='p3'
target def val=3.
. $SE HOME/simenv get sh
target name='p4'
target def val=4.
. $SE HOME/simenv get sh
# run the model
cp ../land sea mask.coarsed .
../world sh $p1 $p2 $p3 $p4
# read model results and output them to SimEnv
../world shput
# clear and remove directory
rm -f *
cd ..
rmdir run$run char
# always perform at end
. $SE HOME/simenv end sh
                                                       Example file: world_sh.run
```

Example 15.5 Model interface at shell script level – model shell script world_sh.run



15.2.7 GAMS Model

The SimEnv version comes with a coupled GAMS model **gams_model.gms** and all associated files that fully correspond with the GAMS example model at http://www.gams.com/docs/gams/Tutorial.pdf. Modifications for SimEnv are marked in **bold**.

```
SETS
        canning plants / SEATTLE, SAN-DIEGO /
  Ι
  J
        markets
                         / NEW-YORK, CHICAGO, TOPEKA / ;
PARAMETERS
  A(I) capacity of plant i in cases
    /
         SEATTLE
                     350
         SAN-DIEGO
                     600
  B(J) demand at market j in cases
         NEW-YORK 325
     /
         CHICAGO
                     300
         TOPEKA
                     275 / ;
* - Before using parameter (here: dem ny and dem ch) as SimEnv experiment
   targets they have to be declared as GAMS model parameters
   default values from above.
* - Then insert $include <model> simenv get.inc
  simenv get.inc is generated automatically based on <model>.edf
* - and assign adjusted targets to model variables
  PARAMETERS
  dem ny /325.0/;
  dem ch /300.0/;
  $include gams_model_simenv_get.inc
  A("SEATTLE") = dem_ny;
  A("SAN-DIEGO") = dem ch;
TABLE D(I,J) distance in thousands of miles
               NEW-YORK CHICAGO TOPEKA
  SEATTLE
                  2.5
                               1.7
                                            1.8
  SAN-DIEGO
                   2.5
                                1.8
                                             1.4 ;
SCALAR F freight in dollars per case per thousand miles /90/
* get the model status as a model output
  modstat is set to transport.modelstat ;
PARAMETER C(I,J) transport cost in thousands of dollars per case ;
  C(I,J) = F * D(I,J) / 1000;
VARIABLES
  X(I,J) shipment quantities in cases
   Ζ
          total transportation costs in thousands of dollars ;
POSITIVE VARIABLE X ;
EOUATIONS
             define objective function
  COST
   SUPPLY(I) observe supply limit at plant i
  DEMAND(J) satisfy demand at market j ;
             Z = E = SUM((I,J), C(I,J) * X(I,J));
COST ..
             SUM(J, X(I,J)) = L = A(I);
SUPPLY(I) ..
DEMAND(J) .. SUM(I, X(I, J)) =G= B(J);
MODEL TRANSPORT /ALL/ ;
SOLVE TRANSPORT USING LP MINIMIZING Z ;
```

```
* After solving the equations $include simenv_put.inc
* has to be inserted.
* simenv_put.inc is generated automatically by SimEnv
* based on <model>.edf and <model>.gdf
* Additional GAMS statements are possible after the $include statement
modstat = transport.modelstat
$include gams_model_simenv_put.inc
* Only if sub-models sub_m1 and sub_m2 are coupled (see Example 5.5):
* $call "gams ../sub_m1.gms ll= lo=2 lf=gams_model.nlog dp=0";
* $call "gams ../sub_m2.gms ll= lo=2 lf=gams_model.nlog dp=0";
```

Example 15.6 Model interface for GAMS models – model gams_model.gms



15.2.8 Post-Processor User-Defined Operator

Implementation of the user-defined operator mat_mul in the file usr_opr_mat_mul.f:

```
function icheck user def operator()
  declare fields to hold extents and coordinates
С
   dimension iext1(9), iext2(9)
   dimension ico blk1(9), ico blk2(9)
   dimension ico beg1(9), ico beg2(9)
c get dimensionality idimens, extents iext,
c coordinate block number ico blk and
c begin number for coordinates ico_beg in coordinate block ico_blk
  idimens1=iget dim arg(1,iext1)
   idimens2=iget dim arg(2,iext2)
   iok=iget co arg(1,ico blk1,ico beg1)
  iok=iget co arg(2,ico blk2,ico beg2)
c get check modus for coordinates
   ichk modus=iget co chk modus()
   if (idimens1.ne.2.or.idimens2.ne.2) then
 wrong dimensionalities
С
      ierror=1
   else
      if(iext1(2).ne.iext2(1)) then
С
  wrong extents
         ierror=2
      else
         if(ico blk1(2).eq.ico blk2(1)) then
С
  coordinates identical
            if (ico beg1(2).eq.ico beg2(1)) then
               iret=31
            else
               iret=33
            endif
         else
С
   differing coordinates
            iret=32
            if (ichk modus.eq.1) then
  check only for weak coordinate
С
               do j=0, iext1(2)-1
  get coordinate values
С
                  iretv1=iget_coord_val(
     #
                         ico blk1(2),ico beg1(2)+j,value1)
                  iretv2=iget_coord_val(
     #
                         ico blk2(1),ico beg2(1)+j,value2)
С
  iret=33: differing coordinate values
                  if(value1.ne.value2) iret=33
               enddo
            endif
         endif
         ierror=0
         if(ichk modus.eq.2) then
            if(iret.gt.31) ierror=3
         elseif(ichk modus.eq.1) then
            if(iret.gt.32) ierror=3
         endif
      endif
   endif
```

```
if(ierror.eq.0) then
     iext1(2)=iext2(2)
     ico blk1(2)=ico blk2(2)
     ico beg1(2)=ico beg2(2)
     iok=iput_struct_res(0,idimens1,iext1,ico_blk1,ico_beg1)
   endif
c return error code
   icheck user def operator=ierror
   return
   end
   function icompute_user_def_operator(res)
c SimEnv operator results are always of type real*4
  real*4 res(1)
c auxiliary variables
   integer*4 iext1(9),iext2(9)
  real*8 value8
c get dimensionality idimens and extents iext for both arguments
   idimens=iget dim arg(1,iext1)
   idimens=iget dim arg(2,iext2)
c perform matrix multiplication
  m=0
   do k=1, iext2(2)
      ioffs2=(k-1) *iext2(1)
      do i=1, iext1(1)
         ioffs1=i
   res(i,k) = sum(arg1(i,l) * arg2(l,k))
С
         value8=0.
         indi defined=0
         do l=1, iext1(2)
            ial=ioffs1+(l-1)*iext1(1)
            ia2=ioffs2+1
            fac1=arg1(ia1)
            fac2=arg2(ia2)
            if(is undef(fac1)+is undef(fac2).eq.0) then
               indi defined=1
               value8=value8+fac1*fac2
            endif
         enddo
         m=m+1
         if(indi defined.eq.0) then
            res(m) = set undef()
         else
            res(m)=clip undef(value8)
         endif
      enddo
   enddo
 return error code
С
  icompute user def operator=0
  return
  end
                                                   Example file: usr_opr_mat_mul.f
```

Example 15.7 Post-processor user-defined operator module – operator mat_mul



15.2.9 Post-Processor Result Import Interface

In Example 15.8 an implementation of an interface to import ASCII post-processor output from SimEnv can be found. A corresponding interface to import IEEE compliant post-processor output is documented as the file read_default_file_ieee.f.

```
subroutine read result file ascii(model name, res nmb)
   character model name*20, res nmb*2
   real*4, pointer, dimension(:) :: coord values
   real*4, pointer, dimension(:) :: result values
   integer*4 idim, iext(9)
   character result expr*512, result desc*128, result unit*32
   character coord name*20
   open(unit=1,file=trim(model name)//`inf'//res nmb//`.ascii',
 #
        form=`formatted', status=`old')
   open(unit=2,file=trim(model name)//`res'//res nmb//`.ascii',
        form=`formatted',status=`old')
 #
   iostat=0
   do while (iostat.eq.0)
      read(1, `(a512)',iostat=iostat) result_expr
      if(iostat.eq.0) then
         read(1, '(a128)') result desc
         read(1, '(a32)') result unit
         read(1, '(10i8)') idim, (iext(i), i=1, 9)
         length result=1
         do i=1,idim
            length_result=length_result*iext(i)
            read(1, '(a20)') coord name
            allocate(coord_values(iext(i)))
            ibeg=1
            do while (ibeg.le.iext(i))
                iend=min0(ibeg+9,iext(i))
                read(1, '(10g12.6)') (coord values(j),j=ibeg,iend)
                ibeg=iend+1
            enddo
            further processing of coordinate values
С
С
             . . .
            deallocate (coord values)
         enddo
         allocate(result values(length result))
         ibeg=1
         do while (ibeg.le.length result)
            iend=min0(ibeg+9,length result)
             read(2, '(10g12.6)') (result values(j),j=ibeg,iend)
            ibeg=iend+1
         enddo
         further processing of result values
С
С
         deallocate (result values)
      endif
   enddo
   close(unit=1)
   close(unit=2)
   return
   end
                                                      Example file: read result file.f
                                         together with subroutine read_result_file_ieee
```

Example 15.8 ASCII compliant post-processor result import interface



15.3 Compilation of Post-Processor Built-In Operators and Operator Arguments

15.3.1 Post-Processor Built-In Operators (in Thematic Order)

arg	general numerical argument
const_arg	constant argument
char_arg	character argument

Name	Meaning	See Tab. at page
Elemental operators		66
arg1 + arg2	addition	
arg1 - arg2	subtraction	
arg1 * arg2	multiplication	
arg1 / arg2	division	
arg1 ** arg2	exponentiation	
+ arg	identity	
- arg	negation	
(arg)	parentheses	
Basi	c operators	67
abs(arg)	absolute value	
dim(arg1,arg2)	positive difference	
exp(arg)	exponential function	
int(arg)	truncation value	
log(arg)	natural logarithm	
log10(arg)	decade logarithm	
mod(arg1,arg2)	remainder	
nint(arg)	round value	
sign(arg)	sign of value	
sqrt(arg)	square root	
	onometric operators	67
sin(arg)	sine	
cos(arg)	cosine	
tan(arg)	tangent	
cot(arg)	cotangent	
asin(arg)	arc sine	
acos(arg)	arc cosine	
atan(arg)	arc tangent	
acot(arg)	arc cotangent	
sinh(arg)	hyperbolic sine	
cosh(arg)	hyperbolic cosine	
tanh(arg)	hyperbolic tangent	
coth(arg)	hyperbolic cotangent	

Name	Meaning	See Tab. at page	
Advanced	operators	72	
classify(const_arg1, const_arg2,const_arg3,arg)	classification of arg into const_arg1 classes		
clip(char_arg,arg)	clip arg according to char_arg		
cumul(char_arg,arg)	cumulates arg according to char_arg		
flip(char_arg,arg)	flip arg according to char_arg		
get_experiment(char_arg1, char_arg2,char_arg3,arg)	include an other experiment		
get_table_fct(char_arg,arg)	table function with linear interpolation of table char_a	rg for position arg	
if(char_arg,arg1,arg2,arg3)	general purpose conditional if-construct		
mask(char_arg,arg1,arg2)	mask elements of argument arg1		
matmul(arg1,arg2)	matrix multiplication		
nr_of_runs()	number of single runs of the current experiment		
rank(char_arg,arg)	rank of arg according to char_arg		
run(char_arg,arg)	values of arg for a single run selected by char_arg		
transpose(char_arg,arg)	transpose arg according to char_arg		
undef()	undefined element		
	n and moment operators for arguments	69	
avg(arg)	argument arithmetic mean of values		
avgg(arg)	argument geometric mean of values		
avgh(arg)	argument harmonic mean of values		
avgw(arg1,arg2)	argument weighted mean of values		
count(char_arg,arg)	count number of values		
hgr(char_arg,const_arg1, const_arg2,const_arg3, arg)	argument histogram of values		
max(arg)	argument maximum of values		
maxprop(arg)	index of the element where the maximum is reached	the first time	
min(arg)	argument minimum of values		
minprop(arg)	index of the element where the minimum is reached t	he first time	
sum(arg)	argument sum of values		
var(arg)	argument variance of values		
Multiple ag	gregation and moment operators for arguments	69	
max_n(arg1,,argn)	maximum per element		
maxprop_n(arg1,,argn)	argument position (1 n) where the maximum is rea	ched the first time	
min_n(arg1,,argn)	minimum per element		
minprop_n(arg1,,argn)	argument position (1 n) where the minimum is read	ched the first time	
Dimension	-related aggregation and moment operators for arg	juments 70	
avg_l(char_arg,arg)	dimension-related argument arithmetic means of valu	ies	
avgg_l(char_arg,arg)	dimension-related argument geometric means of valu	les	
avgh_l(char_arg,arg)	dimension-related argument harmonic means of value		
avgw_l(char_arg,arg1,arg2)	dimension-related argument weighted means of value		
count_l(char_arg1,char_arg2, arg)	dimension-related count numbers of values		
hgr_l(char_arg1,char_arg2, const_arg1,const_arg2, const_arg3,arg)	dimension-related argument histograms of values		
max_l(char_arg,arg)	dimension-related argument maxima of values		
max_i(chai_aig,aig) maxprop_l(char_arg,arg)	dimension-related argument position (1 n) where the	ne maximum is	
	reached the first time		

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Name	Meaning	See Tab. at page	
min_l(char_arg,arg)	dimension-related argument minima of values		
minprop_l(char_arg,arg)	dimension-related argument position (1 n) where t reached the first time	he minimum is	
sum_l(char_arg,arg)	dimension-related argument sums of values		
var_l(char_arg,arg)	dimension-related argument variances of values		
Multi-run o	perators (behavioural analysis)	79	
behav(char_arg,arg)	general purpose operator for navigating and aggrega space	ating in the experiment	
Multi-run o	perators (Monte Carlo analysis and optimization)	82	
avg_e(arg)	run ensemble mean		
avgg_e(arg)	run ensemble geometric mean		
avgh_e(arg)	run ensemble harmonic mean		
avgw_e(arg1,arg2)	run ensemble weighted mean		
cnf(const_arg,arg)	positive distance of confidence line from mean avg_e	e(arg)	
cor(arg1,arg2)	correlation coefficient between arg1 and arg2		
count_e(char_arg,arg)	run ensemble count number of values		
cov(arg1,arg2)	covariance between arg1 and arg2		
ens(arg)	whole Monte Carlo run ensemble		
hgr_e(char_arg1,const_arg1, const_arg2,const_arg3,arg)	heuristic probability density function		
krt(arg)	kurtosis (4 th moment)		
max_e(arg)	run ensemble maximum		
maxprop_e(arg)	run number where the maximum is reached the first	time	
med(arg)	median		
min_e(arg)	run ensemble minimum		
minprop_e(arg)	run number where the minimum is reached the first ti	ime	
qnt(const_arg,arg)	quantile of arg		
reg(arg1,arg2)	linear regression coefficient to forecast arg2 from arg	1	
rng(arg)	range = max_e(arg) - min_e(arg)	, ,	
skw(arg)	skewness (3 rd moment)		
stat_full(const_arg1, const_arg2,const_arg3, const_arg4,arg5)	basic statistical summaries		
stat_red(const_arg1, const_arg2,arg3)	basic statistical summaries		
sum_e(arg)	run ensemble sum		
var_e(arg)	run ensemble variance		
	perators (local sensitivity analysis)	84	
lin_abs(char_arg,arg)	absolute linearity measure		
lin_rel(char_arg,arg)	relative linearity measure		
sens_abs(char_arg,arg)	absolute sensitivity measure		
sens_rel(char_arg,arg)	relative sensitivity measure		
sym_abs(char_arg,arg)	absolute symmetry measure		
sym_abs(char_arg,arg) sym_rel(char_arg,arg)	relative symmetry measure		
sym_rei(unai_aiy,aiy)	i ciauve symmetry measure		

 Tab. 15.6
 Post-processor built-in operators (in thematic order)

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15.3.2 Post-Processor Built-In Operators (in Alphabetic Order)

arggeneral numerical argumentconst_argconstant argumentchar_argcharacter argument

Name	Name Meaning		See Tab. at page	
arg1 + arg2	addition	elemental	66	
arg1 - arg2	subtraction	elemental	66	
arg1 * arg2	multiplication	elemental	66	
arg1 / arg2	division	elemental	66	
arg1 ** arg2	exponentiation	elemental	66	
+ arg	identity	elemental	66	
- arg	negation	elemental	66	
(arg)	parentheses	elemental	66	
abs(arg)	absolute value	basic	67	
acos(arg)	arc cosine	trigonom.	67	
acot(arg)	arc cotangent	trigonom.	67	
asin(arg)	arc sine	trigonom.	67	
atan(arg)	arc tangent	trigonom.	67	
avg(arg)	argument arithmetic mean of values	aggr./mom.	69	
avg_e(arg)	run ensemble mean	Monte C.	82	
avg_l(char_arg,arg)	dimension-related argument arithmetic means of values	aggr./mom.	70	
avgg(arg)	argument geometric mean of values	aggr./mom.	69	
avgg_e(arg)	run ensemble geometric mean	Monte C.	82	
avgg_l(char_arg,arg)	dimension-related argument geometric means of values	aggr./mom.	70	
avgh(arg)	argument harmonic mean of values	aggr./mom.	69	
avgh_e(arg)	run ensemble harmonic mean	Monte C.	82	
avgh_l(char_arg,arg)	dimension-related argument harmonic means of values	aggr./mom.	70	
avgw(arg1,arg2)	argument weighted mean of values	aggr./mom.	69	
avgw_e(arg1,arg2)	run ensemble weighted mean	Monte C.	82	
avgw_l(char_arg,arg1,arg2)	dimension-related argument weighted means of values	aggr./mom.	70	
behav(char_arg,arg)	general purpose operator for navigating and aggre- gating in the experiment space	behav.	79	
classify(const_arg1, const_arg2,const_arg3,arg)	classification of arg into const_arg1 classes	advanced	72	
clip(char_arg,arg)	clip arg according to char_arg	advanced	72	
cnf(const_arg,arg)	positive distance of confidence line from mean avg_e(arg)	Monte C.	82	
cor(arg1,arg2)	correlation coefficient between arg1 and arg2	Monte C.	82	
cos(arg)	cosine	trigonom.	67	
cosh(arg)	hyperbolic cosine	trigonom.	67	
cot(arg)	cotangent	trigonom.	67	
coth(arg)	hyperbolic cotangent	trigonom.	67	
count(char_arg,arg)	count number of values	aggr./mom.	69	
count_e(char_arg,arg)	run ensemble count	Monte C.	82	
count_l(char_arg1,char_arg2, arg)	dimension-related count numbers of values	aggr./mom.	70	

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Name	Meaning	Туре	See Tab. at page
cov(arg1,arg2)	covariance between arg1 and arg2	Monte C.	82
cumul(char_arg,arg)	cumulates arg according to char_arg	advanced	72
dim(arg1,arg2)	positive difference	basic	67
ens(arg)	whole Monte Carlo run ensemble	Monte C.	82
exp(arg)	exponential function	basic	67
flip(char_arg,arg)	flip arg according to char arg	advanced	72
get_experiment(char_arg1, char_arg2,char_arg3,arg)	include an other experiment	advanced	72
get_table_fct(char_arg,arg)	table function with linear interpolation of table char_arg for position arg	advanced	72
hgr(char_arg1,const_arg1, const_arg2,const_arg3,arg)	argument histogram of values	aggr./mom.	69
hgr_e(char_arg1,const_arg1, const_arg2,const_arg3,arg)	heuristic probability density function	Monte C.	82
hgr_l(char_arg1,char_arg2, const_arg1,const_arg2, const_arg3,arg)	dimension-related argument histograms of values	aggr./mom.	70
if(char_arg,arg1,arg2,arg3)	general purpose conditional if-construct	advanced	72
int(arg)	truncation value	basic	67
krt(arg)	kurtosis (4 th moment)	Monte C.	82
lin_abs(char_arg,arg)	absolute linearity measure	sensitivity	84
lin_rel(char_arg,arg)	relative linearity measure	sensitivity	84
log(arg)	natural logarithm	basic	67
log10(arg)	decade logarithm	basic	67
mask(char_arg,arg1,arg2)	mask elements of argument arg1	advanced	72
matmul(arg1,arg2)	matrix multiplication	advanced	72
max(arg)	argument maximum of values	aggr./mom.	69
max_e(arg)	run ensemble maximum	Monte C.	82
max_l(char_arg,arg)	dimension-related argument maxima of values	aggr./mom.	70
max_n(arg1,,argn)	maximum per element	aggr./mom.	69
maxprop(arg)	index of the element where the maximum is reached the first time	aggr./mom.	69
maxprop_e(arg)	run number where the maximum is reached the first time	Monte C.	82
maxprop_l(char_arg,arg)	dimension-related argument position (1 n) where the maximum is reached the first time	aggr./mom.	70
maxprop_n(arg1,,argn)	argument position (1 n) where the maximum is reached the first time	aggr./mom.	69
med(arg)	median	Monte C.	82
min(arg)	argument minimum of values	aggr./mom.	69
min_e(arg)	run ensemble minimum	Monte C.	82
min_l(char_arg,arg)	dimension-related argument minima of values	aggr./mom.	70
min_n(arg1,,argn)	minimum per element	aggr./mom.	69
minprop(arg)	index of the element where the minimum is reached the first time	aggr./mom.	69
minprop_e(arg)	run number where the minimum is reached the first time	Monte C.	82
minprop_l(char_arg,arg)	dimension-related argument position (1 n) where the minimum is reached the first time	aggr./mom.	70
minprop_n(arg1,,argn)	argument position (1 n) where the minimum is reached the first time	aggr./mom.	69
mod(arg1,arg2)			67
nint(arg)	round value	basic basic	67

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Name	Meaning	Туре	See Tab. at page
nr_of_runs()	number of single runs of the current experiment	advanced	72
qnt(const_arg,arg)	quantile of arg	Monte C.	82
rank(char_arg,arg)	rank of arg according to char_arg	advanced	72
reg(arg1,arg2)	linear regression coefficient to forecast arg2 from arg1	Monte C.	82
rng(arg)	range = max_e(arg) - min_e(arg)	Monte C.	82
run(char_arg,arg)	values of arg for a single run selected by char_arg	advanced	72
sens_abs(char_arg,arg)	absolute sensitivity measure	sensitivity	84
sens_rel(char_arg,arg)	relative sensitivity measure	sensitivity	84
sign(arg)	sign of value	basic	67
sin(arg)	sine	basic	67
sinh(arg)	hyperbolic sine	trigonom.	67
skw(arg)	skewness (3 rd moment)	Monte C.	82
sqrt(arg)	square root	trigonom.	67
stat_full(const_arg1, const_arg2,const_arg3, const_arg4,arg5)	basic statistical summaries	Monte C.	82
stat_red(const_arg1, const_arg2,arg3)	basic statistical summaries	Monte C.	82
sum(arg)	argument sum of values	aggr./mom.	69
sum_e(arg)	run ensemble sum	Monte C.	82
sum_l(char_arg,arg)	dimension-related argument sums of values	aggr./mom.	70
sym_abs(char_arg,arg)	absolute symmetry measure	sensitivity	84
sym_rel(char_arg,arg)	relative symmetry measure	sensitivity	84
tan(arg)	tangent	trigonom.	67
tanh(arg)	hyperbolic tangent	trigonom.	67
transpose(char_arg,arg)	transpose arg according to char_arg	advanced	72
undef()	undefined element	advanced	72
var(arg)	argument variance of values	aggr./mom.	69
var_e(arg)	run ensemble variance	Monte C.	82
var_l(char_arg,arg)	dimension-related argument variances of values aggr./mom		70

 Tab. 15.7
 Post-processor built-in operators (in alphabetical order)

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15.3.3 Character Arguments of Post-Processor Built-In Operators

Tab. 15.8 summarises for built-in operators character argument values. User-defined operators can not have pre-defined character argument values.

Operator	Argument number	Argument value (without quotation marks, pre-defined values are case-insensitive)	Re- mark
avg_l	1	sequence of digits 0 and	(**)
avgg_l	1	sequence of digits 0 and 1	(**)
avgh_l	1	sequence of digits 0 and 1	(**)
avgw_l	1	sequence of digits 0 and 1	(**)
behav	1	(not pre-defined, case insensitive)	(*)
clip	1	(not pre-defined, case insensitive)	
count	1	[all def undef]	
count_e	1	[all def undef]	
count_l	1	sequence of digits 0 and 1	(**)
count_l	2	[all def undef]	
cumul	1	sequence of digits 0 and 1	(**)
flip	1	sequence of digits 0 and 1	(**)
get_experiment	1	(not pre-defined, case sensitive)	
get_experiment	2	(not pre-defined, case insensitive)	(*)
get_experiment	3	(not pre-defined, case sensitive)	
get_table_fct	1	(not pre-defined, case sensitive)	
hgr	1	[bin_no bin_mid]	
hgr_e	1	[bin_no bin_mid]	
hgr_l	1	sequence of digits 0 and 1	(**)
hgr_l	2	[bin_no bin_mid]	
if	1	[< <= > >= = != def undef]	
lin_abs	1	(not pre-defined, case insensitive)	(*)
lin_rel	1	(not pre-defined, case insensitive)	(*)
mask	1	[< <= > >= = !=]	
max_l	1	sequence of digits 0 and 1	(**)
maxprop_l	1	sequence of digits 0 and 1	(**)
min_l	1	sequence of digits 0 and 1	(**)
minprop_l	1	sequence of digits 0 and 1	(**)
rank	1	[tie_plain tie_min tie_avg]	
run	1	[run number not pre-defined]	
sens_abs	1	(not pre-defined, case insensitive)	(*)
sens_rel	1	(not pre-defined, case insensitive)	(*)
sum_l	1	sequence of digits 0 and 1	(**)
sym_abs	1	(not pre-defined, case insensitive)	(*)
sym_rel	1	(not pre-defined, case insensitive)	(*)
transpose	1	sequence of digits 1 to 9 (
var_l	1	sequence of digits 0 and 1	(**)

Tab. 15.8

Character arguments of post-processor built-in operators (*)

- Character argument can be empty
- (**) The length of the character argument from a sequence of digits corresponds with the dimensionality of the non-character and non-constant argument under investigation.

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15.3.4 Constant Arguments of Post-Processor Built-In Operators

Operator	Argument number	Argument type	Argument value restriction
classify	1	integer	[0 ≥2]
classify	2	real	arg2 = arg3 = 0. or
classify	3	real	arg2 < arg3
cnf	1	real	[0.001 0.01 0.05 0.1]
hgr	2	integer	[0 ≥4]
hgr	3	real	arg3 = arg4 = 0. or
hgr	4	real	arg3 < arg4
hgr_e	2	integer	[0 ≥4]
hgr_e	3	real	arg3 = arg4 = 0. or
hgr_e	4	real	arg3 < arg4
hgr_l	3	integer	[0 ≥4]
hgr_l	4	real	arg4 = arg5 = 0. or
hgr_l	5	real	arg4 < arg5
qnt	1	real	0. ≤ arg1 ≤ 100.
stat_full	1	real	[0.001 0.01 0.05 0.1]
stat_full	2	real	arg1 < arg2
stat_full	3	real	$0. \le \arg 3 < \arg 4 \le 100.$
stat_full	4	real	
stat_red	1	real	[0.001 0.01 0.05 0.1]
stat_red	2	real	arg1 < arg2

Tab. 15.8 summarises for built-in operators constant argument values.

Tab. 15.9

Constant arguments of post-processor built-in operators

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15.4 Glossary

The glossary defines and/or explains terms in that sense they are used in this User Guide. An arrow \rightarrow refers to another term in the glossary.

- Adjustment: Numerical modification of a \rightarrow target during an \rightarrow experiment. Adjustments are related to an \rightarrow experiment type and are described in the experiment description \rightarrow user-defined file.
- ASCII: The American Standard Code for Information and Interchange developed by the American National Standards Institute (http://www.ansi.org) is used in SimEnv to store information in → user-defined files and on request in result output files.
- Behavioural analysis: → Experiment type to inspect behaviour of a → model in a space, spanned up by → targets. The target space is scanned in a deterministic manner, applying pre-defined → adjustments of the targets with a flexible scanning strategy for target sub-spaces.
- Coordinate coord: Each → dimension of a → variable and each → operand of an → operator in a → result with a → dimensionality greater than 0 a coordinate is assigned to. A coordinate has a unique name and strictly monotonic ordered coordinate values. The number of coordinate values corresponds with the → extent for this dimension. Consequently, each model variable with a dimensionality greater than 0 resides at a assigned (multi-dimensional) → grid. Assignments for variables is done in the model description → user-defined file.
- Coupling: → model interface
- Data type: The type of a → variable as declared in the → model and the corresponding model description → user-defined file. SimEnv data types are byte, short, int, float, and double.
- **Default value**: The nominal (standard) numerical value of an experiment \rightarrow target. The default value is specified in the experiment description \rightarrow user-defined file and for \rightarrow the model interface at the language level also in the model code.
- Dimension: → dimensionality
- Dimensionality dim: The number of dimensions of a model → variable or of an → operator result in model output post-processing. In the model description → user-defined file each variable a dimensionality is assigned to that corresponds with the dimensionality of the related model output field in the model source code. Dimensionality 0 corresponds to a scalar, dimensionality 1 to a vector, dimensionality 2 to a matrix.
- **Environment variable**: At UNIX operating system level the so called environment is set up as an array of operating-system and user-defined environment variables that have the form Name=Value. The Value of a Name can be addressed by \$Name. In SimEnv use of environment variables in directory strings is forbidden.
- Experiment: Performing simulation runs with a → model in a co-ordinated manner by applying → experiment types and running the model in a run ensemble, i.e., a series of single simulation runs.

Experiment target: → target

- Experiment type: Pre-defined multi-run simulation experiment. In the process of experiment preparation (defining an experiment by describing it in the experiment description → user-defined file) → targets are assigned to an experiment type and experiment-specific → adjustments and other information are assigned to the targets. Currently available experiment types are → behavioural analysis, → Monte Carlo analysis, and → local sensitivity analysis.
- Extent ext: The number of values for a dimension (from the → dimensionality) of a model → variable or of an → operator result in model result post-processing. Extents are always greater than 1. Model variables and operator results of dimensionality 0 do not have an extent.

Expression: → result expression



Fortran storage model: A rule how to map the elements of a multi-dimensional data field to a 1-dimensional vector and vice versa. A data field field(1:ext₁, 1:ext₂, ..., 1:ext_{dim-1}, 1:ext_{dim}) of → dimensionality dim and → extents ext₁, ext₂, ..., ext_{dim-1}, ext_{dim} is mapped in Fortran in the following way on a 1-dimensional vector vector (1:ext₁* ext₂* ... * ext_{dim-1}* ext_{dim})

```
      ipointer = 0 \\ do i_{dim} = 1 , ext_{dim} \\ do i_{dim-1} = 1 , ext_{dim-1} \\ ... \\ do i_2 = 1 , ext_2 \\ do i_1 = 1 , ext_1 \\ ipointer = ipointer + 1 \\ vector(ipointer) = field(i_1, i_2, ..., i_{dim-1}, i_{dim}) \\ enddo \\ enddo \\ ... \\ enddo \\ enddo
```

For a two-dimensional matrix this storage model corresponds to a column by column storage of the matrix to the vector, starting with the first column and for each column starting with the first row.

- **GAMS**: The General Algebraic Modeling System (http://www.gams.com) is a high-level modeling system for mathematical programming problems. It consists of a language compiler and a stable of integrated high-performance solvers. GAMS is tailored for complex, large scale modeling applications, and allows to build large maintainable models that can be adapted quickly to new situations.
- **Grid**: Regular topological structure for a model \rightarrow variable or an \rightarrow operator result in post-processing, spanned up as the Cartesian product of the assigned \rightarrow coordinates to the variable or the operator result.
- IEEE: SimEnv can use on demand for storage of model and post-processor output the Institute of Electrical and Electronics Engineers (http://www.ieee.org) standard number 754 for binary storage of floating point numbers.
- Load Leveler: The load leveler LoadL is a job management system that handles compute resources at IBM's p655 cluster at PIK.
- Local sensitivity analysis: → Experiment type with incremental → adjustments of → targets in the neighbourhood of the → default values of the targets. A local sensitivity analysis in SimEnv is always performed independently for all targets involved. During post-processing sensitivity, linearity, and symmetry measures can be determined.
- **Macro**: An abbreviation for a unique \rightarrow result expression to apply during \rightarrow post-processing. Macros can be embedded into result expressions and are plugged into the expression during its evaluation and computation. Macros are described in the macro description \rightarrow user-defined file.
- Model: A model is a deterministic or stochastic algorithm, implemented in one or a number of computer programs that transforms a sequence of input values (→ targets) into a sequence of output values (→ variables). Normally, inputs are parameters, driving forces, initial values, or boundary values to the model, outputs are state variables of the model. For many cases, the model will be state deterministic, time and space dependent. For SimEnv, the model, its targets and variables are coupled in the process of → interfacing the model to SimEnv.

Model coupling: → model interface

Model interface: Interfacing a → model to SimEnv means coupling it to SimEnv and enabling finally experimenting with a model within SimEnv. There are coupling interfaces at programming language level for C/C++, Fortran, → Python, and → GAMS. Additionally, models can be interfaced at the → shell script level by using shell script syntax elements. For all interface techniques the interfaced model itself has to be wrapped into a shell script.

Model output post-processing operator: → operator

Model output variable: → variable

- Monte Carlo analysis: → Experiment type with pre-single run perturbations of experiment → targets. Each perturbed target a → probability density function pdf with function parameters is assigned to. During the → experiment → adjustments of the targets are realizations from the pdf's using random number techniques. In experiment post-processing statistical measures can be derived from model output of the run ensemble. A prominent statistical measure is the heuristic pdf (histogram) of a model → variable and its relation to the pdf's of the targets.
- NetCDF: Network Common Data Form is an interface for array-oriented data access and a library that provides an implementation of the interface. The NetCDF library also defines a machine-independent format for representing scientific data. Together, the interface, library, and format support the creation, access, and sharing of scientific data. The NetCDF software was developed at the Unidata Program Center in Boulder, Colorado (http://www.unidata.ucar.edu). NetCDF is freely available. SimEnv follows for model output and post-processing output storage the NetCDF Climate and Forecast (CF) metadata convention 1.0-beta4

(http://www.cgd.ucar.edu/cms/eaton/cf-metadata/index.html) and extends it.

- **OpenDX:** The **Open D**ata Explorer OpenDX (http://www.opendx.org) is a uniquely full-featured open source project and software package for the visualization of scientific, engineering and analytical data: Its open system design is built on a standard interface environment. The data model provides users with great flexibility in creating visualizations. OpenDX is based on IBM's Visualization Data Explorer.
- **Operand**: Argument of an \rightarrow operator in SimEnv model output post-processing. An operand can be a model \rightarrow variable, an experiment \rightarrow target, a constant, a character string, \rightarrow a macro and an operator.
- **Operator**: Computational algorithm how to transform the values of a sequence of → operands into the values of the operator result during model output post-processing. An operator transforms → dimensionality, → extents, and → coordinates from the operands into the corresponding information for the operator result. There are built-in elemental, basic, and advanced operators as well as built-in operators related to specific → experiment types. Additionally, SimEnv offers specification of user-defined operators according to an operator interface. User-defined operators are announced to the system in the operator description → user-defined file.
- Optimization: → Experiment type to minimize a cost function (objective function) over a bounded → target space. In SimEnv a simulated annealing strategy (check chapter 4.5 for explanation) is used to optimize the cost function that is formed from model → variables. Often the cost function represents a distance between model output and reference data to find an optimal point in the target space that fits best the model behaviour with respect to the reference data.

Parallel Operating Environment: → POE

- **POE:** The **P**arallel **O**perating **E**nvironment POE on IBM's p655 cluster at PIK supplies services to allocate nodes, assign jobs to nodes and launch jobs.
- **Post-processing:** The work step of processing model output data from the whole run ensemble after performing a simulation \rightarrow experiment. SimEnv post-processing enables navigation in the \rightarrow target space that is sampled by an experiment as well as construction of additional output functions by declaration and computation of \rightarrow results.
- **Probability density function pdf**: A probability density function serves to represent a probability distribution in terms of integrals. A probability distribution assigns to every interval of real numbers a probability.
- **Python**: Python (http://www.python.org) is a portable, interpreted, interactive, object-oriented programming language. It incorporates modules, exceptions, dynamic typing, and very high level dynamic data types, and classes.
- **Result:** In SimEnv \rightarrow post-processing a result (synonym: output function) is derived from model output of the \rightarrow experiment and from reference data. A result is specified by a result expression, optionally prefixed by a result description and a result unit string.
- **Result expression:** A chain of \rightarrow operators from built-in or user-defined operators applied to model output \rightarrow variables and/or reference data. A result expression is a part of a \rightarrow post-processing \rightarrow result.



Shell script: A sequence of UNIX operating system commands stored in an → ASCII file. A shell script is interpreted and executed by a command line interpreter, the so-called shell. SimEnv demands the Korn shell ksh.

Simulation: Performing \rightarrow experiments with \rightarrow models

Target: Element of the input set of a → model. Targets are manipulated numerically during an → experiment. Targets can be addressed in model output post-processing and they have there a → dimensionality of zero.

Target adjustment: → adjustment

- User-defined files: A set of → ASCII files to describe → model-, → experiment-, → operator-, → macro-, and → GAMS model specific information and to determine general SimEnv settings. All user-defined files follow the same syntax rules.
- Variable: Element of the output set of a → model that is stored in a SimEnv model output format. Variables are defined in the model description → user file and they are output from the model to SimEnv data structures. Each variable has a unique → data type, a → dimensionality, → extents and an assigned → grid. Normally, a variable consists of a series of values, forming a field.
- White spaces: → ASCII characters space (blank) and horizontal tabulator used in → user-defined files or within result expressions in model output post-processing.

Working directory: The directory, a SimEnv service was started from.

