READEX Tool Suite – User Guide

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1 Introduction

The READEX tool suite has been developed to optimise the execution of HPC applications on Exascale systems for energy efficiency. This is achieved by analysing and modelling the dynamic behaviour of an HPC application in a step known as *design-time analysis*. This step results in identifying the optimal values for different hardware, system software and application parameters that are available for tuning. Currently, READEX is capable of tuning two hardware parameters (CPU core and uncore frequencies), a system software parameter (number of OpenMP threads) and application parameters that have been exposed by the user of the tool suite or the application owner. Following this, the optimal values that have been identified for different regions of an application are set during the production run of the application in a step known as *runtime application tuning*.

This document describes how to use the READEX tool suite according a simple workflow:

- 1. Instrument the application with Score-P. (Section 2)
- 2. Perform design-time analysis of application to create tuning model. (Section 3)
- 3. Use the tuning model during the production run of the application for runtime tuning. (Section 4)

For a better understanding of the READEX approach we recommend to have a look at Deliverable D2.3. The tools in the READEX tool suite are accessible by installing the tool suite as explained in the READEX website.

2 Application instrumentation

2.1 Build application with Score-P

The READEX tool suite is based on instrumenting an application with Score-P. Instrumentation inserts measurement probes into the source code of the application. This can be done by the compiler, by other software tools, or manually. Detailed documentation on Score-P and the instrumentation features can be found at www.score-p.org.

1. Modify the application's makefile for instrumentation with Score-P. Prepend the compilation with the **scorep** command. For example,

Replace MPICXX = mpic++ -fopenmp by MPICXX = scorep --mpp=mpi mpic++ -fopenmp

The **scorep** command switches on compiler instrumentation of program functions as well as instrumentation of MPI routines and OpenMP regions.

Use --mpp=mpi for MPI applications and --mpp=none for non-MPI applications.

- 2. Build the application. Note that Score-P and the application have to be built with the same compiler.
- 3. Run the application like the uninstrumented version.

Outcome: Compiler instrumentation of the application is performed; upon application execution, Score-P creates a profile (profile.cubex) file in the scorep-<xyz> directory at the execution location.

2.2 Filtering

The probes inserted in the application through instrumentation add overhead to the application execution and thus can make any measurements and tuning efforts wasted time. Therefore, it is essential to make sure that the instrumentation overhead is below a certain limit. This section focuses on giving you advice on the support in Score-P for reducing the measurement overheads. To measure the overhead, first measure the execution without instrumentation and then measure it with instrumentation.

To reduce the overhead from instrumentation to an acceptable level,

- 1. First try to reduce the overhead with runtime and compile time filtering as described in Sections A.1 and A.2, respectively.
- 2. You may also remove MPI and OpenMP region instrumentation overhead as described in Section A.3.
- 3. Then switch on the energy measurements which (sometimes) may cause significant performance overhead. Verify the overhead again. For instance, RAPL can be used for energy measurement, which has less overhead than HDEEM. Note that the energy measurements from RAPL may not be precise enough. For instance, an execution time less than 40 ms (that is 40 ms function execution time and 1 ms sampling rate) may result in approximately 2.5% error.

2.3 Phase region instrumentation

4. If the overhead is still too high, consider manual instrumentation of those regions that are relevant for the READEX tool suite as described in Section A.5.

Do not proceed to energy tuning if the overhead is too high, e.g. more than 5%.

2.3 Phase region instrumentation

Specify the phase region: Manually annotate the phase region of the application as shown below:

```
SCOREP_USER_REGION_DEFINE( REGION_HANDLE )
// loop starts
SCOREP_USER_OA_PHASE_BEGIN( REGION_HANDLE, "PHASE_REGION_NAME", SCOREP_USER_REGION_TYPE_COMMON )
// loop body (phase region)
SCOREP_USER_OA_PHASE_END( REGION_HANDLE )
// loop ends
```

A phase region is a repetitive, single-entry and exit region, typically the body of the main progress loop of the application. If the phase region is not known beforehand, it may be useful to look at the profile.cubex file generated after running the scorep-autofilter tool with a performance analysis tool like CUBE.

Example The for-loop body in Integrate::run() is annotated as a phase region as shown in the example in Section D.3.

2.4 Application tuning parameter instrumentation

As mentioned earlier, the READEX tool suite also allows tuning application parameters. This is optional and the application parameters to be tuned should be identified and exposed by the tool suite user and/or by the application owner. This requires some additional manual code annotation and instrumentation to pinpoint the parts of the code that can be exploited as application tuning parameters and annotate them with certain API functions.

This is enabled in READEX using the ATP (Application Tuning Parameter) library and the procedure for this is described in Section B.1.

3 Design-time Analysis (DTA)

3.1 Tuning Potential Analysis

The first step in the DTA is to detect and analyze the dynamism of the application using **readex-dyn-detect**. The tool automatically identifies the significant regions that are subject to the READEX tuning methodology and generates a report on the potentially exploitable dynamism in these regions.

The readex-dyn-detect tool requires a single phase region, which is to be instrumented as described earlier in Section 2.3.

Perform the following steps to use readex-dyn-detect:

- 1. Build the application with instrumentation by scorep. Add --online-access --user for the manually annotated phase region.
- 2. Run the application with the following environment variables set:

```
export SCOREP_PROFILING_FORMAT=cube_tuple
export SCOREP_METRIC_PAPI=PAPI_TOT_INS,PAPI_L3_TCM
export SCOREP_FILTERING_FILE=<filter_file_name_with_extension>
```

This will create a tupled profile.cubex file in the scorep-<xyz> directory at the execution location.

3. Apply the readex-dyn-detect tool on the profile.cubex file as follows:



If the readex configuration file already exists, only the list of significant regions will be updated according to the outcome of **readex-dyn-detect**. It the file is missing, the template file will be copied from the installation directory and updated with the significant regions.

The command line options have the following meaning:

- -t This threshold specifies the minimal mean execution time of regions that are to be considered as significant regions. Use a value larger than 0.1 (100 ms). (default 0.1)
- -p Name of the phase region as given in the instrumentation.
- -c This is the required minimal standard deviation of the compute intensities of significant regions with a weight above the given threshold, such that intra-phase dynamism due to compute intensity variation is reported. (default 10%)
- -v This is the required minimal standard deviation of the execution time of instances of significant regions in percent of the mean region's execution time, such that intra-phase dynamism is reported. It is also used to decide whether inter-phase dynamism exists. Only if the standard variation of the phase time in percent of the mean phase time is greater, inter-phase dynamism is reported. (default 10%)

3.2 Configure DTA

- -w This threshold specifies the minimal weight of a region such that any dynamism due to time variation or compute intensity variation is reported. (default 10%)
- -r This is the desired name (without the file name extension) for the READEX configuration file to be created by readex-dyn-detect.
- 4. The results of readex-dyn-detect are summarized in readex_config.xml in the execution directory, which is used as an input to PTF. An example of readex_config.xml is available in <PTF_installation_path>/templates/readex_config.xml.default.

Alternatively, the **readex_config.xml** file may be manually created from this template and used as input for PTF without applying **readex-dyn-detect** if the significant regions are already known.

Note: readex-dyn-detect currently ignores MPI and shared memory regions in the significant regions analysis.

READEX distinguishes between two types of dynamism in an application:

- Inter-phase dynamism: This occurs when each phase (execution instance) of a phase region in the application exhibits different characteristics. This results in different values for the measured objective values and thus may require different configurations to be applied for the tuning parameters.
- Intra-phase dynamism: This occurs when each runtime situation (execution instance) of the significant regions in a phase region exhibits different characteristics and thus may need different configurations to be applied for the tuning parameters.

Outcome: The readex_config.xml file containing the tuning potential summary, the list of significant regions, and the intra-phase and inter-phase dynamism due to variation in the execution time and compute intensity.

Section D.6 presents an example.

3.2 Configure DTA

The next step of the DTA is to update the readex_config.xml file that the readex-dyn-detect tool has generated with additional criteria for the design-time analysis experiments performed by the Periscope Tuning Framework (PTF). The steps to update the readex_config.xml file are as follows:

1. Specify the tuning parameters: READEX currently supports tuning two hardware parameters (processor core frequency and uncore frequency) and a system software parameter (number of OpenMP threads). A minimum of one of these tuning parameter must be specified. Specify the ranges (minimum, maximum, step size, and default) for the processor core frequency and the uncore frequency in MHz. The default frequencies are given by the system and are defined during the installation. Section C describes how to retrieve the available CPU core and uncore frequencies in a system.

For OpenMP threads, specify the lower bound and the step size to increment to the next value. The upper bound is given by the psc_frontend command.

3.2 Configure DTA

Example

<tuningparameter></tuningparameter>
<frequency></frequency>
<min_freq>1200</min_freq>
<max_freq>2400</max_freq>
<freq_step>200</freq_step>
<default> 2500</default>
<uncore></uncore>
<min_freq>1000</min_freq>
<max_freq>3000</max_freq>
<freq_step>200</freq_step>
<default> 3000</default>
<openmpthreads></openmpthreads>
<lower_value>1</lower_value>
<step>2</step>

Note that application parameters may be optionally tuned as introduced in Section 2.4 and explained in Section B.1.

2. Specify the objectives: Specify at least one objective from Energy, Execution Time, CPU Energy, Energy Delay Product, Energy Delay Product Squared, CPUEnergy, Total Cost of Ownership (TCO). The normalized version of each of the objectives can also be specified. It expresses the energy consumption per instruction and can be used for applications where the amount of computation in a phase varies without changing the phase characteristics ("more of the same"). The plugin measures the objective values for all the specified objectives, but tunes the application only for the objective that is specified first.

Example

<pre><objectives></objectives></pre>				
<objective>Energy</objective>				
<objective>NormalizedEnergy</objective>				
<objective>Time</objective>				
<objective>NormalizedTime</objective>				
<objective>EDP</objective>				
<objective>NormalizedEDP</objective>				
<objective>ED2P</objective>				
<objective>NormalizedED2P</objective>				
<objective>CPUEnergy</objective>				
<objective>NormalizedCPUEnergy</objective>				
<objective>TCO</objective>				
<objective>NormalizedTCO</objective>				

To compute TCO, the CostPerJoule and CostPerCoreHour also needs to be specified.

<Configuration> <CostPerJoule>0.00000008</CostPerJoule> <CostPerCoreHour>1.0</CostPerCoreHour> </Configuration>

3. Specify the energy metrics: Specify the energy plugin name and associated metric names. For hdeem_sync_plugin, it is possible to measure the energy for the whole node or/and two CPUs respectively. The energy metrics should be specified under cope>

3.2 Configure DTA

Example

```
cperiscope>
    <metricPlugin>
        <name>hdeem_sync_plugin</name>
     </metricPlugin>
     <metrics>
        <node_energy>hdeem/BLADE/E</node_energy>
        <cpu0_energy>hdeem/CPU0/E</cpu0_energy>
        <cpu1_energy>hdeem/CPU1/E</cpu1_energy>
     </metrics>
```

To specify the RAPL counter energy plugin x86_energy_sync_plugin, use the configuration as follows:

Example

```
cmetricPlugin>
    <metricPlugin>
    <metricPlugin>
    <metricPlugin>
    <metrics>
        <node_energy>x86_energy/BLADE/E</node_energy>
        <cpu0_energy>x86_energy/CORE0/E</cpu0_energy>
        <cpu1_energy>x86_energy/CORE1/E</cpu1_energy>
        </metrics>
```

4. Specify a search algorithm: Specify a search algorithm from exhaustive, random or individual. For the random search strategy, specify the number of samples (scenarios) that the plugin should limit to. For the individual search, specify the number of tuning parameter values to *keep* in the search space. The search algorithm should be specified under cpriscope>

Example

```
<searchAlgorithm>
<name>exhaustive</name>
<name>random</name>
<samples>2</samples>
<name>individual</name>
<keep>2</keep>
</searchAlgorithm>
</periscope>
```

The search algorithm together with the tuning parameter specification determines the tuning time. Exhaustive search leads to the biggest number of configurations that are tested in subsequent program phases. The individual strategy reduces the number significantly since not the cross product of all tuning parameters is investigated, but the parameters are optimized independently. With the random strategy, the number of experiments can be specified.

5. Specify the tuning model file name: The generated tuning model file name can also be specified under <periscope> </periscope>.

Example

```
ctuningModel>
    <file_path>./tuning_model.json</file_path>
    </tuningModel>
```

Optionally, if the Application Tuning Parameter (ATP) library is used, then the details for the ATP library should be included in the READEX configuration file as outlined in Section B.2.

3.3 Tuning application configuration parameters

The READEX tool suite provides an additional optional step before generating the tuning model. Applications frequently have *Application Configuration Parameters (ACP)* that might effect the performance and energy consumptions. These parameters can be tuned for the given objective with PTF. It provides a tuning plugin which takes a specification of the ACPs with their possible values and searches for the best settings by restarting the application for each experiment.

The tuning plugin is **readex_configuration**. It relies on a configuration file specifying the input files, the configuration parameters and their possible values. The plugin first reads the configuration files and then generates the possible configurations based on the defined search algorithm. For each configuration, the template file of an input file is copied, the configuration variables are replaced by the selected values, and the application is restarted.

Note that names of configurations variables may be composed of the letters 'A'-'Z', 'a-'z', '0'-'9', and '_'.

Be aware that only a single phase is executed for each experiment which significantly reduces the tuning time. In case you want to omit some phases before starting the measurements or you want it to be based on multiple subsequent phases, please use the optional PTF arguments --delay and --iterations. At the end of the search, the optimal configuration is written into the input files and output into the configured results file.

The tuning plugin and its configuration file are selected via command line arguments as shown in the example below. Both configuration files are to be given, i.e., readex_config.xml and appConfigParams.cfg.

6

. . .

```
psc_frontend --apprun=....
--config-file=readex_config.xml
--tune=readex_configuration --readex-app-config=appConfigParams.cfg
...
```

The plugin takes from readex_config.xml the configuration for the energy measurements, all other configurations are taken from the plugin specific configuration file. The syntax of the specification in the plugin specific configuration file is given below:

```
5 //First specify the input file with its template file
```

3.4 Tuning Model Creation

```
//PTF copies the template file to the input file and
6
7
     //replaces the ACP name with the current value
     //-----
8
9
                       FileSpec FileList | FileSpec
     FileList:
10
^{11}
     FileSpec:
                      FileSpecification TemplateFileSpecification ParameterList
     FileSpecification: FILENAME STRING ';'
12
     TemplateFileSpecification: TEMPLATEFILENAME STRING ';'
13
14
     //-----
                   15
     //In each input file a number of ACP can be provided
16
17
     //For each ACP specify the name and the values.
     //The first value is taken as default.
18
     //-----
19
                  ------
20
     ParameterList:
                      ParameterSpecification ParameterList | ParameterSpecification
21
     ParameterSpecification: PARAMETER STRING VALUESSTR TpRange ';'
22
23
     TpRange:
                       IntRange | StringRange
24
                       '[' INT ',' INT ']' | '[' INT ',' INT ',' INT ']'
25
     IntRange:
     StringRange:
                       '[' StringList ']'
26
                       STRING ',' StringList | STRING
27
     StringList:
                         ['-' 'a'-'z' 'A'-'Z' '0'-'9' '_'
     //STRING:
28
                          ·~· ·*· ·@· ·, · ·(· ·)· · · · · · ·/· ·=· ]+
29
     11
30
31
32
     //-----
33
     //Specification of general configuration parameters
     //---
          ------
34
35
                       ConfigStmt ConfigList | ConfigStmt
36
     ConfigList:
37
     ConfigStmt:
                       SearchAlgSpecification ';'
38
39
                       | IndividualKeepSpecification ';'
                       | SampleCountSpecification ';'
40
                       | ResultsFileSpecification ';'
41
                       | ObjectiveSpecification ';'
42
43
     SearchAlgSpecification:
                                SEARCHALG STRING
44
     IndividualKeepSpecification: INDIVIDUALKEEP INT
45
46
     SampleCountSpecification:
                               SAMPLECOUNT '=' INT
47
     ResultsFileSpecification:
                                RESULTSFILE STRING
48
49
     ObjectiveSpecification:
                                OBJECTIVE ObjectiveName
50
                      ENERGY | NORMALIZEDENERGY
51
     ObjectiveName:
                       | CPUENERGY | NORMALIZEDCPUENERGY
52
                       | EDP | NORMALIZEDEDP
53
                       | ED2P | NORMALIZEDED2P
54
55
                       | TIME | NORMALIZEDTIME
                       | TCO | NORMALIZEDTCO
56
57
```

Outcome: The results file containing the optimal configuration for the ACPs and the input files where the ACPs were replaced by the optimal setting.

3.4 Tuning Model Creation

After updating the **readex_config.xml** file for use by PTF, use the following steps to perform design-time analysis using PTF as explained using a slurm job script for the miniMD application as an example.

1. Build the application with instrumentation as discussed in Section 2.3 (scorep --user --online-access) for the instrumented phase region. Additionally, you may optionally

3.4 Tuning Model Creation

use the Score-P options that are required to specify compile-time filtering, MPP and thread instrumentation options. Refer to the Score-P documentation for this.

- 2. Set the number of nodes requested to at least 2, and allocate enough memory to fit the application. In general, if N > 1 nodes are allocated for this job, then PTF will use one node for the tool's agents and the remaining N-1 nodes for the application processes.
- 3. Use the substrate plugin and parameter control plugins compatible with Score-P and PTF as shown in lines 16 and 17.
- 4. Load the Score-P plugin to be used for energy measurements and set the corresponding Score-P environment variables as shown in lines 20-24. The SCOREP_SUBSTRATE_PLUGIN variable is used to specify the substrate plugins to be used, which for READEX should include rrl. The SCOREP_RRL_PLUGINS are a comma-separated list of parameter tuning plugin names to be used by RRL (cpu_freq_plugin, uncore_freq_plugin, OpenMPTP, etc.). The SCOREP_METRIC_PLUGINS is used to set the metric plugin used to perform energy measurements.
- 5. Apply PTF on the application with the psc_frontend command as shown in lines 28– 36. Specify the instrumented phase region name for the option --phase and the readex configuration file for --config-file. Specify the readex_intraphase plugin for --tune.

The options --info and --selective-info are only used for debug messages, and are not mandatory. For more debug output, set the --info=<max_info_level> between 2 and 7, and --selective-info=<comma_separated_list_of_information_levels>. For more information about other options, see psc_frontend --help.

This will produce a tuning model in the execution directory under the name specified in the readex_config.xml file, or tuning_model.json if unspecified.

If readex-dyn-detect reports inter-phase dynamism for the application, the user is advised to proceed with inter-phase tuning, as described in Section 3.5. Note that the ATPs are disabled for inter-phase tuning.

```
### Batch system (SLURM, PBS, ...) directives ###
1
     ### number of nodes requested (at least 2); 1 for PTF and remaining for application run
2
3
     ### ...
     ### memory requested for application run
4
     ### ...
5
6
     ###
7
8
     ###
9
     ### Load READEX tool suite modules, or set paths to executables and libraries
10
     ###
11
     echo "run PTF begin."
12
13
     export SCOREP_SUBSTRATE_PLUGINS=rrl
14
     export SCOREP_RRL_PLUGINS=<list of RRL tuning plugins>
15
     export SCOREP_RRL_VERBOSE="WARN"
16
17
18
     export SCOREP_METRIC_PLUGINS=<energy_plugin_name; for eg. HDEEM, RAPL, ...>
     export SCOREP_METRIC_PLUGINS_SEP=";'
19
     ###
20
     ### set Score-P metric energy plugin environment variables; for eg. SCOREP_METRIC_HDEEM_SYNC_PLUGIN_*
21
22
     ###
23
24
     export SCOREP_MPI_ENABLE_GROUPS=ENV
25
     psc_frontend --apprun="<application executable and command-line arguments>"
26
                   --mpinumprocs=<number of MPI processes>
27
```

```
28 --ompnumthreads=<maximum number of OpenMP threads>
29 --phase=<phase region name>
30 --tune=<readex_intraphase | readex_interphase>
31 --config-file=<READEX configuration file name; for eg. readex_config.xml>
32 --info=<2 .. 7>
33 --selective-info=AutotuneAll,AutotunePlugins
34
35 echo "run PTF done."
```

Outcome:

- A printed summary of the created scenarios, the properties found in each scenario, the optimum and the worst scenarios for the phase, the measured objective values for the phase in each scenario, the best configuration for each rts, the static and dynamic energy savings for the rts's, and the static energy savings for the whole phase.
- A tuning_model.json file containing the list of rts's that were tuned by the plugin, the scenarios into which they are classified, and the best configuration for each scenario.

3.5 Inter-Phase Tuning

To exploit the inter-phase dynamism, only the following must be changed in addition to the steps described in Sections 3.2 and 3.4.

Configure DTA For inter-phase tuning, the search algorithm specified by the user is ignored, and the plugin sets the **random** search strategy by default. Specify the number of experiments/-points to analyze during clustering under the **samples** tag. Higher values enable a better analysis of the dynamism between the phases and result in a more refined tuning model.

Tuning Model Creation Specify the readex_interphase plugin for --tune.

Outcome The readex_interphase tuning plugin clusters similar phases using the DBSCAN algorithm, and determines the best configuration for each cluster, which will be applied to all the phases belonging to that cluster. It also determines the best configuration for individual rts's in the cluster.

- A printed summary of the created scenarios, the properties found in each scenario, the measured objective values for the phase in each scenario, per-cluster results showing the optimum scenario for all the phases of the cluster as well as the best configuration for each rts of the cluster, the static and dynamic energy savings for the rts's, and the static energy savings for the whole phase.
- A tuning_model.json file containing the list of clusters generated by the clustering algorithm, the set of phases belonging to each cluster, the ranges of the features that were used for clustering, the list of rts's that were tuned by the plugin, the scenarios into which they are classified, and the best configuration for each scenario.

3.6 Visualization of the Tuning Model

3.6 Visualization of the Tuning Model

The visualization tool of the tuning model is constructed based on the JavaScript library D3.js. The tool can be built as the app for macOS, Linux, and Windows, using electron-packager. The latest source of the visualization tool of the tuning model can be cloned by git clone https://periscope.in.tum.de/git/visualizationOfTM.git. This source already contains electron-packager source and configured to build as apps. Electron Packager is a command line tool and Node.js library that bundles Electron-based application source code with a renamed Electron executable and supporting files into folders ready for distribution.

Mandatory Library: The tool requires Node.js library to be installed. The library can be downloaded from https://nodejs.org.

Usage: The tool requires two mandatory files which are generated at the end of DTA. The files are: tuning_model.json and rts.xml. The first one is the tuning model file and the latter contains the execution time information of the rts's. The tool can be started with the following command in the source directory:

1

npm start

First, select the tuning model file tuning_model.json and the file with the rts's rts.xml. You have to select both files in the dialog. The tool checks the extension and assumes that a file with .json is the tuning model and the extension .xml identifies the rts file.

The tool will then generate the forced layout view of the tuning model. This will look as in Figure 1. For each scenario a circle is generated. The size of the circle represents the weight, i.e., the aggregated execution times of all rts's of the scenario as percent of the phase time. The thickness of the lines represents the similarity of two connected scenarios. Hovering over a circle triggers a tool tip that gives detailed information about the scenario. Clicking on a scenario opens individual circles for each rts in this scenario.

More details about the tool can be found in deliverable D2.2.



Figure 1: Forced layout graph of the tuning model

4 Runtime Application Tuning (RAT)

4.1 Production Run with Tuning Model

The following steps describe how to use RRL to tune the application during its production run.

- 1. If Application Tuning Parameters are exploited in the application then the ATP related instrumentation functions should remain in the code.
- 2. For the application run tuned with RRL, use the application built for analysis with PTF as described in Section 3. If the readex_interphase tuning plugin was used for DTA, perform the following steps before proceeding with RAT:
 - To perform runtime tuning after the inter-phase analysis, the cluster prediction library must be invoked. The SCOREP_USER_PARAMETER macro must be added to the source code, with a call to the predict_cluster function of the cluster prediction library.

For C/C++ applications:

```
#include <cluster_prediction.h>
// loop starts
SCOREP_USER_OA_PHASE_BEGIN( REGION_HANDLE, "PHASE_REGION_NAME", SCOREP_USER_REGION_TYPE_COMMON )
SCOREP_USER_PARAMETER_INT64( "Cluster", predict_cluster() )
// loop body (phase region)
SCOREP_USER_OA_PHASE_END( REGION_HANDLE )
// loop ends
```

For Fortran applications:

```
SCOREP_USER_REGION_DEFINE( REGION_HANDLE )
SCOREP_USER_PARAMETER_DEFINE(cluster)
// loop starts
SCOREP_USER_OA_PHASE_BEGIN( REGION_HANDLE, "PHASE_REGION_NAME", SCOREP_USER_REGION_TYPE_COMMON )
SCOREP_USER_PARAMETER_INT64( "Cluster", predict_cluster() )
// loop body (phase region)
SCOREP_USER_OA_PHASE_END( REGION_HANDLE )
// loop ends
```

- Add the linker flags in the Makefile and rebuild the application.
- 3. Set the number of nodes to run the application, and allocate enough memory to fit the application (lines 1-6). Here, the number of nodes required is the same as the number of nodes on which to run the application.
- 4. For the RRL-tuned run of the application (lines 68–106) perform the following steps:
 - (a) Disable Score-P profiling and tracing (lines 13 and 14), set the Score-P substrate plugins to **rrl**, RRL plugins to the tuning plugins to use and the tuning model to the file generated by PTF (lines 15–17).
 - (b) Depending on whether the calibration mechanism is enabled or disabled for RRL, set the environment variable SCOREP_FORCE_CFG_FILES to true or false, respectively, to enable or disable generation of Score-P configuration file for use by RRL (line 23).
 - (c) Run the RRL-tuned version of the application (line 26) using the application executable built for PTF.

^{1 ###} Batch system (SLURM, PBS, ...) directives ###

^{2 ###} number of nodes requested for application run

^{3 ### ...}

```
### memory requested for application run
4
     ### ...
5
6
     ###
7
     ###
8
     ### Load READEX tool suite modules, or set paths to executables and libraries
9
10
     ###
11
     ### start RRL-tuned run
12
     export SCOREP_ENABLE_PROFILING="false"
13
     export SCOREP_ENABLE_TRACING="false"
14
15
     export SCOREP_SUBSTRATE_PLUGINS="rrl"
     export SCOREP_RRL_PLUGINS=<list of RRL tuning plugins>
16
     export SCOREP_RRL_TMM_PATH=<tuning model file name>
17
     export SCOREP_MPI_ENABLE_GROUPS=ENV
18
19
     ### If RRL calibration mechanism is enabled, then set SCORE_FORCE_CFG_FILES to true.
20
     ### This will create Score-P configuration files in a Score-P directory for the application run.
21
22
     ### If RRL calibration mechanish is disabled, then set SCORE_FORCE_CFG_FILES to false.
23
     export SCOREP_FORCE_CFG_FILES=false
24
25
     ### run RRL-tuned application
26
     # code to start application run with command-line arguments
     ### end RRL-tuned run
27
```

4.2 Visualise Configuration Switching

There are two ways for visualising the configuration switching:

- 1. The first is a Score-P metric plugin which is implemented as part of RRL and it shows the RRL perspective to the switching, i.e. the configurations applied by RRL. It can be used during DTA and RAT.
- 2. The other way is to use the Score-P APAPI metric plugin to trace metrics such as "APAPI_TOT_CYC" to view the CPU frequency and metric plugins such as "upe_plugin" to view the uncore frequency. These metrics show what actually happens in the processor. They can just be applied during RAT.

By comparing the output of both visualization ways mentioned above, it can be easily verified if the settings applied by RRL correspond to what actually happened in the processor.

Using the Score-P metric plugin for viewing the configuration set by RRL When tracing is enabled, Score-P generates OTF2 traces which can be viewed in Vampir. Hence, to get the metrics in trace, the first step is to enable tracing as follows.

export SCOREP_ENABLE_TRACING=true

1. Set the environment variable "SCOREP_METRIC_PLUGINS" to specify the metric plugin for visualization of tuning parameters settings in Vampir.

export SCOREP_METRIC_PLUGINS="scorep_substrate_rrl"

2. Set the environment variable "SCOREP_METRIC_SCOREP_SUBSTRATE_RRL" to specify the tuning parameters which need to be added to trace. For the hardware and software tuning parameters, names of the parameter control plugins (PCPs) are used. The list of

4.2 Visualise Configuration Switching 4 RUNTIME APPLICATION TUNING (RAT)

Tuning Parameter	Parameter Control Plugin
CPU Frequency	cpu_freq_plugin
Uncore Frequency	uncore_freq_plugin
Number of Threads	
Scheduling Type	OpenMPTP
Schedule chunk size	
Energy Performance Bias (EPB)	epb_plugin
MPIR_CVAR_REDUCE_SHORT_MSG_SIZE	${ m mpit}_{-}{ m plugin}$

Table 1: Tuning Parameters and their correspoding PCPs

tuning parameters with their corresponding PCPs are listed in Table 1. If the user wants to load all the hardware and software tuning parameters, it can be done by simply setting the environment variable to "*".

Application Tuning Parameters (ATPs) need to be explicitly specified. To load ATPs, the value should be set equal to 'ATP/<atp_name>' where atp_name is the name of the ATP. The prefix 'ATP/' is required to recognize the ATPs.

export SCOREP_METRIC_SCOREP_SUBSTRATE_RRL="ATP/<atp_name>, <pcp_name>"

For example, the environment variables to specify the metric plugin and request the processor core frequency and uncore frequency to be traced can be set as follows:

```
export SCOREP_METRIC_PLUGINS="scorep_substrate_rrl"
export SCOREP_METRIC_SCOREP_SUBSTRATE_RRL="cpu_freq_plugin,uncore_freq_plugin"
```

An example trace showing the different configurations of core and uncore frequency applied during RAT is given in Figure 2. The Score-P tracing functionality is enabled and the metric plugin is employed during the RAT phase for Blasbench benchmark. The tuning parameters in Figure 2 are named as CPU_FREQUENCY and UNCORE_FREQUENCY. Figure 2 shows the configurations which have been applied at runtime. To confirm that these configurations are actually set in the processor, Score-P APAPI and uncore metric plugins, which are explained next, can be used.

Using the Score-P APAPI and uncore metric plugins To use the APAPI and uncore metric plugins, and to trace the processor core and uncore frequencies, please add the following lines to your script:

```
module load scorep-uncore
module load scorep-apapi
export SCOREP_ENABLE_TRACING=true
export SCOREP_MABLE_PROFILING=false
export SCOREP_METRIC_PLUGINS="apapi_plugin,upe_plugin"
export SCOREP_METRIC_APAPI_PLUGIN="PAPI_TOT_CYC"
export SCOREP_METRIC_APAPI_INTERVAL_US=10000
export SCOREP_METRIC_UPE_PLUGIN="hswep_unc_cbo0::UNC_C_CLOCKTICKS"
export UPE_INTERVAL_US=10000
export SCOREP_EXPERIMENT_DIRECTORY=<location_for_trace_file>
```

The generated trace file will be placed in the folder specified by the environment variable SCOREP_EXPERIMENT_DIRECTORY. This can be viewed using Vampir.



4.2 Visualise Configuration Switching 4 RUNTIME APPLICATION TUNING (RAT)

Figure 2: Vampir trace showing the different configurations of CPU_FREQUENCY and UN-CORE_FREQUENCY applied at runtime for the Blasbench benchmark

Figure 3 shows the trace for the APAPI_TOT_CYC and UNC_C_CLOCKTICKS traced using the APAPI and uncore metric plugins respectively along with the tuning parameters CPU_FREQUENCY and UNCORE_FREQUENCY shown in Figure 2.

Both the traces presented in Figure 2 and Figure 3 are obtained in the same run of Blasbench benchmark. It can be easily visualized in Figure 3 that the APAPI_TOT_CYC trace confirms the trace of CPU_FREQUENCY. The UNC_C_CLOCKTICKS trace in Figure 3 also confirms that the value of UNCORE_FREQUENCY is set as instructed by RRL.

Details about the plugins can be found at: https://github.com/score-p/scorep_plugin_ apapi and https://github.com/score-p/scorep_plugin_uncore.



Figure 3: Vampir trace showing the APAPI_TOT_CYC and UNC_C_CLOCKTICKS recorded using the Score-P APAPI and uncore plugin respectively

A Filtering and Manual Instrumentation

A.1 Runtime Filtering

The first way to reduce the instrumentation overhead is to suppress the measurements done by Score-P for instrumented regions. This is called runtime filtering of regions. READEX provides the **scorep-autofilter** tool that inspects a generated profile and creates a filter file for guiding runtime filtering. This file includes the names of too fine-granular regions that are dominated by the measurement overhead.

1. Apply the scorep-autofilter tool on the profile.cubex file as follows:

```
scorep-autofilter -t <region_granularity_threshold_in_sec>
    -f <filter_file_name_without_extension>
    <path_to_cubex_file>/profile.cubex
```

Choose a value to use as a threshold, for example 100 ms (-t 0.1), for regions to be instrumented. This will create a filter file with .filt extension. The user of the tool-suite can decide the value of the threshold depending on the amount of instrumentation overhead that they wish to retain for the analysis of regions in the application. The higher the threshold value, the lower will be the instrumentation overhead, but also the number of regions accessible to the READEX tool suite.

2. It is advisable but not required to rerun the application and scorep-autofilter to detect additional fine granular regions that were missed in the previous step because their execution time was increased by the measurement overhead of nested regions. This requires that the environment variable SCOREP_FILTERING_FILE is to be set to the filter file name (including the .filt extension) before rerunning the application.

Apply scorep-autofilter to the new profile. Be careful not to overwrite the current filter file. Copy the newly found region names into the original filter file.

Repeat this step until no more regions were found.

Outcome: A filter file with .filt extension containing the application regions that Score-P will not measure.

Section D.2 presents an example.

A.2 Compile-time Filtering

Runtime filtering only suppresses the measurements while the overhead for the probes is still there. You can apply the filter file also during instrumentation of the application to suppress the insertion of probes for the given regions. Please check the Score-P user manual for details on how to perform compile-time filtering. It is advisable that the user do this whenever possible since each existing instrumentation interrupts the program flow during its execution.

In order to apply compile time filtering using the Intel compiler, an additional option needs to specified for scorep-autofilter:

```
scorep-autofilter -t <region_granularity_threshold_in_sec>
        [-f <filter_file_name_without_extension>]
        [-i <intel_filter_file_name_without_extension>]
        <path_to_cubex_file>/profile.cubex
```

A.3 Filtering OpenMP and MPI regionsFILTERING AND MANUAL INSTRUMENTATION

This will create a filter file that can be used by the Intel compiler to suppress instrumentation of regions. This filter file can be passed to the compiler using the option -tcollect-filter=<intel_filter_file_name>.

A.3 Filtering OpenMP and MPI regions

You can remove instrumentation of MPI routines and OpenMP regions as follows:

• Filtering OpenMP regions: To skip the instrumentation of OpenMP regions, the option --thread=none should be used. As a side-effect, no instrumented regions should occur inside of parallel regions. Otherwise, a runtime error will occur. Instead of switching off instrumentation of all OpenMP regions, you can also disable regions selectively via

```
--opari="--disable=omp:single,master,atomic,critical,barrier"
```

This will instrument parallel regions and nested instrumented regions would be handled as expected by Score-P.

• Filtering MPI regions: To disable measurements for MPI routines, you can add the following line to your batch script:

export SCOREP_MPI_ENABLE_GROUPS=ENV

It suppresses measurements for all MPI routines except MPI_Init, MPI_Finalize and other environment routines. These are required for Score-P to work correctly.

A.4 Energy Measurements

Due to potential overhead of energy measurements for application profiling with Score-P, it is necessary to check the overhead when the energy measurements are switched on.

For energy measurements on the Taurus cluster at TU Dresden, load the HDEEM module compatible with the compiler that was used to build the READEX tool suite.

```
module load scorep-hdeem/sync-xmpi-gcc6.3
(or)
module load scorep-hdeem/sync-hdeem2.2.5-intelmpi-intel2017
```

Load the scorep-hdeem sync plugin that is compatible with the Score-P built for the READEX tool suite, and set the following environment variables:

```
export SCOREP_METRIC_PLUGINS=hdeem_sync_plugin
export SCOREP_METRIC_HDEEM_SYNC_PLUGIN_CONNECTION="INBAND"
export SCOREP_METRIC_HDEEM_SYNC_PLUGIN_VERBOSE="WARN"
export SCOREP_METRIC_HDEEM_SYNC_PLUGIN_STATS_TIMEOUT_MS=1000
```

If the overhead for HDEEM measurements for the application regions is more than a few percent, you need to reduce measurement and instrumentation even further, may be by using manual instrumentation as explained in Section A.5.

A.5 Manual Instrumentation A FILTERING AND MANUAL INSTRUMENTATION

A.5 Manual Instrumentation

If none of the other filtering methods is successful in reducing the overhead to an acceptable level, then manually annotate regions where most of the computation time is spent. You can find these regions with a standard profiler. It is also recommended to instrument the parents of all the significant regions up until the main caller in the hierarchy. This is an optional step which will allow the annotated regions to be used as identifiers for runtime situations.

- 1. Build the application with additional options to disable compiler instrumentation (--nocompiler) and to enable user region instrumentation (--user).
- 2. Manually annotate coarse granular application regions or any other regions that are of interest for tuning using SCOREP_USER_REGION_DEFINE inside the function definition as shown below:

SCOREP_USER_REGION_DEFINE(REGION_HANDLE)
SCOREP_USER_REGION_BEGIN(REGION_HANDLE, "REGION_NAME", SCOREP_USER_REGION_TYPE_COMMON)
// application region
SCOREP_USER_REGION_END(REGION_HANDLE)

Note: You also have to instrument the main routine.

Section D.4 presents an example.

B Application Tuning Parameter (ATP) Library

As explained earlier, it is also possible to optionally exploit application level tuning using the READEX tool suite. This requires some additional manual code annotation and instrumentation to pinpoint the parts of the code that can be exploited as application tuning parameters and annotate them with certain API functions. Note that the ATP library can only be used to exploit intra-phase dynamism.

B.1 Instrumentation for ATP library

- 1. Include the atplib.h header file in the source code.
- 2. Declare the parameter in the source code using ATP_PARAM_DECLARE function. Each parameter must contain a unique name, type, default value, and domain name (uses default domain if domain name is NULL):

ATP_PARAM_DECLARE("PARAM_NAME", ATP_PARAM_TYPE_RANGE, DEFAULT_VALUE, "DOMAIN_NAME");

Available ATP parameter types are:

- ATP_PARAM_TYPE_RANGE defines a range with min, max and step values
- ATP_PARAM_TYPE_ENUM defines an array of all possible values
- 3. Add values to the parameter using ATP_ADD_VALUES. The second parameter is an array of values added to the parameter, the third parameter is the number of values added.

ATP_ADD_VALUES("PARAM_NAME", {1,5,1}, 3, "DOMAIN_NAME");

- If parameter type is range, the number of values should be 3 and the values array should contain {min_value, max_value, step}.
- If the parameter type is enum, then the values array should contain all the possible values that the parameter can have, and the number of values parameter indicates how many values are in this array.
- 4. Add the call for parameter value assignment. This assigns the parameter value to control_variable. The value is assigned by RRL. In case no value is available to RRL, the default parameter value defined in ATP is used:

ATP_PARAM_GET("PARAM_NAME", &control_variable, "DOMAIN_NAME");

5. Add constraint to the parameters of domain "DOMAIN_NAME" (optional):

ATP_CONSTRAINT_DECLARE("CONSTRAINT_NAME", "expr", "DOMAIN_NAME");

- The constraint is expressed in the form of a character string "expr" which contains a logical expression of how parameters in this domain are constrained (see example in Section D.5).
- Any ATP parameters declared in the application can be used in the constraint as long as they belong to the same domain as the constraint.
- Multiple constraints can be defined for the same domain.

- B.2 Using the ATP Library B APPLICATION TUNING PARAMETER (ATP) LIBRARY
 - If the domain name is not specified (NULL) the constraint will apply to parameters in the default domain.

Section D.5 presents an example.

B.2 Using the ATP Library

- 1. Build the application by linking with the ATP library (-latp).
- 2. Specify a search algorithm for the ATP library from among exhaustive_atp and individual_atp strategies. This is done by adding sections in the READEX configuration file (readex_config.xml) used as input for PTF during DTA as shown below:

For the individual strategy, the *keep* factor is always 1. Updating/extending the READEX configuration file was explained in detail in Section 3.2.

- 3. Running the application: there are two phases for running the application with ATP:
 - parameter collection phase parameters, constraints and explorations defined in application are collected and saved for the tuning system to explore.
 - parameter exploration phase declaration functions are turned off and the tuning system can explore the parameter combinations by providing parameter values through the ATP_PARAM_GET function.

There are two ATP modes available that allow to enable which phase will be used in the application, although the parameter collection phase needs to be run at least once for the application to allow parameter collection and ATP configuration file creation.

- DTA mode:
 - Includes both ATP phases.
 - ATP_EXECUTION_MODE environment variable should be set to DTA.
 - The name and location of the ATP description file can be set by the ATP_DESCRIPTION_FILE environment variable. If this variable is not set then the ATP description file will be created in the current working directory as ATP_description_file.json.
 - Starts with parameter collection phase: parameter, constraint and exploration declaration functions are executed only once.
 - The second time the same parameter declaration is executed, it triggers the end of parameter collection phase, generates the ATP description file and begins the exploration phase.
 - ATP_PARAM_GET assigns parameter values decided by RRL. In the first phase, the default value is used.

• RAT mode:

- Only the parameter exploration phase is running.
- The ATP_EXECUTION_MODE environment variable should be unset or set to RAT.
- Declaration functions are shut down, only <code>ATP_PARAM_GET</code> function is working.
- Details of parameters are loaded from the ATP description file.

C Retrieving CPU Frequencies

In Linux-based systems,

• the available CPU core frequencies can be retrieved as follows:

```
cat /sys/devices/system/cpu/cpu0/cpufreq/scaling_available_frequencies
```

For Intel processors, make sure that intel_pstate is not loaded.

• the minimum and maximum values for the CPU uncore frequency can be retrieved as follows:

```
sudo modprobe msr
# Minimum frequency
echo "'sudo rdmsr 0x620 --bitfield 14:8 --decimal'00"
# Maximum frequency
echo "'sudo rdmsr 0x620 --bitfield 6:0 --decimal'00"
```

Alternatively, Likwid can be used to retrieve the CPU core and uncore frequencies as follows:

• the available CPU core frequencies (in GHz) can be retrieved as follows:

likwid-setFrequencies -1

• the available CPU uncore frequency (including the minimum and maximum values) can be retrieved as follows:

likwid-setFrequencies -p

Alternatively, x86_adapt can be used to retrieve the CPU uncore frequencies as follows:

```
# Minimum frequency (in 100 MHz)
x86a_read -n -i Intel_UNCORE_MIN_RATIO
# Maximum frequency (in 100 Mhz)
x86a_read -n -i Intel_UNCORE_MAX_RATIO
```

D Examples

D.1 Modules on Taurus

The tools in the READEX tool suite are accessible through modules created either by the continuous integration process or the beta release of the tool suite. Users in the p_readex group may use either, while those in p_readextest can only use the beta release.

Depending on the choice of compilers used for the application (GCC or Intel), load one of these modules to use the READEX tools that are required to analyse and tune an application at the different steps in the workflow.

D.1.1 Continuous integration

Load the continuous integration modules on Taurus as follows:

• For gcc/6.3.0 and bullxmpi/1.2.8.4:

```
module use /projects/p_readex/modules
module load readex/ci_readex_bullxmpi1.2.8.4_gcc6.3.0
```

• For intel/2017.2.174 and intelmpi/2017.2.174:

```
module use /projects/p_readex/modules
module load readex/ci_readex_intelmpi2017.2.174_intel2017.2.174
```

D.1.2 Beta release

Load the beta release modules on Taurus as follows:

• For gcc/6.3.0 and bullxmpi/1.2.8.4:

module load readex/beta_gcc6.3.0_bullxmpi1.2.8.4

• For intel/2017.2.174 and intelmpi/2017.2.174:

module load readex/beta_intel2017.2.174_intelmpi2017.2.174

D.2 Runtime Filtering

Apply scorep-autofilter as follows:

scorep-autofilter -t 0.1 -f scorep scorep-*/profile.cubex

The file scorep.filt contains the region names to be filtered enclosed between SCOREP_REGION_NAMES_BEGIN and SCOREP_REGION_NAMES_END, as shown below:

```
SCOREP_REGION_NAMES_BEGIN
EXCLUDE
Atom::Atom()
Atom::~Atom()
...
SCOREP_REGION_NAMES_END
```

D.3 MiniMD Phase Region Annotation

On the Taurus cluster at TU Dresden, a script to repeat the identification of too fine-granular regions for the miniMD application is available in

```
/projects/p_readextest/miniMD/run_saf.sh
```

and is executed as

sh run_saf.sh

For different applications, run_saf.sh can be reused by updating the line to execute the application. This script requires do_scorep_autofilter_single.sh that is present in the same directory.

D.3 MiniMD Phase Region Annotation

```
void Integrate::run(Atom &atom, Force* force, Neighbor &neighbor,
                    Comm & comm, Thermo & thermo, Timer & timer)
{
 int i, n;
 comm.timer = &timer;
 timer.array[TIME_TEST] = 0.0;
 int check_safeexchange = comm.check_safeexchange;
 mass = atom.mass;
 dtforce = dtforce / mass;
  #pragma omp parallel private(i,n)
   SCOREP_USER_REGION_DEFINE(R1)
   for(n = 0; n < ntimes; n++)
    {
      SCOREP_USER_OA_PHASE_BEGIN(R1, "INTEGRATE_RUN_LOOP", 2)
     #pragma omp barrier
     x = \&atom.x[0][0];
     v = &atom.v[0][0];
     f = &atom.f[0][0];
     xold = &atom.xold[0][0];
     nlocal = atom.nlocal;
      initialIntegrate();
      #pragma omp barrier
      #pragma omp master
      timer.stamp();
      if((n + 1) % neighbor.every)
      {
        #pragma omp barrier
        comm.communicate(atom);
        #pragma omp master
        timer.stamp(TIME_COMM);
        #pragma omp barrier
      }
      else
      {
       {
          if(check_safeexchange)
          {
            #pragma omp master
              double d_max = 0;
```

3

r

```
for(i = 0; i < atom.nlocal; i++)</pre>
        ſ
          double dx = (x[3 * i + 0] - xold[3 * i + 0]);
          if(dx > atom.box.xprd) dx -= atom.box.xprd;
          if(dx < -atom.box.xprd) dx += atom.box.xprd;</pre>
          double dy = (x[3 * i + 1] - xold[3 * i + 1]);
          if(dy > atom.box.yprd) dy -= atom.box.yprd;
          if(dy < -atom.box.yprd) dy += atom.box.yprd;</pre>
          double dz = (x[3 * i + 2] - xold[3 * i + 2]);
          if(dz > atom.box.zprd) dz -= atom.box.zprd;
          if(dz < -atom.box.zprd) dz += atom.box.zprd;</pre>
          double d = dx * dx + dy * dy + dz * dz;
          if(d > d_max) d_max = d;
        7
        d_max = sqrt(d_max);
        if((d_max > atom.box.xhi - atom.box.xlo) || \
           (d_max > atom.box.yhi - atom.box.ylo) || \
           (d_max > atom.box.zhi - atom.box.zlo))
          printf("Warning: Atoms move further than your subdomain size, \
                  which will eventually cause lost atoms.
\n" \
          "Increase reneighboring frequency or choose a different processor grid \
          "Maximum move distance: %lf; Subdomain dimensions: %lf %lf %lf ^{,} \
          d_max, atom.box.xhi - atom.box.xlo, \
          atom.box.yhi - atom.box.ylo, \
          atom.box.zhi - atom.box.zlo);
      }
    }
    #pragma omp master
    timer.stamp_extra_start();
    comm.exchange(atom);
    comm.borders(atom);
    #pragma omp master
    ſ
      timer.stamp_extra_stop(TIME_TEST);
      timer.stamp(TIME_COMM);
    7
    if(check_safeexchange)
      for(int i = 0; i < 3 * atom.nlocal; i++) atom.xold[i] = atom.x[i];</pre>
  #pragma omp barrier
  neighbor.build(atom);
  #pragma omp barrier
  #pragma omp master
  timer.stamp(TIME_NEIGH);
force->evflag = (n + 1) % thermo.nstat == 0;
force->compute(atom, neighbor, comm, comm.me);
#pragma omp master
timer.stamp(TIME_FORCE);
if(neighbor.halfneigh && neighbor.ghost_newton)
{
  comm.reverse_communicate(atom);
  #pragma omp master
  timer.stamp(TIME_COMM);
v = \&atom.v[0][0];
f = \&atom.f[0][0];
nlocal = atom.nlocal;
#pragma omp barrier
finalIntegrate();
#pragma omp barrier
if (thermo.nstat) thermo.compute (n + 1, atom, neighbor, force, timer, comm);
```

```
SCOREP_USER_OA_PHASE_END(R1)
}
//end OpenMP parallel
}
```

This example is also available on the Taurus cluster at TU Dresden in

```
/projects/p_readextest/miniMD/integrate.cpp
```

D.4 Manual Instrumentation

```
main()
{
    ...
    integrate.run(...);
    ...
}
void Integrate::run(...)
{
    SCOREP_USER_REGION_DEFINE( REGION_HANDLE )
    SCOREP_USER_REGION_BEGIN( REGION_HANDLE, "REGION_NAME", SCOREP_USER_REGION_TYPE_COMMON )
    // application region
    SCOREP_USER_REGION_END( REGION_HANDLE )
}
```

Example For the miniMD application, manually annotate ForceLJ::compute_halfneigh() and its parents Integrate::run() and main() as significant regions as shown in the following files respectively, which are available on the Taurus cluster at TU Dresden:

```
/projects/p_readextest/miniMD/force_lj.cpp
/projects/p_readextest/miniMD/integrate.cpp
/projects/p_readextest/miniMD/ljs.cpp
```

D.5 Application Tuning Parameter (ATP) Instrumentation

```
void foo(){
    int atp_cv;
    ATP_PARAM_DECLARE("solver", ATP_PARAM_TYPE_RANGE, 1, "DOM1");
    int solver_values[3] = {1,5,1};
       //{1,5,1} means a range with a minimum value of 1, a maximum one of 5 and an increment of 1
    ATP_ADD_VALUES("solver", solver_values, 3, "DOM1");
ATP_PARAM_GET("solver", &atp_cv, "DOM1");
    switch (atp_cv){
       case 1:
          // choose algorithm 1
          break;
      case 2:
          // choose algorithm 2
          break;
      . . .
 }
    int atp_ms;
    ATP_PARAM_DECLARE("mesh", ATP_PARAM_TYPE_RANGE, 40, "DOM1");
    int mesh_values[3] = {0,120,10};
```

D.6 Tuning Potential Analysis

1. The miniMD application with manually annotated phase region is built for readex-dyn-detect as follows:

make openmpi PREP="scorep --online-access --user --thread=none"

2. When miniMD is run with in2.data as its input file and readex-dyn-detect is applied on the resulting tupled profile.cubex as follows, the function ForceLJ::compute_halfneigh() is identified as the significant region.

readex-dyn-detect -t 0.001 -p INTEGRATE_RUN_LOOP -c 10 -v 10 -w 10 scorep-<xyz>/profile.cubex

Here, readex-dyn-detect takes the granularity for the region as 1 ms with -t 0.001. The option -p INTEGRATE_RUN_LOOP is given to the tool to identify the phase region from the profile.cubex call tree. The three options -c 10 -v 10 -w 10 define thresholds for the compute intensity variation (absolute value), time deviation in % of the mean region time and weight of the region (%) which is execution time w.r.t. phase time.

On the Taurus cluster at TU Dresden, a script to perform steps 1 and 2 for the miniMD application is available in

/projects/p_readextest/miniMD/run_rdd.sh

and is executed as

sh run_rdd.sh

For different applications, run_rdd.sh can be reused by updating the line to execute the application. This is to be run from the location with the application's executable and the filter file name considered to be scorep.filt.

The following lines are printed as part of the output by readex-dyn-detect for miniMD:

D.6 Tuning Potential Analysis

```
\mathbf{2}
 3
 4
 \mathbf{5}
 6
 7
 8
 9
10
11
12
13
14
15
16
17
18
19
20
^{21}
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
```

1

. . .

```
Significant regions are:
void Comm::borders(Atom&)
void ForceLJ::compute_halfneigh(Atom&, Neighbor&, int) [with int EVFLAG = 0; int GHOST_NEWTON = 1]
void ForceLJ::compute_halfneigh(Atom&, Neighbor&, int) [with int EVFLAG = 1; int GHOST_NEWTON = 1]
void Neighbor::build(Atom&)
Significant region information
                                                           Time Dev.(%Reg) Ops/L3miss
                                                                                          Weight(%Phase)
Region name
                             Min(t)
                                              Max(t)
void Comm::borders(Atom&)
                             0.001
                                                0.001
                                                                 2.6
                                                                                109
                                                                                                 0
                                                                                                68
void ForceLJ::compute_hal
                                                0.014
                                                                 2.9
                                                                                 97
                             0.013
void ForceLJ::compute_hal
                             0.016
                                                0.016
                                                                 0.0
                                                                                 91
                                                                                                 1
void Neighbor::build(Atom
                             0.047
                                                0.048
                                                                 0.7
                                                                                332
                                                                                                23
Phase information
Min
                     Max
                                           Mean
                                                                Dev.(% Phase)
                                                                                      Dyn.(% Phase)
0.0138626
                     0.0664566
                                           0.020337
                                                                72.731
                                                                                      258.612
. . .
SUMMARY:
Inter-phase dynamism due to variation of the execution time of phases
No intra-phase dynamism due to time variation
Intra-phase dynamism due to variation in the compute intensity of the following important significant
regions
void ForceLJ::compute_halfneigh(Atom&, Neighbor&, int) [with int EVFLAG = 0; int GHOST_NEWTON = 1]
void Neighbor::build(Atom&)
```

The printed output above for the miniMD application can be divided into three parts:

First, lines 2–7 list the names of the significant regions computed from the detection algorithm. For details of the algorithm, please see deliverable D2.1.

Secondly, lines 10-26 show the profile statistic output for the detected significant regions and phase region. This section consists of two parts. The significant region information presents the minimum and the maximum of the execution time for each significant region as well as the aggregated execution time for the region. It also prints the time deviation in % with respect to its mean value. The Ops/L3miss column prints the absolute compute intensity value. In the last column, Weight(%Phase), is the execution time with respect to phase time.

After that, the tool summarises the statistics information for the phase region. It shows the minimum, maximum, and mean values of the execution time spent on the phase region as well as the aggregated execution time for the phase. The Dev. (% Phase) column prints the time deviation w.r.t. the phase mean execution time. The last column, Dyn. (% Phase), prints the variation between minimum and maximum execution time w.r.t. the mean execution time of the phase.

Finally, the tool prints the summary results of the dynamism analysis (lines 28–40). First, if the standard deviation of the phase is larger than the variation threshold, then the tool indicates having inter-phase dynamism due to variation of the execution time of phases. Otherwise, the

D.7 Tuning Model Creation

application does not have inter-phase dynamism. For miniMD, the variation is larger than the threshold. So the tool detects inter-phase dynamism for miniMD.

The tool compares Weight (%Phase) with the given threshold given by the user. If a significant region has enough weight and its time deviation w.r.t. region is more than the time deviation threshold given via -v, the tool detects intra-phase dynamism for these significant region(s) due to time variation. For miniMD, there are two significant regions having weights larger than the given threshold (> 10%):

void ForceLJ::compute_halfneigh(Atom&, Neighbor&, int) [with int EVFLAG = 0; int GHOST_NEWTON = 1]
void Neighbor::build(Atom&)

But neither of them has a time deviation greater than 10%. So the tool does not detect intraphase dynamism due to time deviation for miniMD.

The tool computes the variation of the compute intensity for the set of detected significant regions having a minimum weight of 10%. For miniMD the variation value is larger than the provided threshold of compute intensity specified with -c. So the tool detects intra-phase dynamism due to the variation in the compute intensity characteristic and lists the region names that exhibit intra-phase dynamism.

D.7 Tuning Model Creation

On the Taurus cluster at TU Dresden, a batch job script to apply PTF for design-time analysis and create a tuning model for the miniMD application is available in

/projects/p_readextest/miniMD/run_ptf.sh

and is submitted as

sbatch run_ptf.sh

For different applications, run_ptf.sh can be reused by updating the command to run the application in --apprun. This script is to be run from the location with the application's executable.

```
#!/bin/sh
1
2
     #SBATCH --time=5:00:00
                               # walltime
3
     #SBATCH --nodes=2 # number of nodes requested; 1 for PTF and remaining for application run
4
     #SBATCH --tasks-per-node=8 # number of processes per node for application run
\mathbf{5}
     #SBATCH --cpus-per-task=1
6
     #SBATCH --exclusive
\overline{7}
     #SBATCH --partition=haswell
8
     #SBATCH --mem-per-cpu=2500M
9
                                     # memory per CPU core
     #SBATCH -J "miniMD_PTF" # job name
10
     #SBATCH -A p_readex
11
12
13
     echo "run PTF begin."
14
15
     NP=8 # check against --ntasks and tasks-per-node
16
17
     module purge
18
     module use /projects/p_readex/modules
```

```
#module load readex/beta_gcc6.3
19
     module load readex/ci_readex_bullxmpi1.2.8.4_gcc6.3.0
20
21
     INPUT_FILE=in3.data #in.lj.miniMD
22
     PHASE=INTEGRATE_RUN_LOOP
23
^{24}
     export LD_LIBRARY_PATH=$LD_LIBRARY_PATH:/usr/local/lib
25
26
     export SCOREP_SUBSTRATE_PLUGINS=rrl
27
     export SCOREP_RRL_PLUGINS=cpu_freq_plugin,uncore_freq_plugin
28
     export SCOREP_RRL_VERBOSE="WARN"
29
30
     module load scorep-hdeem/sync-xmpi-gcc6.3
31
     export SCOREP_METRIC_PLUGINS=hdeem_sync_plugin
32
     export SCOREP_METRIC_PLUGINS_SEP=";
33
     export SCOREP_METRIC_HDEEM_SYNC_PLUGIN_CONNECTION="INBAND"
34
     export SCOREP_METRIC_HDEEM_SYNC_PLUGIN_VERBOSE="WARN"
35
     export SCOREP_METRIC_HDEEM_SYNC_PLUGIN_STATS_TIMEOUT_MS=1000
36
37
38
     export SCOREP_MPI_ENABLE_GROUPS=ENV
39
40
     psc_frontend --apprun="./miniMD_openmpi_ptf -i $INPUT_FILE"
41
                   --mpinumprocs=$NP
                   --ompnumthreads=1
42
43
                   --phase=$PHASE
                   --tune=readex_intraphase
44
45
                   --config-file=readex_config.xml
                   --force-localhost
46
                   --info=7
47
48
                   --selective-info=AutotuneAll,AutotunePlugins
49
     echo "run PTF done."
50
```

To use the RAPL counter energy plugin change from lines 22–24 with the following:

```
1module load scorep_plugin_x86_energy2export SCOREP_METRIC_PLUGINS=x86_energy_sync_plugin3export SCOREP_METRIC_X86_ENERGY_SYNC_PLUGIN=*/E4export SCOREP_METRIC_PLUGINS_SEP=";"5export SCOREP_METRIC_X86_ENERGY_SYNC_PLUGIN_CONNECTION="INBAND"6export SCOREP_METRIC_X86_ENERGY_SYNC_PLUGIN_VERBOSE="WARN"7export SCOREP_METRIC_X86_ENERGY_SYNC_PLUGIN_STATS_TIMEOUT_MS=1000
```

D.8 Production Run with Tuning Model

On the Taurus cluster at TU Dresden, a batch job script is available in

/projects/p_readextest/miniMD/run_rrl.sh

and is submitted as

sbatch run_rrl.sh

For different applications, run_rrl.sh can be reused by updating the command to run the application in lines 41 and 81. This script is to be run from the location with the application's executable. The following script can be used to tune the application during its production run with RRL and compare the execution time and energy consumption with an untuned run of the application.

```
#!/bin/sh
1
2
     #SBATCH --time=2:00:00
3
     #SBATCH --nodes=1
4
     #SBATCH --ntasks=8
\mathbf{5}
     #SBATCH --tasks-per-node=8
6
     #SBATCH --cpus-per-task=1
7
     #SBATCH --exclusive
8
     #SBATCH --partition=haswell
9
     #SBATCH --mem-per-cpu=2500M
10
11
     #SBATCH -J "miniMD_rrl"
     #SBATCH -A p_readex
12
13
     module use /projects/p_readex/modules
14
     module load readex/ci_readex_bullxmpi1.2.8.4_gcc6.3.0
15
16
     energy_label="Energy"
17
18
     rm -rf host_names.out
19
     srun -N 1 -n 1 --ntasks-per-node=1 -c 1 hostname >> host_names.out
20
21
     #####
22
     # application-specific setup here
     INPUT_FILE=in3.data #in.lj.miniMD
23
24
     #####
25
     export LD_LIBRARY_PATH=$LD_LIBRARY_PATH:/usr/local/lib
26
27
     28
     export SCOREP_ENABLE_PROFILING="false"
29
     export SCOREP_ENABLE_TRACING="false"
30
     export SCOREP_SUBSTRATE_PLUGINS=""
31
     export SCOREP_RRL_PLUGINS=""
32
33
     export SCOREP_RRL_TMM_PATH=""
     export SCOREP_MPI_ENABLE_GROUPS=ENV
34
35
36
     # start measurements
     srun -N 1 -n 1 --ntasks-per-node=1 -c 1 clearHdeem
37
     srun -N 1 -n 1 --ntasks-per-node=1 -c 1 startHdeem
38
     start_time=$(($(date +%s.N)/100000))
39
40
     # run untuned application
     srun ./miniMD_openmpi_plain -i $INPUT_FILE
41
42
     # stop measurements
43
     stop_time=$(($(date +%s%N)/1000000))
     srun -N 1 -n 1 --ntasks-per-node=1 -c 1 stopHdeem
44
45
     srun -N 1 -n 1 --ntasks-per-node=1 -c 1 sleep 5
46
     exec < host_names.out</pre>
     while read host name: do
47
48
      srun -N 1 -n 1 --ntasks-per-node=1 -c 1 --nodelist=$host_name checkHdeem >> hdeem.out
49
     done
50
     # aggregate energy measurements from HDEEM
51
     energy_total=0
52
53
     if [ -e hdeem.out ]; then
54
       exec < hdeem.out</pre>
       while read max max_unit min min_unit average average_unit energy energy_unit; do
55
         if [ "$energy" == "$energy_label" ]; then
56
57
          read blade max_val min_val average_val energy_val
           energy_total=$(echo "${energy_total} + ${energy_val}" | bc)
58
        fi
59
       done
60
       time_total=$(echo "${stop_time} - ${start_time}" | bc)
61
62
       echo ""
       echo "Untuned run: Total time = $time_total ms, Total energy = $energy_total J"
63
64
       rm -rf hdeem.out
65
     fi
     66
67
68
     export SCOREP_ENABLE_PROFILING="false"
69
```

D.8 Production Run with Tuning Model

```
70
     export SCOREP_ENABLE_TRACING="false"
      export SCOREP_SUBSTRATE_PLUGINS="rrl"
71
      export SCOREP_RRL_PLUGINS="cpu_freq_plugin,uncore_freq_plugin"
72
73
      export SCOREP_RRL_TMM_PATH="tuning_model.json"
      export SCOREP_MPI_ENABLE_GROUPS=ENV
74
75
      # start measurements
76
      srun -N 1 -n 1 --ntasks-per-node=1 -c 1 clearHdeem
77
      srun -N 1 -n 1 --ntasks-per-node=1 -c 1 startHdeem
78
      start_time=$(($(date +%s<sup>7</sup>N)/1000000))
79
      # run RRL-tuned application
80
81
      srun ./miniMD_openmpi_ptf -i $INPUT_FILE
      # stop measurmenents
82
      stop_time=$(($(date +%s%N)/1000000))
83
      srun -N 1 -n 1 --ntasks-per-node=1 -c 1 stopHdeem
84
      srun -N 1 -n 1 --ntasks-per-node=1 -c 1 sleep 5
85
86
      exec < host_names.out</pre>
87
      while read host_name; do
       srun -N 1 -n 1 --ntasks-per-node=1 -c 1 --nodelist=$host_name checkHdeem >> hdeem.out
88
89
      done
90
91
      # aggregate energy measurements from HDEEM
      energy_total=0
92
      if [ -e hdeem.out ]; then
93
94
        exec < hdeem.out</pre>
        while read max max_unit min min_unit average average_unit energy energy_unit; do
95
          if [ "$energy" == "$energy_label" ]; then
96
97
            read blade max_val min_val average_val energy_val
            energy_total=$(echo "${energy_total} + ${energy_val}" | bc)
98
          fi
99
100
        done
        time_total=$(echo "${stop_time} - ${start_time}" | bc)
101
102
        echo ""
        echo "RRL-tuned run: Total time = $time_total ms, Total energy = $energy_total J"
103
104
        rm -rf hdeem.out
105
      fi
      106
```