# RunMCv4.0 Manual

#### S. Chekanov

November 28, 2005

HEP division, Argonne National Laboratory, 9700 S.Cass Avenue, Argonne, IL 60439 USA

Main reference: Comm. Phys. Commun. 173 (2005) p.115-198

# 1 Introduction

RunMC is an object-oriented framework aimed to run various FORTRAN Monte Carlo (MC) programs for simulation of the particle production in high-energy collisions between elementary particles. It allows to fill 1D and 2D histograms, as well as to create ntuples (ROOT trees). The program is written in C++, and interfaced with ROOT [1,2] and CLHEP [3,4] library classes. To include user-defined histograms, you need a minimum coding. It comes already with a number of histograms, which can be used to look at different event characteristics for many high-energy processes. One can also write a ROOT event tree with selected events for further analysis. This application is open-source licensed, and at this moment, it is available on the LINUX platforms with g++ and f77 compilers.

# 2 Installation under LINUX

Download the file "runmcv[version].tgz" from http://www.desy.de/~chekanov/runmc ([version] denotes the current RunMC version). Then "gunzip" and "untar" this file, i.e. "gunzip runmc[version].tar.gz; tar -vxf runmcv[version].tar". This creates the directory RunMC[version]. To install the RunMC program from the source file, you need: g++, g77 compilers. Also, you have to have "sed", "zip/unzip" and "acroread" to read the user manual. Normally, all these packages are already installed on Linux PCs.

Before further installation, you should have:

- 1. ROOT C++ package, http://root.cern.ch/;
- 2. CERNLIB library, http://wwwasd.web.cern.ch/wwwasd/cernlib/version.html;
- 3. CLHEP class library http://wwwinfo.cern.ch/asd/lhc++/clhep/;
- 4. JAVA virtual machine (JRE);

Check JAVA by typing "java -version" . If JAVA is installed, you should see

```
java version "1.5.0_05"
Java(TM) 2 Runtime Environment, Standard Edition (build 1.5.0_05-b05)
Java HotSpot(TM) Client VM (build 1.5.0_05-b05, mixed mode, sharing)
```

or a similar mesage. If not, install recent java JRE (Java Runtime Environment) from the SUN web page http://www.java.com/en/download/manual.jsp.

The simplest way to compile C++ source of the main package is:

- cd RunMC[version]/build i.e. go to the directory containing the package's source code, and edit "install.sh" file: (a) specify the location of the CERNLIB library (default location is CERN\_DIR = /cern/pro/lib); (b) specify the location of the CLHEP library, library itself and the include files;
- RUNMC environmental variable should be set to the RunMC[version] directory, such that path gives the location to the RunMC directory. Then, add \$RUNMC/bin to the PATH variable:

sh family:

export RUNMC=<path>/RunMC
export PATH=\$RUNMC/bin:\$PATH

csh family:

```
setenv RUNMC <path>/RunMC
setenv PATH ${RUNMC}/bin:${PATH}
```

Type "echo \$RUNMC" in the bash shell, to be sure that you set this variable properly. It is also advisable to check your ROOT installation before compiling the RunMC. Type "cd \$ROOTSYS/test" and compile ROOT examples by typing "make". If ROOT is installed correctly, you should be able to compile all examples in the "test" directory.

To compile RunMC, go to the "\$RUNMC/build" directory and type "./install.sh" to compile the package. This script makes some most elementary checks of your settings. The installation takes 3-10 minutes depending on CPU. After the first compilation, next compilations should take significantly less time, since all Monte Carlo libraries should be installed already.

You can remove the program binaries and object files from the source code directory by typing "make cleanall". To remove only object files, do "make clean". To remove the Monte-Carlo-related files, type "make cleanmc".

Here, "\$RUNMC" denotes the installation directory of the RunMC project, and will not be mentioned again; all the directories discussed below are assumed to be located in this directory.

Go to the "proj" directory and type "runmc" to start RunMC GUI. As long as you set \$RUNMC, you can execute "runmc" from any directory.

#### 2.1 Installation problems

This could be due to incorrectly installed ROOT or CLHEP library Check ROOT as: cd "\$RUNMC/build/test; ./check\_root.sh" Check CLHEP as: cd "\$RUNMC/build/test; ./check\_clhep.sh" If by running these scripts you will see errors, then reinstall these libraries

#### 3 Installation under Windows

At present, RunMC can be installed under on Windows with Cygwin/X11.

• Install recent version of Cygwin (free package, see http://www.cygwin.com). Install the base packages, and, in addition, the following packages: zip, unzip, rxvt, g++, g77, make, X11, X11lib (X11 library for X11 development, this is necessary for ROOT installation). Use "UNIX" file system for the CYGWIN installation;

- Install ROOT. How to build ROOT under Cygwin is documented on the ROOT web page.
- Download the main RunMC, runmc[version].tar.gz, and gunzip, and untar the package as described on the RunMC installation page;
- Now set up the bash environmental variables in ".bashrc":

```
# set RUNMC variables
export RUNMC=/home/sergei/work/RunMC  # location of RunMC
export RUNMCwin32='cygpath -da \$RUNMC'  # do not modify
export PATH=$RUNMC/bin:\$PATH
```

if this is done, do "source .bashrc";

In order to use RunMC together with ROOT, first you should start X-windows on Cygwin Read more details how to start X-windows on Cygwin. The best prompt on Cygwin is RXVT, so do not forget to install it.

Next, download the additional package "runmcv[version]-win32.tar.gz" from the installation RunMC web page. This package contains compiled CLHEP and CERNLIB libraries. Do "gunzip runmc[version]-win32.tar.gz; tar -vxf runmc[version]-win32.tar (note the package may have a version number, therefore, use appropriate name for [version]). This will create "build-win32" directory; Move it to \$RUNMC directory, "mv build[version]win32 \$RUNMC/" Now, go to \$RUNMC/build-win32 and type "./install.sh". This should compile the RunMC.

Note that if CYGWIN does not have "libtermcap" library installed, CASCADE Monte Carlo will not be compliled. CASCADE library assumes that this library is located here "/usr/lib/termcap/libtermcap.a"

#### 4 Quick start

Go to the user directory "\$RUNMC/proj" and start RunMC by typing "runmc". You will see JAVA-based GUI (JRunMC). Start the Monte Carlo simulation by selecting. The status panel should show MC models installed on your system. They all should have "idle" state. Select from the top menu "RMC project" and "open RMC". You will see RMC modules which represent particular calculations. Using mouse double click, one can view the information on these modules. Select "default" file and click on "Load". Then click on "Exit". You will see the yellow progress bar indicating the compilation progress. You also should see which MC model is running (red message "running" on the status panel). At the end, another window should show what is inside of this physics project. Then click on "run" on the main window. You should see ROOT canvas with 3 filled histograms. Then you can save these histogram using the ROOT menu.

The same can be done with any RMC project files.

### 5 Starting JRunMC

When you execute "runmc" (which is a link to jrunmc bash script), JRunMC.jar executable (located in "main/jrunmc") reads executables located in "bin" directory and check their status ("idle" or "running"). Then JRunMC builds a database from the physics project files (RMC) which are located in "archive" directory of the main installation RunMC directory. Each \*.rmc file is a gzipped user directory "proj". This directory contains user files with physics calculations, thus files in this directory can be modified.

To read information on each \*.rmc file, JRunMC reads XML file "user-comment.xml'. You can manually unzip \*.rmc file and look what is inside. One RMC file is always unzipped - the user directory "proj" - which corresponds to the current user calculation.

#### 6 JRunMC user interface in more details

At present, the following MC models are included to RunMC: PYTHIA6.3 [5], HERWIG 6.5 [6], ARIADNE 4.12 [7], LEPTO 6.5 [8], AROMA 2.2 [9], CASCADE 1.2 [10], PHO-JET 1.05 [11], RAPGAP 3.1 [12]. When JRunMC is used, one can choose one of these MC programs. The final state used in your calculation (or/and in the RUNMC output trees) can be: all stable particles, "partons", "charged particles", "all particles" (which is equivalent to the HEPEVT record), "stable+all" (fill both stable particles and, in addition, complete HEPEVT record), and "user-defined". The option "stable+all" allows to reconstruct some quantity from the final state (PP or PJ vectors, see the definition of the "heplist" class), and at the same time one can preselect events using information on all generated particles (HP vector, which is a copy of HEPEVT event record). Obviously, this option is the slowest. The last option is used to fill the final state in the "proj" directory.

Select the number of events to be processed, type of initial particles, the nucleon structure function (PDF). You also can set kinematic cuts applied on the final state in the transverse momentum (PT) or pseudorapidity (Eta) of particles, if particles were selected, or on jets, if your choice was jet reconstruction. If, instead of the "particles" option, one selects "hadronic jets", all kinematic cuts will be applied on the jets, without any kinematic constrains on the particles used to form jets. For DIS, one can set up the event kinematics in Q2, Bjorken x, y, or W.

Then, you can select the histograms to be filled by clicking on the "Histogram" from the "Ouput" panel. You should see the window "Variables". Normally, left window is empty - this means that the user did not specify the variables in "proj/user-name.txt" file. Howver, one can still use some system variables by selecting the option "system variables" below the left pannel.

Use mouse double-click to set the variable. This will add a new histogram to the histogram window (on the right). Then one should set the minimum and maximum range of the histograms, as well as the number of bins. You can also select two-dimensional (2D) histograms using the option from the right side.

You can select up to 8 one-dimensional histograms at the same time (or 4 2D histograms), but in fact, the user can specify up to 500 1D histograms (or 250 2D histograms), but only 8 can be viewed during the run (see below). If you click on "more histograms", you will get a spreadsheet "Histograms", where you can define more than 8

histograms (use the "Variable" window to do this). This spreadsheet should allow you to delete, copy, paste selected histograms.

Then, by clicking on "Options", you can select the size Canvas size, update interval for histograms (the larger number, the less CPU consuming run), color scheme (for 1D or 2D histograms separately, use plain style to look scientific!), and a renice level. Note that the log-scale option at present works only if you set a single histogram for on-line viewing (of course, when histograms will created, one can use ROOT to set log scale). You can also specify whether you want to fill the histograms for events, or use different normalizations. One can normalize histograms to unity, but also one can calculate differential cross sections (in "nb" or "pb") after taking into account the bin size and the luminosity for this MC run. Note that the histogram-normalization option works only for 1D histograms for a moment.

Now you can select the output using "Output" pannel. Three types of the output files are possible:

- The histogram file output. It has the extension ".root";
- The HEPEVT event record with output file, which has the extension ".htup"; The HEPEVT output was slightly modified, to make it simpler. Read Monte Carlo manuals for definitions of the variables. Note: the output file can be very big!
- The RUNMC event record. The output file has the extension ".rtup". This is a reduced HEPEVT output. The output contains only particles selected from the RunMC GUI: all particles, stable particles, charged stable particles;

No any kinematic cut is applied for such output, i.e. you cannot set any cut on the final state from RunMC GUI. To keep the output file small, only particle ID, charge (times 3), mother, daughter and Px, Py, Pz, E of particles are filled. The size of the output file is by a factor 4 smaller than that for the complete HEPEVT record. Note that double precision values were replaced by float values, to keep output small.

Now specify the name of your project on the main window, and save it by clicking on "File - save MC". Another type of projects (which called "RMC", and has the same extension) will be explained latter. The option "Save MC" saves the current status of RunMC GUI into the file project;.mc in the directory "proj". Note, that even if you will not name your project, RunMC will save the present project in the default "analmc.mc" file automatically.

If you do want to preselect events for your RunMC tree, you can do this using three methods:

1. Set appropriate values in the subroutines which initialize the MC programs. All user files are in the "proj" directory (read about this below). FORTRAN subroutines which initialize Monte Carlo models are located in the "proj/ini" directory. For every MC model, there is a separate subroutine, in which new parameters can be specified. (You have to read MC manuals for this, and use FORTRAN style of coding, of course). Then do "make" from the "proj" directory. It will rebuild all MC binary files in the "bin" directory. If you do this, the original MC HEPEVT record will contain the preselected events you want. The event class of RUNMC will also contain only preselected events, since this class is derived from the HEPEVT.

Read about the "proj" directory in more details below. Note that this option is not preferable, but should be used if nothing else can help. Instead, use the steering cards (see below).

- 2. Another way to change MC parameters is to use the steering cards. Examples of the steering cards are located in the "example/steer/" directory. Copy necessary steering card to a directory were you execute JRunMC (i.e. where you type "runmc". The steering cards will be loaded automatically. Note than in the parameter settings, the functions in "proj/ini" directory are always the last one, i.e. parameters in steering cards can be overwritten by the settings in the "proj/ini" functions. The MC settings is the most complicated part, please carefully check the output log file.
- 3. One can preselect specific events, while generating fully inclusive events. Again, go to the "proj" directory, and put your user calculations in "user-select.cpp". By default, the function returns "0", i. e. it takes the event. If this function will return any other number, the event will not be taken. Note that this only affects the RUNMC tree (i.e. file "project\_name.rtup"). The event selection done in the "user-select.cpp" file does not change events in the original HEPEVT record, as well as in the simplified HEPEVT tree (i.e. the file "project\_name.htup"), which is simply a copy of the FORTRAN HEPEVT common block. In addition, such selection has no affect on the event class used to make histograms on-line. The event class "main/inc/heplist.h" is described below. After a modification in "proj/user-select.cpp", type "make" to recompile the project. Read about the "proj" directory below.

Now select predefined histograms, and set minimum and maximum values and the number of bins. To start the run, click "run'. This will check all your setting, as well as will display the output file options. After 1-5 seconds, you will see the ROOT canvas with histograms updated in real time. Note that for some models, like AROMA or CASCADE, this takes more time since these programs do some initial integrations before they start looping over events. You will see a blue progress bar indicating how many events were processed.

To verify log file generated by RunMC, go to the "Control" panel and click on "view log". This executes "JRunMC Notepad" - an editor with the log file (".runmc.log"). The log file can also be found in your current directory, look at "project\_name.log" file.

In addition to the log file, there a file with possible errors "project\_name.err". If it is not empty, you will be notified by JRunMC.

You can stop the run at any time by clicking on "Stop". (You have to wait for 3-10 sec after this, since the program needs to finish all calculations. This however depends on type of calculations.). When run is finished, you will see a pop-up window with the run information.

After the run finished, you should see a new button "View output" (below "More histograms" button), which has light color. Click on this button: you should see ROOT Browser with the opened file project;.root.

Then, using the ROOT built-in graphics editor, you can modify the canvas, i.e. change scale, put/remove labels etc. To exit, you should click on "Exit". It is advisable to quite the ROOT canvas fist, before you exit RUNMC. All files can be saved as a ROOT

file, "project\_name.root". If you do not want to look at the histograms latter using "project\_name.root file", go to "Select output" to change this setting.

If something goes wrong, click on "Cancel". It kills all the runing programs associated with the RunMC program. Note: if you are running some RunMC executables in the background, they will also be killed without saving the results!).

# 7 Saving and loading the RunMC project file

You can save all you current JRunMC settings to a project file (by default, it has has the name "default.mc"). The extension "mc" is important. Go to the menu "File" and save this file as "MC" file.

You can restore this project using the same "File" menu ("Read MC" item);

There is also a fast way to load project without using the "File" menu. In the shell prompt, type "runmc project.mc", where "project.mc" is the name of your input file.

# 8 The RMC physics projects

You cannot do complicated studies just by using JRunMC. First of all, only a small number of histograms are available from JRunMC. Monte Carlo settings also cannot be changed flexibly enough. Note that if you will copy steering cards to the directory where you execute "runmc", you can change some Monte Carlo parameters by modifying the steering cards. Examples of the steering cards are located in "example/steer" directory.

As was mentioned above, the user can include personal histograms and modifications to the MC settings by changing subroutines in the "proj" directory. This directory contains:

- 1. At least one "project.mc" file which defines the status of RunMC. This file can be loaded to RunMC GUI via "File-read MC", or using the command line: "runmc project.mc"
- 2. One file "user-comment.xml". It has comments on the structure of this projects, and is shown automatically when an RMC project is loaded. It has some useful information on how to run this project (in XML format). This file can be edited the menu "RMC project RMC info editor". You can view HTML formated information using "RMC project RMC info". In fact, the user can edit the file "user-comment.xml" using any text editor.
- 3. Directory "ini" with "dummy" subroutines for each MC model. Here you can set MC parameters using FORTRAN codings (read MC manuals how to do this. All MC manuals are located in "main/mcarlo" directory).
- 4. , "user-init.cpp", "user-end.cpp", "user-run.cpp" project initialisation, analysis and terminations subroutines. The RMC file "par-had-jetLHC.mc" illustrates how to use these files.
- 5. "user-select.cpp" here you can set any cuts you want on hadronic final state or original HEPEVT record (if option "stable+all" used).

- 6. "user\_afill.cpp" defines user variables for each event, each particle (1-particle density), or each 2-particle combinations (2-particle density). The same file is used to fill histograms for jets. Read comments in this file. One may also look at a very similar system file "main/src/afill.cpp".
- 7. This directory can contain steering cards for MC models. Settings in these cards can be overwritten by subroutines in "ini" directory.
- 8. "Makefile" and "Makefile\_in". You need this to compile your project by typing "make". The same make files are used when you load RMC project using "RMC project open RMC";
- 9. "user-name.txt" file which attributes names to your personal histograms which will be automatically included to RunMC during compilation. All names here should be exactly as in the "user\_afill.cpp" file.

The structure of the "user-name.txt" is:

```
# this is just a comment
#
variable1 describe it here
variable2 describe it here
....
```

Use "#" to add a comment. Do not use spaces in the definition of the variable names!.

All the subroutines defined above can be modified, and all executable for MC models can be recompiled by typing "make" from the "proj" directory.

In fact, the directory "proj" is "unpacked" form of the so called "RMC" file (or project), which contains everything you need to do the analysis. At this moment, this directory has only dummy functions. Go to this directory "proj", type "runmc" and save this directory from "RMC project - save RMC". You will see the message "archive/¡project¿.rmc created" (of course, ¡project¿.rmc' could be any name which you type in JRunMC).

The file *project*<sub>i</sub>.rmc is nothing but the "proj" directory, archived and compressed using the "zip" utility. This file is stored in the "archive" directory.

In the directory "archive" you can find a number of files with the extension "rmc". These files contain various physics projects (MC settings+histogram definitions as in the "proj" directory), saved exactly as you did before. This is a list of the present projects: Every time when you start JRunMC, the program read XML file inside such file and build a database.

- 1. "default.rmc" just "proj" directory with dummy functions; "default1.rmc", "default2.rmc" are few examples of how to fill ROOT histograms.
- 2. "dis\_kinematics.rmc" project for DIS related studies;
- 3. "dis\_strange.rmc" project to study strange particles in DIS;
- 4. "event\_shapes.rmc" project to study the event shapes;

- 5. "charm\_dis.rmc" project to study D\* cross sections in DIS;
- 6. "jets\_HERA.rmc" project to study jets at HERA (using the longitudinally invariant algorithm in the Breit frame);
- 7. "jets\_LHC.rmc" project to study jets at LHC; It also illustrates how one can preselect events in the user function "user-select.cpp".
- 8. "jets+charm\_LHC.mc" project to study jets at LHC. It also requires at least one charm particle in an event. This example illustrates how one can preselect events in "user-select.cpp" function. Note: option "stable+all" are used for the final state (fills both PP and HP vectors from heplist class).
- 9. "par-had-jetLHC.rmc"- project to study jets at LHC. This calculates jets at parton and hadron levels, and takes the ratio (the so-called hadronisation correction). This example is very important: it illustrates how to set user-defined calculations via "user-run.cpp" file, without using the internal RunMC functions. Also, the user can set some histogram options "user-init.cpp".
- 10. "view3d.rmc"- project to study top production at NLC. It creates a 3D view of the event, following to a very similar approach of the example tree2.C file in \$ROOT-SYS/tutorials. In this example, RUNMC canvas is modified by the user. The settings for MC are done in "ini" directory

Try to load one of this projects to "proj" directory for testing. Click "RMC project - open RMC". Select one of the files with the extension. Use doble click to read about the selected project. To load the RMC project, click on "Load". You can get recent RMC files from http://www.desy.de/~chekanov/runmc.

When you click on "Load", this automatically recreates "proj" directory by copying new files from the selected RMC file. You will see a yellow message "Wait" and yellow status bar indicating the compilation progress: it starts compilations of all MC models using new files from "proj".

Note that if you had important files in "proj", and you have forgotten to save them before loading a new RMC project, such files will not be lost. Check the "main/tmp" directory. It should contain old "proj" directory (in zip format) with the name "proj\_date.tmp". Note that files which are one week old are removed automatically.

When the compilation process finished, you can see that there are new histogram in RunMC. A new message will appear which contains information on this project (from the "proj/user-comment.xml" file). Say "OK" to close this window. Then, press histogram bottom, and study the list of the histograms. Now you can do necessary modification via RunMC and can start the run. (Note: if histograms in JRunMC were not updated, try to restart "runmc").

You can read comments on the current project using the the menu "RMC Projects-RMC info". This information be modified by using "RMC Projects-RMC info editor"

The new "proj" file should contain at least one file with the extension "mc" which defines the status of RunMC GUI (which can be loaded as "runmc project.mc" or using "File-read MC" option.

To debug files in "proj" directory, just type "make" from the shell prompt.

You can study the RMC project which is currently in "proj" using "RMC project-RMC browser". This executes appropriately modified Jedit editor.

The user can define and fill external variables in "user-run.cpp". Load the "parhad-jetLHC.rmc" project to see how this was done. In this example, several histogram files are defined in the user initialization subroutine, and final states are defined in the "user-run.cpp" file. In this subroutine, jets are reconstructed using an external function "ktjet.cpp", and histograms are filled separately for hadrons and partons. The final state in RunMC GUI was set to "user-defined" final state, since the final state is filled by a user.

# 9 Analyzing ROOT output ntuples and files off-line

RunMC allows to analyse the ntuple files (with the extension "rtup" and "htup") off-line. Just select "Ntuple HEPEVT" or "Ntuple RUNMC", and load the selected files. Always select only one file. Then, if you want to chain over all ntuples, select this option on RunMC GUI. Do not forget to click "check ntuple files". This will show you the list of the files currently selected. All calculations for those files should be done exactly as for real Monte Carlo calculations. (Note that RUNMC ntuple format cannot be used for many calculations since this type of ntuples does not have much information on the event structure).

Once you have the output files, "project.root", "project.rtup", or "project.htup", you can analyze the output using ROOT TBrowser. The output files can be analyzed latter, calculating complicated event characteristics off-line. Look at the "projexampleproj\_hepevt" directory. It contains a demo program which reads "project.htup", and prints all values for each event on the screen. Analogously, "proj/example/proj\_runmc" contains a demo program which reads "README" in these directories for more details.

### 10 RunMC event structure

To include an user-defined histogram or additional calculation, you need to know the event structure. The event structure is defined in the "main/inc/heplist.h" include file.

```
//\ {\rm Class}\ {\rm characterizing}\ {\rm a}\ {\rm particle}
```

```
public:
                                  // 4-momenta
HepLorentzVector p;
HepLorentzVector v;
                                  // vertex
double
                                  // mass
                  mass;
                                  // position in HEPEVT
int
                      pos;
                                  // identity (particle data group)
                      id;
int
                                  // charge times 3
int
                  charge3;
                  status;
                                  // status of entry
int
int
                  mother;
                                  // mother
int
                  mother2;
                                  // point to second mother
int
                  daughter;
                                  // daughter
```

daughter2; // point to second daughter int}; // Class characterizing a jet (under construction) class jet{ public: // 4-momenta HepLorentzVector p; int in; // number of particles in this jet }; // Event record for Monte Carlo events class heplist { public: heplist(); // constructor ~heplist(); // destructor HepLorentzVector beam1; // beam 1 // PID of this particle int id1; // beam 2 HepLorentzVector beam2; int id2; // ID of this particle HepLorentzVector boson; // Px,Py,Pz,E of exchange boson int idboson; // ID of boson HepLorentzVector lepton; // Px,Py,Pz,E of outgoing lepton(DIS) int idlepton; // ID of lepton // Information on separate particles vector<particle> PP; // vector with particles // Information on separate particles (options stable+all) // contains complete HEPEVT record vector<particle> HP; // vector with particles // information on event double PTtot , PZtot, Etot; // event PT, PZ, Etot // information on jets vector<jet> PJ; // vector with jets void fill\_HEPEVT(); // fill all particles from HEPEVT (stable+all option) void fill\_All(); // fill all particles from HEPEVT void fill\_ALLstable(); void fill\_Partons(); // fill stable particles from HEPEVT // fill partons from HEPEVT void fill\_CHARGEDstable(); // fill partons from HEPEVT int set\_jetPtEta(); // set Pt and Eta cuts on particles int set\_particlePtEta(); // set Pt and Eta cuts on jets // transformations for jets int set\_jetLABtoHCMframe(); // move jets to the HCM frame

```
// move jets to the lab from HCM frame
    int
          set_jetHCMtoLABframe();
    int
          set_jetLABtoBREITframe();
                                       // move jets to Breit frame
          set_jetBREITtoLABframe();
                                       // move jets to lab frame from Breit frame
    int
// for sepate particles transformation
          set_particleLABtoHCMframe();
                                          // move jets to the HCM frame
    int
          set_particleHCMtoLABframe();
    int
                                          // move jets to the lab from HCM frame
    int
          set_particleLABtoBREITframe(); // move jets to Breit frame
          set_particleBREITtoLABframe(); // move jets to lab frame from Breit frame
    int
          set_JADEjets();
                                          // move to the JADE jets
    int
          set_DURHAMjets();
                                          // move to the Durham jets
    int
          set_KTjets();
                                          // move to the KT long invariant
    int
                                                                             jets
 };
```

```
11 Structure of HEPEVT and RUNMC outputs
```

If you decide to write the entire event on the hard disk, you can choose between the HEPEVT record or reduced RUNMC record. Let's remind that the HEPEVT record can only be changed by the FORTRAN initialization subroutine "proj/ini" (for each MC separately). But you can do complicated selections of events for a reduced RUNMC record (in "proj/user-select.cpp" file, see above the description, and read the "proj/user-select.cpp" file).

Use the RunMC GUI option "Select output". The HEPEVT output file has "thepevt" tree. It has the following structure:

```
int NEVHEP;
                                 // read HEPEVT output
                                 // NHEP - total number of particles
int NHEP;
int ISTHEP[hepevt_local_max];
                                 // ISTHEP(j)
int IDHEP[hepevt_local_max];
                                 // IDHEP(j)
int CHA3[hepevt_local_max];
                                 // charge times 3
                                 // JMOHEP(1,j) in fortran
int JMOHEP1[hepevt_local_max];
                                 // JMOHEP(2,j) in fortran
int JMOHEP2[hepevt_local_max];
int JDAHEP1[hepevt_local_max];
                                 // JDAHEP(1,j) in fortran
int JDAHEP2[hepevt_local_max];
                                 // JDAHEP(2,j) in fortran
double PHEP_X[hepevt_local_max]; // PHEP(1,j)
double PHEP_Y[hepevt_local_max]; // PHEP(2,j)
double PHEP_Z[hepevt_local_max]; // PHEP(3,j)
double PHEP_E[hepevt_local_max]; // PHEP(4,j) energy
double PHEP_M[hepevt_local_max]; // PHEP(5,j) mass
double VHEP_X[hepevt_local_max]; // VHEP(1,j)
double VHEP_Y[hepevt_local_max]; // VHEP(2,j)
double VHEP_Z[hepevt_local_max]; // VHEP(3,j)
double VHEP_T[hepevt_local_max]; // VHEP(4,j)
```

The RUNMC output contains only particles which you select via the RunMC GUI (all stable, all charged partons etc.). The RUNMC tree is called "trunmc". It has the foolowing structure:

```
const int hepevt_runmc_max =4000;
    int NHEP:
                                     // total number of particles
    int IDHEP[hepevt_runmc_max];
                                     // particle ID
    int ISTCHA[hepevt_runmc_max];
                                     // 100*ISTHEP+(10 + charge times 3)
    int JMOHEP[hepevt_runmc_max];
                                     // JMOHEP(1,j)
                                                    in fortran
    int JDAHEP[hepevt_runmc_max];
                                     // JDAHEP(1,j) in fortran
    float PX[hepevt_runmc_max];
                                     // PHEP(1,j)
    float PY[hepevt_runmc_max];
                                     // PHEP(2,j)
    float PZ[hepevt_runmc_max];
                                     // PHEP(3,j)
    float E[hepevt_runmc_max];
                                     // PHEP(4,j) energy
```

See the example "mrun.cxx" in the "example/proj\_runmc" directory which can read "project.rtup" file. Note it uses "float" instead of "double" precision numbers to save the disk space (and there is no reason to think that you really need double precision momenta of final-state particles even for LHC energies).

# 12 How to include user-defined histograms

The easiest, just load any "RMC" project from "archive" directory using "RMC project - open RMC". You will see changes in "proj" directory, and can study functions copied from RMC. Below are some additional details:

You can include additional histograms in a few steps: Go to the user project directory, "proj", and edit "user-name.txt" file. You have to put the line:

Variable description of this variable

The variable names should be defined as following:

- Should not have spaces in it;
- Any calculation which does not involve looping over all particles/jets should not have the symbol "@" at the beginning;
- If you calculate a single-particle density (looping over all particles in an event), your variable name should have "@" at the beginning. Note that "@" will be removed automatically for histogram titles etc.
- If you calculate two-particle density, i.e. calculate a variable for each particle pair, your variable name should have "@@" at the beginning.

Once you defined the title of your calculation, put the description of the calculation after a space. You can put up to 500 histogram definitions. Each definition should be on a new line.

There are a few types of histograms:

1. A histogram for a value characterizing the entire event. You have to define the variable "\*getval" (the star in front is important!) to do this in the function "user\_afill.cpp". It is very important that the name in "if" statement of this function should match the histogram name in "user-name.txt". You can include as many calculations as you like, but they all should be included using "if-else" statements.

- 2. A histogram representing a variable for each particle. Do this exactly as before, only specify particle counting number in your variable in "user\_afill.cpp" function. See examples in the RMC files. The title of a calculation should contain "@" as explained above.
- 3. A histogram for each particle pair. Do this exactly as before, but specify additional 2 arguments in this variable representing particle number in "user\_afill.cpp" function. The name of the calculation should contain "@@" as explained above.
- 4. A histogram for each jet. In this case use additional variable "jet" string in "if" statement of the function "user\_afill.cpp".

When you did this, you have to recompile the project, type "make clean" and then "make".

You can find a number of examples in "main/src/afill.cpp". You should not modify this file. Better, load one of the physics projects from "archive" as described before.

Note that the name convention described above is not needed when a user fills the histograms in "user-run.cpp" file. See "par-had-jetLHC.rmc" project. In this example, the final state and well as histograms are filled without using RunMC directly. In addition, user can initialize the calculations in "user-init.cpp"

If you now save the project as "File - save MC", you will see that your histograms are added to the end of the file "project.mc".

In total, 500 histograms can be included to your project. Instead of using the spreadsheet "Histograms", one can use any your favourite editor and correct the file "project.mc". Go to the end of this file. For each new histogram, you need to add 2 lines:

H-NAME "Name of the histogram" "comment or definition" H-INFO "Dimension" Xmin Xmax "No of bins" "weight"

where "Dimension" means "1" or "2", i.e. 1D or 2D histogram, Xmin and Xmax are the minimum and maximum values for histogram, and "No of bins" is the number of bins (should be always integer). Xmin and Xmax, weight are real numbers. Note: do not use more than 20 characters to for the title of your histogram.

If you want to use "2D" histograms, set "2" after "H-INFO". The number of histogram with this option should be always even, otherwise RunMC will fail to calculate how many 2D histograms you want and will report an error.

You can also change any option in "project.mc" by hand. To learn about this, use RunMC GUI and study what it does when you create the "project.mc" file.

### 13 User-defined settings for MC models

At this moment, all Monte Carlo models are initialised (via the RunMC GUI) for fully inclusive events (AROMA- for charm events). You can change this if you will include your Monte Carlo settings into the files located in "proj/ini". You have to use FORTRAN coding for this. Please refer to Monte Carlo manuals on how to specify your new Monte Carlo settings (they are in "main/mcarlo/" directories. The names of the subroutines in "proj/ini" are self-explanatory. The subroutine "set-lepto-ariadne.f" corresponds to LEPTO with ARIADNE (for ep DIS only!). When you did this,go to the "proj" directory and recompile it by typing "make". Note: PYTHIA does not work well for DIS, use it for photoproduction.

Another way to change MC parameters is to use steering cards (i.e initialisation files). Examples of the steering cards are located in "example/steer/" directory. Copy necessary steering card to a directory were you run RunMC GUI (i.e. where you execute "runmc"). The steering cards will be included automatically. Note that in the parameters settings, the functions in "proj/ini" directory are always the last one, i.e. parameters in steering cards can be overwritten by the settings in "proj/ini functions.

The steering cards can be loaded and edited using RunMC steering card editor. Use "Settings- Steering card". You should see a spreadsheet-type of editor for the steering-cards, where you can type new parameters for MC models. Note: you loaded RMC file, which may contain some steering cards, it is important to start RunMC from the "proj" directory, otherwise the RunMC steering-card editor will never find the current RMC steering card.

Note that the steering card for HERWIG is still rather simple, i.e. only a few parameters were implemented so far. Look at the "main/mcarlo/herwig-65/RUNMC-herwig.steer.f" to check which parameters can be included. Therefore, the best way to change the HERWIG parameters is to use more flexible function in "proj/ini".

### 14 How to include user-defined RMC projects

There are three functions in the "proj" directory: "user-init.cpp" (initialisation), "userrun.cpp" (for event calculation), "user-end.cpp" (for job termination). You can access the HEPLIST class for every events and do any calculation you want to, including filling of a ntuple. If you have an external function you want to link to the project, you should add an additional line in the file "Makefile\_in" with the name of new file. Then you have to recompile the project from the "proj" directory.

You can also include user-defined labraries, use variable "\$(PROJ\_LIBS)" in Make-file\_in to do this. Look at the example "hztool.rmc".

The example "par-had-jetLHC.rmc" illustrates how to fill user-defined histograms. It does not used the final state option from RunMC GUI (it is set to "user-defined"). Instead, the final states (partons and hadrons) are filled inside "user-run.cpp". Histograms are also filled inside this function. Drawing options and some protection conditions are redefined in the "user-init.cpp" function.

You can save and restore the "proj" directory using "RMC project - save RMC". This will create a file with the extention "rmc" in the "archive" directory. You can restore this file latter using "RMC project - read RMC".

It is very easy to make a personal RMC project. First, load the default RMC project file using "RMC project - open RMC" and select "default.rmc". Then start "RMC browser". Now you can modify any file in the "proj" directory. Alternatively, just go to "proj" and edit the RMC source using any editor.

#### 15 How to access some MC parameters

During MC running, sometimes you need to access MC parameters and plot them. To do this, you should copy them in the array REAL rUSER(50) (from "getevkin.inc"). This

should be done in the main loop of the FORTRAN program.

For example, for PYTHIA6, copy necessary variables to the rUSER array in "main/mcarlo/pythia-62/RUNMC-pythia6.f" after the line "nev=nev+1". Then you can access this variable in any of your function (like "user\_afill.cpp" under "proj") if you specify the include file "#include<getevkin.h>" and put the line "extern getevkincommon getevkin\_;", You variable should be seen as "getevkin\_.rUSER[0]", "getevkin\_.rUSER[1]" etc. Look at the example in "dis\_kinematics.rmc" where the kinematic variables Q2 etc were taken directly from the LEPTO model.

Similarly, you can do the same with any MC model.

#### 16 How to use RunMC without GUI

You can run any MC model without the GUI. You need GUI only to create "name.mc" file and to be able to communicate with the MC. But this can be done without the graphical interface as well.

Before doing this, it is advisable first to run JRunMC and create a project file. Go to "Options" and select "Do not update" and "Do not show plots". Then you can save this project (the default name is "analmc.mc").

Next step is to run this MC model. Look at "bin". This directory has binary files for each MC model. These files read the input file ".runmc.in" located in your current directory. This is just a link to your input project file, say "project.mc". One can edit this file, to set additional settings you want. Then, execute the MC binary file in shell. For example, to run ARIADNE, just type: "runmc.ariadne ¿ runmc.log"

It is likely that you will get the message in the log file:

PROCESS PID=2331 renice 10 2331

2331: old priority 0, new priority 10 output file: analmc

\$RUNMC/pipes/pipe\_<date>.stop has 1

where "date" is a string containing the time when your job was started. This means that the communication file "pipe\_<date>.stop" forbids this run since it has "1". To start the run, you should replace "1" in "pipe\_<date>.stop" by "0". If you set "0" in this file, you can run this MC. If you will want to stop the MC run, again set "1" to "pipes/pipe\_date.stop". After 3-5 seconds, the run should be terminated and all histograms will be saved. Since all pipe files are not identical, you can run as many jobs as you like at the same time.

If you get the message that "pipe\_date.stop" is not find, you should create this file in the "pipes" directory and make sure that the name of this file was set correctly in the "project.mc" file (it should not contain any extension like ".stop"!).

The easiest way to learn this – use first RunMC GUI and analyse changes in the project file "project.mc" file.

In addition to the "pipe\_<date>.stop" file, each run has another file called "pipe\_<date>.events", which contains the number of generated events.

The RUNMC will clean the "pipes" directory from the old files. At present, it will clean pipe files which more than 20 days old.

# 17 How to load RMC projects without GUI

First, remove your current "proj" directory. Copy a RMC file to the top \$RUNMC directory, for example "cp \$RUNMC/archive/default.rmp \$RUNMC/". Then unzip it, "unzip default.rmp". This will create new "proj" directory with new files. Then recompile the project by executing "make" from the "proj" directory.

# 18 Random seeds for MC models

Every time when you start new MC run, seeds for random numbers used by the MC models are different. Seeds are calculated from many numbers which include the current time, job id etc. Just check few first lines in "project.log" file to see that the numbers are indeed different every time you star the run.

# 19 Specific issues how to run MC

### 19.1 PYTHIA, HERWIG, LEPTO, AROMA models

For these models, just include your settings to \$RUNMC/proj/ini" and recompile the project from "proj". In addition to this, you can use steering cards in your current directory. Examples of the steering cards are in "example/steer/(MC model)" for each MC model. Note: settings defined in "proj/ini" are set at the very end, i. e. your parameters defined in the steering cards can be overwritten via files in "proj/ini".

#### 19.2 CASCADE model

For this model, you need a steering card and a file with unintegrated gluon densities. Unintegrated densities can be found in http://www-h1.desy.de/~jung/cascade. You should copy one of it to your current directory and rename it to "ccfm.dat". One file with example unintegrated PDF is in the "main/mcarlo/cascade-12/steer" directory.

RunMC contains example steering cards for several processes in "main/mcarlo/cascade-12/steer". You should copy one steering card to your current directory and rename it to "cascade.cards". The file "cascade.cards" should be in the same directory where you execute "runmc".

#### 19.3 PHOJET model

This model is not supported by the author, therefore, the PHOJET model was implemented as it is, with a minimal coding necessary for RunMC. The cards files for initialisations, located in "example/steer/PHOJET", are not standard and are different from other Monte Carlo models. Because of this reason, it is impossible to edit these steering cards using RunMC GUI. This model also requires input PDF file, "fitpar.dat" (located in "main/mcarlo/phojet/files").

In addition, there is a rather different concept to fill histograms. In case when PHOJET steering cards are used, RunMC first fills a temporary binary file "runmc.tmp" with events, and only then it start to read it and fill histograms. Thus, one may notice some

delay at the very beginning of the run, but this compensated by a very fast histogram filling. Make sure that you have enough disk space (50k events usually require 60M for HERA experiment).

# 20 How to share RunMC physics projects

All user-defined settings and calculations (including those necessary to update RunMC GUI) are in the "proj" directory. You should create RMC file of from this directory using RunMC GUI "RMC project - save RMC". Then you can send just created RMC file located in the "archive" directory.

# 21 Possible problems

#### 21.1 ROOT canvas corrupted:

ROOT canvas with updated histograms is corrupted when overlapped with another window. Answer: you have to activate "BackingStore" in X11: (see PAW howto) LINUX: It depends which daemon is in charge of putting up the login box on the X-Windows screen, and start the X-Window session at login time. It can be xdm, kdm (provided with the KDE environment) or gdm. If it is xdm or kdm the line starting the X server in the file /usr/lib/X11/xdm/Xservers should not contain the option "-bs" it should look something like:

```
:0 local /usr/X11R6/bin/X +bs
```

For XFree86 4.0.1, the Screen section of the file /etc/X11/XF86Config (or /etc/X11/XF86Config-4) should be something like:

Section "Screen"

. Option "backingstore" . . EndSection

# 22 Tabular Summary

Program name:	RunMC
Version:	4.0
Date of last version:	September, 2005
Author:	Sergei Chekanov
Size:	11M (Linux/Windows-Cygwin), 12M addition (Windows-Cygwin)
Operating system:	Linux, Windows/Cygwin
Additional packages needed:	CLHEP, ROOT, CERNLIB with PDFLIB (only for Linux)
Program requirements:	g77, g++, make, zip/unzip, sed, X11
Programming language:	C++, C, Fortran, bash
Availability:	http://www.desy.de/~chekanov/runmc

### References

- R. Brun, F. Rademakers, ROOT: An object oriented data analysis framework, Nucl. Instrum. Meth. A389 (1997) 81. URL http://root.cern.ch/
- [2] R. Brun, F. Rademakers, P. Canal, M. Goto, Root status and future developments, ECONF C0303241 (2003) MOJT001.
- [3] L. Lönnblad, CLHEP: A project for designing a C++ class library for high-energy physics, Comput. Phys. Commun. 84 (1994) 307.
- [4] M. Fischler, A. Pfeiffer, CLHEP new developments and directionsThe proceedings of International Conference on Computing in High Energy Physics and Nuclear Physics (CHEP 2000), Padova, Italy, 7-11 Feb 2000. URL http://wwwasd.web.cern.ch/wwwasd/lhc++/clhep/
- [5] T. Sjöstrand, L. Lönnblad, S. Mrenna, Pythia 6.2: Physics and manual.
- [6] G. Corcella, et al., HERWIG 6.5 release note.
- [7] L. Lönnblad, ARIADNE version 4: A program for simulation of qcd cascades implementing the color dipole model, Comput. Phys. Commun. 71 (1992) 15.
- [8] G. Ingelman, A. Edin, J. Rathsman, LEPTO 6.5 a monte carlo generator for deep inelastic lepton-nucleon scattering, Comput. Phys. Commun. 101 (1997) 108.
- [9] G. Ingelman, J. Rathsman, G. Schuler, AROMA 2.2 a monte carlo generator for heavy flavour events in *ep* collisions, Comput. Phys. Commun. 101 (1997) 135.
- [10] H. Jung, The CCFM monte carlo generator CASCADE, Comput. Phys. Commun. 143 (2002) 100.
- [11] R. Engel, PHOJET: manual, Univ. Siegel preprint 95-05. URL http://www-ik.fzk.de/~engel/phojet.html
- [12] H. Jung, Hard diffractive scattering in high energy ep collisions and the monte carlo generator RAPGAP, Comm. Phys. Commun. 86 (1995) 147.

# Index

ARIADNE, 5 AROMA, 5

CASCADE, 5 CLHEP, 2

directory archive, 5 directory proj/ini, 6

file user-name.txt, 9 file project.mc, 8 file user-comment.xml, 8 file user-end.cpp, 8 file user-init.cpp, 8 file user-name.txt, 5 file user-run.cpp, 8 file user-select.cpp, 8 file user\_afill.cpp, 9

#### HERWIG, 5

installation problems, 3

JRunMC, 4

LEPTO, 5

PHOJET, 5

RAPGAP, 5 RMC, 5, 8 ROOT, 2 RUNMC environmental variable, 2

steering cards, 7