## ICOOL User’s Guide

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## 1. Introduction

ICOOL is a 3-dimensional tracking program that was originally written to study ionization cooling of muon beams [1,2]. The program simulates particle-by-particle propagation through materials and electromagnetic fields. Particles are tracked and regions are described using "accelerator" coordinates. The program was written with low energy ( $1 \mathrm{MeV} / \mathrm{c}--1 \mathrm{GeV} / \mathrm{c}$ ) muons in mind, but tracking of electrons, pions, kaons, and protons is also possible. The physics processes available include decays, delta rays, multiple scattering, energy loss and straggling.

Information is input to the code and written out via ASCII data files.
The incident beam particles can be generated from uniform or Gaussian distributions or read from an input file. The code can read its own output, so simulations can be staged. The particles are tracked through a sequence of fixed-length longitudinal regions. In general a region is cylindrical in shape and may be subdivided radially. Every region has a specified material and field type associated with it. Groups of regions can be grouped in cells and a separate cell field can be superimposed over the region fields when tracking is done. In general the program takes user-defined steps along a reference axis. For each step it updates the particle position, time and momentum, taking into account the local field, and corrects the particle's momentum for energy loss and multiple scattering in the step. There is an option for letting the program make adaptive step sizes.

ICOOL uses analytic and other methods to compute field strengths at a given location. There are in general several model levels for each field type, which determine the approximation used to calculate the field.

The program always generates an output log file. In addition, depending on control variable settings, it may generate several other output files. A number of internal diagnostics, such as histograms and scatterplots, are conveniently available. There are options for saving information about each particle after every region (or step) in a postprocessor file.

This guide gives an overview of the main features of ICOOL from a functional point of view. Many examples have been included. However, not all capabilities of the code are discussed here. A complete description of all commands can be found in the ICOOL Reference Manual, icoolman.pdf.

### 1.1 File structure

The source code for ICOOL is broken up logically into the following files.

| file | description |
| :--- | :--- |
| icommon.inc | common blocks |
| icool.for | Monte Carlo superstructure |
| ifld.for | field routines |
| iint.for | interaction routines |
| idiag.for | internal diagnostics |
| imath.for | mathematical routines |
| imsoft.for | Windows-specific code |

When running a job ICOOL uses the file naming convention for0 $x x$. dat. File numbers 01 to 19 are reserved by the program. Some common execution files are listed below.

| file | type | description |
| :--- | :--- | :--- |
| for001.dat | input | command file |
| for002.dat | output | log file |
| for003.dat | input | beam file |
| for004.dat | output | beam file |
| for007.dat | output | region summary table |
| for008.dat | internal | overflow particle file |
| for009.dat | output | postprocessor file |
| for011.dat | output | beam envelope file |

Every job must have at least a command file for001.dat. Sections 2-7 of this guide are mainly concerned with a description of the contents of this file. Every job produces at least a $\log$ file for002.dat. Section 8 of this guide describes a number of useful utility programs that can be used with the output files from an ICOOL job.

### 1.2 Basic job structure

An ICOOL job is run by first setting up the command file for001.dat and any other required files for0xx.dat, and then invoking the executable by typing icool. The basic parts of the command file are

1. title
2. control variables (\&cont)
3. beam definition (\&bmt)
4. interactions definition (\&ints)
5. internal diagnostics (\&nhs, \&nsc, \&nzh, \&nrh, \&nem, \&ncv)
6. region definition (SECT ...)

A simple example of a complete command file follows.

```
Drift space example with a scatterplot of x vs. Y
&cont npart=500 / ! control variables
&bmt / ! beam definition
1 1. 1 ! Gaussian definition of muon beam
0.0.0.0.0.0.200 ! means
3e-3 3e-3 0.01 0.005 0.005 0.010 ! sigmas
0 ! no imposed beam correlations
&ints / ! use default interactions
&nhs / ! no histograms
&nsc nscat =1 / ! define 1 scatter plot
-0.10 5e-3 40 1 2 -0.10 10e-3 20 2 2
&nzh / ! no z-histories
&nrh / ! no r-histories
&nem / ! no 2-D emittance calculations
&ncv / ! no covariance calculations
SECTION ! start problem definition
SREGION ! define a region
1.00 1 0.003 ! length, 1 radial subregion, step
1 0. 0.10 ! radial extent
NONE ! no associated field
0.0.0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
VAC ! vacuum material
CBLOCK ! cylindrical block geometry
0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
ENDSECTION ! end of problem definition
```

Note that the default units used in ICOOL are meters, seconds, $\mathrm{GeV} / \mathrm{c}$, tesla and MV/m.

## 2. Basic concepts

### 2.1 Comments

Comments may be inserted anywhere in the input file for001. A comment is anything on a line that follows the ! character. Blank lines may also be inserted to improve clarity in the input file.

```
! this whole line is a comment
SREGION ! so is the remainder of this line
```

"Comments and blank lines can help format a control file so that it is more understandable to human readers. The computer subroutines that read and parse control files don't want to see the comments and blank lines. A subroutine in ICOOL removes comments and blank lines before the command processor parses them.

## Rules for commenting a control file

A comment is any string of printable characters whose leftmost character is an exclamation point.
! This is a comment.
! So is this.
! Additional exclamation points !! in a comment don't matter.
A whitespace is either a space or a horizontal tab.
A blank line either has no characters (except for the end-of-line terminator) or no characters except whitespaces. Blank lines may be placed anywhere in the control file. They are ignored by the command parser.

A comment line contains a comment, preceded by zero or more whitespaces. Comment lines may be placed anywhere in the control file. They are ignored by the command parser.

An end-of-line comment is a comment placed to the right of a normal input line. End-ofline comments may be placed at the end of any input line, separated from the data ICOOL is to parse by zero or more whitespaces.
1.2342 .345 !This is an end-of-line comment.
3.4564.567!So is this (valid but hard to read).

RING!This is valid even though "RING" will be read into a 6 -character field.
RING !This is a more readable end-of-line comment.

## The Use of Informal Comments

Though it is not forbidden, users are strongly discouraged from using "informal comments" --
those without leading exclamation points that are imagined to be out of view of the parser. There are numerous ways to go wrong. For example, free format reads can extract data from more than one input line; an informal comment on the end of any but the last line will generally produce an error. An informal comment following a text string must come with enough preceding spaces to fill the input field with spaces. All of these potential problems are avoided by using formal comments with leading exclamation points." $\{$ S.B. $\}$

### 2.2 Beam definition

The beam particles used for a simulation may be defined internally or read in from a file, depending on the value of the control variable BGEN. Internal generation uses the namelist command \$BMT. The input beam can consist of one or more types of particle, each with its own production distribution. The distributions can be either gaussian or part of a uniform circular segment. A number of correlations can be imposed on the distribution, including angular momentum and longitudinal momentum-transverse amplitude. Beams with both signs of charge can be generated using the variable BMALT. ICOOL uses an internal particle numbering system

| code | particle |
| :--- | :--- |
| $\pm 1$ | electron |
| $\pm 2$ | muon |
| $\pm 3$ | pion |
| $\pm 4$ | kaon |
| $\pm 5$ | proton |
| 6 | deuteron |
| 7 | $\mathrm{He}^{3}$ |
| 8 | $\mathrm{Li}^{7}$ |

where the sign indicates the charge of the particle.

```
Example: generate a muon beam with a gaussian distribution
&bmt /
1 2 1. 1
0. 0. 0. 0. 0. 0.200
0.010 0.010 0.010 0.010 0.010 0.010
0
```

Externally produced beam files are read from the file for003.dat.

### 2.3 Regions

The basic unit in a problem description is called a region. This command describes the field and material in a given longitudinal area. The basic structure is

SREGION

| SLEN | NRREG | ZSTEP |
| :--- | :--- | :--- |
| IRREG | RLOW | RHIGH |
| FTAG |  |  |
| FPARM |  |  |
| MTAG | (MTAG2) |  |
| MGEOM |  |  |
| GPARM |  |  |

Each region begins with an SREGION command. There is no specific end command for a region.

SLEN is the axial length of the region. In a curved region this length follows the bent path. A region can be broken up into up to 4 radial subdivisions. NRREG is the number of radial subdivisions. ZSTEP gives the stepsize for this region. The following commands are repeated for each radial subdivision. IRREG enumerates the radial regions. RLOW is the minimum radius and RMAX is the maximum radius for this subdivision. FTAG is a character string identifying the field. This is followed by 15 parameters FPARM that give the specific details for the field. MTAG is a character string that identifies the material in the region. For a wedge an optional second material may be given that specifies the material exterior to the wedge. MGEOM is a character string specifying the geometry of the material. This is followed by 10 parameters GPARM that give specific details about the geometry.

```
Example: Simple solenoid with no material
    SREGION
    0.20 1 0.01
    1 0. 0.30
    SOL
    1.0.0.0.1.0 0.0.0.0.0. 0. 0. 0. 0. 0.
    VAC
    CBLOCK
    0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
```

```
Example: stepped beryllium window
SREGION 
750e-6 1 1 1 % 1 0.00 0.25
NONE
0
BE
CBLOCK
```



```
750e-6 2 100e-6
1 0.00 0.20
ONONE N
VAC
CBLOCK
0
NONE N
BE
CBLOCK
0
```


### 2.4 Repeats

The simplest form of looping can be done using the REPEAT structure. The basic form of this structure is

```
REPEAT
NREP
    whatever you want to repeat
ENDREPEAT
```

NREP is the number of times you want the commands repeated. REPEAT structures cannot be nested. They are often used to describe a string of identical RF cavity cells.

```
Example: String of 4 pillbox rf cavities
    REPEAT
    4
    SREGION
    0.20 1 0.01
    1 0. 0.30
    ACCEL
    2. 201.25 11.0 0.0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
    VAC
    CBLOCK
    0
    ENDR
```


### 2.5 Cells

The CELL structure can also be used for looping, but the main use is to define the lattice field for a channel. The basic structure is

CELL
NCELLS
CELLFLIP
CFTAG
CFPARM
whatever is inside the cell

## ENDCELL

NCELLS is the number of cells. A cell has an associated field given by the parameters CFTAG and CFPARM. Each repetition of the cell uses the same field, unless CELLFLIP is true, in which case alternate cells have reversed polarity.

```
Example: 15 cells of an alternating solenoid lattice. The cell field is
a model 3 SHEET. The second SHEET parameter specifies the input data is
in the file for020.dat.
    CELL
    15
    .true.
    SHEET
    3 20 0.01 0.01 1.51 0.11 15.0
    SREGION
    1.50 1 0.01
    10.0.30
    NONE
    0. 0 0 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
    VAC
    CBLOCK
    0
    ENDC
```

ICOOL can define the magnetic field on a cylindrical grid made up from a sum of individual coils or current sheets. It is possible to define this grid over a short portion of the field lattice and then reuse the same grid over and over. The procedure is to define the grid from $\mathrm{z}=0$ to one grid point beyond the end of one cell. The length of the cell is either the period of the magnetic field, or half the period if the field alternates in direction. One then defines the sheets, for example, which create the field starting one cell before the cell of interest and extending to one cell beyond it, i.e. over 3 cells. ICOOL gets the field it needs from the $z$ distance from the start of the cell it is currently in, so this same grid can be used at any $z$ location in the lattice.

### 2.6 Reference particle

When running with RF cavities it is customary to define a reference particle. The reference particle is used to automatically set the phase of each cavity to 0 -crossing of the electric field when the reference particle passes the center of the cavity. The reference particle is initialized using the pseudocommand

REFP
REFPAR PZOREF TOREF GRADREF PHMODREF
where REFPAR describes the particle type, PZ0REF is the initial momentum and T0REF is the initial time. PHMODREF $=\{3,4\}$ describes how the reference particle moves. If PHMODREF $=3$ the reference particle moves with constant velocity. If it equals 4 , then the particle slows down because of energy loss in matter and speeds in RF cavities at a rate given by GRADREF. The control variable PHASEMODEL must be set to 3 to enable this feature. The REFP command should immediately follow the SECT command in for001.dat. The relative phasing of the real particles with respect to the reference particle can be changed for example by adjusting T0REF or the phase offsets for the RF cavities. A second reference particle defined by the pseudocommand REF2 is used for ACCEL model 10.

```
Example: use a muon traveling with constant momentum 200 MeV/c as a
reference particle
    SECT
    REFP
    2 0.200 0. 0. 3
```

The reference particle is designated as particle 0 in the $\log$ output. A second reference particle, if any, is denoted as particle -1 in the log file output.

If a problem is restarted using an existing beam file that has a non-zero value for $\mathrm{P}_{\mathrm{Z}}$ for the reference particle momentum, the reference particle information for the problem is initialized using information from the file. A REFP command must still be used to activate the reference particle logic for the new job. Normally the REFP command immediately follows the SECTION command. The reference particle can be redefined later in a problem, if desired, by using another REFP command. The reference particle data is initialized using the input file data. The momentum is reset by the REFP command if the parameter PZOREF is not equal to 0 . The time is reset by the REFP command is the parameter TOREF is not equal to 0 .

When a reference particle has been defined and DIAGREF=true, the internal diagnostics in ICOOL compute time and longitudinal momentum as the difference from the reference particle value at a given location. In addition, when the control variable PHASEMODEL is $2-4$, the phases of the rf cavities are initialized from the reference particle. The reference particle is started on-axis with no divergence. Stochastic processes, such as multiple scattering and straggling are temporarily turned off. The time when the reference particle crosses the center of the cavity is used to initialize the cavity electric field to 0 (zero crossing). When PHASEMODEL=3 the reference particle is assumed to move with
constant velocity (determined from the parameter PZOREF). When PHASEMODEL=4 the reference particle is tracked through all regions except RF cavities. The reference particle velocity drops due to the mean $\mathrm{dE} / \mathrm{dx}$ in any material that may be present. It is assumed that the reference particle gains energy linearly in the cavity region. The amount of energy it gains is determined by the parameter GRADREF. You can use (1) the mean value of the initial time position or (2) the parameter PHASE0 for the ACCEL field type to adjust the phase of the non-reference particles.

### 2.7 Reference orbit

Tracking is done in ICOOL in the Frenet-Serret coordinate system. This is a right-handed system where $s$ is tangent to the reference orbit, $y$ is vertical and $x$ is the third orthogonal coordinate. In a circular orbit $x$ is in the radial direction. We define the reference orbit to be that path where the transverse coordinates $x$ and $y$ and the transverse momenta $\mathrm{p}_{\mathrm{x}}$ and $p_{y}$ always remain zero. The shape of the reference orbit in a global Cartesian coordinate system is determined by the curvature parameter. It is important to understand that the reference orbit is not necessarily the same path taken by the RF reference particle described in the previous section.

### 2.8 Material

Every region has an associated material type and geometry. ICOOL has a number of predefined material types, such as LH for liquid hydrogen. The standard material geometry is the cylindrical block CBLOCK. However, a number of wedge-shaped geometries are also available for studies of emittance exchange.

```
Example: liquid hydrogen absorber in a drift region
    SREGION
    0.20 1 0.01
    1 0.0.30
    NONE
    0 0 0 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
    LH
    CBLOCK
```



```
Example: lithium hydride wedge absorber in a drift region. The wedge has
an opening angle of }1\mp@subsup{5}{}{\circ}\mathrm{ and has the apex oriented vertically.
    SREGION
    0.20 1 0.01
    1 0. 0.30
    NONE
    0 0 0 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
    LIH
    WEDGE
    15.0.05 0.10 90.0.10 0.10 0 0 0 0
```


### 2.9 Postprocessor output file

ICOOL can save information about the particles at production and about the particles and their electromagnetic field environment at various $z$ locations in the problem in the file FOR009.DAT for later analysis by the user. Particle information at production is included in the file by setting the control variable OUTPUT1 true. Note that data for the electromagnetic field environment at the production point is NOT included in these records because ICOOL doesn't compute fields until after the particles move away from the production plane. This deficiency can be effectively remedied though by simply defining the first region to be a very thin drift space or by entering the BZ field manually using the control variable BZFLDPRD.

The user can generate particle and field information at the end of a given ICOOL region by preceding the region definition with the OUTPUT pseudocommand. Alternatively, particle and field information at the end of every region can be generated by setting the control variable NTUPLE true.

Particle and field information inside of regions can be generated by setting the control variable RTUPLE true. This variable should be used with caution, since it is capable of generating an enormously large file. The variable RTUPLE works together with the control variable RTUPLEN, which specifies the number of steps inside the region to skip before generating output.

The precision (and length) of the data in the output file can be controlled using the control variable F9DP.

### 2.10 Restarting a job

It is often convenient to save the beam and reference particle at some location as the starting conditions for a subsequent job. This can be done using the control variables FSAV, IZFILE and FSAVSET. If FSAV is true, all particles that reach the region number IZFILE are written to the file for004.dat. If FSAVSET is true, the data is modified to have $\mathrm{z}=0$ and times relative to the reference particle. This file has the same format as the ICOOL beam input file for003.dat, so the job can be restarted by copying the file for004 to for003 and then running ICOOL with the control variable BGEN false.

### 2.11 Name substitution

Parameters in for001 may be referred to symbolically using name substitution. The basic structure is
\&SUB name value
Following this command any occurrence of the text string \&name is replaced with its value. This feature can be used for example to change a parameter in many regions at the same time, or to conveniently change a parameter at the top of the file when doing an optimization study.

```
Example: define region length symbolically
    &SUB slen 0.20
    SREGION
    &slen 1 0.01
```

A variant allows scaling of a class of previously-defined symbolic names.
\&SCL NameString type value
NameString is a set of characters. All previously defined names that begin with this character string will be scaled. The type of scaling is determined by type $=\{*,+\}$. The amount of scaling is determined by value.

Example: scale all parameters starting with the letter $s$ to $99 \%$ of their value
\&SUB step1 0.01
\&SUB step2 0.015
\&SCL s * 0.99
SREGION
0.20 1 \&step1

### 2.12 Keyboard input during execution

ICOOL recognizes two keyboard inputs while it is executing.
p pauses execution until the Enter key is pressed
x stops execution with the current particle and region and executes any end of run diagnostics.

## 3. Control variables

There are a number of variables that can be used to control an ICOOL simulation. Many of these variables have default values that are used if the variable is not defined in a given job.

### 3.1 Global simulation

BGEN is used to control whether ICOOL generates the beam particles internally or from the external beam file for003.dat.

BUNCHCUT sets the maximum time difference allowed between a particle and the reference particle. It is used to stop tracking particles that fall out of an RF bunch.

DECTRK determines whether the charged daughter particle from a decay is tracked.
NPART sets the number of particles used in the simulation.
NPSKIP sets the number of beam particles in an external file to skip before the simulation starts.

NSECTIONS sets the number of times the particles are tracked through an ICOOL section. A section consists of all region commands between the BEGS and ENDS commands. This is frequently used to simulate multiple turns around a ring.

```
Example: simulate 10 turns around a ring following an injector lattice
    &cont npart=1 nsections=10 /
    ........................ .
    SECT
            (the injector lattice description goes here)
    BEGS
            (the ring description goes here)
            (only this part is repeated)
    ENDS
```

PHASEMODEL is used to specify how the phases of the RF cavities are determined. In jobs with a reference particle this parameter determines how the reference particle moves. If the value is 3 , the reference particle moves with a constant velocity, while if the value is 4 , it loses energy in material and gains energy in RF cavities. If the value is 5 , the RF cavity information is read in from a file.

RFPHASE specifies an external file of RF cavity information that can be used with phasemodel 5.

TIMELIM sets the maximum time allowed for the simulation.

### 3.2 Analysis and diagnostics

BETAPERP sets the value of the transverse beta function for calculation of the transverse amplitude parameter $\mathrm{A}^{2}$.

BZFLDPRD gives the value of the solenoid field at production. This needed for the postprocessor file and emittance calculations at production.

DIAGREF controls whether time and $p_{z}$ are specified relative to $t$ the RF reference particle for ICOOL internal diagnostics.

FORCERP is used to force the reference particle back to the axis when phasemodel=4.
GOODTRACK determines whether failed tracks in a beam input file are processed.
PHANTOM forces a particle to maintain its initial transverse coordinates after every step. This can be used to examine field components at fixed transverse positions.

RFDIAG is used to write diagnostic information for every RF cavity.
RNSEED is used to set the initial random number seed.
RUN_ENV is used to run ICOOL in a paraxial beam envelope mode, instead of doing particle tracking. This only works for straight solenoid-focused channels. Quantities are only calculated for planes that are written to the postprocessor file, for009.dat. The initial beam parameters are taken from the ICOOL beam input. Envelope parameters are written to the file for011.dat.

```
Example: examine the paraxial behavior of a straight channel using the
beam envelope mode.
    Run channel in beam envelope mode
    &cont npart=1 ntuple=.true. run_env=.true. /
    &bmt /
    1 2 1. 1
    0. 0. 0. 0. 0. 0.200
    0.010 0. 0. 0.010 0. 0.010
```

This will automatically run a single particle along the axis, and evolve the envelope equations. OUTPUT must be enabled to the postprocessor file for009.dat. Envelope parameters are output to the file for011.dat.
"Currently, this only works if the beam parameters are given from the BMT input lines. Only the first beam type is used. This applies even if bgen is set to .false.; the particle data in for003.dat is ignored... There must be nonzero sigma x and sigma Px ; in addition, the average Pz must be $>0$.

The envelope equations assume that there are no dispersive regions or shaped absorbers,
the bunch is well matched into the rf bucket, and the reference orbit stays on axis. In addition, the envelope equations will not work well for induction linacs or bunching sections.
The only source of longitudinal emittance growth considered is the slope of the $\mathrm{dE} / \mathrm{ds}$ curve. The size of the rf bucket is ignored, but this can be put in after the fact (see below). Besides paraxial equations of motion, beam scraping is included where the beam is scraped against the widest aperture for each region.

## How to interpret the results of the envelope equations

The parameter Psi (listed as "gauss" in the output file) represents how severely the beam is scraped; when Psi is large, only the extreme tails are missing; when Psi is small, the beam distribution is almost flat until the cutoff. The envelope equations give as results z , $\mathrm{Bz},<\mathrm{t}\rangle,\langle\mathrm{pz}>$, transverse emittance e_N, beta, alpha, and dimensionless canonical angular momentum, transverse scraping parameter Phi (listed as "gauss" in output file). There are also parameters which do NOT take into account losses due to falling out of rf bucket, in particular the total number N_0, and the longitudinal emittance e_L0.
Dimensionless canonical angular momentum is $<$ Lcanon $>/\left(2 \mathrm{mc} \mathrm{e}_{-} \mathrm{N}\right)$. Particle number and longitudinal emittance ignores losses due to particles falling out of rf bucket. Thus, if rf bucket separatrix corresponds to amplitude A_RF, then after those losses are taken into account the particle number is $\mathrm{N}=\mathrm{N} \_0$ G0(A_RF/e_L0).

The longitudinal emittance taking into account losses from falling out of the rf bucket is $\mathrm{e}_{-} \mathrm{L}=\mathrm{e}_{-} \mathrm{L} 0 \mathrm{G} 1\left(\mathrm{~A}_{-} \mathrm{RF} / \mathrm{e}_{-} \mathrm{L} 0\right) / \mathrm{G} 0\left(\mathrm{~A}_{-} \mathrm{RF} / \mathrm{e}_{-} \mathrm{L} 0\right)$.

The transverse "scale length" (not given in output file) is given by A_G = e_N G1(Psi)/G2(Psi).

The fraction of particles within the transverse acceptance C_perp is
f_perp $=$ G1(C_perp/A_G) / G1(Psi),
or just 1 when Cperp > Psi A_G.
The fraction of particles with the longitudinal acceptance C_L is
f_L = G0(C_L/e_L0), if C_L < A_RF, or G0(A_RF/e_L0) if C_L > A_RF.

The total number of particles within the 6D acceptance defined by C_perp, C_L, is given by N_6D = N_0 f_perp f_L

Key functions:
$\mathrm{G} 0(\mathrm{x})=1-\exp (-\mathrm{x} / 2)$
$\mathrm{G} 1(\mathrm{x})=1-(1+\mathrm{x} / 2) \exp (-\mathrm{x} / 2)$
$\mathrm{G} 2(\mathrm{x})=1-\left(1+\mathrm{x} / 2+\mathrm{x}^{\wedge} 2 / 8\right) \exp (-\mathrm{x} / 2) "\{\mathrm{G} . \mathrm{P}\}$

SUMMARY is used to write a file containing a list of all the regions (and pseudoregions) defined in the job.

### 3.3 Print

Print diagnostics are written to the $\log$ file for002.dat.
NPRNT sets the number of particles for which print diagnostics is desired.
PRLEVEL specifies the amount of information to print for each particle.
PRNMAX sets the maximum number of steps to print out inside a given region.

### 3.4 Beam restart

FSAV specifies whether beam information at a given location is saved to the file for004.dat.

IZFILE specifies the location where beam information should be saved.
FSAVSET tells whether the information saved in for004. dat should have the z position and time reset.

### 3.5 Postprocessor output

These commands control the amount of information written to the postprocessor output file for009.dat.

F9DP sets the number of decimal places to use for floating point variables.
NTUPLE determines whether to write output after every region.
OUTPUT1 determines whether production information is written to the file.
RTUPLE is used to write output after a fixed number of steps inside a region.
RTUPLEN sets the number of steps to skip between outputs when using RTUPLE.

### 3.6 Accuracy

EPSF sets the tolerance on the fractional change on field variations, energy loss or multiple scattering when using variable stepping.

EPSREQ sets the required tolerance on tracking when using variable stepping.
EPSSTEP sets the desired tolerance for reaching the end of a region.
PZMINTRK sets the minimum value of $\mathrm{P}_{\mathrm{Z}}$ a track can have, after which tracking is stopped.

SCALESTEP sets a factor that modifies all the step sizes in a problem simultaneously. This only works when not using variable stepping.

STEPMAX specifies the maximum step size to use for variable stepping.
STEPMIN specifies the minimum step size to use for variable stepping.
STEPRK specifies whether to use the Runge-Kutta or the Boris algorithm for stepping.
"The Boris integrator is a space-stepping integration scheme similar to the time-based scheme used in plasma physics (see "Plasma Physics via Computer Simulation", Birdsall and Langdon, p. 356).This scheme is second-order accurate, compared to fourth-order for the default Runge-Kutta, so a smaller stepsize may be required. However, the Boris integrator is less computationally intensive and so may give significantly shorter run times. Adaptive stepsize control is not yet implemented for the Boris integrator." $\{$ P.S. $\}$

The Boris integrator is enabled by setting the control variable STEPRK=false. It only works for straight regions. The program automatically switches to Runge-Kutta integration in curved regions. The underlying algorithm is described in MC note 229.

VARSTEP specifies whether to use fixed or variable (adaptive) step sizes in tracking.

### 3.7 Moving grid

A solenoidal channel can be defined in ICOOL from a single list of coil properties. A second file is used to define the field grid partitions where the magnetic field must be calculated. The program fills the first grid partition, tracks all the particles to the end of the grid, then recomputes the field values for the second grid partition, tracks all the particles, and so forth.

MAGCONF specifies a file containing the coil information.
MAPDEF specifies a file containing the field grid partition data.

### 3.8 Spin

The muon spin is followed in pion decay, tracking in fields and interactions in matter.
SPIN determines whether spin calculations are processed. It also generates muon polarization from pion decay.

SPINMATTER determines whether depolarization effects in matter are simulated.
SPINTRK controls whether spin variables are tracked.
Setting the control variable SPIN true enables the handling of spin in ICOOL. At present there are two ways of getting initial spin information: (1) the user may read in a beam input file with spin information already present or (2) muon spin from pion decay may be calculated inside the program. If the control variable SPINTRK $=0$, the muon helicity is stored in the variable POL(3) when the pion decays and is unchanged as the particle is tracked.

If the control variable $\operatorname{SPINTRK}=1$, the BMT equation is used to track the particle spin in the muon rest frame through electric and magnetic fields. The spin tracking needs a 3component, normalized Cartesian vector to represent the spin. For muons from pion decay the array POL contains the spin vector in the muon rest frame.

Muon depolarization effects in matter are controlled by the variable SPINMATTER. If SPINMATTER $=1$ the Rossmanith model of depolarization is used, which relates the amount of depolarization to the energy loss in material regions. If SPINMATTER=2 quantum mechanical formulae are use for the probability of spin flip due to elastic and Coulomb scattering. The elastic scattering effect is more important and requires that delta rays interactions be enabled (LDRAY=true).

### 3.9 Neutrino production

Neutrino production information can be saved from muon, pion and kaon decays.
NEUTRINO controls whether neutrino production information is written to a file.
NNUDK sets the number of neutrinos to produce at each muon, pion or kaon decay location.

NUTHMAX sets the maximum polar angle for neutrinos written to the file.

NUTHMIN sets the minimum polar angle for neutrinos written to the file.
It is possible to make files of neutrino production data using four control variables. The variable NEUTRINO specifies the output file, if any, that should be used. Neutrinos can
be generated from muon, pion, and kaon decays. A large number (NNUDK) of independent neutrinos can be generated at each decay point. The user can select which of the generated events are written to the output file using the polar angle variables NUTHMIN and NUTHMAX. Neutrinos have a particle type that uniquely identifies their flavor and their parent particle.

```
Example: save information on neutrino production from pion decay in a
file, e.g. for030.dat. Produce }100\mathrm{ neutrinos at each decay point. Only
accept neutrinos produced within 100 mrad of the axis.
    Save neutrino production data
    $cont npart=1000 neutrino=30 nnudk=100 nuthmax=0.100 $
    $bmt $
    1 3 1. 1
    0. 0. 0. 0. 0. 0.200
    0.010 0.010 0.010 0.010 0.010 0.010
```


### 3.10 Miscellaneous

FFCR inserts form feed and carriage return characters into the log file for002.dat so that there are two plots per printed page.

NEIGHBOR specifies whether fields from neighboring regions can overlap when using ICOOL soft-edge field models. The control variable NEIGHBOR can be used to model the contribution to a given region of the tails from the fields from neighboring regions. This is used with the soft-edged, analytic delta hyperbolic tangent model of the field available for the field types \{BSOL , DIP , HDIP , QUAD , ROD , SEX , SOL , SQUA \}. If NEIGHBOR is true, the program looks to see if the regions immediately preceding and following the current region have a field type in the above set and, if so, add the field to the field at the current location.

Example: consider the field in two adjacent 1 m long solenoid regions. Allow the fringe fields to overlap. Overlapping soft edge solenoid fields. \&cont npart=1 neighbor=.true. /

```
&zhist nzhist=1 /
```

10.0 .02857 70 0. 0. 33
SREGION
1.0010 .01
10.0 .20
SOL
2. 1.0 0.8 0.1 1 0.2 0. 0. 0. 0. 0. 0. 0. 0. 0.
VAC
CBLOCK
0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
SREGION
$1.00 \quad 1 \quad 0.01$
10.0 .20
SOL
2. 1.0 0.8 0.1 1 0.2 0. 0. 0. 0. 0. 0. 0. 0. 0.
VAC
CBLOCK
0. 0. 0. 0. 0. 0. 0. 0. 0. 0.


TERMOUT specifies whether output to the terminal is permitted. This is used by some optimization programs.

## 4. Interaction variables

### 4.1 Global controls

Information about interactions is entered with the INTS namelist. The interactions with matter used in a simulation are controlled by setting the logical interaction variables given in the following table.

| interaction variable | default | level | description |
| :--- | :--- | :--- | :--- |
| LDEDX | T | DELEV | mean energy loss |
| LSTRAG | T | STRAGLEV | straggling |
| LSCATTER | T | SCATLEV | multiple scattering |
| LDRAY | T | (none) | delta ray production |
| LDECAY | T | DECLEV | decay |
| LELMS | F | (none) | scattering \& straggling |
| LSAMCS | F | (none) | scattering \& straggling |
| LINTERACT | F | INTLEV | nuclear interactions |
| LSPACE | F | SPACELEV | space charge |

The level variable is used to select a particular model for a given interaction.

### 4.2 Energy loss

PDELEV4 gives the value of momentum to use in the level 4 model.

### 4.3 Multiple scattering

FACFMS is a factor to correct the characteristic angle squared $\chi_{\mathrm{C}}{ }^{2}$ in the Moliere theory.
FACMMS is a factor to correct the characteristic angle squared $\chi_{\mathrm{A}}{ }^{2}$ in the Moliere theory.

### 4.4 Correlated scattering and straggling

There are three methods for simulating correlated scattering and straggling in particle interactions. The first is to turn on delta rays by setting LDRAY=true. Two other variables control the transition energy between continuous, uncorrelated scattering and straggling and correlated delta ray (knock-on electron) interactions.

DCUTE is the kinetic energy of electrons, above which delta rays are simulated discretely.

DCUTM is the kinetic energy of muons and other heavy particles, above which delta rays are simulated discretely.

The second method is to use the ELMS algorithm [3] by setting LELMS=true. This only applies to liquid hydrogen. Interactions in other materials use the models specified by SCATLEV and STRAGLEV.

ELMSCOR determines whether ELMS runs with correlations or not.
The third method is to use the SAMCS algorithm [4].

### 4.5 Decays

FASTDECAY is a diagnostic variable that forces muons, pions and kaons to decay immediately.

### 4.6 Nuclear interactions

The following parameters are used for the Wang model of pion production in proton interactions,
$\mathrm{d}^{2} \sigma / \mathrm{dp} \mathrm{d} \Omega=\mathrm{A} \mathrm{p}_{\mathrm{MAX}} \mathrm{x}(1-\mathrm{x}) \exp \left\{-\mathrm{Bx}^{\mathrm{C}}-\mathrm{Dp}_{\mathrm{T}}\right\}$
where $\mathrm{x}=\mathrm{p}_{\mathrm{L}} / \mathrm{p}_{\mathrm{MAX}}$.
WANGA is the Wang A parameter.
WANGB is the Wang B parameter.
WANGC is the Wang C parameter.
WANGD is the Wang D parameter.
WANGPMX is the Wang $p_{\text {MAX }}$ parameter.
WANGFMX is the maximum value for the Wang differential cross section.

### 4.7 Space charge

FRFBUNSC is the RF frequency for space charge model 4.
PARBUNSC is the number of muons per bunch.

## 5. Internal diagnostics

ICOOL contains a number of internal diagnostics. These can be used to check if the program is behaving "reasonably". For simple jobs these diagnostics may all that is needed. The postprocessor file information can be used for more detailed analysis. Apart from the region summary table, all internal diagnostics are written to the log file for002.dat.

### 5.1 Region summary table

ICOOL identifies physical regions and pseudoregions by an integer region number. The region number can be found from a convenient summary table that lists all the regions that have been defined for the job. It is useful to check this table when first setting up a problem to be sure that it has been set up in the intended way. The region summary table is created by setting the control variable SUMMARY true. The data is written to the file for007.dat. The figure shows the beginning of a region summary table.


The region number is given in the first column (labeled IZ).

### 5.2 Data print out

Data about individual particle tracking can be printed out using the control variables NPRNT, PRLEVEL, and PRNMAX. NPRNT sets the number of tracks to print out. The amount of information is controlled using PRLEVEL. This ranges from only giving values at the end of regions to listing particle and field data after every step inside a region. PRNMAX can be used to limit the number of steps that is printed inside regions.

| ${ }_{5}$ Command Prompt - browse for002.dat |  | $\square$ |
| :---: | :---: | :---: |
| ompleted SETUP: 16 20 8 |  |  |
| INITIAL |  |  |
|  |  |  |
| XP, PP: 0.6170E-03-0.8512E-03 0.3000E-02 0.2643E-03 0.7779E-02 0.1882E+00 |  |  |
|  |  |  |
|  |  |  |
|  |  |  |
| XP, PP: 0.6254E-03-0.6032E-03 | -0.6032E-03 0.9000E-02 0.2643E-03 | 7779E-02 0.1882E+00 |
|  |  |  |
|  |  |  |
|  |  |  |
| XP, PP: 0.6339E-03-0.3553E-03 | -0.3553E-03 0.1500E-01 0.2643E-03 | $0.7779 \mathrm{E}-020.1882 \mathrm{E}+00$ |
|  |  |  |
| XP, PP: 0.6381E-03-0.2313E-03 | -0.2313E-03 0.1800E-01 0.2643E-03 | $0.7779 \mathrm{E}-020.1882 \mathrm{E}+00$ |
| KNT, TP, HDID, HF, HE, HS: 7 O.110E-09 0.300E-02 0.3G日E-02 0.300E-02 0.300E-02 |  |  |
| XP, PP: 0.6423E-03-0.1073E-03 | -0.1073E-03 0.2100E-01 0.2643E-03 | $0.7779 \mathrm{E}-020.1882 \mathrm{E}+00$ |
|  |  |  |
|  |  |  |
| KNT, TP, HDID, HF, HE, HS: 9 0.133E-09 0.300E-02 0.300E-02 0.300E-02 0.300E-62 |  |  |
| XP, PP: 0.6507E-03 0.1407E-03 | $0.1407 \mathrm{E}-03 \mathrm{0}$-2700E-01 0.2643E-03 | $0.7779 \mathrm{E}-020.1882 \mathrm{E}+00$ |
| KNT, TP, HDID, HF, HE, HS: 10 0.144E-09 0.300E-02 0.300E-02 0.300E-02 0.300E-02 |  |  |
| XP,PP: 0.6549E-03 0.2646E-03 0.3000E-01 0.2643E-03 0.7779E-02 |  |  |
|  |  |  |
|  |  |  |

The figure shows part of the step-by-step print out using NPRNT=1 and PRLEVEL=2.
A message is written in the LOG file FOR002.DAT for each particle that fails to successfully reach the end of the problem. The following information is given in the message. IFLAG is a code indicating the cause of the failure. IEVT is the "event" number. This number increases sequentially up to the number of particles requested for the simulation. IPNUM is the "particle" number. In general this is set to 1 . However, when decays are enabled, this number increments by 1 each time the tracked particle decays. IPFLG is a particle-specific code that may be implemented in the future. JSRG is the ICOOL region number where the failure occurred.

### 5.3 Histograms

Histograms can be produced of particle coordinates, field components, etc. The quantity to be histogrammed is specified using the region number and an ID number from the following table.

| ID | variable | ID | variable | ID | variable | ID | variable |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| 1 | x | 11 | $\mathrm{x}^{\prime}$ | 21 | $\mathrm{p}_{\mathrm{T}}$ | 31 | $\mathrm{~B}_{\mathrm{X}}$ |
| 2 | y | 12 | $\mathrm{y}^{\prime}$ | 22 |  | 32 | $\mathrm{~B}_{\mathrm{Y}}$ |
| 3 | z | 13 | $\theta$ | 23 |  | 33 | $\mathrm{~B}_{\mathrm{Z}}$ |
| 4 | $\mathrm{p}_{\mathrm{X}}$ | 14 | r | 24 |  | 34 | $\mathrm{E}_{\mathrm{X}}$ |
| 5 | $\mathrm{p}_{\mathrm{Y}}$ | 15 | $\phi$ | 25 |  | 35 | $\mathrm{E}_{\mathrm{Y}}$ |
| 6 | $\mathrm{p}_{\mathrm{Z}}$ | 16 | $\mathrm{p}_{\mathrm{R}}$ | 26 |  | 36 | $\mathrm{E}_{\mathrm{Z}}$ |
| 7 | ct | 17 | $\mathrm{p}_{\phi}$ | 27 |  | 37 | $\mathrm{~S}_{\mathrm{X}}$ |
| 8 | p | 18 | $\mathrm{~L}_{\mathrm{Z}}$ | 28 | $\mathrm{~A}^{2}$ | 38 | $\mathrm{~S}_{\mathrm{Y}}$ |
| 9 | E | 19 | $\mathrm{~L}^{2}$ | 29 | $\mathrm{r}^{2}$ | 39 | $\mathrm{~S}_{\mathrm{Z}}$ |
| 10 | KE | 20 | s | 30 | helicity | 40 | phase |

By default histograms are automatically scaled to prevent overflows, although this can be changed using the histogram variable HAUTO. The contents of histograms can be written to the log file using the histogram variable HCPRN.

Example: make a histogram of x at production
\&nhs nhist=1 /
$\begin{array}{ccccc}-0.02 & 1 E-3 & 40 & 1\end{array}$


The ID number -1 can be used to obtain the production characteristics of tracks that reach the end of the simulation. The ID number may also be set to an ICOOL error flag,
e.g. -23 for particles which an lost due to a radial limit, in order to obtain production characteristics of tracks that failed.

### 5.4 Scatterplots

Scatterplots can be produced of pairs of particle coordinates, field components, etc. The quantities to be plotted are specified using pairs of variable ID numbers and region numbers. By default scatterplots are automatically scaled to prevent overflows, although this can be changed using the scatter plot variable SAUTO.



Density is represented by the plot symbol in the sequence $\{+, 2-9, a-z, A-Z, *\}$.

### 5.5 Z-history

A z-history is a plot of a quantity as a function of the longitudinal coordinate $z$ (or $s$ in a curved region). Z-histories can be produced of particle coordinates, field components, etc. The quantity to be plotted is specified using the variable ID number. By default zhistories are automatically scaled to prevent overflows, although this can be changed using the z-history variable ZAUTO. The contents of z-histories can be written to the log file using the $z$-history variable ZHPRN.

```
Example: make a z-history of }\mp@subsup{B}{z}{}\mathrm{ over the first 6 m of a lattice
&nzh nzhist=4 /
(first 3 definitions)
2 0. 0.08571 70 0. 0. 33
```



### 5.6 R-history

An r-history is a longitudinal plot of the minimum, maximum, mean and standard deviation of all particles in a distribution at the end of regions. The quantity to be plotted is specified using the variable ID number. By default r-histories are automatically scaled to prevent overflows, although this can be changed using the r-history variable RHAUTO. The contents of r-histories can be written to the log file using the r-history variable RHPRN.

```
Example: plot an r-history of the kinetic energy over regions 3 to 210.
&nrh nrhist=1 /
3 3 0. 0. 10
```



### 5.7 Emittances

2D emittances can be printed out at selected region numbers. A number of emittance variables \{DISCORR, PXYCORR, IPZCOR, SIGMACUT, SIG_CUT, EDETAIL\} can be used to control what corrections and correlations are used in the calculations.

```
Example: compute the 2D emittances for 16 regions. Correct for
solenoidal field.
&nem nemit=16 pxycorr=.true. /
    38
5 7 8
```

| Méc Command Prompt - browse asol.f02 |  |  |  |  |  |  |  | - $\square \times$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Total number of initial good particles $=$ Total number of final good particles $=$ |  |  |  |  | $99000$ |  |  |  |
| ***********************) |  |  | Weighted NORMALIZED 2-D EMITTANGES |  |  | ******************* |  |  |
| des | No | ${ }^{\mathrm{N}}$ | emitX [m] | emity [m] | emitz | Tr | fET | fEL |
| 24 | 1000. | 1800. | 0.1582E-02 | 0.1564E-02 | $2{ }^{0}$ 0.1624E-02 | 1.006 | 1.006 | 1.006 |
| 110 | 1699. | 996. | 0.1420E-02 | -1458E--13 | - $0.1626 \mathrm{E}-02$ | 0.996 | 0. 891 | 1.051 |
| 46 | 998. | 997. | $0.1336 \mathrm{E}-62$ | 0.1334E-02 | 0.1765E-02 | 6.997 | 0.849 | 1.656 |
| 182 218 | 998. | 993. | O.1256E-G2 | O.1240E-02 | $2{ }^{\text {O }}$-1798E-02 | 0.993 | 0.793 | 1.107 <br> 1.115 <br> 1 |
| 254 | 998. | 994. | 0.1150E-62 | 0.1134E-02 |  | 0.994 | -. 726 | 1.232 |
| 296 | 997. | 990. | $0.1693 \mathrm{E}-62$ | $0.1041 \mathrm{E}-02$ | 0.2071E-02 | 0.994 | 0.678 | 1.225 |
| 326 362 | 997. | 988. | 0.9883E-03 | -.9505E-03 | $3{ }^{\text {a }}$ | 0.984 0.988 | 0.625 | 1.238 1.371 1 |
| 398 <br> 4 <br> 18 | 997. | 988. | $0.8993 \mathrm{E}-03$ | 0. 9431 E - -03 | 0.2334E-02 | 0.988 | 0. 588 | 1.437 |
| 434 | 996. | ${ }_{985}^{980}$. | 6.8372E-63 | C.8833E-03 | 3 $\begin{aligned} & \text { O.2282E-62 } \\ & 3\end{aligned}$ | 0.989 | 0.547 | 1.405 1.558 1.5 |
| 566 | 995. | 981. | 0.7761E-03 | Q. $7789 \mathrm{E}-03$ | 3 0.2561E-02 | 0.981 | 0.494 | 1.572 |
| 542 | 994. | 978. | 6. $27.7036 \mathrm{E}-03$ | 0. $7303 \mathrm{E}-03$ | 3 $0.2521 \mathrm{E}-02$ <br> 0.2752 E  <br> 02  | 0.978 0.979 | 0.468 | 1.552 |
|  | 994. | 979. | $0.7086 \mathrm{E}-\mathrm{b3}$ | ๑.7060E-63 | 0.2757E-घ2 | 0.979 | 0.450 | 1.697 |

The ICOOL log file contains two sets of emittance calculations. The first set of emittances use 2-dimensional calculations for the $x-P_{x}, y-P_{y}$, and $z-P_{z}$ phase spaces. The emittance definition removes the slope in each of these 2-dimensional spaces. In addition a control variable (PXYCORR) is provided to compute the emittance using canonical momentum variables inside solenoidal fields (Larmor frame). Another control variable (IPZCOR) is provided to remove the Pz-amplitude correlation in the longitudinal phase space calculation. The 6-dimensional emittance calculated by this method is the product of the three 2-dimensional emittances.

The second set of emittance calculations ("LBNL") is derived from the computed covariance matrix. The transverse emittances comes from the determinant of a 4dimensional covariance matrix and the 6 -dimensional emittance is the determinant of a 6 dimensional covariance matrix. This method essentially removes all non-diagonal correlations from the final emittance values. In addition, other quantities (e.g. beta functions) following from the covariance matrix are computed

The program computes dispersions statistically as

$$
\begin{aligned}
& \mathrm{D}=\mathrm{r}(\mathrm{x}, \mathrm{Pz}) \sigma(\mathrm{x})<\mathrm{Pz}>/ \sigma(\mathrm{Pz}) \\
& \mathrm{D}^{\prime}=\mathrm{r}(\mathrm{Px}, \mathrm{Pz}) \sigma(\mathrm{Px}) / \sigma(\mathrm{Pz})
\end{aligned}
$$

where
$\mathrm{r}(\mathrm{u}, \mathrm{v})=(<\mathrm{uv}>-<\mathrm{u}><\mathrm{v}>) /(\sigma(\mathrm{u}) \sigma(\mathrm{v}))$

### 5.8 Covariances

Second- and third-order covariances among the particle variables can be printed out for selected region numbers. The figure shows part of the covariance print out at a given plane.

```
Example: compute emittances and other correlations from the covariance
matrix
&ncv ncovar=2 /
    1 578
(the figure shows some of the output)
```



### 5.9 Magnetic field maps

ICOOL can copy an internally created magnetic field map to an external file. This can be done for 2D maps using the GRID command or by using SHEET model 3. An internally created 3D map generated from the BACKGROUND command can also be written to an external file.

### 5.10 RF diagnostics

Diagnostic information about particle and field variables in RF cavities can be generated by using the control variable RFDIAG.

## 6. Pseudoregion commands

### 6.1 Simulation

APERTURE imposes a rectangular or elliptical constraint on all particles transverse coordinates at this location.

```
Example: insert a symmetric rectangular collimator with a half-opening
of 15 cm in x.
    APERTURE
    2 -0.15 0.15 1
```

CUTV applies a cut on an ICOOL variable.

```
Example: only accept particles with p greater than 200 MeV/c.
    CUTV
    6 2 0.200
```

DENP applies a transverse density profile.

```
Example: associate a quadratic density profile along the x direction
with beryllium material.
    DENP
    BE 1 0.5 0.3 0.1 0.
```

DENS adjusts the density of a material.

```
Example: use gaseous hydrogen at a pressure of 20 atm as a material.
    DENS
    GH 20.
```

DVAR makes an additive change to one of the particles variables.

```
Example: offset all particles x coordinates by -1 cm. Variables are
specified using ICOOL's internal variable numbering, e.g. 1 means x. The
third parameter controls whether the change also affects the reference
particle.
        DVAR
        1 -0.01 0
```

EDGE takes into account fringe field focusing for hard-edge magnet models.

```
Example: model a 1.2 T hard-edge solenoid including effects of the end
fields.
    EDGE
    SOL
    1 -1.2 0. 0. 0.
    SREGION
    1.00 1 0.01
    1 0.00 0.30
    SOL
```



```
    VAC
    CBLOCK
    0
    EDGE
    SOL
    1 1.2 0. 0. 0.
```

REFP specifies reference particle parameters.

```
Example: define a 200 MeV/c muon that moves with constant velocity as
the reference particle.
    REFP
    2 0.200 0. 0. 3
```

REF2 specifies the parameters for a second reference particle. This is only used for the ACCEL field model 10 for the adiabatic buncher.

RESET changes the times of all particles to the current value of the reference particle time.

ROTATE changes the angles of the coordinate system.

```
Example: bend particles vertically by rotating coordinates \(90^{\circ}\) around \(z\),
followed by a normal 0.2 T dipole field.
    ROTATE
    90. 30
    SREGION
    \(1.00 \quad 1 \quad 0.01\)
    \(10.00 \quad 0.30\)
    \(\begin{array}{lllllllllllllll}\text { DIP } & & & 0.2 & 0 & 0.200 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0\end{array} 0\)
    \(\begin{array}{lllllllllllllll}1 & 0.2 & 0 & 0.200 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0\end{array}\)
    VAC
    CBLOCK
    \(\begin{array}{llllllllll}0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0\end{array}\)
```

TAPER is used to set up a tapered lattice. It is used in conjunction with the BLOCK(4) and BSOL(5) field models.

TRANSPORT performs a matrix transformation on all particles.

### 6.2 Postprocessor file

OUTPUT enables writing of particle information to for009. dat in the following region.

```
Example: generate postprocessor output for the next region
    OUTPUT
    SREGION
    0.20 1 0.01
    1 0. 0.30
    QUAD
    1. 1.0 0. 0.0. 0.0.0.0.0. 0. 0. 0. 0. 0.
    VAC
    CBLOCK
    0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
```


### 6.3 Field grid

GRID defines a new magnetic field grid. This can be done either by reading in a file of current elements or by reading in an existing field grid.

```
Example: define a new field grid #2 using the data in an external file
of current sheets, e.g. for032.dat. Let the grid have 1 cm spacing in z
and r and extend over 3 m in z and 30 cm in r. Save the field values on
the grid in file for043.dat.
    GRID
    2
    SHEET
```



```
Example: define a new field grid #3 using an existing grid in the
external file for050.dat.
    GRID
    3
    USER
```



### 6.4 Error analysis

DISP randomly displaces particle transverse coordinates.
TILT randomly rotates all particles' coordinates in 3D.
RKICK makes random magnetic kicks on all particles.
Example: apply random dipole kicks to all particles at this location. Let the standard deviation of the error field be 0.1 T and let the kick be applied with random azimuthal orientations.

RKICK
$\begin{array}{lllll}\text { DIP } 0 . & 0.1 & 0 & -1\end{array}$

### 6.5 Background fields

The background field construct sets up a static magnetic field on a 3-D cartesian grid. A new definition is begun using the BACKGROUND command. The definition is terminated with the ENDB command. One of the parameters ZTOTALBKG specifies the total incremental distance from the present location that this background can be used. Following the BACKGROUND command the user can enter a series of BFIELD
commands, which are used to construct the field on the background grid. Two of the parameters of this command, ZMINBKG and ZMAXBKG, specify the region of the grid where this field will be used. In general ZMAXBKG $=$ ZTOTALBKG. The field type specified by the BFIELD command can either be an internally defined ICOOL field or STUS, which allows a user-specified static field to be read in from a file. In order for higher order interpolation to work properly, it is necessary to make the background grid at least 1 grid point larger than the region where you actually want to track particles. Otherwise the particles will see a zero field at the last grid point.

The background field can be used in two ways. First the field can be defined as a sum of predefined ICOOL fields. Second the user can read in an external file containing a 3D field grid. The background field, if any, is superposed on the ICOOL cell and region fields.

BACKGROUND begins a background field definition.
BFIELD defines a field contribution to the total background field.
ENDB ends the definition of a background field.

```
Example: define a new field for the following 3 m consisting of a 0.5 T
solenoid and a 0.2 T/m quadrupole field.
    BACKGROUND
        .false. 0.200 3.00
        -0.20 0.02 21 
    BFIELD
    0. 0.20 0.00 3.00
    SOL
```



```
    BFIELD
    0. 0.20 0.00 3.00
    QUAD
```



```
    ENDB
```

Example: use an external field grid, e.g. for044.dat, as a background field for the next 3 m . BACKGROUND

```
.false. 0.200 3.00
    -0.20 0.02 21 -0.20 0.02 21 0. 0.02 151 
    BFIELD
    0. 0.20 0.00 3.00
    STUS
```


ENDB

### 6.6 Miscellaneous

DUMMY inserts a dummy placeholder in a job. This can be used to reserve a place for an OUTPUT command without changing the region numbers used for ICOOL's internal diagnostics.

## 7. Field description

ICOOL uses three methods for specifying the field environment of the tracked particles.
(1) Each region has a field associated with it. For simple problems this may provide a sufficient description of the field. (2) Groups of regions can be associated together into cells. Each cell has a field associated with it that applies to all the regions in the cell. This can be used for example to describe RF cavity regions inside a solenoid field over the cell. (3) A background field grid can be defined over a specified region of $s$ from any combination of region fields or from a user-supplied field grid. This can be used, for example, to build a complicated field pattern or to apply error fields to some part of a lattice. The actual field seen by a particle at a given location is the superposition of all of these component fields.

### 7.1 Hard-edge fields

Hard-edge fields drop abruptly to 0 at the ends. The simplest hard-edge fields are uniform longitudinally over the entire length of the region. These are listed in the following table.

| Longitudinally uniform hard-edge fields |  |  |
| :--- | :--- | :--- |
| type | description | model |
| ACCEL | RF cavity | 1 (mode=0) |
| BROD | bent axial rod | 1 |
| BSOL | bent solenoid | 1 |
| DIP | dipole | 1,3 |
| EFLD | electric field | 1 |
| HDIP | horizontal dipole | 1 |
| QUAD | quadrupole | 1 |
| ROD | axial current rod | 1 |
| SEX | sextupole | 1 |
| SOL | solenoid | 1 |
| SQUA | skew quadrupole | 1 |

```
Example: 20 cm long, 1 T/m quadrupole field
    SREGION
    0.20 1 0.01
    1 0. 0.30
    QUAD
    1. 1.0 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
    VAC
    CBLOCK
    0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
```

There is in addition a hard edge dipole model with rotated pole faces.


### 7.2 Linear longitudinal variation

There are several simple models where the longitudinal field variation is piecewise linear.

| Longitudinally non-uniform hard-edge fields |  |  |
| :--- | :--- | :--- |
| type | description | model |
| FOFO | alternating solenoid | 1 |
| ROD | axial current rod | 2,3 |
| SOL | solenoid | 1 |

```
Example: 1 m long, 1 T solenoid with 10 cm end regions
        SREGION
        1.00 1 0.01
        1 0. 0.30
        SOL
        1. 1.0 0.80 0.10 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
        VAC
        CBLOCK
        0.0.0.0. 0. 0. 0. 0. 0. 0.
```


### 7.3 Soft edge fields

Soft edge fields in ICOOL use a $\Delta \tanh (\mathrm{s})$ longitudinal field dependence.

| Soft-edge fields |  |  |
| :--- | :--- | :--- |
| type | description | model |
| BSOL | bent solenoid | 2 |
| DIP | dipole | 2 |
| HDIP | horizontal dipole | 2 |
| QUAD | quadrupole | 2 |
| ROD | axial current rod | 4 |
| SEX | sextupole | 2 |
| SOL | solenoid | 2 |
| SQUA | skew quadrupole | 2 |

Example: 60 cm long, 1 T solenoid with 10 cm symmetric fall-off at the ends
SREGION
$1.00 \quad 1 \quad 0.01$
10.0 .10

SOL
2. 1.0 0.60 0.20 3. 0.10 0. 0. 0. 0. 0. 0. 0. 0. 0.

VAC
CBLOCK
0.0.0.0.0.0.0.0.0.0. 0 .

### 7.4 Current elements

ICOOL can produce a field from a set of current elements.

| Fields from current elements |  |  |
| :--- | :--- | :--- |
| type | description | model |
| BLOCK | current block | 1,2 |
| BLOCK | tapered current | 4 |
| COIL | current loop | 1,2 |
| SHEET | current sheet | $1-4$ |
| SOL | solenoid | $3-5$ |
| HELIX | helical winding | 2,3 |

```
Example: field from 20 cm long, 30 cm radius cylindrical current block
    SREGION
    1.00 1 0.01
    1 0. 0.30
    BLOCK
    1.0.40 0.30 0.40 0.20. 100.0.0.0.0. 0.0.0.0.0.
    VAC
    CBLOCK
    0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
Example: field from an external file, e.g. for022.dat, containing sheet
parameters. Prepare a grid and interpolate field values from the grid.
    SREGION
    1.00 1 0.01
    1 0. 0.30
    SHEET
    3. 22 0.01 0.01 1.01 0.31 10. 2. 0.0. 1.0.0.0.0.
    VAC
    CBLOCK
    0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
```


### 7.5 B on-axis

The field in a region can be determined if either the field or multipole coefficients are given on-axis.

| Fields from B on-axis |  |  |
| :--- | :--- | :--- |
| type | description | model |
| SOL | solenoid | 9 |
| BSOL | bent solenoid | 3,4 |
| BSOL | tapered lattice | 5 |
| HELIX | helix | 4 |

Example: combined dipole plus solenoid field from an external file, e.g. for023.dat, of on-axis multipole coeffients. Use as a cell field.

CELL
1
.false.
BSOL
$\begin{array}{llllllllllllllll}4 & 23 & 0.200 & 4 & 0.80 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 0\end{array}$
ENDC

### 7.6 Field grids

The field may be taken from a field grid in an external file. The first four types give essentially equivalent ways to input a 2D r-z field grid.

| Field grids |  |  |
| :--- | :--- | :--- |
| type | description | model |
| BLOCK | $\mathrm{r}-\mathrm{z}$ | 3 |
| COIL | $\mathrm{r}-\mathrm{z}$ | 3 |
| SHEET | $\mathrm{r}-\mathrm{z}$ | 5 |
| SOL | $\mathrm{r}-\mathrm{z}$ | 6 |
| STUS | $\mathrm{x}-\mathrm{y}-\mathrm{s}$ | 1,2 |

```
Example: find the field from an r-z grid that has been previously stored
in grid #3 using a GRID command.
    SREGION
        1.00 1 0.01
        1 0.0.30
        SHEET
        5. 3 2 0 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
        VAC
        CBLOCK
        0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
Example: find the field from a user-supplied static 3D field grid in an
external file, e.g. for023.dat.
        SREGION
        1.00 1 0.01
        1 0.0.30
        STUS
        1. 23 0 0.200 1. 2. 0. 0. 0. 0. 0. 0. 0. 0. 0.
        VAC
        CBLOCK
        0. 0. 0. 0. 0. 0. 0. 0. 0. 0.
```


### 7.7 Miscellaneous static fields

| Other static fields |  |  |
| :--- | :--- | :--- |
| type | description | model |
| FOFO | alternating solenoid | 2 |
| HELIX | helix | 1 |
| HORN | toroidal horn | 1,2 |
| WIG | wiggler | 1,2 |
| ROD | axial current rod | 5 |
| SOL | solenoid | 7 |

```
Example: 10 cells of an alternating solenoid lattice with peak field of
1 T and magnetic period of 1 m. Use as a cell field.
    CELL
        10
    .true.
    FOFO
    2 1.0 0. 1.00 0. 0 0 0 0 0 0 0 0 0 0 0 0
    ENDC
```


### 7.8 Time-varying fields

| Time-varying fields |  |  |
| :--- | :--- | :--- |
| type | description | model |
| ACCEL | E Z only | 1 (mode 1) |
|  | RF cavity | $2-4,9-13$ |
|  | induction linac | $6-8$ |
|  | external RF file | 5 |
| KICK | kicker | $1,2,3$ |

Example: 201 MHZ pillbox RF cavity SREGION
$0.30 \quad 1 \quad 0.01$
10.0 .55

ACCE
2. 201. 10000 0. 0. 0. 0. 0. 0. 0. 0. 0. 0 .

VAC
CBLOCK
0. 0. 0. 0. 0. 0. 0. 0. 0. 0.

## 8. ICOOL utilities

We list here some of the useful utility programs that operate on ICOOL files.

### 8.1 ERRSUM2

This program creates a summary of error messages by reading the ICOOL log file for002.dat. It displays a short summary on the terminal and writes a more detailed description to the file errsum2.dat.

### 8.2 ECALC9F

This is the standard emittance calculation program for ICOOL. It was written by Gregg Penn. It reads the ICOOL file for009.dat. The calculation is controlled with the input file ecalc9f.inp. The output of the program is written to the file ecalc9f.dat. ECALC9F assumes azimuthal symmetry in the fields and is most suited to solenoidal channels. Two variations of the code, ECALCXY and EMITCALC, are more suited for channels containing dipoles or quadrupoles.

### 8.3 ENDOF9

This program reads the ICOOL file for009.dat and produces a new file endof9.dat in the for009 format that only contains information from tracks that make it to the end of the simulation.

### 8.4 NOTEND9

This program is the inverse of ENDOF9. It reads the ICOOL file for009.dat and produces a new file notend9.dat in the for009 format that only contains information from tracks that do not make it to the end of the simulation. This could be useful, for example, in understanding the acceptance of a channel.

### 8.5 EXTPAR9

This program reads the ICOOL file for009. dat and produces a new file extpar9.dat in the for009 format that only contains information on specified tracks. The calculation is controlled with the input file extpar9.inp.

### 8.6 EXTREG9

This program reads the ICOOL file for009.dat and produces a new file extreg9.dat in the for009 format that only contains information at specified regions. The calculation is controlled with the input file extreg9.inp.

### 8.7 BUNCH9

This program reads the ICOOL file for009.dat. It produces a short summary of the amount of bunching on the terminal and a more detailed analysis in the file bunch9.dat. Several auxiliary files are also produced. The calculation is controlled with the input file bunch9.inp.

## 9. Other useful utilities

### 9.1 HistoRoot

The HistoRoot program, which is distributed in conjunction with G4Beamline, can be used to examine for009 output.

## 10. References

[1] R.C. Fernow, ICOOL: A simulation code for ionization cooling of muon beams, Proc. 1999 Particle Accelerator Conference, New York, 1999, p. 3020.
[2] R.C. Fernow, Recent developments on the muon-facility design code ICOOL, Proc. 2005 Particle Accelerator Conference, Knoxville, 2005, p. 2651.
[3] W. Allison et al, Ab initio liquid hydrogen muon cooling simulations with ELMS, J. Phys. G: Nucl. Part. Phys. 34 (2007) 679-685.
[4] Sergei Striganov, On the theory and simulation of multiple Coulomb scattering of heavy charged particles, FERMILAB-Conf-04/056-AD, April 2004.

