# USER’S GUIDE

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Conventions

The accelerator and/or beam line to be studied is described as a sequence of beam elements placed sequentially along a reference orbit. The reference orbit is the path of a charged particle having the central design momentum of the accelerator through idealised magnets with no fringe fields (see Figure 1).

The reference orbit consists of a series of straight line segments and circular arcs. It is defined under the assumption that all elements are perfectly aligned. The accompanying tripod of the reference orbit spans a local curvilinear right-handed coordinate system \( (x,y,s) \) The local \( s \)-axis is the tangent to the reference orbit. The two other axes are perpendicular to the reference orbit and are labelled \( x \) (in the bend plane) and \( y \) (perpendicular to the bend plane).

- Closed Orbit
- Global Reference System
- Local Reference System
- Sign Conventions for Magnetic Fields
- Variable
  - Canonical Variables Describing Orbits
  - Normalised Variables and other Derived Quantities
- Physical Units

Figure 1: Local Reference System

\[ \text{hansg} \ May \ 8, \ 2001 \]
Closed Orbit

Due to various errors like misalignment errors, field errors, fringe fields etc., the closed orbit does not coincide with the reference orbit. It also changes with the momentum error. The closed orbit is described with respect to the reference orbit, using the local reference system \((x, y, s)\). It is evaluated including any nonlinear effects.

MAD also computes the betatron and synchrotron oscillations with respect to the closed orbit. Results are given in the local \((x, y, s)\)-system defined by the reference orbit.

\[\text{hansg} \quad \text{January 24, 1997}\]
Global Reference System

The global reference orbit of the accelerator is uniquely defined by the sequence of physical elements. The local reference system \((x, y, s)\) may thus be referred to a global Cartesian coordinate system \((X, Y, Z)\) (see Figure 1). The positions between beam elements are numbered \(0,\ldots,i,\ldots,n\). The local reference system \((x_i, y_i, s_i)\) at position \(i\), i.e. the displacement and direction of the reference orbit with respect to the system \((X, Y, Z)\) are defined by three displacements \((X_i, Y_i, Z_i)\) and three angles \((\theta_i, \phi_i, \psi_i)\). The above quantities are defined more precisely as follows:

- **X**: Displacement of the local origin in \(X\)-direction.
- **Y**: Displacement of the local origin in \(Y\)-direction.
- **Z**: Displacement of the local origin in \(Z\)-direction.
- **THETA**: Angle of rotation (azimuth) about the global \(Y\)-axis, between the global \(Z\)-axis and the projection of the reference orbit onto the \((Z, X)\)-plane. A positive angle THETA forms a right-hand screw with the \(Y\)-axis.
- **PHI**: Elevation angle, i.e. the angle between the reference orbit and its projection onto the \((Z, X)\)-plane. A positive angle PHI correspond to increasing \(Y\). If only horizontal bends are present, the reference orbit remains in the \((Z, X)\)-plane. In this case PHI is always zero.
- **PSI**: Roll angle about the local \(s\)-axis, i.e. the angle between the intersection \((x, y)\)- and \((Z, X)\)-planes and the local \(x\)-axis. A positive angle PSI forms a right-hand screw with the \(s\)-axis.

The angles \((\theta_i, \phi_i, \psi_i)\) are not the Euler angles. The reference orbit starts at the origin and points by default in the direction of the positive \(Z\)-axis. The initial local axes \((x, y, s)\) coincide with the global axes \((X, Y, Z)\) in this order. The six quantities \((X_0, Y_0, Z_0, \theta_0, \phi_0, \psi_0)\) thus all have zero initial values by default. The program user may however specify different initial conditions.

Internally the displacement is described by a vector \(V\) and the orientation by a unitary matrix \(W\). The column vectors of \(W\) are the unit vectors spanning the local coordinate axes in the order \((x, y, s)\). \(V\) and \(W\) have the values:

\[
V = \begin{pmatrix} X \\ Y \\ Z \end{pmatrix}, \quad W = \Theta \Phi \Psi
\]

where
The reference orbit should be closed and it should not be twisted. This means that the displacement of the local reference system must be periodic with the revolution frequency of the accelerator, while the position angles must be periodic modulo(2 π) with the revolution frequency. If PSI is not periodic module(2 π), coupling effects are introduced. When advancing through a beam element, MAD computes $V_i$ and $W_i$ by the recurrence relations

$$V_i = W_{i-1}R_i + V_{i-1}, \quad W_i = w_{i-1}S_i.$$ 

The vector $R_i$ is the displacement and the matrix $S_i$ is the rotation of the local reference system at the exit of the element $i$ with respect to the entrance of the same element. The values of $R_i$ and $S_i$ are listed in the:

- straight reference system

for each physical element type.

![Figure 1: Global Reference System](image)

hansg, January 24, 1997
Local Reference Systems

Reference System for Straight Beam Elements

In straight elements the local reference system is simply translated by the length of the element along the local \( s \)-axis. This is true for

- Drift space
- Quadrupole
- Sextupole
- Octupole
- Solenoid
- CRAB CAVITY
- RF cavity
- Electrostatic separator
- Closed orbit corrector
- Beam position monitor

The corresponding \( R \) and \( S \) are

\[
R = \begin{pmatrix} 0 \\ 0 \\ L \end{pmatrix}, \quad S = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{pmatrix}.
\]

A rotation of the element about the \( s \)-axis has no effect on \( R \) and \( S \), since the rotations of the reference system before and after the element cancel.

Figure 1: Reference System for Straight Beam Elements
Reference System for Bending Magnets

Bending magnets have a curved reference orbit. For both rectangular and sector bending magnets

\[ R = \begin{pmatrix} \rho (\cos \alpha - 1) \\ 0 \\ \rho \sin \alpha \end{pmatrix}, \quad S = \begin{pmatrix} \cos \alpha & 0 & -\sin \alpha \\ 0 & 1 & 0 \\ \sin \alpha & 0 & \cos \alpha \end{pmatrix}, \]

where \( \alpha \) is the bend angle. A positive bend angle represents a bend to the right, i.e. towards negative \( x \) values. For sector bending magnets, the bend radius is given by \( \rho \), and for rectangular bending magnets it has the value

\[ \rho = L / 2 \sin(\alpha/2). \]

If the magnet is rotated about the \( s \)-axis by an angle \( \psi \), \( R \) and \( S \) are transformed by

\[ R^* = \mathbf{T} R, \quad S^* = \mathbf{T} S \mathbf{T}^{-1}. \]

where \( \mathbf{T} \) is the orthogonal rotation matrix

\[ \mathbf{T} = \begin{pmatrix} \cos \psi & -\sin \psi & 0 \\ \sin \psi & \cos \psi & 0 \\ 0 & 0 & 1 \end{pmatrix}. \]

The special value \( \psi = \pi/2 \) represents a bend down.

![Figure 2: Reference System for Rectangular Bends; The signs of the pole-face rotations are positive as shown.](image-url)
Elements which do not Change the Local Reference

**MARKER** elements do not affect the reference orbit. They are ignored for geometry calculations.
Drift Space

label: DRIFT,L=real;

A DRIFT space has one real attribute:

- L: The drift length (default: 0 m)

Examples:

DR1:   DRIFT,L=1.5;
DR2:   DRIFT,L=DR1[L];

The length of DR2 will always be equal to the length of DR1. The straight reference system for a drift space is a cartesian coordinate system.

hansg  January 24, 1997
A QUADRUPOLE has four real attributes:

- **L**: The quadrupole length (default: 0 m).
- **K1**: The normal quadrupole coefficient
  
  \[ K_1 = 1/(B \rho) \left( \frac{\partial B_y}{\partial x} \right) \]

  The default is 0 m**(-2). A positive normal quadrupole strength implies horizontal focusing of positively charged particles.

- **K1S**: The skew quadrupole coefficient
  
  \[ K_{1s} = 1/(2 B \rho) \left( \frac{\partial B_x}{\partial x} - \frac{\partial B_y}{\partial y} \right) \]

  where \((x,y)\) is now a coordinate system rotated by \(45^\circ\) around \(s\) with respect to the normal one. The default is 0 m**(-2). A positive skew quadrupole strength implies defocusing (!) of positively charged particles in the \((x,s)\) plane rotated by \(45^\circ\) around \(s\) (particles in this plane have \(x = y > 0\)).

- **TILT**: The roll angle about the longitudinal axis (default: 0 rad, i.e. a normal quadrupole). A positive angle represents a clockwise rotation. A TILT=\(\pi/4\) turns a positive normal quadrupole into a negative skew quadrupole.

  **Please note that contrary to MAD8 one has to specify the desired TILT angle, otherwise it is taken as 0 rad. This was needed to avoid the confusion in MAD8 about the actual meaning of the TILT attribute for various elements.**

  **Note also that** \(K_1/K_{1s}\) **can be considered as the normal or skew quadrupole components of the magnet on the bench, while the TILT attribute can be considered as an tilt alignment error in the machine. In fact, a positive** \(K_1\) **with a tilt=0 is equivalent to a positive** \(K_{1s}\) **with positive tilt=+\(\pi/4\).**

Example:

\[ \text{QF: QUADRUPOLE, L=1.5, K1=0.001;} \]

The straight reference system for a quadrupole is a cartesian coordinate system.
**Sextupole**

(label: `SEXTUPOLE,L=real,K2=real,K2S=real,TILT=real;`)

A SEXTUPOLE has four real attributes:

- **L**: The sextupole length (default: 0 m).
- **K2**: The normal sextupole coefficient
  
  \[ K_2 = \frac{1}{(B \rho)} (\frac{\partial^2 B_y}{\partial x^2}). \]
  
  (default: 0 m**(-3)).
- **K2S**: The skew sextupole coefficient
  
  \[ K_{2S} = \frac{1}{2(B \rho)} (\frac{\partial^2 B_x}{\partial x^2} - \frac{\partial^2 B_y}{\partial y^2}). \]
  
  where (x,y) is now a coordinate system rotated by -30° around s with respect to the normal one. (default: 0 m**(-3)). A positive skew sextupole strength implies defocussing (!) of positively charged particles in the (x,s) plane rotated by 30° around s (particles in this plane have x > 0, y > 0).
- **TILT**: The roll angle about the longitudinal axis (default: 0 rad, i.e. a normal sextupole). A positive angle represents a clockwise rotation. A TILT=\(\pi/6\) turns a positive normal sextupole into a negative skew sextupole.

Please note that contrary to MAD8 one has to specify the desired TILT angle, otherwise it is taken as 0 rad. This was needed to avoid the confusion in MAD8 about the actual meaning of the TILT attribute for various elements.

Note also that \(K_2/K_{2S}\) can be considered as the normal or skew sextupole components of the magnet on the bench, while the TILT attribute can be considered as an tilt alignment error in the machine. In fact, a positive \(K_2\) with a tilt=0 is equivalent to a positive \(K_{2S}\) with positive tilt=\(+\pi/6\).

Example:

```
S: SEXTUPOLE,L=0.4,K2=0.00134;
```

The straight reference system for a sextupole is a cartesian coordinate system.

\[hans\] \[frs\] August 28, 2003
Octupole

label: OCTUPOLE,L=real,K3=real,K3S=real,TILT=real;

An OCTUPOLE has four real attributes:

- **L**: The octupole length (default: 0 m).
- **K3**: The normal octupole coefficient
  \[ K_3 = \frac{1}{(B \rho)} \left( \frac{\partial^3 B_y}{\partial x^3} \right). \]
  (default: 0 m**(-4)).
- **K3S**: The skew octupole coefficient
  \[ K_{3S} = \frac{1}{(2B \rho)} \left( \frac{\partial^3 B_x}{\partial x^3} - \frac{\partial^3 B_y}{\partial y^3} \right). \]
  where \((x,y)\) is now a coordinate system rotated by \(-22.5^\circ\) around \(s\) with respect to the normal one.
  (default: 0 m**(-4)). A positive skew octupole strength implies defocussing (!) of positively charged particles in the \((x,s)\) plane rotated by \(22.5^\circ\) around \(s\) (particles in this plane have \(x > 0, y > 0\)).
- **TILT**: The roll angle about the longitudinal axis (default: 0 rad, i.e. a normal octupole). A positive angle represents a clockwise rotation. A TILT=\(\pi/8\) turns a positive normal octupole into a negative skew octupole.

Please note that contrary to MAD8 one has to specify the desired TILT angle, otherwise it is taken as 0 rad. This was needed to avoid the confusion in MAD8 about the actual meaning of the TILT attribute for various elements.

**Note also that** \(K_3/K_{3S}\) can be considered as the normal or skew quadrupole components of the magnet on the bench, while the TILT attribute can be considered as an tilt alignment error in the machine. In fact, a positive \(K_3\) with a tilt=0 is equivalent to a positive \(K_{3S}\) with positive tilt=+\(\pi/8\).

Example:

O3: OCTUPOLE, L=0.3, K3=0.543;

The straight reference system for an octupole is a cartesian coordinate system. Octupoles are normally treated as thin lenses, except when tracking by Lie-algebraic methods.
**Solenoid**

**label:** SOLENOID, \( L=\text{real}, \ KS=\text{real}; \) *(thick version)*  
**label:** SOLENOID, \( L=0, \ KS=\text{real}, \ KSI=\text{real}; \) *(thin version)*

A SOLENOID has two (three) real attributes:

- **L:** The length of the solenoid (default: 0 m)
- **KS:** The solenoid strength \( K_s \) (default: 0 rad/m). For positive KS and positive particle charge, the solenoid field points in the direction of increasing \( s \).
- **KSI:** The solenoid integrated strength \( K_s \times L \) (default: 0 rad). This additional attribute is needed only when using the thin solenoid, where \( L=0! \)
- **\( KNL \) & **\( KSL:** Take note that one can specify multipole coefficients but they have no effect in MAD-X proper but are used for solenoids with multipoles in PTC.

Example:

```
SOLO: SOLENOID, \( L=2., \ KS=0.001; \)
THINSOLO: SOLENOID, \( L=0, \ KS=0.001, \ KSI=0.002; \)
```

The **straight reference system** for a solenoid is a cartesian coordinate system.

*hansg* January 27, 1977
CRAB Cavity

label: CRABCAVITY, L=real, VOLT=real, LAG=real, FREQ=real,
 rv1=integer, rv2=integer, rv3=integer, rv4=integer,
 rph1=integer, rph2=integer, lagf=real;

An CRABCAVITY has ten real attributes and seven integer attributes:

- **L**: The length of the cavity (default: 0 m)
- **VOLT**: The peak RF voltage (default: 0 MV). The effect of the cavity is
  \[
  \delta(p_x) = VOLT \times \sin(\phi - \omega \times t) \\
  \delta(E) = -VOLT \times \omega \times x \times \cos(\phi - \omega \times t) \\
  \text{(where, } \phi = \sin(2 \pi \times (LAG - HARMON \times f_0 t)))
  \]
- **LAG**: The initial phase lag [2pi] (default: 0).
- **FREQ**: The frequency [MHz] _fenergy_ (no default). Note that if the RF frequency is not given, it is computed from the harmonic number and the revolution frequency _f_0 as before. However, for deflecting structures this makes no sense, and the frequency is mandatory.
- **RV1**: Number of initial turns with zero voltage (default: 0).
- **RV2**: Number of turns to ramp voltage from zero to nominal (default: 0).
- **RV3**: Number of turns with nominal voltage (default: VOLT).
- **RV4**: Number of turns to ramp voltage from nominal to zero (default: 0).
- **RPH1**: Number of initial turns with nominal phase (default: 0).
- **EPHASE**: Value of the final crab RF phase [2pi] with respect to nominal value (default: 0).
- **RPH2**: Number of turns to ramp phase [2pi] from nominal to specified value (default: 0).
- **HARMON**: The harmonic number _h_ (no default). Only if the frequency is not given.

Please take note, that the following MAD8 attributes: BETRF, PG, SHUNT and TFILL are currently not implemented in MAD-X!

Note that crab cavities are only implemented for tracking purposes. TWISS will ignore any effect of the crab cavity.

A cavity requires the particle energy **ENERGY** and the particle charge **CHARGE** to be set by a **BEAM** command before any calculations are performed.

Example:

```
BEAM, PARTICLE=PROTON, ENERGY=7000.0;
CAVITY: CRABCAVITY, L=10.0, VOLT=5.0, LAG=0.0, FREQ=400,
rv1=0, rv2=50, rv3=1000, rv4=50, rph1=100, rph2=500, lagf=0.125;;
```
The straight reference system for a cavity is a cartesian coordinate system.

R. Calaga  September 2010
BEAM: Set Beam Parameters

Many commands in MAD-X require the setting of various quantities related to the beam in the machine. Therefore, MAD-X will stop with a fatal error if an attempt is made to expand (USE) a sequence for which no BEAM command has been issued before.

The quantities are entered by a BEAM command:

```
BEAM, PARTICLE=name, MASS=real, CHARGE=real,
    ENERGY=real, PC=real, GAMMA=real,
    EX=real, EXN=real, EY=real, EYN=real,
    ET=real, SIGT=real, SIGE=real,
    KBUNCH=integer, NPART=real, BCURRENT=real,
    BUNCHED=logical, RADIATE=logical, BV=integer, SEQUENCE=name;
```

Warning: BEAM updates, i.e. it replaces attributes explicitly mentioned, but does not return to default values for others! To reset to beam value defaults, use RESBEAM. The particle restmass and charge are defined by:

- **PARTICLE**: The name of particles in the machine. MAD knows the restmass and the charge for the following particles:
  - POSITRON: The particles are positrons (MASS=$m_e$, CHARGE=1),
  - ELECTRON: The particles are electrons (MASS=$m_e$, CHARGE=-1),
  - PROTON: The particles are protons (MASS=$m_p$, CHARGE=1),
  - ANTI proton: The particles are anti-protons (MASS=$m_p$, CHARGE=-1),
  - POSMUON: The particles are positive muons (MASS=$m_{mu}$, CHARGE=1),
  - NEGMUON: The particles are negative muons (MASS=$m_{mu}$, CHARGE=-1).

Therefore neither restmass nor charge can be modified for these predefined particles. On the other hand, for ions and all other user-defined particles the name, restmass, and charge can be entered independently.

By default the total particle energy is 1 GeV. A different value can be defined by one of the following:

- **ENERGY**: The total energy per particle in GeV. If given, it must be greater than the particle restmass.
- **PC**: The momentum per particle in GeV/c. If given, it must be greater than zero.
- **GAMMA**: The ratio between total energy and rest energy of the particles: $\Gamma = E / m_0$. If given, it must be greater than one. If the restmass is changed a new value for the energy should be entered. Otherwise the energy remains unchanged, and the momentum PC and the quantity GAMMA are recalculated. The emittances are defined by:
- **EX**: The horizontal emittance $E_x$ (default: 1 m).
- **EY**: The vertical emittance $E_y$ (default: 1 m).
- **ET**: The longitudinal emittance $E_t$ (default: 1 m). The emittances can be replaced by the normalised emittances and the energy spread:
  - **EXN**: The normalised horizontal emittance [m]: $E_{xn} = 4 \left( \Gamma^2 - 1 \right)^{1/2} E_x$ (ignored if $E_x$ is given).
  - **EYN**: The normalised vertical emittance [m]: $E_{yn} = 4 \left( \Gamma^2 - 1 \right)^{1/2} E_y$ (ignored if $E_x$ is given).
- **SIGT**: The bunch length $c \sigma(t)$ in [m].
- **SIGE**: The relative energy spread $\sigma(E)/E$ in [1].

Certain commands compute the synchrotron tune $Q_s$ from the RF cavities. If $Q_s$ is non-zero, the relative energy spread and the bunch length are

\[
\sigma(E)/p_0 c = (2\pi Q_s E_t / \eta C)^{1/2},
\]

\[
c \sigma(t) = (\eta C E_t / 2\pi Q_s)^{1/2}.
\]

$C$ is the machine circumference, and $\eta = \Gamma^{-2} - \Gamma$ (transition) $^{-2}$.

The order of precedence in the parameter evaluation is given below:

```
particle->(mass+charge)
energy->pc->gamma->beta
ex->exn
ey->eyn
current->npart
et->sigt->sige
```

where any item to the left takes precedence over the others.

Finally, the BEAM command accepts

- **KBUNCH**: The number of particle bunches in the machine (default: 1).
- **NPART**: The number of particles per bunch (default: 0).
- **BCURRENT**: The bunch current (default: 0 A).
- **BUNCHECKED**: A logical flag. If set, the beam is treated as bunched whenever this makes sense.
- **RADIATE**: A logical flag. If set, synchrotron radiation is considered in all bipolar magnets.
- **BV**: an integer specifying the direction of the particle movement in a beam line; either +1 (default), or -1. For a detailed explanation see under bv flag.
- **SEQUENCE**: this attaches the beam command to a specific sequence; if the name is omitted, the BEAM command refers to the default beam always present. Sequences without attached beam use this default beam. When updating a beam, the corresponding sequence name, if any, must always be mentioned.
The BEAM command changes only the parameters entered. The command RESBEAM resets all beam data to their beam value defaults.

Examples:

BEAM, PARTICLE=ELECTRON,ENERGY=50,EX=1.E-6,EY=1.E-8,SIGE=1.E-3;
... BEAM, RADIATE;
... RESBEAM;
BEAM, EX=2.E-5,EY=3.E-7,SIGE=4.E-3;

The first command selects electrons, and sets energy and emittances. The second one turns on synchrotron radiation. The last two select positrons (by default), set the energy to 1 GeV (default), clear the synchrotron radiation flag, and set the emittances to the values entered.

Some program modules of MAD-X may also store data into a beam data block. Expressions may refer to data in this beam data block using the notation

BEAM->attribute-name

or

BEAM%sequence->attribute-name.

This notation refers to the value of attribute-name found in the default BEAM resp. the beam belonging to the sequence given. This can be used for receiving or using values, e.g.

value,beam%lhcb2->bv;

or for storing values in the beam (this does NOT trigger an update of dependent variables !), e.g.

beam->charge=-1;

The current values in the BEAM bank can be obtained by the command

show,beam;

resp.

show,beam%sequence;

hansg 11.9.2000
RF Cavity

label: RFCAVITY, L=real, VOLT=real, LAG=real, HARMON=integer, FREQ=real;

An RFCAVITY has eight real attributes and one integer attribute:

- **L**: The length of the cavity (DEFAULT: 0 m)
- **VOLT**: The peak RF voltage (DEFAULT: 0 MV). The effect of the cavity is
  \[ \text{delta}(E) = \text{VOLT} \times \sin(2\pi \times (\text{LAG} - \text{HARMON} \times f_0 \times t)) \].
- **LAG**: The phase lag [2pi] (DEFAULT: 0).
- **FREQ**: The frequency [MHz] (no DEFAULT). Note that if the RF frequency is not given, it is computed from the harmonic number and the revolution frequency \( f_0 \) as before. However, for accelerating structures this makes no sense, and the frequency is mandatory.
- **HARMON**: The harmonic number \( h \) (no DEFAULT). Only if the frequency is not given.

Please take note, that the following MAD8 attributes: BETRF, PG, SHUNT and TFILL are currently not implemented in MAD-X!

Note as well that twiss is 4D only. As a consequence the TWISS parameters in the plane of non-zero dispersion may not close as expected. Therefore, it is best to perform TWISS in 4D only, i.e. with cavities switched off. If 6D is needed one has to use the `ptc_twiss` command.

The RFCAVITY has attributes that will only become active in PTC:

- **n_bessel** (DEFAULT: 0):
  Transverse focussing effects are typically ignored in the cavity in MAD-X or even PTC. This effect is being calculated to order \( n_{bessel} \), with \( n_{bessel}=0 \) disregarding this effect and with a correct treatment when \( n_{bessel} \) goes to infinity.
- **no_cavity_totalpath** (DEFAULT: no_cavity_totalpath=false):
  flag to choose if in a cavity the transit time factor is considered (no_cavity_totalpath=false) or if the particle is kept on the crest of RF voltage (no_cavity_totalpath=true).

A cavity requires the particle energy (\texttt{ENERGY}) and the particle charge (\texttt{CHARGE}) to be set by a \texttt{BEAM} command before any calculations are performed.

Example:
The straight reference system for a cavity is a cartesian coordinate system.

hansg January 24, 1997
ELSEPARATOR: Electrostatic Separator

label: ELSEPARATOR, L=real, EX=real, EY=real, TILT=real;

An ELSEPARATOR (electrostatic separator) has four real attributes:

- **L**: The length of the separator (default: 0 m).
- **EX**: The horizontal electric field strength (default: 0 MV/m). A positive field increases \( p_x \) for positive particles.
- **EY**: The vertical electric field strength (default: 0 MV/m). A positive field increases \( p_y \) for positive particles.
- **TILT**: The roll angle about the longitudinal axis (default: 0 rad). A positive angle represents a clockwise rotation of the electrostatic separator.

A separator requires the particle energy (ENERGY) and the particle charge (CHARGE) to be set by a BEAM command before any calculations are performed.

Example:

```
BEAM, PARTICLE=POSITRON, ENERGY=50.0;
SEP: ELSEPARATOR, L=5.0, EY=0.5;
```

The straight reference system for a separator is a cartesian coordinate system.

*frs* August 28, 2003
Closed Orbit Correctors

Three types of closed orbit correctors are available:

- **HKICKER**, a corrector for the horizontal plane,
- **VKICKER**, a corrector for the vertical plane,
- **KICKER**, a corrector for both planes.

<table>
<thead>
<tr>
<th>Label</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>HKICKER</td>
<td>$L=\text{real}, K\text{ICK}=\text{real}, T\text{ILT}=\text{real}$</td>
</tr>
<tr>
<td>VKICKER</td>
<td>$L=\text{real}, K\text{ICK}=\text{real}, T\text{ILT}=\text{real}$</td>
</tr>
<tr>
<td>KICKER</td>
<td>$L=\text{real}, HK\text{ICK}=\text{real}, VK\text{ICK}=\text{real}, T\text{ILT}=\text{real}$</td>
</tr>
</tbody>
</table>

The type **KICKER** should not be used when an orbit corrector kicks only in one plane.

The attributes have the following meaning:

- **L**: The length of the closed orbit corrector (default: 0 m).
- **KICK**: The kick angle for either horizontal or vertical correctors. (default: 0 rad).
- **HKICK**: The horizontal kick angle for a corrector in both planes (default: 0 rad).
- **VKICK**: The vertical kick angle for a corrector in both planes (default: 0 rad).
- **TILT**: The roll angle about the longitudinal axis (default: 0 rad). A positive angle represents a clockwise rotation of the kicker.

A positive kick increases $p_x$ or $p_y$ respectively. This means that a positive horizontal kick bends to the left, i.e. to positive $x$ which is opposite of what is true for bends.

It should be noted that the kick values assigned to an orbit corrector like above are not overwritten by an orbit correction using the **CORRECT** command. Instead the kicks computed by an orbit correction and the assigned values are added when the correctors are used.

**Examples:**

| HK1:  | HKICKER, KICK=0.001;             |
| VK3:  | VKICKER, KICK=0.0005;            |
| VK4:  | VKICKER, KICK:=AVK4;             |
| KHV1: | KICKER, HKICK=0.001,VKICK=0.0005;|
| KHV2: | KICKER, HKICK:=AKHV2H,VKICK:=AKHV2V;|

The assignment in the form of a deferred expression has the advantage that the values can be assigned and/or modified at any time (and matched !).

The **straight reference system** for an orbit corrector is a Cartesian coordinate system.
Please note that there is a new feature introduced by Stefan Sorge from GSI. Here his description:

The elements KICKER, HKICKER, and VKICKER can also be used as an exciter providing a sinusoidal momentum kick. The usage in this case is

\[
\text{xykick: KICKER, SINKICK=integer, SINPEAK=real, SINTUNE=real, SINPHASE=real;} \\
x\text{kick : HKICKER, SINKICK=integer, SINPEAK=real, SINTUNE=real, SINPHASE=real;} \\
y\text{kick : VKICKER, SINKICK=integer, SINPEAK=real, SINTUNE=real, SINPHASE=real;}
\]

where a sinusoidal momentum kick \( dpz \) as a function of the revolution number \( n \) given by

\[
dpz(n)=\text{SINPEAK} \times \sin(2\times\pi\times\text{SINTUNE}\times n + \text{SINPHASE}), \quad pz=px,py
\]

is provided. So, the variables are

SINKICK - integer, must be set to 1 to switch on the sinusoidal signal, default: 0.

SINPEAK - amplitude of the bending angle (rad), default: 0 rad.

SINTUNE - frequency of the signal times the revolution frequency. Hence, the phase per revolution is \( 2\times\pi\times\text{SINTUNE} \), default: 0.

SINPHASE - initial phase, default: 0 rad.

KICKER generates a kick in horizontal and a kick vertical direction, where both are synchron, HKICKER generates a horizontal kick, and VKICKER generates a vertical kick.

The momentum kick of a kicker has only a single frequency. An element having a finite bandwidth can approximately created by defining thin kickers with all amplitudes SINPEAK, frequencies SINTUNE, and initial phases SINPHASE desired and putting them at the same position \( s \) in the accelerator.

From S.Sorge@gsi.de

\[\text{hansi frs August 28, 2003}\]
Beam Position Monitors

A beam monitor acts on the beam like a drift space. In addition it serves to record the beam position for closed orbit corrections. Four different types of beam position monitors are recognised:

- **HMONITOR.** Monitor for the horizontal beam position,
- **VMONITOR.** Monitor for the vertical beam position,
- **MONITOR.** Monitor for both horizontal and vertical beam position.
- **INSTRUMENT.** A place holder for any type of beam instrumentation. Optically it behaves like a drift space; it returns *no beam observation*. It represent a class of elements which is completely independent from drifts and monitors.

```plaintext
label: HMONITOR,   L=real;
label: VMONITOR,   L=real;
label: MONITOR,    L=real;
label: INSTRUMENT, L=real;
```

A beam position monitor has one real attribute:

- **L:** The length of the monitor (default: 0 m). If the length is different from zero, the beam position is recorded in the centre of the monitor.

Examples:

```plaintext
MH: HMONITOR, L=1;
MV: VMONITOR;
```

The [straight reference system](#) for a monitor is a cartesian coordinate system.

_hansg_ June 17, 2002
Bending Magnets

Two different type keywords are recognised for bending magnets, they are distinguished only by the reference system used:

- **RBEND** is a rectangular bending magnet. It has parallel pole faces and is based on a curved reference system; its length is the straight length as in the Figure but internally the arc length is being used. To define an RBEND with the arc length as length (straight line shorter than input for compatibility with MAD8 version up to version 8.23.06 including), the option RBARC=FALSE has to be set.

- **SBEND** is a sector bending magnet. Its pole faces meet at the centre of curvature of the curved reference system.

They are defined by the commands:

```plaintext
SBEND,  L=real,ANGLE=real,TILT=real,K0=real,K0S=real,K1=real,E1=real,E2=real, 
        FINT=real,FINTX=real,HGAP=real,K2=real,H1=real,H2=real;
```

```plaintext
RBEND,  L=real,ANGLE=real,TILT=real,K0=real,K0S=real,K1=real,E1=real,E2=real, 
        FINT=real,FINTX=real,HGAP=real,K2=real,H1=real,H2=real;
```

For both types, the following first-order attributes are permitted:

- **L**: The length of the magnet (default: 0 m). For a rectangular magnet the length is measured along a straight line as in the Figure (internally the arc length is used), while for a sector magnet it is the arc length of the reference orbit. To define an RBEND with the arc length (shorter straight length), the option RBARC=FALSE has to be set.

- **ANGLE**: The bend angle (default: 0 rad). A positive bend angle represents a bend to the right, i.e. towards negative x values.

- **TILT**: The roll angle about the longitudinal axis (default: 0 rad, i.e. a horizontal bend). A positive angle represents a clockwise rotation. A TILT=pi/2 turns a horizontal into a vertical bend, i.e. a positive bend ANGLE denotes a deflection down. Please note that contrary to MAD8 one has to specify the desired TILT angle, otherwise it is taken as 0 rad. This was needed to avoid the confusion in MAD8 about the actual meaning of the TILT attribute for various elements.

- Please take note that K₀ and K₀s are left in the data base but are no longer used for the MAP of the bends (but see below for what K₀ is being used), instead ANGLE and TILT are used exclusively. We believe that this will allow for a clearer and unambiguous definition, in particular in view of the upcoming integration of MAD-X with PTC which will allow a more
general definition of bends. However, it is required to specify k0 to assign RELATIVE field errors to a bending magnet since k0 is used for the normalization and NOT the ANGLE. (see EFCOMP).

- K1: The quadrupole coefficient
  
  $K_1 = (1 / B \rho) \left( \frac{\partial B_y}{\partial x} \right)$.  
  
  The default is 0 $m^{-2}$. A positive quadrupole strength implies horizontal focusing of positively charged particles.

- E1: The rotation angle for the entrance pole face (default: 0 rad).

- E2: The rotation angle for the exit pole face (default: 0 rad).

- FINT: The field integral whose default value is 0.

- FINTX: Allows (FINTX > 0) to set FINT at the element exit different from its entry value. In particular useful to switch it off (FINTX=0).

- HGAP: The half gap of the magnet (default: 0 m).

The pole face rotation angles are referred to the magnet model for rectangular bend and sector bend respectively. The quantities FINT and HGAP specify the finite extent of the fringe fields as defined in [SLAC-75]. There they are defined as follows:

$$F_{\text{INT}} = \int_{-\infty}^{\infty} \frac{B_y(s) (B_0 - B_y(s))}{g \cdot B_0^2} ds, \quad g = 2 \cdot \text{HGAP},$$

The default values of zero corresponds to the hard-edge approximation, i.e. a rectangular field distribution. For other approximations, enter the correct value of the half gap, and one of the following values for FINT:

- Linear Field drop-off: 1/6
- Clamped "Rogowski" fringing field: 0.4
- Unclamped "Rogowski" fringing field: 0.7
- "Square-edged" non-saturating magnet: 0.45

Entering the keyword FINT alone sets the integral to 0.5. This is a reasonable average of the above values. The following second-order attributes are permitted:

- K2: The sextupole coefficient $K_2 = (1 / B \rho) (\partial^2 B_y / \partial x^2)$.

- H1: The curvature of the entrance pole face (default: 0 m$^{-1}$).

- H2: The curvature of the exit pole face (default: 0 m$^{-1}$). A positive pole face curvature induces a negative sextupole component; i.e. for positive H1 and H2 the centres of curvature of the pole faces are placed inside the magnet.

Examples:

```
BR: RBEND, L=5.5, ANGLE=+0.001; // Deflection to the right
BD: SBEND, L=5.5, K0S=+0.001/5.5; // Deflection up
BL: SBEND, L=5.5, K0=-0.001/5.5; // Deflection to the left
BU: SBEND, L=5.5, K0S=-0.001; // Deflection down
```
Marker.

label: MARKER;

The simplest element which can occur in a beam line is the MARKER. It has no effect on the beam, but it allows one to identify a position in the beam line, for example to apply a matching constraint.

Example:

m27: marker;

hansg June 6, 2002
Sign Conventions for Magnetic Fields

The MAD program uses the following Taylor expansion for the field on the mid-plane $y=0$, described in [SLAC-75]

$$B_y(x, 0) = \sum_{n=0}^{\infty} \frac{B_n x^n}{n!}$$

Note the factorial in the denominator. The field coefficients have the following meaning:

- $B_0$: Dipole field, with a positive value in the positive $y$ direction; a positive field bends a positively charged particle to the right.
- $B_1$: Quadrupole coefficient

$$B_1 = \frac{\partial B_y}{\partial x};$$

a positive value corresponds to horizontal focussing of a positively charged particle.
- $B_2$: Sextupole coefficient

$$B_2 = \frac{\partial^2 B_y}{\partial x^2}.$$  
- $B_3$: Octupole coefficient

$$B_3 = \frac{\partial^3 B_y}{\partial x^3}.$$  

Using this expansion and the curvature $h$ of the reference orbit, the longitudinal component of the vector potential to order 4 is:

$$A_\nu = B_0 \left( z - \frac{h x^2}{3(1 + a e)} \right) + B_1 \left( \frac{1}{2} (x^2 - y^2) - \frac{h}{6} x^3 + \frac{h^2}{24} (4x^2 - y^2) + \ldots \right) + B_2 \left( \frac{1}{6} (x^2 - 6y^2) - \frac{h}{5} (x^4 - y^4) + \ldots \right) + B_3 \left( \frac{1}{36} (x^4 - 6x^2y^2 + y^6) + \ldots \right) + \ldots$$

Taking curl $A$ in curvilinear coordinates, the field components can be computed as
It can be easily verified that both curl $\mathbf{B}$ and div $\mathbf{B}$ are zero to the order of the $B_3$ term. Introducing the magnetic rigidity $B_{\rho}$, the multipole coefficients are computed as

$$K_n = \frac{eB_n}{p_s} = \frac{B_n}{B_{\rho}}.$$
For each variable the physical units are listed in square brackets.

**Canonical Variables Describing Orbits**

MAD uses the following canonical variables to describe the motion of particles:

- X: Horizontal position $x$ of the (closed) orbit, referred to the ideal orbit [m].
- PX: Horizontal canonical momentum $p_x$ of the (closed) orbit referred to the ideal orbit, divided by the reference momentum: $PX = \frac{p_x}{p_0}$, [1].
- Y: Vertical position $y$ of the (closed) orbit, referred to the ideal orbit [m].
- PY: Vertical canonical momentum $p_y$ of the (closed) orbit referred to the ideal orbit, divided by the reference momentum: $PY = \frac{p_y}{p_0}$, [1].
- T: Velocity of light times the negative time difference with respect to the reference particle: $T = -c t$, [m]. A positive T means that the particle arrives ahead of the reference particle.
- PT: Energy error, divided by the reference momentum times the velocity of light: $PT = \frac{\Delta E}{p_s c}$, [1]. This value is only non-zero when synchrotron motion is present. It describes the deviation of the particle from the orbit of a particle with the momentum error DELTAP.
- DELTAP: Difference of the reference momentum and the design momentum, divided by the reference momentum: $DELTAP = \frac{\Delta p}{p_0}$, [1]. This quantity is used to normalize all element strengths.

The independent variable is:

- S: Arc length $s$ along the reference orbit, [m].

In the limit of fully relativistic particles ($\gamma \gg 1$, $v = c$, $p c = E$), the variables T, PT used here agree with the longitudinal variables used in [TRANSPORT]. This means that T becomes the negative path length difference, while PT becomes the fractional momentum error. The reference momentum $p_s$ must be constant in order to keep the system canonical.

**Normalised Variables and other Derived Quantities**

- XN: The normalised horizontal displacement
  
  $XN = x_n = Re(E_1 T S Z)$, [sqrt(m)].
- PXN: The normalised horizontal transverse momentum
  \[ PXN = x_n = \text{Im}(E_1^T S Z), \text{[sqrt(m)]}. \]
- WX: The horizontal Courant-Snyder invariant
  \[ WX = \sqrt{x_n^2 + p_{xn}^2}, \text{[m]}. \]
- PHIX: The horizontal phase
  \[ PHIX = -\frac{\text{atan}(p_{xn} / x_n)}{2 \pi} \text{[1]}. \]
- YN: The normalised vertical displacement
  \[ YN = x_n = \text{Re}(E_2^T S Z), \text{[sqrt(m)]}. \]
- PYN: The normalised vertical transverse momentum
  \[ PYN = x_n = \text{Im}(E_2^T S Z), \text{[sqrt(m)]}. \]
- WY: The vertical Courant-Snyder invariant
  \[ WY = \sqrt{y_n^2 + p_{yn}^2}, \text{[m]}. \]
- PHIY: The vertical phase
  \[ PHIY = -\frac{\text{atan}(p_{yn} / y_n)}{2 \pi} \text{[1]}. \]
- TN: The normalised longitudinal displacement
  \[ TN = x_n = \text{Re}(E_3^T S Z), \text{[sqrt(m)]}. \]
- PTN: The normalised longitudinal transverse momentum
  \[ PTN = x_n = \text{Im}(E_3^T S Z), \text{[sqrt(m)]}. \]
- WT: The longitudinal invariant
  \[ WT = \sqrt{t_n^2 + p_{tn}^2}, \text{[m]}. \]
- PHIT: The longitudinal phase
  \[ PHIT = +\frac{\text{atan}(p_{tn} / t_n)}{2 \pi} \text{[1]}. \]

in the above formulas \( Z \) is the phase space vector
\[ Z = (x, p_x, y, p_y, t, p_t)^T. \]
the matrix $S$ is the “symplectic unit matrix”

$$
S = \begin{pmatrix}
0 & 1 & 0 & 0 & 0 & 0 \\
-1 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & 0 & 0 \\
0 & 0 & -1 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 1 & 0 \\
0 & 0 & 0 & 0 & -1 & 0 \\
\end{pmatrix}
$$

and the vectors $E_i$ are the three complex eigenvectors.

**Linear Lattice Functions (Optical Functions)**

Several MAD commands refer to linear lattice functions. Since MAD uses the canonical momenta $(p_x, p_y)$ instead of the slopes $(x', y')$, their definitions differ slightly from those in [Courant and Snyder]. Notice that in MAD-X PT substitutes DELTAP as longitudinal variable. Dispersive and chromatic functions are hence derivatives with respects to PT. Being PT=BETA*DELTAP, where BETA is the relativistic Lorentz factor, those functions must be multiplied by BETA a number of time equal to the order of the derivative. The linear lattice functions are known to MAD under the following names:

- **BETX**: Amplitude function $\beta_x$, [m].
- **ALFX**: Correlation function $\alpha_x$, [1]:
  
  $$
  ALFX = \alpha_x = - \frac{1}{2} \left( \frac{\partial \beta_x}{\partial s} \right).
  $$

- **MUX**: Phase function $\mu_x$, [2pi]:
  
  $$
  MUX = \mu_x = \text{integral} \left( \frac{ds}{\beta_x} \right).
  $$

- **DX**: Dispersion $D_x$ of $x$, [m]:
  
  $$
  DX = D_x = \left( \frac{\partial x}{\partial \text{PT}} \right).
  $$

- **DPX**: Dispersion $D_{px}$ of $p_x$, [1]:
  
  $$
  DPX = D_{px} = \left( \frac{\partial p_x}{\partial \text{PT}} \right) / p_x.
  $$

- **BETY**: Amplitude function $\beta_y$, [m].
- **ALFY**: Correlation function $\alpha_y$, [1]:
  
  $$
  ALFY = \alpha_y = - \frac{1}{2} \left( \frac{\partial \beta_y}{\partial s} \right).
  $$
MUY: Phase function $\mu_y$, $[2\pi]$.

$$MUY = \mu_y = \int \frac{ds}{\beta_y}.$$ 

DY: Dispersion $D_y$ of $y$, $[m]$:

$$DY = D_y = \frac{\delta y}{\delta PT}.$$ 

DPY: Dispersion $D_{px}$ of $p_x$, $[1]$:

$$DPY = D_{py} = \frac{\delta p_y}{\delta PT} / p_s.$$ 

R11, R12, R21, R22: Coupling Matrix

ENERGY: The total energy per particle in GeV. If given, it must be greater then the particle mass.

### Chromatic Functions

Several MAD commands refer to the chromatic functions. $(p_x, p_y)$ instead of the slopes $(x', y')$, their definitions differ slightly from those in [Montague]. Notice that in MAD-X PT substitutes DELTAP as longitudinal variable. Dispersive and chromatic functions are hence derivatives with respects to PT. Being $PT = BETA*DELTAP$, where $BETA$ is the relativistic Lorentz factor, those functions must be multiplied by $BETA$ a number of time equal to the order of the derivative. The chromatic functions are known to MAD under the following names:

Please note that this option is needed for a proper calculation of the chromaticities in the presence of coupling!

WX: Chromatic amplitude function $W_x$, $[1]$:

$$WX = W_x = \sqrt{a_x^2 + b_x^2},$$

$$a_x = \frac{\delta \beta_x}{\delta PT} / \beta_x,$$

$$b_x = \frac{\delta \alpha_x}{\delta PT} - (\alpha_x / \beta_x) \cdot \frac{\delta \beta_x}{\delta PT}.$$ 

PHIX: Chromatic phase function $\Phi_x$, $[2\pi]$:

$$PHIX = \Phi_x = \text{atan}(a_x / b_x).$$ 

DMUX: Chromatic derivative of phase function $\mu_x$, $[2\pi]$:

$$DMUX = (\delta \mu_x / \delta PT).$$ 

DDX: Chromatic derivative of dispersion $D_x$, $[m]$: 

36
DDX = 1/2 * (del^2 x / del PT^2).

- **DDPX**: Chromatic derivative of dispersion $D_{px}$, [1]:
  
  $$DDPX = 1/2 * (del^2 p_x / del PT^2) / p_s.$$ 

- **WY**: Chromatic amplitude function $W_y$, [1]:
  
  $$WY = W_y = \sqrt{a_y^2 + b_y^2},$$
  
  $$a_y = (del \beta_y / del PT) / \beta_y,$$
  
  $$b_y = (del \alpha_y / del PT) - (\alpha_y / \beta_y) * (del \beta_y / del PT).$$

- **PHIY**: Chromatic phase function $\Phi_y$, [2pi]:
  
  $$PHIY = \Phi_y = \text{atan}(a_y / b_y).$$

- **DMUY**: Chromatic derivative of phase function $\mu_y$, [2pi]:
  
  $$DMUY = (del \mu_y / del PT).$$

- **DDY**: Chromatic derivative of dispersion $D_y$, [m]:
  
  $$DDY = 1/2 * (del^2 y / del PT^2).$$

- **DDPY**: Chromatic derivative of dispersion $D_{py}$, [1]:
  
  $$DDPY = 1/2 * (del^2 p_y / del PT^2) / p_s.$$ 

### Variables in the SUMM Table

After a successful TWISS command a summary table is created which contains the following variables:

- **LENGTH**: The length of the machine, [m].
- **ORBIT5**: The T ($= c t$, [m]) component of the closed orbit.
- **ALFA**: The momentum compaction $\alpha_p$, [1].
- **GAMMATR**: The transition energy $\gamma_{\text{transition}}$, [1].
- **Q1**: The horizontal tune $Q_1$, [1].
- **DQ1**: The horizontal chromaticity $dQ_1$, [1].
DQ1 = dq1 = (del Q1 / del PT).

- BETXMAX: The largest horizontal beta_x, [m].
- DXMAX: The largest horizontal dispersion [m].
- DXRMS: The r.m.s. of the horizontal dispersion [m].
- XCOMAX: The maximum of the horizontal closed orbit deviation [m].
- XRMS: The r.m.s. of the horizontal closed orbit deviation [m].
- Q2: The vertical tune Q2 [1].
- DQ2: The vertical chromaticity dq2, [1]:
  DQ2 = dq2 = (del Q2 / del PT).
- BETYMAX: The largest vertical beta_y, [m].
- Dymax: The largest vertical dispersion [m].
- Dyrms: The r.m.s. of the vertical dispersion [m].
- YCOMAX: The maximum of the vertical closed orbit deviation [m].
- YCORS: The r.m.s. of the vertical closed orbit deviation [m].
- DELTAP: Energy difference, divided by the reference momentum times the velocity of light, [1]:
  DELTAP = delta(E) / p \_x c.

Notice that in MAD-X PT substitutes DELTAP as longitudinal variable. Dispersive and chromatic functions are hence derivatives with respects to PT. Being PT=BETA*DELTAP, where BETA is the relativistic Lorentz factor, those functions must be multiplied by BETA a number of time equal to the order of the derivative.

**Variables in the TRACK Table**

The command RUN writes tables with the following variables:

- X: Horizontal position x of the orbit, referred to the ideal orbit [m].
- PX: Horizontal canonical momentum p_x of the orbit referred to the ideal orbit, divided by the reference momentum.
- Y: Vertical position y of the orbit, referred to the ideal orbit [m].
- PY: Vertical canonical momentum p_y of the orbit referred to the ideal orbit, divided by the reference momentum.
- T: Velocity of light times the negative time difference with respect to the reference particle, [m].
positive $T$ means that the particle arrives ahead of the reference particle.

- **PT**: Energy difference, divided by the reference momentum times the velocity of light, [1].

When tracking Lyapunov companions (not yet implemented), the TRACK table defines the following dependent expressions:

- **DISTANCE**: the relative Lyapunov distance between the two particles.
- **LYAPUNOV**: the estimated Lyapunov Exponent.
- **LOGDIST**: the natural logarithm of the relative distance.
- **LOGTURNS**: the natural logarithm of the turn number.

Throughout the computations MAD uses international (SI, Système International) units. These units are summarised in the Units table.

**Table 1: Physical Units**

<table>
<thead>
<tr>
<th>Description</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length</td>
<td>m (metres)</td>
</tr>
<tr>
<td>Angle</td>
<td>rad (radians)</td>
</tr>
<tr>
<td>Quadrupole coefficient</td>
<td>m**(-2)</td>
</tr>
<tr>
<td>Multipole coefficient, 2n poles</td>
<td>m**(-n)</td>
</tr>
<tr>
<td>Electric voltage</td>
<td>MV (Megavolts)</td>
</tr>
<tr>
<td>Electric field strength</td>
<td>MV/m</td>
</tr>
<tr>
<td>Frequency</td>
<td>MHz (Megahertz)</td>
</tr>
<tr>
<td>Phase angles</td>
<td>2 pi</td>
</tr>
<tr>
<td>Particle energy</td>
<td>GeV</td>
</tr>
<tr>
<td>Particle mass</td>
<td>GeV/c**2</td>
</tr>
<tr>
<td>Particle momentum</td>
<td>GeV/c</td>
</tr>
<tr>
<td>Beam current</td>
<td>A (Amperes)</td>
</tr>
<tr>
<td>Particle charge</td>
<td>e (elementary charges)</td>
</tr>
<tr>
<td>Impedances</td>
<td>MOhm (Megohms)</td>
</tr>
<tr>
<td>Emittances</td>
<td>π m mrad</td>
</tr>
<tr>
<td>RF power</td>
<td>MW (Megawatts)</td>
</tr>
<tr>
<td>Higher mode loss factor</td>
<td>V/pc</td>
</tr>
</tbody>
</table>

*Hansg June 17, 2002*
Command Format

- Statements
- Comments
- Identifiers or Label
- Command Attribute
  - Name or String Attribute
  - Logical Attribute
  - Integer Attribute
  - Real Expression built from operator and operand
  - A Deferred Expression is evaluated every time it is used
  - Constraint
  - Variable Name
- Wild Card Pattern

hansg May 8, 2001
EUROPEAN ORGANIZATION FOR NUCLEAR RESEARCH

Statements

and

Comments

Input for MAD-X follows in broad lines the new MAD-9 format, i.e. free format with commas ",," as separators; however, outside {...} enclosures blanks may be used as separators. Blank input lines do not affect program execution. The input is not case sensitive except for strings enclosed in " ".

The input file consists of a sequence of commands, also known as statements. A statement may occupy any number of input lines. It must be terminated by a semicolon, except if it contains a block of statements itself, like in

\[
\text{if (a < 3) \{a=b^2; b=2*b+4;\}}
\]

Several statements may be placed on the same line. When a "!" or "/" is found on an input line, the remaining characters of the line are skipped. A line "/" starts a comment region, it ends with a "*/" line. The general format for a command is (items enclosed in /rep/ ... /rep/ can be repeated any number of times, including zero):

\[
\text{label: keyword /rep/,attribute/rep/;}
\]

It has three parts:

- A \text{[label]} is required for a definition statement. It gives a name to the stored command.
- A \text{[keyword]} identifies the action desired.
- The \text{[attributes]} are normally entered in the form "attribute-name=attribute-value" and serve to define data for the command, where:
  - \text{[attribute-name]} selects the attribute,
  - \text{[attribute-value]} gives it a value.

If a value has to be assigned to an attribute, the attribute name is mandatory. For logical attributes it is sufficient to enter the name only. The attribute is then given a default value taken from the command dictionary.

Example: TILT attribute for various magnets.

The command attributes can have one of the following types:
Any integer or real attribute can be replaced by a real expression expressions are normally deferred (see deferred expression), except in the definition of constants where they are evaluated immediately. When a command has a label MAD-X keeps it in memory. This allows repeated execution of the same command by just entering EXEC label. This construct may be nested. For an exhaustive list of valid declarations of constants or variables see declarations.
Identifiers or Labels

A keyword begins with a letter and consists of letters and digits. The MAD-X keywords are protected; using one of them as a label results in a fatal error.

hansg May 8, 2001
String Attributes

A string attribute makes alphanumeric information available, e.g. a title or a file name. It can contain any characters, enclosed in single (’) or double (”) quotes, except for quotes of the type used as its delimiter.

Examples:

```
TITLE,’This is a title for the program run "test"';
system("ln -fns some-lengthy-directory-name local-link");
```

hansg  June 17, 2002
Real Attributes

Most attributes are of type REAL and are treated internally as double precision values. They may be set by integer values, real values, or expressions. Example:

```
ddd:drift,l=1.2345;
dddd:drift,l=ddd->1-0.3;
```

hansg  May 8, 2001
Selection Statements

The elements, or a range of elements, in a sequence can be selected for various purposes. Such selections remain valid until cleared (in difference to MAD-8); it is therefore recommended to always start with a

\[ \text{select,flag}=\ldots,\text{clear}; \]

before setting a new selection.

\[ \text{SELECT,FLAG}=\text{name},\text{RANGE}=\text{range},\text{CLASS}=\text{class},\text{PATTERN}=\text{pattern}[,\text{FULL}][,\text{CLEAR}]; \]

where the name for FLAG can be one of ERROR, MAKETHIN, SEQEDIT or the name of a twiss table which is established for all sequence positions in general.

Selected elements have to fulfill the \text{RANGE}, \text{CLASS} and \text{PATTERN} criteria.

Any number of SELECT commands can be issued for the same flag and are accumulated (logically ORed). In this context note the following:

\[ \text{SELECT,FLAG}=\text{name},\text{FULL}; \]

selects all positions in the sequence for this flag. This is the default for all tables and makethin, whereas for ERROR and SEQEDIT the default is "nothing selected".

SAVE: A SELECT,FLAG=SAVE statement causes the selected sequences, elements, and variables to be written into the save file. A class (only used for element selection), and a pattern can be specified. Example:

\[ \text{select,flag}=\text{save, class=variable, pattern}="\text{abc}.*"; \]
\[ \text{save, file}=?\text{mysave}; \]

will save all variables (and sequences) containing "abc" in their name, but not elements with names containing "abc" since the class "variable" does not exist (astucieux, non ?).

SECTORMAP: A SELECT,FLAG=SECTORMAP statement causes sectormaps to be written into the file "sectormap" like in MAD-8. For the file to be written, a flag SECTORMAP must be issued on the TWISS command in addition.

TWISS: A SELECT,FLAG=TWISS statement causes the selected rows and columns to be written into the Twiss TFS file (former OPTICS command in MAD-8). The column selection is done on the same select. See as well example 2. Example 1:
TITLE,'Test input for MAD-X';

option,rbarc=false; // use arc length of rbends
beam; ! sets the default beam for the following sequence
option,-echo;
call file=fv9.opt;  ! contains optics parameters
call file="fv9.seq"; ! contains a small sequence "fivecell"
OPTION,ECHO;
SELECT,FLAG=SECTORMAP,clear;
SELECT,FLAG=SECTORMAP,PATTERN="^m.*";
SELECT,FLAG=TWISS,clear;
SELECT,FLAG=TWISS,PATTERN="^m.*",column=name,s,betx,bety;
USE,PERIOD=FIVECELL;
twiss,file=optics,sectormap;
stop;

This produces a file [sectormap] and a twiss output file (name = optics):

<table>
<thead>
<tr>
<th>TYPE</th>
<th>%05s &quot;TWISS&quot;</th>
</tr>
</thead>
<tbody>
<tr>
<td>PARTICLE</td>
<td>%08s &quot;POSITRON&quot;</td>
</tr>
<tr>
<td>MASS</td>
<td>%le 0.000510998902</td>
</tr>
<tr>
<td>CHARGE</td>
<td>%le 1</td>
</tr>
<tr>
<td>E0</td>
<td>%le 1</td>
</tr>
<tr>
<td>PC</td>
<td>%le 0.99999986944</td>
</tr>
<tr>
<td>GAMMA</td>
<td>%le 1956.95136738</td>
</tr>
<tr>
<td>KBUNCH</td>
<td>%le 1</td>
</tr>
<tr>
<td>NPART</td>
<td>%le 0</td>
</tr>
<tr>
<td>EX</td>
<td>%le 1</td>
</tr>
<tr>
<td>EY</td>
<td>%le 1</td>
</tr>
<tr>
<td>ET</td>
<td>%le 0</td>
</tr>
<tr>
<td>LENGTH</td>
<td>%le 534.6</td>
</tr>
<tr>
<td>ALFA</td>
<td>%le 0.00044339992938</td>
</tr>
<tr>
<td>ORBIT5</td>
<td>%le -0</td>
</tr>
<tr>
<td>GAMMAIR</td>
<td>%le 47.4900022541</td>
</tr>
<tr>
<td>Q1</td>
<td>%le 1.25413071556</td>
</tr>
<tr>
<td>Q2</td>
<td>%le 1.25485338377</td>
</tr>
<tr>
<td>Q1</td>
<td>%le 1.05329608302</td>
</tr>
<tr>
<td>Q2</td>
<td>%le 1.04837000224</td>
</tr>
<tr>
<td>DXMAX</td>
<td>%le 2.17763211131</td>
</tr>
<tr>
<td>DYMAY</td>
<td>%le 0</td>
</tr>
<tr>
<td>XCOMAX</td>
<td>%le 0</td>
</tr>
<tr>
<td>YCOMAX</td>
<td>%le 0</td>
</tr>
<tr>
<td>BETXMAX</td>
<td>%le 177.70993499</td>
</tr>
<tr>
<td>BETYMAX</td>
<td>%le 177.671582415</td>
</tr>
<tr>
<td>XCORMS</td>
<td>%le 0</td>
</tr>
<tr>
<td>YCORMS</td>
<td>%le 0</td>
</tr>
<tr>
<td>DXRMS</td>
<td>%le 1.66004270906</td>
</tr>
<tr>
<td>DYRMS</td>
<td>%le 0</td>
</tr>
<tr>
<td>DELTAP</td>
<td>%le 0</td>
</tr>
<tr>
<td>TITLE</td>
<td>%20s &quot;Test input for MAD-X&quot;</td>
</tr>
<tr>
<td>ORIGIN</td>
<td>%16s &quot;MAD-X 0.20 Linux&quot;</td>
</tr>
<tr>
<td>DATE</td>
<td>%08s &quot;07/06/02&quot;</td>
</tr>
<tr>
<td>TIME</td>
<td>%08s &quot;14.25.51&quot;</td>
</tr>
</tbody>
</table>

* NAME S BETX BETY
| %s | %le | %le |
| "MSCBH" | 4.365 | 171.6688159 | 33.31817319 |
| "MB" | 19.72 | 108.1309095 | 58.58680717 |
Example 2:

Addition of variables to (any internal) table:

```
select,flag=table,column=name,s,betx,...,var1,var2,...; or
select,flag=table,full,column=var1,var2,...; ! default col.s + new
```

will write the current value of var1 etc. into the table each time a new line is added; values from the same (current) line can be accessed by these variables, e.g.

```
var1:= sqrt(beam->ex*table(twiss,betx));
```

in the case of table above being "twiss". The plot command accepts the new variables.
Remark: this replaces the "string" variables of MAD-8.

This example demonstrates as well the usage of a user defined table.

beam, ex=1.e-6, ey=1.e-3;
// element definitions
mb: rbend, l=14.2, angle=0, k0=bang/14.2;
mq: quadrupole, l=3.1, apertype=ellipse, aperture=(1,2);
qft: mq, l=0.31, kl=kqf, tilt=pi/4;
qf.1: mq, l=3.1, kl=kqf;
qf.2: mq, l=3.1, kl=kqf;
qf.3: mq, l=3.1, kl=kqf;
qf.4: mq, l=3.1, kl=kqf;
qf.5: mq, l=3.1, kl=kqf;
qd.1: mq, l=3.1, kl=kqd;
qd.2: mq, l=3.1, kl=kqd;
qd.3: mq, l=3.1, kl=kqd;
qd.4: mq, l=3.1, kl=kqd;
qd.5: mq, l=3.1, kl=kqd;
bph: hmonitor, l=1.bpm;
bpv: vmonitor, l=1.bpm;
cbh:hkicker;
cbv:vkicker;
cbh.1: cbh, kick=achbh1;
cbh.2: cbh, kick=achbh2;
cbh.3: cbh, kick=achbh3;
cbh.4: cbh, kick=achbh4;
cbh.5: cbh, kick=achbh5;
cbv.1: cbv, kick=acbvl;
cbv.2: cbv, kick=acbv2;
cbv.3: cbv, kick=acbv3;
cbv.4: cbv, kick=acbv4;
cbv.5: cbv, kick=acbv5;
!mscbh: sextupole, l=1.1, k2=ksf;
mscbh: multipole, knl={0,0,0,ksf}, tilt=-pi/8;
mscbv: sextupole, l=1.1, k2=ksd;
!mscbv: octupole, l=1.1, k3=ksd, tilt=-pi/8;
// sequence declaration
fivecell: sequence, refer=centre, l=534.6;
   qf.1: qf.1, at=1.550000e+00;
   qft: qft, at=3.815000e+00;
   cbh.1: cbh.1, at=4.365000e+00;
   mb: mb, at=1.262000e+01;
   mb: mb, at=2.828000e+01;
   mb: mb, at=4.394000e+01;
   bpv: bpv, at=5.246000e+01;
   qd.1: qd.1, at=5.501000e+01;
   mscbv: mscbv, at=5.727500e+01;
   cbv.1: cbv.1, at=5.782500e+01;
   mb: mb, at=6.608000e+01;
   mb: mb, at=8.174000e+01;
   mb: mb, at=9.740000e+01;
   bph: bph, at=1.059200e+02;
   qf.2: qf.2, at=1.084700e+02;
mscbh:mscbh, at=1.107350e+02;
cbh.2:cbh.2, at=1.112850e+02;
mb:mb, at=1.195400e+02;
mb:mb, at=1.352000e+02;
mb:mb, at=1.508600e+02;
bpv:bpv, at=1.593800e+02;
qd.2:qd.2, at=1.619300e+02;
mscbv:mscbv, at=1.641950e+02;
cbv.2:cbv.2, at=1.647450e+02;
mb:mb, at=1.730000e+02;
mb:mb, at=1.886600e+02;
mb:mb, at=2.043200e+02;
bph:bph, at=2.128400e+02;
qf.3:qf.3, at=2.153900e+02;
mscbh:mscbh, at=2.176550e+02;
cbh.3:cbh.3, at=2.182050e+02;
mb:mb, at=2.264600e+02;
mb:mb, at=2.421200e+02;
mb:mb, at=2.577800e+02;
bpv:bpv, at=2.663000e+02;
qd.3:qd.3, at=2.688500e+02;
mscbv:mscbv, at=2.711150e+02;
cbv.3:cbv.3, at=2.716550e+02;
mb:mb, at=2.799200e+02;
mb:mb, at=2.955800e+02;
mb:mb, at=3.112400e+02;
bph:bph, at=3.197600e+02;
qf.4:qf.4, at=3.223100e+02;
mscbh:mscbh, at=3.245750e+02;
cbh.4:cbh.4, at=3.251250e+02;
mb:mb, at=3.338000e+02;
mb:mb, at=3.490400e+02;
mb:mb, at=3.647000e+02;
bpv:bpv, at=3.732200e+02;
qd.4:qd.4, at=3.757700e+02;
mscbv:mscbv, at=3.780350e+02;
cbv.4:cbv.4, at=3.785850e+02;
mb:mb, at=3.868400e+02;
mb:mb, at=4.025000e+02;
mb:mb, at=4.181600e+02;
bph:bph, at=4.266800e+02;
qf.5:qf.5, at=4.292300e+02;
mscbh:mscbh, at=4.314950e+02;
cbh.5:cbh.5, at=4.320450e+02;
mb:mb, at=4.403000e+02;
mb:mb, at=4.559600e+02;
mb:mb, at=4.716200e+02;
bpv:bpv, at=4.801400e+02;
qd.5:qd.5, at=4.826900e+02;
mscbv:mscbv, at=4.849550e+02;
cbv.5:cbv.5, at=4.855050e+02;
mb:mb, at=4.937600e+02;
mb:mb, at=5.094200e+02;
mb:mb, at=5.250800e+02;
bph:bph, at=5.336000e+02;
end:marker, at=5.346000e+02;
endsequence;
// forces and other constants

1.bpm:=-.3;
bang:=.509998807401e-2;
kqf:=.872651312e-2;
kqd:=-.872777242e-2;
ksf:=.0198492943;
ksd:=-.039621283;
acbv1:=1.e-4;
acbh1:=1.e-4;
!save,sequence=fivecell,file,mad8;

s := table(twiss,bpv[5],betx);
myvar := sqrt(beam->ex*table(twiss,betx));
use, period=fivecell;
select,flag=twiss,column=name,s,myvar,apertype;
twiss,file;

create,table=mytab,column=dp,mq1,mq2;
mq1:=table(summ,q1);
mq2:=table(summ,q2);
while ( n < 11)
{
    n = n + 1;
dp = 1.e-4*(n-6);
twiss,deltap=dp;
    fill,table=mytab;
}
write,table=mytab;
plot,haxis=s,vaxis=aper_1,aper_2,colour=100,range=#s/cbv.1,notitle;

prints the following user table on output:

| NAME | %05s "MYTAB" |
| TYPE | %04s "USER" |
| TITLE | %08s "no-title" |
| ORIGIN | %16s "MAD-X 1.09 Linux" |
| DATE | %08s "10/12/02" |
| TIME | %08s "10.45.25" |

<table>
<thead>
<tr>
<th>DP</th>
<th>MQ1</th>
<th>MQ2</th>
</tr>
</thead>
<tbody>
<tr>
<td>$%le</td>
<td>%le</td>
<td>%le</td>
</tr>
<tr>
<td>-0.0005</td>
<td>1.242535951</td>
<td>1.270211135</td>
</tr>
<tr>
<td>-0.0004</td>
<td>1.242495534</td>
<td>1.270197018</td>
</tr>
<tr>
<td>-0.0003</td>
<td>1.242452432</td>
<td>1.270185673</td>
</tr>
<tr>
<td>-0.0002</td>
<td>1.242406653</td>
<td>1.270177093</td>
</tr>
<tr>
<td>-0.0001</td>
<td>1.242358206</td>
<td>1.270171269</td>
</tr>
<tr>
<td>0</td>
<td>1.242307102</td>
<td>1.27016819</td>
</tr>
<tr>
<td>0.0001</td>
<td>1.242253353</td>
<td>1.270167843</td>
</tr>
<tr>
<td>0.0002</td>
<td>1.242196974</td>
<td>1.270170214</td>
</tr>
<tr>
<td>0.0003</td>
<td>1.24213798</td>
<td>1.270175288</td>
</tr>
<tr>
<td>0.0004</td>
<td>1.242076387</td>
<td>1.270183048</td>
</tr>
<tr>
<td>0.0005</td>
<td>1.242012214</td>
<td>1.270193477</td>
</tr>
</tbody>
</table>

and produces a twiss file with the additional column myvar, as well as a plot file with the aperture values plotted.
Example of joining 2 tables with different lengths into a third table making use of the length of either table as given by `table("your_table_name", tablelength)` and adding names by the "_name" attribute.

```plaintext
title, "summing of offset and alignment tables";
set, format="13.6f";

readtable, table-align, file="align.ip2.b1.tfs"; // measured alignment
readtable, table-offset, file="offset.ip2.b1.tfs"; // nominal offsets

n_elem = table(offset, tablelength);
create, table-align_offset, column=_name, s_ip, x_off, dx_off, ddx_off, y_off, dy_off, ddy_off;

calcul(elem_name) : macro = {
    x_off = table(align, elem_name, x_ali) + x_off;
    y_off = table(align, elem_name, y_ali) + y_off;
}

one_elem(j_elem) : macro = {
    setvars, table-offset, row=j_elem;
    exec, calcul(tabstring(offset, name, j_elem));
    fill, table-align_offset;
}

i_elem = 0;
while (i_elem < n_elem) { i_elem = i_elem + 1;
    exec, one_elem($i_elem); }

write, table-align_offset, file="align_offset.tfs";

stop;

hansg
May 8, 2001
```

Range and Class Selection Format

- **RANGE**: A range can be defined starting at a given element and ending at another element, both elements included. Two forms exist:
  
  range=position;  
  range=position1/position2;

  In the first case, only one element is selected; in the second case, one or several elements are selected. NOTE: position1 must not be behind position2 in the sequence.

"position" is composed of the element name and an optional occurrence count in the sequence:

mq.ir5.l6..1          ! no occurrence count given
mb[17]                ! occurrence count given

There are two predefined MAD indices:

- S. The start of the beam line expanded by USE,
- E. The end of the beam line expanded by USE.

If, in the USE statement, only a range is selected:

use,period=lhcbl,range=ir1/ir5;

then "#s" and "#e" refer to the start and end of the expanded range, of course.

Examples for ranges:

...,range=#s;           ! first element
...,range=#s/#e;        ! full expansion range
...,range=mb[5]/#e;     ! from mb 5 to end
...,range=mq.ir5.l6..1; ! first occurrence of element mq.ir5.l6..1

- **CLASS**: The single name of a class (no occurrence counts). A class is the name of an element (or basic type) from which other elements have been derived. Example:

mq:quadrupole;
q1:mq;
q2:mq;
q1..a:q1;
q2..b:q2;

makes classes from mq, q1, and q2. Selection class="mq" will actually select q1, q2, q1..a, and q2..b in the above example.

hansg  June 17, 2002
**Sectormap output**

The flag "sectormap" on the Twiss command (together with an element selection via select.flag=sectormap,...) causes a file "sectormap" to be written.

For each user-selected element, it contains the user-selected coefficients of the kick vector $K$ (6 values), of the first-order map $R$ (6 x 6 values) and of the second-order map $T$ (6 x 6 x 6 values)

The sector file is the output of a standard TFS table, which means that the set of columns of interest may be selected through a MAD-X command such as the following:

```
select,flag=my_sect_table,column=name,pos,kl,r11,rl66,t111;
```

Each line of the sectormap file contains for each selected element, the set of chosen $K$,$R$,$T$ matrix coefficients:

<table>
<thead>
<tr>
<th>NAME</th>
<th>TYPE</th>
<th>TITLE</th>
<th>ORIGIN</th>
<th>DATE</th>
<th>TIME</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;MY_SECT_TABLE&quot;</td>
<td>&quot;SECTORMAP&quot;</td>
<td>&quot;no-title&quot;</td>
<td>&quot;MAD-X 3.04.62 Linux&quot;</td>
<td>&quot;18/12/08&quot;</td>
<td>&quot;10.33.58&quot;</td>
</tr>
</tbody>
</table>
* NAME | POS | K1 | R11 | R66 | T111
---|---|---|---|---|---
$ s | %le | %le | %le | %le | %le
"FIVECELL$START" | 0 | 0 | 1 | 1 | 0
"SEQSTART" | 0 | 0 | 1 | 1 | 0
"QF.1" | 3.1 | -1.305314637e-05 | 1.042224745 | 1 | 0
"DRIFT_0" | 3.265 | 7.451656548e-21 | 1 | 1 | 0
"MSCBH" | 4.365 | -1.686090613e-15 | 0.9999972755 | 1 | 0.006004411526
"CBH.1" | 4.365 | 0 | 1 | 1 | 0
"DRIFT_1" | 5.519992305 | -6.675347543e-21 | 1 | 1 | 0
"MB" | 19.72000769 | 2.566889547e-18 | 1.000000091 | 1 | -4.135903063e-25
"DRIFT_2" | 21.17999231 | -1.757758802e-20 | 1 | 1 | 0
"MB" | 35.38000769 | 2.822705549e-18 | 1.000000091 | 1 | -4.135903063e-25
"DRIFT_2" | 36.83999231 | 2.480880093e-20 | 1 | 1 | 0
"MB" | 51.04000769 | 3.006954115e-18 | 1.000000091 | 1 | -4.135903063e-25
"DRIFT_3" | 52.21 | -4.886652187e-20 | 1 | 1 | 0
...
...
...
...
...
...
...
...

Of course, the select statement can be combined with additional options to filter-out the list of elements, such as in the following statement, which for instance only retains drift-type elements:

```
select,flag=my_sect_table,class=drift,column=name,pos,k1,r11,r66,t111;
```

K coefficients range: K1... K6

R coefficients range:

```
R11  ...  R61
R12  ...  R62
...  ...  ...
R61  ...  R66
```
T coefficients range:

\[
\begin{align*}
T_{111} & \quad \ldots \quad T_{611} \\
T_{121} & \quad \ldots \quad T_{621} \\
& \quad \ldots \\
T_{161} & \quad \ldots \quad T_{661} \\
T_{112} & \quad \ldots \quad T_{612} \\
& \quad \ldots \\
T_{166} & \quad \ldots \quad T_{666}
\end{align*}
\]

In the above notation, $R_{ij}$ stands for "effect on the $i$-th coordinate of the $j$-th coordinate in phase-space", whereas $T_{ijk}$ stands for "combined effect on the $i$-th coordinate of both the $j$-th and $k$-th coordinates in phase-space" along the lattice.

The maps are the accumulated maps between the selected elements. They contain both the alignment, and field errors present. Together with the starting value of the closed orbit (which can be obtained from the standard twiss file) this allows the user to track particles over larger sectors, rather than element per element. A typical usage therefore lies in the interface to other programs, such as TRAIN.

\textit{hansg} May 8, 2001
Variable Declarations

In the following, "=" means that the variable at the left receives the current value of the expression at right, but does not depend on it any further, whereas ":=" makes it depend on this expression, i.e. every time the expression changes, the variable is re-evaluated, except for "const" variables.

```
var = expression;
var := expression;
real var = expression;    // identical
real var := expression;    // to above
int var = expression;     // truncated if expression is real
int var := expression;
const var = expression;
const var := expression;
const real var = expression;    // identical
const real var := expression;    // to above
const int var = expression;    // truncated if expression is real
const int var := expression;
```

May 8, 2001
Identifiers or Labels

A label begins with a letter, followed by up to fifteen letters, digits, decimal points (.), or underscores (_). Characters beyond the sixteenth are dropped, but should be avoided, and the resulting sequence must be unique.

A label may refer to a keyword, an element, a beam-line, a sequence, etc. The MAD-X keywords are protected; using one of them as a label results in a fatal error.

hansg May 8, 2001
Command Attributes

The following types of attributes are available in MAD:

- A **name or string attribute** refers to an object, or a string.
- A **logical attribute** selects or deselects an option.
- An **integer attribute** is a counter, as for repetition in a beam line.
- A **real expression** defines a datum for a command, it may be varied in matching. An expression is built of a combination of **operator** and **operand**
- A **constraint** specifies a matching constraint.
- A **variable name** selects a variable to be matched.

*hansg* May 8, 2001
Name or String Attributes

A name or string attribute often selects one of a set of options:

```
use,period=lhc;  // expand the LHC sequence
```

It may also refer to a user-defined object:

```
twiss,file=optics;  // specifies the name of the OPTICS output file
```

It may also define a string:

```
title,"LHC version 6.2";
```

The case of letters is only significant if a string is enclosed in quotes, otherwise all characters are converted to lower at reading. On the other hand, strings that do not contain blanks do not need to be enclosed in quotes. Example:

```
call,file="my.file";
call,file=my.file;
call,file=MY.FILE;
call,file="MY.FILE";
call,file='MY.FILE';
```

In the first three cases, MAD-X will try to read a file my.file, in the last two it will try to read MY.FILE.

hansg May 8, 2001
Logical Attributes

Many commands in MAD require the setting of logical values (flags) to represent the on/off state of an option. A logical value "flag" can be set in two ways:

```plaintext
flag | flag = true
```

It can be reset by:

```plaintext
-flag | flag=false
```

Example:

```plaintext
option,-echo; // switch off copying the input to the standard output
```

The default for a logical flag is normally false, but can be found e.g. for options by the command

```plaintext
help,option;
```

hansg May 8, 2001
Integer Attributes

An integer attribute usually denotes a count. Example:

myline:line=(-3*(a,b,c));

In this case, a literal integer is requested; however, in the following

rfc:rfcavity,harmon=12345;

or

rfc:rfcavity,harmon=num;

"num" may be an integer variable, a real variable, or an expression (in the two latter cases, the value is truncated).

hansg  May 8, 2001
Real Expressions

To facilitate the definition of interdependent quantities, any real value and integer value can be entered as an arithmetic expression. When a value used in an expression is redefined by the user or changed in a matching process, the expression is reevaluated. Expression definitions may be entered in any order. MAD evaluates them in the correct order before it performs any computation. At evaluation time all operands used must have values assigned.

An expression is built from a combination of operator and operand and it may contain random generators.

Operators in Arithmetic Expressions

An expression can be formed using the following operators:

Arithmetic operators

- + Addition,
- - Subtraction,
- * Multiplication,
- / Division,
- ^ Exponentiation.

Ordinary functions

- sqrt(x) square root,
- log(x) natural logarithm,
- log10(x) logarithm base 10,
- exp(x) exponential,
- sin(x) trigonometric sine,
- cos(x) trigonometric cosine,
- tan(x) trigonometric tangent,
- asin(x) arc sine,
- acos(x) arc cosine,
- atan(x) arc tangent,
- sinh(x) hyperbolic sine,
- cosh(x) hyperbolic cosine,
- tanh(x) hyperbolic tangent,
- abs(x) absolute value;
random number generators

- ranf() random number, uniformly distributed in [0,1],
- gauss() random number, gaussian distribution with unit standard deviation,
- tgauss(x) random number, gaussian distribution with unit standard deviation, truncated at x standard deviations;

in the above cases, "x" can be any expression, i.e. contain other functions, variable or constant expressions. To initialize the MAD-X random generator use the \texttt{Eoption} command.

table access functions

- table(x,z): accesses value of the named table column "z" of table "x"; example: table(summ,q1) returns the hor. tune of the Twiss summary table "summ".
- table(x,y,z): accesses value of the named table column "z" for element "y" (first table row with that name) of table "x"; example: table(twiss,mb.12,betx) returns the beta_x at element mb.12 from the Twiss table "twiss". When the element is called with its proper name, as in the example above, the value is returned at the first occurrence of the element of this name. If the value is needed at a occurrence number: NNN, then "[NNN]" has to be appended to the name, e.g. in the above example one obtains the betx of the 23th occurrence of the element "mb.12" by changing the example to: "table(twiss,mb.12[23],betx)". Mind you that the old, but little known, form: "table(twiss,mb.12->23,betx)" continues to work. Lastly, if NNN exceeds the maximum occurrence number the return value is arbitrarily small.

Beware:

- Unnamed Drifts are not included in the table name database, so as to speed up the search for "real" elements. Therefore, those unnamed drifts cannot be found. However, named drifts can be searched for.
- Due to the importance of finding elements in the table for a proper functioning of the MAD-X runs, the programs throws a "fatal_error" if an element cannot be found in the table.

There is a second option of this function with 3 entries

- table(x,z,N_row): accesses the value of the named table column "z" at the "N_row" number of rows of table "x" (row numbers start at 1); example: table(twiss,betx,370) returns the beta_x at row number "370" of the Twiss table "twiss". The return value is zero if "N_row" is outside of the allowed range.

Note that "N_row" has to be a number and not a variable. However, the \texttt{Macro facility} in MAD-X allows one to use a variable instead.

A typical example could look like this, in fact the square root of betx (user defined variable myvar) is added to the twiss table:

\begin{verbatim}
myvar := sqrt(table(twiss,betx));
select,flag=twiss,column=name,s,myvar,betx;
twiss,file;
\end{verbatim}
Or another arbitrary test case of adding k1l taken from the Twiss table:

Define macro:

\[
\text{mycrap}(xx,yy,zz): \text{macro} = \{\text{myval} = \text{table}(xx,yy,zz);\};
\]

Use macro in loop:

\[
i = 0; \\
\text{incval} = 0; \\
\text{while } (i < 100) \{ \\
\text{value},i; \\
\text{exec,mycrap(twiss,k1l,$i);} \\
\text{incval} = \text{incval} + \text{myval}; \\
\text{value},i,\text{myval},\text{incval}; \\
i = i + 1; \\
\}
\]

**Features as of Version 3_03_50**

- **FILL, TABLE=t, ROW=n;**
  
  fill a table row with the present variable values. If ROW is negative or missing a new row is created. If ROW is greater than the number of rows, the last row is selected without creating a new row.

- **SETVARS, TABLE=t, ROW=n;**
  
  set variables according to the column names of the given table and the values of the given row. if ROW is negative, missing or greater than the number of rows, the last row is selected.

- **An example can be found at:** [Special Features](#)

- **The length of a table can be determined by using the attribute "tablelength" via table("your_table_name", tablelength). This is useful when creating a table from existing ones. See an example at:** [user table II](#)

**Operands in Arithmetic Expressions**

An expression may contain the following operands:

**Literal constants**

Numerical values are entered like FORTRAN constants. Real values are accepted in INTEGER or REAL format. The use of a decimal exponent, marked by the letter D or E, is permitted.

Examples:
Symbolic constants

MAD recognizes some mathematical and physical constants. Their names must not be used for user-defined labels.

Additional symbolic constants may be defined to simplify their repeated use in statements and expressions.

\[
\text{CONST name=constant-expression;}
\]

defines a real constant with the name given. An existing symbolic constant can be redefined, but it cannot change in a matching procedure.

Example:

\[
\text{const in = 0.0254;}
\]

<table>
<thead>
<tr>
<th>mad name</th>
<th>symbol</th>
<th>value used</th>
<th>unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>pi</td>
<td>pi</td>
<td>4 * atan(1)</td>
<td>1</td>
</tr>
<tr>
<td>two pi</td>
<td>2 pi</td>
<td>2 * pi</td>
<td>1</td>
</tr>
<tr>
<td>degrad</td>
<td>180/pi</td>
<td>180 / pi</td>
<td>deg/rad</td>
</tr>
<tr>
<td>raddeg</td>
<td>pi/180</td>
<td>180 / pi</td>
<td>rad/deg</td>
</tr>
<tr>
<td>e</td>
<td>e</td>
<td>exp(1)</td>
<td>1</td>
</tr>
<tr>
<td>emass</td>
<td>m_e</td>
<td>.510998902*10(-3)</td>
<td>GeV</td>
</tr>
<tr>
<td>pmass</td>
<td>m_p</td>
<td>.938271998</td>
<td>GeV</td>
</tr>
<tr>
<td>mumass</td>
<td>m_mu</td>
<td>.1056583568</td>
<td>GeV</td>
</tr>
<tr>
<td>clight</td>
<td>c</td>
<td>2.99792458*10**8</td>
<td>m/s</td>
</tr>
<tr>
<td>qelect</td>
<td>elem. charge</td>
<td>1.602176462e-19</td>
<td>As</td>
</tr>
</tbody>
</table>

Parameter labels

Often a set of numerical values depends on a common variable parameter. Such a parameter must be defined as a global parameter. A global parameter always has a current value; however, this value may be re-evaluated or not, depending on the parameter definition:

\[
x = a;
\]

x is set to the current value of a and not changed, even if a changes. This makes assignments such as
x = x + 1;

perfectly valid (this replaces the old SET instruction). The definition of the deferred expression

x := a;

assign the current value of a to x every time x is used, i.e. it is re-evaluated using the latest value of a;
therefore,

x := x + 1;

results in an infinite loop (!) when x is used (error abort). Of course, the following definitions are
equivalent:

x = 0.1;
x := 0.1;

When such a parameter is used in an expression, MAD uses the current value of the parameter if the
expression is deferred:

Example:

x:=1.0;
d1: drift,l = x;
d2: drift,l := 2.0 - x;

When the value of X is changed, the length of the drift d1 remains unchanged, that of d2 is recalculated.

**Element or command attributes**

In arithmetic expressions the attributes of physical elements or commands can occur as operands. They are
named respectively by

- `element-name->attribute-name`
- `command-name->attribute-name`

Values are assigned to attributes in element definitions or commands.

Example:

D1: DRIFT,L=1.0;
D2: DRIFT,L=2.0-D1->L;

D1->L refers to the length L of the drift space D1.

**Expressions and Random Values**

The definition of random machine imperfections requires evaluation of expressions containing random
functions. These are evaluated like any other expression when the expression is used, i.e. only once if a
"=" assignment refers to it, or every time if the assignment is ":="; this latter case is used by the error
generation routines.
Example:

\[ a := 3 \cdot \text{ranf}(); \]

Every time a is used, it gets a random value assigned from a uniform distribution between 0 and 3.

\[ \text{error: } \text{ealign}, \text{range, dx:= sigma*gauss>();} \]

All elements in range are assigned independent random displacements sampled from a Gaussian distribution with standard deviation sigma.

\[ \text{hansg} \ May 8, 2001 \]
EOPTION: Set Error Options

The random generator for MAD is taken from [Knuth]. The error option command specifies different seeds for random values:

EOPTION, SEED=real, ADD=logical;

- **SEED**: Selects a particular sequence of random values. A SEED value is an integer in the range [0...999999999] (default: 123456789). SEED alone continues with the current sequence. See also: Random values. SEED may be an expression.
- **ADD**: If this logical flag is set, an EALIGN or EFCOMP, causes the errors to be added on top of existing ones. If it is not set, new errors overwrite any previous definitions. The default value is TRUE if it is omitted in the EOPTION command. The default value is false if no EOPTION command is used.

Please note a recent modification: the default value for the ADD option is only applied as long as the ADD option has not been set explicitly. Once it was set with EOPTION, it is NOT reset to the default when the ADD option is omitted in subsequent calls to EOPTION.

Example:

EOPTION, SEED=987456321;

Werner Herr 18.6.2002
Program Flow Statements

- **IF**

  if (logical_expression) {statement 1; statement 2; ...; statement n; }

  where "logical_expression" is one of

  expr1 oper expr2
  expr11 oper1 expr12 && expr21 oper2 expr22
  expr11 oper1 expr12 || expr21 oper2 expr22

  and oper one of

  ==     ! equal
  <>     ! not equal
  <      ! less than
  >      ! greater than
  <=     ! less than or equal
  >=     ! greater than or equal

  The expressions are arithmetic expressions of type real. The statements in the curly brackets are executed if the logical expression is true.

- **ELSEIF**

  elseif (logical_expression) {statement 1; statement 2; ...; statement n; }

  Only possible (in any number) behind an IF, or another ELSEIF; is executed if logical_expression is true, and if none of the preceding IF or ELSEIF logical conditions was true.

- **ELSE**

  else {statement 1; statement 2; ...; statement n; }

  Only possible (once) behind an IF, or an ELSEIF; is executed if logical_expression is true, and if none of the preceding IF or ELSEIF logical conditions was true.
For a real life example, see **ELSE example**

**WHILE**

\[
\text{while (logical\_condition)}\{\text{statement 1}; \text{statement 2}; \ldots; \text{statement n}; \}
\]

executes the statements in curly brackets while the logical_expression is true. A simple example (in case you have forgotten the first ten factorials) would be

\[
\begin{align*}
\text{option, -info; } & \text{ ! otherwise you get redefinition warnings} \\
n &= 1; m = 1; & \\
\text{while (}n \leq 10\text{)} & \\
& \{ \\
& \quad m = m \times n; \text{ value, } m; \\
& \quad n = n + 1; \\
& \}
\end{align*}
\]

For a real life example, see **WHILE example**

**MACRO**

\[
\begin{align*}
\text{label: macro} & = \{ \text{statement 1}; \text{statement 2}; \ldots; \text{statement n}; \}; \\
\text{label(arg1,...,argn): macro} & = \{ \text{statement 1}; \text{statement 2}; \ldots; \text{statement n}; \};
\end{align*}
\]

The first form allows the execution of a group of statements via a single command:

\[
\text{exec, label;}
\]

will execute the statements in curly brackets exactly once. This command can be issued any number of times.

The second form allows to replace strings anywhere inside the statements in curly brackets by other strings, or integer numbers prior to execution. This is a powerful construct and should be handled with care.

Simple example:

\[
\begin{align*}
\text{option, -echo, -info; } & \text{ ! otherwise the output is somewhat confusing} \\
\text{simple(xx, yy): macro} & = \{ \ xx = yy^2 + xx; \text{ value, } xx; \}; \\
a &= 3; & \\
b &= 5; & \\
\text{exec, simple(a, b);}
\end{align*}
\]

Somewhat more tricky (a "$" in front of an argument means that the truncated integer value of this argument is used for replacement, rather than the argument string itself).
tricky(xx,yy,zz): macro = {mzz.yy: xx, l = 1.yy, kzz = k.yy};
n=0;
while (n < 3)
{
    n = n+1;
    exec,tricky(quadrupole,$n,1);
    exec,tricky(sextupole,$n,2);
}

Whereas the actual use of the preceding example is NOT recommended, a real life example, showing the full power (!) of macros is to be found under macro usage for the usage, and under macro definition for the definition.

Beware of the following rules:

- Generally speaking: special constructs like IF, WHILE, MACRO will only allow one level of inclusion of another special construct.
- Macros must not be called with numbers, but with strings (i.e. variable names in case of numerical values). i.e.

**NOT**

exec,thismacro($99,$129);

**BUT**

n1=99; n2=219;
exec,thismacro($n1,$n2);

* hansg  June 17, 2002*
Real life example for IF statements, and MACRO usage

! Creates a footprint for head-on + parasitic collisions at IP1+5
! of lhc.6.5; both lhcb1 (for tracking) and lhcb2 (to define the
! beam-beam elements, i.e. weak-strong) are used; there are flags to
! select head-on, left, and right parasitic separately at all IPs.
! The bunch spacing can be given in nanosec and automatically creates
! the beam-beam interaction points at the correct positions.
! It is important to set the correct BEAM parameters, i.e. number
! of particles, emittances, bunch length, energy.

!--- For completeness, all files needed by this job are copied
! to the local directory ldb. The links to the the originals
! in offdb (official database) are commented out.

Option, warn,info,echo;
!System,
"ln -fn /afs/cern.ch/eng/sl/MAD-X/dev/test_suite/foot/V3.01.01 ldb";
!system,"ln -fn /afs/cern.ch/eng/lhc/optics/V6.4 offdb";
Option, -echo,-info,warn;
SU=1.0;
call, file = "ldb/V6.5.seq";
call,file="ldb/slice_new.madx";
Option, echo,info,warn;

!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!! Step 1 +!!!!!!!!!!!!!!!!!!!!
! define beam constants
!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!

b_t_dist  = 25.e-9;                  !--- bunch distance in [sec]
b_h_dist  = clight * b_t_dist / 2 ;  !--- bunch half-distance in [m]
ip1_range = 58.;                     ! range for parasitic collisions
ip5_range = ip1_range;
ip2_range = 60.;
ip8_range = ip2_range;

npara_1 = ip1_range / b_h_dist;     ! # parasitic either side
npara_2 = ip2_range / b_h_dist;
npara_5 = ip5_range / b_h_dist;
npara_8 = ip8_range / b_h_dist;

value,npara_1,npara_2,npara_5,npara_8;

eg   =  7000;
bg   =  eg/pmass;
en   = 3.75e-06;
epsx = en/bg;
epsy = en/bg;
Beam, particle = proton, sequence=lhcb1, energy = eg,
  sigt= 0.077,
  bv = +1, NPART=1.1E11, sige = 1.1e-4,
  ex=epsx, ey=epsy;

Beam, particle = proton, sequence=lhcb2, energy = eg,
  sigt= 0.077,
  bv = -1, NPART=1.1E11, sige = 1.1e-4,
  ex=epsx, ey=epsy;

beamx = beam%lhcb1->ex;  beamy%lhcb1 = beam->ey;
sigz = beam%lhcb1->sigt; sige = beam%lhcb1->sige;

--- split5, 4d
long_a= 0.53 * sigz/2;
long_b= 1.40 * sigz/2;
value,long_a,long_b;

ho_charge = 0.2;

!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!! Step 2 !!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!
! slice, flatten sequence, and cycle start to ip3
!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!

use,sequence=lhcb1;
makethin,sequence=lhcb1;
!save,sequence=lhcb1,file=lhcb1_thin_new_seq;
use,sequence=lhcb2;
makethin,sequence=lhcb2;
!save,sequence=lhcb2,file=lhcb2_thin_new_seq;
!stop;

option,-warn,-echo,-info;
call,file="ldb/V6.5.thin.coll.str";
option,warn,echo,info;

! keep sextupoles
ksf0=ksf; ksd0=ksd;
use,period=lhcb1;
select,flag=twiss.1,column=name,x,y,betx,bety;
twiss,file;
plot,haxis=s,vaxis=x,y,colour=100,noline;

use,period=lhcb2;
select,flag=twiss.2,column=name,x,y,betx,bety;
twiss,file;
plot,haxis=s,vaxis=x,y,colour=100,noline;
seqedit,sequence=lhcb1;
flatten;
endedit;
seqedit,sequence=lhcb1;
cycle,start=ip3.b1;
endedit;
seqedit,sequence=lhcb2;
flatten;
endedit;

seqedit,sequence=lhcb2;
cycle,start=ip3.b2;
endedit;

bbmarker: marker; /* for subsequent remove */

!+++++++++++++++++++++++++ Step 3 ++++++++++++++++++++++
!       define the beam-beam elements
!+++++++++++++++++++++++++++++++++++++++++++++++++++++
!===============================================================================
! read macro definitions
option,-echo;
call,file="ldb/bb.macros";
option,echo;

!
!===============================================================================
! this sets CHARGE in the head-on beam-beam elements.
! set +1 * ho_charge for parasitic on, 0 for off

  on_ho1  = +1 * ho_charge; ! ho_charge depends on split
  on_ho2  = +0 * ho_charge; ! because of the "by hand" splitting
  on_ho5  = +1 * ho_charge;
  on_ho8  = +0 * ho_charge;

!
!===============================================================================
! set CHARGE in the parasitic beam-beam elements.
! set +1 for parasitic on, 0 for off

  on_lr1l = +1;
  on_lr1r = +1;
  on_lr2l = +0;
  on_lr2r = +0;
  on_lr5l = +1;
  on_lr5r = +1;
  on_lr8l = +0;
  on_lr8r = +0;

!
!===============================================================================
! define markers and savebetas
assign,echo=temp.bb.install;
!--- ip1
if (on_ho1 <> 0)
{
  exec, mkho(1);
  exec, sbhomk(1);
}
if (on_lr1l <> 0 || on_lr1r <> 0)
{
  n=1; ! counter
  while (n < npara_1)
  {

}
exec, mkl(1,$n);
exec, sbl(1,$n);
n=n+1;
}

if (on_lr1r <> 0 || on_lr1l <> 0)
{
    n=1; ! counter
    while (n < npara_1)
    {
        exec, mkr(1,$n);
        exec, sbr(1,$n);
        n=n+1;
    }
}

!--- ip5
if (on_ho5 <> 0)
{
    exec, mkho(5);
    exec, sbhomk(5);
}

if (on_lr5l <> 0 || on_lr5r <> 0)
{
    n=1; ! counter
    while (n < npara_5)
    {
        exec, mkl(5,$n);
        exec, sbl(5,$n);
        n=n+1;
    }
}

if (on_lr5r <> 0 || on_lr5l <> 0)
{
    n=1; ! counter
    while (n < npara_5)
    {
        exec, mkr(5,$n);
        exec, sbr(5,$n);
        n=n+1;
    }
}

!--- ip2
if (on_ho2 <> 0)
{
    exec, mkho(2);
    exec, sbhomk(2);
}

if (on_lr2l <> 0 || on_lr2r <> 0)
{
    n=1; ! counter
    while (n < npara_2)
    {
        exec, mkl(2,$n);
        exec, sbl(2,$n);
        n=n+1;
    }
}
if (on_lr2r <> 0 || on_lr2l <> 0)
{
    n=1; ! counter
    while (n < npara_2)
    {
        exec, mkr(2,$n);
        exec, sbr(2,$n);
        n=n+1;
    };
}
!--- ip8
if (on_ho8 <> 0)
{
    exec, mkho(8);
    exec, sbhmk(8);
}
if (on_lr8l <> 0 || on_lr8r <> 0)
{
    n=1; ! counter
    while (n < npara_8)
    {
        exec, mkl(8,$n);
        exec, sbl(8,$n);
        n=n+1;
    };
}
if (on_lr8r <> 0 || on_lr8l <> 0)
{
    n=1; ! counter
    while (n < npara_8)
    {
        exec, mkr(8,$n);
        exec, sbr(8,$n);
        n=n+1;
    };
}
assign, echo=terminal;
call, file=temp.bb.install;
system, "rm temp.bb.install";
!
!===========================================================
!   install bb markers
assign, echo=temp.bb.install;
!--- ip1
if (on_ho1 <> 0)
{
    exec, inho(mk,1);
}
if (on_lr1l <> 0 || on_lr1r <> 0)
{
    n=1; ! counter
    while (n < npara_1)
    {
        exec, inl(mk,1,$n);
        n=n+1;
    };
}
if (on_lr1r <> 0 || on_lr1l <> 0)
{
    n=1; ! counter
    while (n < npara_1)
    {
        exec, inr(mk,1,$n);
        n=n+1;
    }
}
!--- ip5
if (on_lr5l <> 0 || on_lr5r <> 0)
{
    n=1; ! counter
    while (n < npara_5)
    {
        exec, inl(mk,5,$n);
        n=n+1;
    }
}
if (on_lr5r <> 0 || on_lr5l <> 0)
{
    n=1; ! counter
    while (n < npara_5)
    {
        exec, inr(mk,5,$n);
        n=n+1;
    }
}
!--- ip2
if (on_lr2r <> 0 || on_lr2l <> 0)
{
    n=1; ! counter
    while (n < npara_2)
    {
        exec, inl(mk,2,$n);
        n=n+1;
    }
}
if (on_lr2l <> 0 || on_lr2r <> 0)
{
    n=1; ! counter
    while (n < npara_2)
    {
        exec, inr(mk,2,$n);
        n=n+1;
    }
}
!--- ip8
if (on_lr8r <> 0 || on_lr8l <> 0)
if (on_lr8l <> 0 || on_lr8r <> 0)
{
    n=1; ! counter
    while (n < npara_8)
    {
        exec, inl(mk,8,$n);
        n=n+1;
    }
}
if (on_lr8r <> 0 || on_lr8l <> 0)
{
    n=1; ! counter
    while (n < npara_8)
    {
        exec, inr(mk,8,$n);
        n=n+1;
    }
}
assign,echo=terminal;
seqedit,sequence=lhcb2;
call,file=temp.bb.install;
endedit;
system, "rm temp.bb.install";

!-------------------------------------------------------------
!--- get beta functions at bb in all four IPs
use,period=lhcb2;
!select,flag=twiss,class=bbmarker,column=name,s,x,y;
twiss, sequence=lhcb2; !,file;
!--- separation for halo collisions at IP2
on_sep2 = 2.118 * sqrt(epsx * r2ip2->betx) / 0.0007999979093;
value,on_sep2;
!-------------------------------------------------------------
!   define bb elements
assign,echo=temp.bb.install;
!--- ip1
if (on_ho1 <> 0)
{
    exec, bbho(1);
}
if (on_lr1l <> 0)
{
    n=1; ! counter
    while (n < npara_1)
    {
        exec, bbl(1,$n);
        n=n+1;
    }
}
if (on_lr1r <> 0)
{
    n=1; ! counter
    while (n < npara_1)
{  exec, bbr(1,8n);
  n=n+1;
}

!--- ip5
if (on_ho5 <> 0) 
{  
exec, bbho(5);
}

if (on_lr5l <> 0) 
{
  n=1; ! counter
  while (n < npara_5) 
  {  
    exec, bbl(5,8n); 
    n=n+1;
  }
}

if (on_lr5r <> 0) 
{
  n=1; ! counter
  while (n < npara_5) 
  {  
    exec, bbr(5,8n); 
    n=n+1;
  }
}

!--- ip2
if (on_ho2 <> 0) 
{  
exec, bbho(2);
}

if (on_lr2l <> 0) 
{
  n=1; ! counter
  while (n < npara_2) 
  {  
    exec, bbl(2,8n); 
    n=n+1;
  }
}

if (on_lr2r <> 0) 
{
  n=1; ! counter
  while (n < npara_2) 
  {  
    exec, bbr(2,8n); 
    n=n+1;
  }
}

!--- ip8
if (on_ho8 <> 0) 
{  
exec, bbho(8);
}

if (on_lr8l <> 0)
{  
    n=1; ! counter  
    while (n < npara_8)  
    {  
        exec, bbl(8,$n);  
        n=n+1;  
    }  
}  
if (on_lr8r <> 0)  
{  
    n=1; ! counter  
    while (n < npara_8)  
    {  
        exec, bbr(8,$n);  
        n=n+1;  
    }  
}  
assign,echo=terminal;  
call,file=temp.bb.install;  
system, "rm temp.bb.install";  
!  
!============================================================================  
!   install bb elements  
assign,echo=temp.bb.install;  
!--- ip1  
if (on_ho1 <> 0)  
{  
    exec, inho(bb,1);  
}  
if (on_lr1l <> 0)  
{  
    n=1; ! counter  
    while (n < npara_1)  
    {  
        inl(bb,1,$n);  
        n=n+1;  
    }  
}  
if (on_lr1r <> 0)  
{  
    n=1; ! counter  
    while (n < npara_1)  
    {  
        inr(bb,1,$n);  
        n=n+1;  
    }  
}  
!--- ip5  
if (on_ho5 <> 0)  
{  
    exec, inho(bb,5);  
}  
if (on_lr5l <> 0)  
{  
    n=1; ! counter  
    while (n < npara_5)  
    {  

exec, inl(bb,5,$n);
 n=n+1;
);
}
if (on_lr5r <> 0)
{
 n=1; ! counter
 while (n < npara_5)
 {
   exec, inr(bb,5,$n);
   n=n+1;
   
  

  

};
}!--- ip2
if (on_ho2 <> 0)
{
  exec, inho(bb,2);
}
if (on_lr2l <> 0)
{
 n=1; ! counter
 while (n < npara_2)
 {  
   exec, inl(bb,2,$n);
   n=n+1;
   
  

  

};
}
if (on_lr2r <> 0)
{
 n=1; ! counter
 while (n < npara_2)
 {  
   exec, inr(bb,2,$n);
   n=n+1;
   
  

  

};
}!--- ip8
if (on_ho8 <> 0)
{
  exec, inho(bb,8);
}
if (on_lr8l <> 0)
{
 n=1; ! counter
 while (n < npara_8)
 {  
   exec, inl(bb,8,$n);
   n=n+1;
   
  

  

};
}
if (on_lr8r <> 0)
{
 n=1; ! counter
 while (n < npara_8)
 {  
   exec, inr(bb,8,$n);
   n=n+1;
   
  

  

};


assign, echo=terminal;
select, flag=seqedit, class=bbmarker;
seqedit, sequence=lhcb2;
remove, element=selected;
endedit;
select, flag=seqedit, clear;
seqedit, sequence=lhcb1;
call, file=temp.bb.install;
endedit;

!!--- Now the beam-beam element installation is complete
system, "rm temp.bb.install";

seqedit, sequence=lhcb1;
cycle, start=ipl;
endedit;

use, period=lhcb1;
!twiss, sequence=lhcb1;
!
! make footprint
!

option, trace;
small=0.05;
big=sqrt(1.-small^2);
track;
xsmall; ysmall;
value, xs, ys;
start, fx=xsmall, fy=ys; // zero amplitude
nsigmax=6;
n=1; // sigma multiplier
m=0; // angle multiplier
while (n <= nsigmax)
{
    angle = 15*m*pi/180;
    if (m == 0) {xs=n*big; ys=n*xsmall;}
    elseif (m == 6) {xs=n*xsmall; ys=n*big;}
    else
    {
        xs=n*cos(angle);
        ys=n*sin(angle);
    }
    value, xs, ys;
    start, fx=xsmall, fy=ys;
    m=m+1;
    if (m == 7) { m=0; n=n+1;}
};
dynap, fastune, turns=1024;
endtrack;


Real life example of MACRO definitions

bbho(nn): macro = {
    !--- macro defining head-on beam-beam elements; nn = IP number
    print, text="bbipnnl2: beambeam, sigx=sqrt(rnnipnnl2->betx*epsx),";
    print, " sigy=sqrt(rnnipnnl2->bety*epsy),";
    print, " xma=rnnipnnl2->x, yma=rnnipnnl2->y,";
    print, " charge:=on_honn;";
    print, text="bbipnnl1: beambeam, sigx=sqrt(rnnipnnl1->betx*epsx),";
    print, " sigy=sqrt(rnnipnnl1->bety*epsy),";
    print, " xma=rnnipnnl1->x, yma=rnnipnnl1->y,";
    print, " charge:=on_honn;";
    print, text="bbipnn:   beambeam, sigx=sqrt(rnnipnn->betx*epsx),";
    print, " sigy=sqrt(rnnipnn->bety*epsy),";
    print, " xma=rnnipnn->x, yma=rnnipnn->y,";
    print, " charge:=on_honn;";
    print, text="bbipnnr1: beambeam, sigx=sqrt(rnnipnnr1->betx*epsx),";
    print, " sigy=sqrt(rnnipnnr1->bety*epsy),";
    print, " xma=rnnipnnr1->x, yma=rnnipnnr1->y,";
    print, " charge:=on_honn;";
    print, text="bbipnnr2: beambeam, sigx=sqrt(rnnipnnr2->betx*epsx),";
    print, " sigy=sqrt(rnnipnnr2->bety*epsy),";
    print, " xma=rnnipnnr2->x, yma=rnnipnnr2->y,";
    print, " charge:=on_honn;";
};

mkho(nn): macro = {
    !--- macro defining head-on markers; nn = IP number
    print, text="mkipnnl2: bbmarker;"
    print, text="mkipnnl1: bbmarker;"
    print, text="mkipnn:   bbmarker;"
    print, text="mkipnnr1: bbmarker;"
    print, text="mkipnnr2: bbmarker;"
};

inho(xx,nn): macro = {
    !--- macro installing bb or markers for head-on beam-beam (split into 5)
    print, text="install, element= xxipnnl2, at=-long_b, from=ipnn;"
    print, text="install, element= xxipnnl1, at=-long_a, from=ipnn;"
    print, text="install, element= xxipnn,   at=1.e-9,   from=ipnn;"
    print, text="install, element= xxipnnr1, at=+long_a, from=ipnn;"
    print, text="install, element= xxipnnr2, at=+long_b, from=ipnn;"
};

sbhomk(nn): macro = {
    !--- macro to create savebetas for ho markers
    print, text="savebeta, label=rnnipnnl2, place=mkipnnl2;"
    print, text="savebeta, label=rnnipnnl1, place=mkipnnl1;"
    print, text="savebeta, label=rnnipnn,  place=mkipnn;"
    print, text="savebeta, label=rnnipnnr1, place=mkipnnr1;"
    print, text="savebeta, label=rnnipnnr2, place=mkipnnr2;"
};
mkl(nn,cc): macro = {
  !--- macro to create parasitic bb marker on left side of ip nn; cc = count
  print, text="mkipnnplcc: bbmarker;";
};

mkr(nn,cc): macro = {
  !--- macro to create parasitic bb marker on right side of ip nn; cc = count
  print, text="mkipnnprcc: bbmarker;";
};

sbl(nn,cc): macro = {
  !--- macro to create savebetas for left parasitic
  print, text="savebeta, label=rnnipnnplcc, place=mkipnnplcc;";
};

sbr(nn,cc): macro = {
  !--- macro to create savebetas for right parasitic
  print, text="savebeta, label=rnnipnnprcc, place=mkipnnprcc;";
};

inl(xx,nn,cc): macro = {
  !--- macro installing bb and markers for left side parasitic beam-beam
  print, text="install, element= xxipnnplcc, at=-cc*b_h_dist, from=ipnn;";
};

inr(xx,nn,cc): macro = {
  !--- macro installing bb and markers for right side parasitic beam-beam
  print, text="install, element= xxipnnprcc, at=cc*b_h_dist, from=ipnn;";
};

bbl(nn,cc): macro = {
  !--- macro defining parasitic beam-beam elements; nn = IP number
  print, text="bbipnnplcc: beambeam, sigx=sqrt(rnnipnnplcc->betx*epsx),";
  print, text="          sigy=sqrt(rnnipnnplcc->bety*epsy),";
  print, text="          xma=rnnipnnplcc->x, yma=rnnipnnplcc->y,";
  print, text="          charge:=on_lrnnl;";
};

bbr(nn,cc): macro = {
  !--- macro defining parasitic beam-beam elements; nn = IP number
  print, text="bbipnnprcc: beambeam, sigx=sqrt(rnnipnnprcc->betx*epsx),";
  print, text="          sigy=sqrt(rnnipnnprcc->bety*epsy),";
  print, text="          xma=rnnipnnprcc->x, yma=rnnipnnprcc->y,";
  print, text="          charge:=on_lrnrr;";
};

\hansg June 17, 2002
Parameter Statements

Relations between Variable Parameters

A relation is established between variables by the statement

\[ \text{parameter-name} = \text{expression}; \]

or

\[ \text{parameter-name} := \text{expression}; \]

The first form evaluates the expression at the right immediately and assigns its value to the parameter. The second form assigns the value by evaluating the expression at right every time the parameter is actually used. This holds as well for element parameters - beware! If you want to modify e.g. the strength of a quadrupole later (e.g. in a match, or by entering a new value for a parameter on which it depends) then the definition has to be

\[
\text{qd:quadrupole,k1:= ak1;}
\]

and not

\[
\text{qd:quadrupole,k1 = ak1;}
\]

In the latter case, k1 will be set to the current value of ak1, and will not change when ak1 changes.

Parameters not yet defined have the value zero.

Example:

\[
\text{gev = 100;}
\]

\[
\text{beam,energy=gev;}
\]

the parameter on the left may appear on the right as well:

\[
\text{x = x+1;}
\]

Increases the value of x by 1. As a result, the SET statement of MAD-8 is no longer necessary and is not implemented.

Circular definitions are allowed in the first form:

\[
\text{a = b + 2;}
\]

\[
\text{b = a * b;}
\]

However, circular definitions in the second form are forbidden:
a := b + 2;
b := a * b;

will result in an error.

**VALUE: Output of Parameters**

The **VALUE** statement

```
VALUE = expression;
```

or

```
VALUE = expression1, expression2, ...;
```

evaluates the current value of "expression" resp. "expression1" etc. and prints the result on the standard output file.

Example:

```
p1 = 5;
p2 = 7;
value, p1*p2-3;
```

After echoing the command, this prints:

```
p1*p2-3 = 32       ;
```

Another example:

```
option, -warn;
while (x < 100) {x = x + 1;}
value, x, x^2, log10(x);
```

After echoing the command, this prints:

```
x = 100        ;
x^2 = 10000      ;
log10(x) = 2      ;
```
Constraints

In matching it is desired to specify equality constraints, as well as lower and upper limits for a quantity. MAD accepts the following forms of constraints:

! equality constraint:
name=expression

! upper limit:
name<expression

! lower limit:
name>expression

! both upper and lower limit for the same name:
name<expression,name>expression

hansg May 8, 2001
Variable Names

A variable name can have one of the formats:

1. parameter-name
2. element-name->attribute-name
3. command-name->attribute-name
4. beam%sequence-name->attribute-name
5. table(table-name,..,..)

The first format refers to the value of the global parameter "parameter-name", the second and third formats refer to the real attribute "attribute-name" of the element "element-name", or the command "command-name". Number four is specific to beams belonging to a particular sequence (for details see sequences and beams). Number five allows extraction of variables from existing tables, as specified in table access.

hansg May 8, 2001
Regular Expressions

Some commands allow selection of items via "regular expression" strings. Such a pattern string must be enclosed in single or double quotes. MAD-X follows regexp (Unix regular expression patterns) for matching. The following features are implemented:

- "^" at the start of the search string: Match following search string at the start of the target string; otherwise the search string can start anywhere in the target string. To search for a genuine "^" anywhere, use "\^".
- "$" at the end of the search string: Match preceding search string at the end of the target string; otherwise the search string can end anywhere in the target string. To search for a genuine "$" anywhere, use "\$".
- ".": Stands for an arbitrary character; to search for a genuine ".", use "\.".
- "[xyz]": Stands for one character belonging to the string contained in brackets (example: "[abc]" means one of a, b, c).
- "[a-ex-z]": Stands for ranges of characters (example: "[a-zA-Z]" means any letter).
- "[^xyz]" (i.e. a "^" as first character in a square bracket): Stands for exclusion of all characters in the list, i.e. "[^a-z]" means "any character but a lower case letter".
- "*": Allows zero or more repetitions of the preceding character, either specified directly, or from a list. (examples: "a*" means zero or more occurrences of "a", "[A-Z]*" means zero or more upper-case letters).
- "backslash-c" (e.g. "\."): Removes the special meaning of character c.

All other characters stand for themselves. Example:

```plaintext
select,flag=twiss,pattern="^d..$" ;
select,flag=twiss,pattern="^k.*qd.*\r1$" ;
```

The first command selects all elements whose names have exactly three characters and begin with the letter "D". The second command selects elements beginning with the letter "K", containing the string "QD", and ending with the string ".R1". The two occurrences of ".*" each stand for an arbitrary number (including zero) of any character, and the occurrence ".\" stands for a literal period.

hansg May 8, 2001
Control Statements

MAD-X consists of a core program, and modules for specific tasks such as twiss parameter calculation, matching, thin lens tracking, and so on.

The statements listed here are those executed by the program core. They deal with the I/O, element and sequence declaration, sequence manipulation, statement flow control (e.g. IF, WHILE), MACRO declaration, saving sequences onto files in MAD-X or MAD-8 format, and so on.

- **Program flow control**
  - IF
  - ELSEIF
  - ELSE
  - WHILE
  - MACRO

- **General control**
  - ASSIGN
  - CALL
  - COGUESS
  - CREATE
  - DUMPSEQU
  - EXEC
  - EXIT
  - FILL
  - HELP
  - OPTION
  - PRINT
  - QUIT
  - READTABLE
  - RETURN
  - SAVE
  - SAVEBETA
• SELECT
• SET
• SHOW
• STOP
• SYSTEM
• TABSTRING
• TITLE
• USE
• VALUE
• WRITE

Beam specification

• BEAM
• RESBEAM

PLOT

• PLOT
• RESPLOT
• SETPLOT

Sequence editing

• SEQEDIT
• FLATTEN
• INSTALL
• MOVE
• REMOVE
• CYCLE
• REFLECT
• ENDEDIT

hansg June 17, 2002
General Control Statements

- **ASSIGN**

  assign, echo = file_name;

  where "file_name" is the name of an output file, or "terminal". This allows switching the echo stream to a file or back, but only for the commands value, show, and print. Allows easy composition of future MAD-X input files. A real life example (always the same) is to be found under footprint \texttt{example}.

- **CALL**

  call, file = file_name;

  where "file_name" is the name of an input file. This file will be read until a "return;" statement, or until end_of_file; it may contain any number of calls itself, and so on to any depth.

- **COGUESS**

  coguess,tolerance=double,x=double,
  px=double,y=double,py=double,t=double,pt=double;

  sets the required convergence precision in the closed orbit search ("tolerance", see as well Twiss command \texttt{tolerance}.

  The other parameters define a first guess for all future closed orbit searches in case they are different from zero.

- **CREATE**

  create,table=table,column=var1,var2,\_name,...;

  creates a table with the specified variables as columns. This table can then be filled, and finally one can write it in TFS format. The attribute "\_name" adds the element name to the table at the specified column, this replaces the undocumented "withname" attribute that was not always working properly.
See the example:

or an example of joining 2 tables of different length in one table including the element name:

- **DELETE**

  delete, sequence = s_name, table = t_name;

  deletes a sequence with name "s_name" or a table with name "t_name" from memory. The sequence deletion is done without influence on other sequences that may have elements that were in the deleted sequence.

- **DUMPSEQ**

  dumpseq, sequence = s_name, level = integer;

  Actually a debug statement, but it may come handy at certain occasions. Here "s_name" is the name of an expanded (i.e. USEd) sequence. The amount of detail is controlled by "level":

  level = 0: print only the cumulative node length = sequence length
  > 0: print all node (element) names except drifts
  > 2: print all nodes with their attached parameters
  > 3: print all nodes, and their elements with all parameters

- **EXEC**

  exec, label;

  Each statement may be preceded by a label; it is then stored and can be executed again with "exec, label;" any number of times; the executed statement may be another "exec", etc.; however, the major usage of this statement is the execution of a macro.

- **EXIT**

  exit;

  ends the program execution.
**FILL**

Every command

```
fill,table=table;
```

adds a new line with the current values of all column variables into the user table created beforehand. This table one can then write in TFS format. See as well the user table example.

**HELP**

```
help,statement_name;
```

prints all parameters, and their defaults of the statement given; this includes basic element types.

**OPTION**

```
option, flag { = true | false };
option, flag | -flag;
```

sets an option as given in "flag"; the part in curly brackets is optional: if only the name of the option is given, then the option will be set true (see second line); a "-" sign preceding the name sets it to "false".

Example:

```
option,echo=true;
option,echo;
```

are identical, ditto

```
option,echo=false;
option,-echo;
```

The available options are:

<table>
<thead>
<tr>
<th>name</th>
<th>default</th>
<th>meaning if true</th>
</tr>
</thead>
<tbody>
<tr>
<td>bborbit</td>
<td>false</td>
<td>the closed orbit is modified by beam-beam kicks</td>
</tr>
<tr>
<td>sympl</td>
<td>false</td>
<td>all element matrices are symplectified in Twiss</td>
</tr>
<tr>
<td>echo</td>
<td>true</td>
<td>echoes the input on the standard output file</td>
</tr>
<tr>
<td>trace</td>
<td>false</td>
<td>prints the system time after each command</td>
</tr>
<tr>
<td>verify</td>
<td>false</td>
<td>issues a warning if an undefined variable is used</td>
</tr>
<tr>
<td>warn</td>
<td>true</td>
<td>issues warnings</td>
</tr>
<tr>
<td>info</td>
<td>true</td>
<td>issues informations</td>
</tr>
<tr>
<td>tell</td>
<td>false</td>
<td>prints the current value of all options</td>
</tr>
<tr>
<td>reset</td>
<td>false</td>
<td>resets all options to their defaults</td>
</tr>
<tr>
<td>rbarc</td>
<td>true</td>
<td>converts the RBEND straight length into the arc length</td>
</tr>
<tr>
<td>thin_foc</td>
<td>true</td>
<td>if false suppresses the 1(rho**2) focusing of thin dipoles</td>
</tr>
<tr>
<td>no_fatal_stop</td>
<td>false</td>
<td>Prevents madx from stopping in case of a fatal error. Use at your own risk.</td>
</tr>
</tbody>
</table>

The option "rbarc" is implemented for backwards compatibility with MAD-8 up to version 8.23.06 included; in this version, the RBEND length was just taken as the arc length of an SBEND with
inclined pole faces, contrary to the MAD-8 manual.

- **PRINT**
  
  `print,text="...";`

  prints the text to the current output file (see ASSIGN above). The text can be edited with the help of a [macro statement](#). For more details, see there.

- **QUIT**
  
  `quit;`

  ends the program execution.

- **READTABLE**
  
  `readtable,file=filename;`

  reads a TFS file containing a MAD-X table back into memory. This table can then be manipulated as any other table, i.e. its values can be accessed, it can be plotted, written out again etc.

- **READMYTABLE**
  
  `readmytable,file=filename,table=internalname;`

  reads a TFS file containing a MAD-X table back into memory. This table can then be manipulated as any other table, i.e. its values can be accessed, it can be plotted, written out again etc. An internal name for the table can be freely assigned while for the command READTABLE it is taken from the information section of the table itself. This feature allows to store multiple tables of the same type in memory without overwriting existing ones.

- **RESBEAM**
  
  `resbeam,sequence=s_name;`

  resets the default values of the beam belonging to sequence s_name, or of the default beam if no sequence is given.
RETURN

return;

ends reading from a "called" file; if encountered in the standard input file, it ends the program execution.

SAVE

save, beam, sequence=sequ1,sequ2,..., file=filename, beam,bare;

saves the sequence(s) specified with all variables and elements needed for their expansion, onto the file "filename". If quotes are used for the "filename" capital and low characters are kept as specified, if they are omitted the "filename" will have lower characters only. The optional flag can have the value "mad8" (without the quotes), in which case the sequence(s) is/are saved in MAD-8 input format.

The flag "beam" is optional; when given, all beams belonging to the sequences specified are saved at the top of the save file.

The parameter "sequence" is optional; when omitted, all sequences are saved.

However, it is not advisable to use "save" without the "sequence" option unless you know what you are doing. This practice will avoid spurious saved entries. Any number of "select,flag=save" commands may precede the SAVE command. In that case, the names of elements, variables, and sequences must match the pattern(s) if given, and in addition the elements must be of the class(es) specified. See here for a SAVE with SELECT example.

It is important to note that the precision of the output of the save command depends on the output precision. Details about default precisions and how to adjust those precisions can be found at the SET Format instruction page.

The Attribute 'bare’ allows to save just the sequence without the element definitions nor beam information. This allows to re-read in a sequence with might otherwise create a stop of the program. This is particularly useful to turn a line into a sequence to seqedit it. Example:

t13:line=(ld16,qt1301,mqn,qt1301,ld17,qt1302,mqn,qt1302,ld18,ison);
DLTL3 : LINE=(delay, t13);
use, period=dltl3;

save,sequence=dltl3,file=t1,bare; // new parameter "bare": only sequ. saved
call,file=t1; // sequence is read in and is now a "real" sequence
// if the two preceding lines are suppressed, seqedit will print a warning
// and else do nothing
use, period=dltl3;
twiss, save, betx=bxa, alfx=alfxa, bety=bya, alfy=alfya;
plot, vaxis=betx, bety, haxis=s, colour:=100;
SEQEDIT, SEQUENCE=dltl3;
remove,element=cx.bhe0330;
remove,element=cd.bhe0330;
ENEDIT;

use, period=dltl3;
twiss, save, betx=bx, alfx=alfxa, bety=bya, alfy=alfya;

SAVEBETA

savebeta, label=label, place=place, sequence=s_name;

marks a place "place" in an expanded sequence "s_name"; at the next TWISS command execution, a beta0 block will be saved at that place with the label "label". This is done only once; in order to get a new beta0 block there, one has to re-issue the command. The contents of the beta0 block can then be used in other commands, e.g. TWISS and MATCH.

Example (after sequence expansion):

savebeta, label=sb1, place=mb[5], sequence=fivecell;
twiss;
show, sb1;

will save and show the beta0 block parameters at the end (!) of the fifth element mb in the sequence.

SELECT

select, flag=flag, range=range, class=class, pattern=pattern, slice=integer, column=s1, s2, s3, ... sn, sequence=s_name, full, clear;

selects one or several elements for special treatment in a subsequent command. All selections for a given command remain valid until "clear" is specified; the selection criteria on the same command are logically ANDed, on different SELECT statements logically ORed.

Example:

select, flag=error, class=quadrupole, range=mb[1]/mb[5];
select, flag=error, pattern="^mqw.*";

selects all quadrupoles in the range mb[1] to mb[5], and all elements (in the whole sequence) the name of which starts with "mqw" for treatment by the error module.

"flag" can be one of the following::
- seqedit: selection of elements for the seqedit module.
- error: selection of elements for the error assignment module.
- makethin: selection of elements for the makethin module that converts the sequence into one with thin elements only.
- sectormap: selection of elements for the sectormap output file from the Twiss module.
- table: here "table" is a table name such as twiss, track etc., and the rows and columns to be written are selected.
For the RANGE, CLASS, PATTERN, FULL, and CLEAR parameters see SELECT.

"slice" is only used by makethin and prescribes the number of slices into which the selected elements have to be cut (default = 1).

"column" is only valid for tables and decides the selection of columns to be written into the TFS file. The "name" argument is special in that it refers to the actual name of the selected element. For an example, see SELECT.

**SHOW**

show,command;

prints the "command" (typically "beam", "beam%sequ", or an element name), with the actual value of all its parameters.

**STOP**

stop;

ends the program execution.

**SYSTEM**

system,"...";

transfers the string in quotes to the system for execution.

Example:

system,"ln -s /afs/cern.ch/user/u/user/public/some/directory short";

**TABSTRING**

tabstring(arg1,arg2,arg3)

The "string function" tabstring(arg1,arg2,arg3) with exactly three arguments; arg1 is a table name (string), arg2 is a column name (string), arg3 is a row number (integer), count starts at 0. The function can be used in any context where a string appears; in case there is no match, it returns _void_.

!
**TITLE**

```
title,"...";
```

inserts the string in quotes as title in various tables and plots.

**USE**

```
use,period=s_name,range=range;
```

expands the sequence with name "s_name", or a part of it as specified in the `range`.

**VALUE**

```
value,exp1,exp2,...;
```

prints the actual values of the expressions given.

Example:

```
a=clight/1000.;
value,a,pmass,exp(sqrt(2));
```

results in

```
a = 299792.458 ;
pmass = 0.938271998 ;
exp(sqrt(2)) = 4.113250379 ;
```

**WRITE**

```
write,table=table,file=file_name;
```

writes the table "table" onto the file "file_name"; only the rows and columns of a preceding `select,flag=table,...;` are written. If no select has been issued for this table, the file will only contain the header. If the FILE argument is omitted, the table is written to standard output.

\[[\text{hansg}]\] June 17, 2002
Set Statements

set,format="...", sequence="...";

The set command allows 2 actions:

1) Format

The first command lets you vary the output precision.

parameter: format = s1, s2, s3

(up to) three strings defining the integer, floating, and string output format for the save, show, value, and table output. The formats can be given in any order and stay valid until replaced. The defaults are:

"10d","18.10g","-18s".

They follow the C convention. The quotes are mandatory. The allowed formats are:

"nd" for integer with n = field width.

"m.nf" or "m.ng" or "m.ne" for floating, m field width, n precision.

"ns" for string output.

The default is "right adjusted", a "-" changes it to "left adjusted". Example:

set,format="22.14e";
changes the current floating point format to 22.14e; the other formats remain untouched.

set,format="s","d","g";
sets all formats to automatic adjustment according to C conventions.

2) Sequence

The second command lets you choose the current sequence without having to use the "USE" command, which would bring you back to a bare lattice without errors. The command only works if the chosen sequence had been activated before with the "USE" command, otherwise a warning will be issued and MAD-X will continue with the unmodified current sequence. This command is particularly useful for commands that do not have the sequence as an argument like "EMIT" or "IBS".

hansg frs June 18, 2003
RESBEAM: reset beam defaults

label: RESBEAM,SEQUENCE=name;

If the sequence name is omitted, the default beam is reset.

Default BEAM Data:
PARTICLE        POSITRON
ENERGY          1 GeV
EX              1 rad m
EY              1 rad m
ET              1 rad m
KBUNCH          1
NPART           0
BCURRENT        0 A
BUNCHEDET       .TRUE.
RADIATE         .FALSE.

hansg Januay 24, 1997
Edit a Beam Line Sequence

With the help of the commands explained below, a sequence may be modified in many ways: the starting point can be moved to another place; the order of elements can be inverted; elements can be inserted one by one, or as a whole group with one single command; single elements, or classes thereof can be removed; elements can be replaced by others; finally, the sequence can be "flattened", i.e. all inserted sequences are replaced by their actual elements, such that a flattened sequence contains only elements. It is good practice to add a flatten; statement at the end of a seqedit operation to ensure a fully operational sequence. And this is particularly useful for the save command to properly save shared sequences and to write out in MAD-8 format.

- **SEQEDIT**

  ```
  seqedit, sequence=s_name;
  ```

  selects the sequence named for editing. The editing is performed on the non-expanded sequence; after having finished the editing, one has to re-expand the sequence if necessary.

- **EXTRACT**

  ```
  extract, sequence=s_name, from=MARKER_1, to=MARKER_2, newname=p;
  ```

  From the sequence named "s_name" is extracted a new sequence with name "p" starting from MARKER_1 and ending at MARKER_2. The new sequence "p" can be USEd as any other sequence. It is declared as "shared" and can therefore be combined E.G. into the cycled original sequence.

- **FLATTEN**

  ```
  flatten;
  ```

  This command includes all sequences in the sequence being edited, if any. The resulting sequence contains only elements.
INSTALL

install, element=name, class=class_name, at=real, from=place|selected;

where the parameters have the following meaning:
- element: name of the (new) element to be inserted (mandatory)
- class: class of the new element to be inserted (mandatory)
- at: position where the element is to be inserted; if no "from" is given, this is relative to the start of the sequence. If "from" is given, it is relative to the position specified there.
- from: either a place (i.e. the name(+occurrence count) of an element already existing in the sequence, e.g. mb[15], or mq.a..i1..4 etc.; or the string "selected"; in this latter case an element of the type specified will be inserted behind all elements in the sequence that are currently selected by one or several [SELECT] commands of the type
  select, flag=seqedit, class=., pattern=., range=.;
- Attention: No element definitions inside seqedit.

MOVE

move, element=name|selected, by=real, to=real, from=place;

- element: name of the existing element to be moved, or "selected", in which case all elements from existing [SELECT] commands will be moved; in the latter case, "by" must be given.
- by: amount by which the element(s) is/are to be moved; no "to" nor "from" in this case.
- to: position to which the element has to be moved; if no from, then this is relative to the start of the sequence; otherwise, it is relative to the place given in "from".
- from: place in the sequence with respect to which the element is to be positioned.

REMOVE

remove, element=name|selected;

- element: name of the existing element to be removed, or "selected", in which case all elements from existing [SELECT] commands will be removed.
- Attention: It is a bad idea to remove all markers from a sequence! In particular the "start=" marker and the new markers added by "cycle" must never be removed from a sequence.

CYCLE

cycle, start=place;

This makes the sequence start at the place given, which must be a marker.
In the case there is a shared sequence in the used sequence, the command FLATTEN has to be used before the command CYCLE. Example:

flatten ; cycle,start=place;

- **REFLECT**
  
  reflect;

  This inverts the order of element in the sequence, starting from the last element.
  
  If there are shared sequences inside the USEd sequence, the command FLATTEN must be used before the command REFLECT. Alternatively each shared sequence must first be reflected. Example:

  flatten ; reflect;

- **REPLACE**
  
  replace,element=name1|selected,by=name2;

  Element with name1 is replaced by element with name2. If name1 is "selected", then all elements selected by SELECT commands will be replaced by the element name2.

- **ENDEDIT**
  
  endedit;

  terminates the sequence editing process. The nodes in the sequence are renumbered according to their occurrence which might have changed during editing.

hansg June 17, 2002
Elements and Markers

- **Element Input Format**
- **Aperture, Geometric**
- **MARKER: Marker Definition**
- **DRIFT: Drift Space**
- **Bending Magnet**
  - RBEND: Rectangular Bending Magnet
  - SBEND: Sector Bending Magnet
  - Dipedge Element
- **QUADRUPOLE**
- **SEXTUPOLE**
- **OCTUPOLE**
- **MULTIPOLE**
- **SOLENOID**
- **Closed Orbit Corrector**
  - HKICKER: Horizontal Orbit Corrector
  - VKICKER: Vertical Orbit Corrector
  - KICKER: Combined Orbit Corrector
- **Transverse Kicker**
- **RFCAVITY**
- **CRABCAVITY**
- **ELSEPARATOR: Electrostatic Separator**
- **Beam Position Monitor**
  - HMONITOR: Horizontal Monitor
  - VMONITOR: Vertical Monitor
  - MONITOR: Combined Monitor
  - INSTRUMENT: Other Beam Instrumentation
- **Collimators**
  - RCOLLIMATOR: Rectangular Collimator
  - ECOLLIMATOR: Elliptic Collimator
- **Coordinate Transformations**
  - YROTATION: Rotation About the Vertical Axis
  - SROTATION: Rotation Around the Longitudinal Axis
- **BEAMBEAM: Beam-Beam Interaction**
- **MATRIX: Arbitrary Element**
- **Editing Element Definitions**
• Element Class

hansg January 24, 1997
Element Input Format

All physical elements are defined by statements of the form

```
label: keyword {,attribute};
```

Example:

```
QF: QUADRUPOLE,L=1.8,K1=0.015832;
```

where

- **label** is a name to be given to the element (in the example QF),
- **keyword** is an element type keyword (in the example QUADRUPOLE).
- **attribute** normally has the form "attribute-name=attribute-value" or "attribute-name:=attribute-value" (except for multipoles).
- **attribute-name** selects the attribute, as defined for the element type keyword (in the example L and K1).
- **attribute-value** gives it a value (in the example 1.8 and 0.015832). The value may be specified by an expression. The "=" assigns the value on the right to the attribute at the time of definition, regardless of any further change of the right hand side; the ":=" re-evaluates the expression at the right every time the attribute is being used. For constant right hand sides, "=" and ":=" are of course equivalent.

Omitted attributes are assigned a default value, normally zero.

A special format is used for a multipole

```
m:multipole, kn= {kn0, kn1, kn2, ..., knmax},
               ks= {ks0, ks1, ks2, ..., ksmax};
```

where kn and ks give the integrated normal and skew strengths, respectively. The commas are mandatory, each strength can be an expression; their position defines the order. example:

```
m:multipole, kn={0,0,0,myoct*lrad}, ks={0,0,0,0,-1.e-5};
```

defines a multipole with a normal octupole, and a skew decapole component.

To know the current maximum order, enter the command

```
help,multipole;
```

and count.
Dipedge Element

A thin element describing the edge focusing of a dipole has been introduced in order to make it possible to track trajectories in the presence of dipoles with pole face angles. Only linear terms are considered since the higher order terms would make the tracking non-symplectic. The transformation of the machine elements into thin lenses leaves dipedge untouched and splits correctly the SBEND’s.

It does not make sense to use it alone. It can be specified at the entrance and the exit of a SBEND. They are defined by the commands:

```
label : dipedge, h=real, e1=real, fint=real, hgap=real, tilt=real;
```

It has zero length and five attributes.

- **H**: Is angle/length or 1/rho (default: 0 m$^{-1}$ - for the default the dipedge element has no effect). (must be equal to that of the associated SBEND)
- **E1**: The rotation angle for the pole face. The sign convention is as for a SBEND $\text{Bending Magnet}$. Note that it is different for an entrance and an exit (default: 0 rad).
- **FINT**: field integral as for SBEND $\text{sector bend}$. Note that each dipedge has its own fint, so fintx is no longer necessary.
- **HGAP**: half gap height of the associated SBEND $\text{Bending Magnet}$
- **TILT**: The roll angle about the longitudinal axis (default: 0 rad, i.e. a horizontal bend). A positive angle represents a clockwise rotation.

*[fr* February 27, 2005]
MULTIPOLE: General Thin Multipole

label: MULTIPOLE, LRAD=real, TILT=real,
      KNL:={..,..,..}, KSL:={..,..,..};

A MULTIPOLE is a thin-lens magnet of arbitrary order, including a dipole:

- LRAD: A fictitious length, which was originally just used to compute synchrotron radiation effects. A non-zero LRAD in conjunction with the \texttt{OPTION thin\_foc} set to a \texttt{true} logical value takes into account of the weak focussing of bending magnets.
- TILT: The roll angle about the longitudinal axis (default: 0 rad). A positive angle represents a clockwise rotation of the multipole element.

Please note that contrary to MAD8 one has to specify the desired TILT angle, otherwise it is taken as 0 rad. We believe that the MAD8 concept of having individual TILT values for each component and on top with default values led to considerable confusion and allowed for excessive and unphysical freedom. Instead, in MAD-X the KNL/KSL components can be considered as the normal or skew multipole components of the magnet on the bench, while the TILT attribute can be considered as an tilt alignment error in the machine.

- KNL: The normal multipole coefficients from order zero to the maximum; the parameters are positional, therefore zeros must be filled in for components that do not exist. Example of a thin-lens sextupole:

\begin{verbatim}
ms:multipole, knl:={0, 0, k2l};
\end{verbatim}

- KSL: The skew multipole coefficients from order zero to the maximum; the parameters are positional, therefore zeros must be filled in for components that do not exist. Example of a thin-lens skew octupole:

\begin{verbatim}
ms:multipole, ksl:={0, 0, 0, k3sl};
\end{verbatim}

Both KNL and KSL may be specified for the same multipole.

A multipole with no dipole component has no effect on the reference orbit, i.e. the reference system at its exit is the same as at its entrance. If it includes a dipole component, it has the same effect on the reference orbit as a dipole with zero length and deflection angle K0L, being the first component of KNL above.

\[\text{hansg, Frank.Schmidt, August 28, 2003}\]
Transverse Kicker

The type TKICKER should be used to create horizontal, vertical or combined transverse kickers physically equivalent to elements of type KICKER, but not used by the closed orbit correction module (see CORRECT command).

Examples of elements that may use the type TKICKER:

- Fast kickers for injection, dump & tune
- Magnetic septa towards beam dump
- Dampers of transverse beam oscillations
- Undulator & Wiggler magnets

For further information on element type TKICKER and its attributes, look at the documentation of the orbit corrector type KICKER.

madx team, September 15, 2011
Collimators

Two types of collimators are defined:

- **ECOLLIMATOR.** Elliptic aperture,
- **RCOLLIMATOR.** Rectangular aperture.

```plaintext
label: ECOLLIMATOR,TYPE=name,L=real,XSIZE=real,YSIZE=real;
label: RCOLLIMATOR,TYPE=name,L=real,XSIZE=real,YSIZE=real;
```

Either type has three real attributes:

- **L**: The collimator length (default: 0 m).
- **XSIZE**: The horizontal half-aperture (default: unlimited).
- **YSIZE**: The vertical half-aperture (default: unlimited).

For elliptic apertures, **XSIZE** and **YSIZE** denote the half-axes respectively, for rectangular apertures they denote the half-width of the rectangle. Optically a collimator behaves like a drift space, but during tracking, it also introduces an aperture limit. The aperture is checked at the entrance. If the length is not zero, the aperture is also checked at the exit.

Example:

```plaintext
COLLIM: ECOLLIMATOR,L=0.5,XSIZE=0.01,YSIZE=0.005;
```

The [**straight reference system**](#) for a collimator is a cartesian coordinate system.

**NOTE:** When a collimator is displaced transversally in order to model an asymmetric collimator, particle losses in tracking are reported with respect to the **displaced** reference system, not with respect to the surrounding beam line.

*hansg* January 24, 1997
Coordinate Transformations

YROTATION: Rotation About the Vertical Axis

\[ x_2 = x_1 \cos(\theta) - y_1 \sin(\theta), \quad y_2 = x_1 \sin(\theta) + y_1 \cos(\theta). \]

It has one real attribute:

- \text{ANGLE:} The rotation angle \( \theta \) (default: 0 rad). It must be a small angle, i.e. an angle comparable to the transverse angles of the orbit.

A positive angle means that the new reference system is rotated clockwise about the local \( y \)-axis with respect to the old system.

Example:

KINK: YROTATION, ANGLE=0.0001;

SROTATION: Rotation Around the Longitudinal Axis

\[ x_2 = x_1 \cos(\psi) - y_1 \sin(\psi), \quad y_2 = x_1 \sin(\psi) + y_1 \cos(\psi). \]

It has one real attribute:

- \text{ANGLE:} The rotation angle \( \psi \) (default: 0 rad)

A positive angle means that the new reference system is rotated clockwise about the \( s \)-axis with respect to the old system.

Example:
ROLL1: SROTATION,ANGLE=\pi/2.;
ROLL2: SROTATION,ANGLE=-\pi/2.;
HBEND: SBEND,L=6.0,ANGLE=0.01;
VBEND: LINE=(ROLL1,HBEND,ROLL2);

The above is a way to represent a bend down in the vertical plane, it could be defined more simply by

VBEND: SBEND,L=6.0,KOS=0.01/6;

hansg June 17, 2002
**BEAMBEAM: Beam-beam Interaction**

The command BEAMBEAM may be inserted in a beam line to simulate a beam-beam interaction point:

```plaintext
label: BEAMBEAM, SIGX=real,SIGY=real,
      XMA=real,YMA=real,CHARGE=real
      BBSHAPE=int,WIDTH=real,BBDIR=int;
```

The beam-beam interaction is represented by a four-dimensional interaction with a thin element, i.e. horizontal and vertical non-linear kicks. The code for this element has been contributed by J.M. Veuillen (1987) and extended by S. Sorge (2007).

- **SIGX**: The horizontal extent of the opposite beam (default: 1 m). Meaning depends on parameter BBSHAPE.
- **SIGY**: The vertical extent of the opposite beam (default: 1 m). Meaning depends on parameter BBSHAPE.
- **XMA**: The horizontal displacement of the opposite beam with respect to the ideal orbit (default: 0 m).
- **YMA**: The vertical displacement of the opposite beam with respect to the ideal orbit (default: 0 m).
- **CHARGE**: The charge of particles in the opposite beam in elementary charges. It is set by default CHARGE=1. So, if you want to describe collisions between beams containing the same particles having a charge different from 1, you have to set CHARGE explicitly in BEAM and in BEAMBEAM.
- **BBSHAPE**: The parameter to choose the radial density shape of the opposite beam (default: 1)
  - **BBSHAPE=1**: Gaussian shape (default), SIGX/SIGY: standard deviation in vertical/horizontal direction.
  - **BBSHAPE=2**: trapezoidal shape, SIGX/SIGY: half width of density profile, i.e. distance from the centre to half edge region with linear decrease of density in horizontal/vertical direction. Still only circular opposite beam possible, i.e. in the calculations SIGX′=SIGY′=(SIGX+SIGY)/2 is used, if SIGX and SIGY have different values.
BBSHAPE=3: hollow-parabolic shape, SIGX/SIGY: distance from the centre to the maximum of the parabolic density profile in vertical/horizontal direction. Still only circular opposite beam possible, i.e. in the calculations SIGX’=SIGY’=(SIGX+SIGY)/2 is used, if SIGX and SIGY have different values.
The restriction to circular opposite beams in the cases BBSHAPE=2,3 appears to be sufficient, because such beam profiles are more important for the description of the interaction between the particle beam and an electron beam of an electron cooler, which are usually circular.

- **WIDTH**: The relative extent of the edge region, absolute value is given by WIDTH*SIGX and WIDTH*SIGY vertical and horizontal direction, respectively. For
  - BBSHAPE=1, WIDTH is meaningless and will be ignored.
  - BBSHAPE=2, WIDTH denotes the full width of the edge region in units of SIGX (or SIGX’ and SIGY’, respectively, if SIGX and SIGY are not equal), i.e. if WIDTH=0.01 and SIGX=5 mm, the edge region has a full width of 0.05 mm. It must be WIDTH < 2.0.
  - BBSHAPE=3, WIDTH denotes the full width at half maximum of the parabolic density profile in units of SIGX (or SIGX’ and SIGY’, respectively, if SIGX SIGY are not equal. It must be WIDTH < SQRT(2.0).

- **BBDIR**: The parameter to choose the direction of motion of the opposite beam relative to the beam considered. It determines the sign of the Lorenz force between the both beams (default: -1):
  - BBDIR=-1: Beams move in the opposite direction as in a collider. Therefore, the Lorenz force enhances the beam-beam interaction.
  - BBDIR=0: Opposite beam does not move, Lorenz force is neglected
  - BBDIR=1: Beams move in the same direction as in an electron cooler. So, the Lorenz force reduces the beam-beam interaction.

Note:
- The particles in the beam considered may have a momentum deviation given by DELTAP
defined in the TRACK command.

- The opposite beam is assumed to have the velocity according to the unperturbed energy of the particles in the beam considered. So, only the direction of motion can be chosen.
- In the case of motion in the opposite direction (BBDIR=-1), the time of interaction between the beams is given by \( \tau = \frac{\text{length}}{2\beta c_{\text{light}}} \), where length is the length of a bunch in the opposite beam. In the case of motion in the same direction (BBDIR=1) as in an electron cooler, this time is given by \( \tau = \frac{\text{length}}{\beta c_{\text{light}}} \), where length is the length of the cooler. So, the factor 1/2 is inserted only for BBDIR=-1 to calculate correct results.

A beam-beam element requires the particle energy (ENERGY) and the particle charge (CHARGE) as well as the number of particles per bunch (NPART) to be set by a BEAM command before any calculations are performed.

Examples of a four-dimensional beam-beam element definition:

Collider regime example:

```
beam,   particle=positron,npart=1.e12,energy=50.0;
bb:     beambeam,sigx=1.e-3,sigy=5.e-4,charge=1.;
```

Electron cooler example:

```
gamma0=1.032;                           ! relativistic factors
beta0=sqrt(1.0-1.0/gamm0/gamm0);      ! relativistic factors
i_e=0.2;                                ! electron current
re_cool=0.01;                           ! electron beam radius
l_cool=5.0;                             ! cooling length
nelect=i_e*l_cool/beta0/clight/qelect;  ! electron number in e-cooler
beam,particle=antiproton,gamma=gamma0,npart=nelect;
bb_ecool:beambeam,sigx=re_cool,sigy=re_cool,bbshape=2,width=0.01,charge=-1,bbdir=1;
```

For the definition of the LHC head-on and parasitic beam-beam elements see beam-beam element examples.

**hans** ssorge, July 13, 2007
The MATRIX permits the definition of an arbitrary transfer matrix. It has four real array attributes:

- **L**: Length of the element, which may be zero.
- **KICKi**: Defines the kick of the element acting on the six phase space coordinates.
- **RMik**: Defines the linear transfer matrix (6x6) of the element.
- **TMikl**: Defines the second-order terms (6x6x6) of the element.

Data values not entered are taken from the identity transformation, kick and second-order terms are zero as default. In the thin-lens tracking module the length of an arbitrary matrix is accepted, however no second-order are allowed to avoid non symplectic tracking runs. In the latter case the tracking run will be aborted.

![Ir] June 25, 2003
Editing Element Definitions

An element definition can be changed in two ways:

- **Entering a new definition:** The element will be replaced in the main beam line expansion.
- **Entering the element name together with new attributes:** The element will be updated in place, and the new attribute values will replace the old ones.

This example shows two ways to change the strength of a quadrupole:

```
QF: QUADRUPOLE,L=1,K1=0.01;  ! Original definition of QF
QF: QUADRUPOLE,L=1,K1=0.02;  ! Replace whole definition of QF
QF,K1=0.02;                   ! Replace value of K1
```

When the type of the element remains the same, replacement of an attribute is the more efficient way.

Element definitions can be edited freely. The changes do not affect already defined objects which belong to its **element class**.

---

* hansg, January 24, 1997
Element Classes

The concept of element classes solves the problem of addressing instances of elements in the accelerator in a convenient manner. It will first be explained by an example. All the quadrupoles in the accelerator form a class QUADRUPOLE. Let us define three subclasses for the focussing quadrupoles, the defocussing quadrupoles, and the skewed quadrupoles:

MQF: QUADRUPOLE, L=LQM, K1=KQF; ! Focussing quadrupoles
MQD: QUADRUPOLE, L=LQM, K1=KQD; ! Defocussing quadrupoles
MQT: QUADRUPOLE, L=LQT; ! Skewed quadrupoles

These classes can be thought of as new keywords which define elements with specified default attributes. We now use these classes to define the real quadrupoles:

QD1: MQD; ! Defocussing quadrupoles
QD2: MQD;
QD3: MQD;
...
QF1: MQF; ! Focussing quadrupoles
QF2: MQF;
QF3: MQF;
...
QT1: MQT, K1S=KQT1; ! Skewed quadrupoles
QT2: MQT, K1S=KQT2;
...

These quadrupoles inherit all unspecified attributes from their class. This allows to build up a hierarchy of objects with a rather economic input structure.

The full power of the class concept is revealed when object classes are used to select instances of elements for various purposes. Example:

select, flag=twiss, class=QUADRUPOLE; ! Select all quadrupoles for the
     ! Twiss TFS file

More formally, for each element keyword MAD maintains a class of elements with the same name. A defined element becomes itself a class which can be used to define new objects, which will become members of this class. A new object inherits all attributes from its class; but its definition may override some of those values by new ones. All class and object names can be used in range selections, providing a powerful mechanism to specify positions for matching constraints and printing.

hansg January 24, 1997
Beam Line Sequences

MAD-X accepts two forms of an accelerator definition: sequences and lines. However, the sequence definition is the only one used internally; lines are converted into sequences when they are USEd. Consequently, only sequences can be saved (written onto a file) by MAD-X.

The corresponding sequence of statements defining a sequence is

```
name: SEQUENCE,REFER=keyword,REFPOS=name,LENGTH=real
label: class, AT=real{,attributes} | class, AT=real | sequ_name, AT=real
...
ENDSEQUENCE
```

where "real" means a real number, variable, or expression.

The first line gives the sequence name, a REFER flag (entry, centre, or exit) which specifies at which part of the element its position along the beam line will be given (default: centre), a REFPOS argument used for sequence insertion, and the total length.

Inside the sequence ... endsequence bracket three types of commands may be placed:

- an element declaration as usual, with an additional "at" attribute giving the element position relative to the start of the sequence; CAUTION: an existing definition for an element with the same name will be replaced, however, defining the same element twice inside the same sequence results in a fatal error, since a unique object (this element) would be placed at two different positions.
- a class name with a position; this causes an instance of the class to be placed at the position given. For uses inside ranges, instances of the same class can be accessed with an occurrence count.
- a sequence name with a position; this causes the sequence with that name to be placed at the position indicated. The entry, centre, or exit of the inserted sequence are placed at the position given, UNLESS a "refpos" (the name of an element in the inserted sequence) is given, in which case the sequence is inserted such that the refpos element is at the insertion point.

When the sequence is expanded in a USE command, MAD generates the missing drift spaces. At this moment, overlapping elements will cause "negative drift length" errors.

For efficiency reasons MAD-X imposes an important restriction on element lengths and positions: once a sequence is expanded, the element positions and lengths are considered as fixed; in order to vary a position or element length, a re-expansion of the sequence becomes necessary. The MATCH command contains a special flag "vlength" to match element lengths.
Example:

! define a default beam (otherwise fatal error)
beam;
! Define element classes for a simple cell:
b:    sbend,l=35.09, angle = 0.011306116;
qf:   quadrupole,l=1.6,k1=-0.02268553;
qd:   quadrupole,l=1.6,k1=0.022683642;
sf:   sextupole,l=0.4,k2=-0.13129;
sd:   sextupole,l=0.76,k2=0.26328;
! define the cell as a sequence:
sequ:  sequence,l=79;
    b1:  b,   at=19.115;
sf1:   sf,  at=37.42;
    qf1: qf,  at=38.70;
    b2:  b,   at=58.255,angle=b1->angle;
sd1:   sd,  at=76.74;
    qd1: qd,  at=78.20;
    endm: marker, at=79.0;
endsequence;
Beam Lines

The accelerator to be studied is known to MAD-X either as a sequence of physical elements called sequence or as a hierarchically structured list of elements called a beam line. A beam line is built from simpler beam lines whose definitions can be nested to any level. A powerful syntax allows to repeat, to reflect, or to replace pieces of beam lines. However, internally MAD-X knows only sequences, and lines as shown below are converted into flat sequences with the same name when they are expanded. Consequently, only sequences can be SAVEd onto a file (see save).

Formally a beam line is defined by a LINE command:

```
label(arg{,arg}): LINE=(member{,member});
```

Label gives a name to the beam line for later reference.

The formal argument list (arg{,arg}) is optional (see below). Each "member" may be one of the following:

- Element label,
- Beam line label,
- Sub-line, enclosed in parentheses,
- Formal argument name,
- Replacement list label.

Beam lines may be nested to any level.

Simple Beam Lines

The simplest beam line consists of single elements:

```
label: LINE=(member{,member});
```

Example:

```
l:      line=(a,b,c,d,a,d);
        use,period=l;
```

The USE command tells MAD to perform all subsequent calculations on the sequence

```
a,b,c,d,a,d
```
Sub-lines

Instead of referring to an element, a beam line member can refer to another beam line defined in a separate command. This provides a shorthand notation for sub-lines which occur several times in a beam line. Lines and sub-lines can be entered in any order, but when a line is expanded, all its sub-lines must be known.

Example:

```
l:     line=(a,b,s,b,a,s,a,b);
s:     line=(c,d,e);
       use,period=1;
```

this example produces the following expansion steps:

1. replace sub-line s:
   ```
   (a,b,(c,d,e),b,a,(c,d,e),a,b)
   ```

2. omit parentheses:
   ```
   a,b,c,d,e,b,a,c,d,e,a,b
   ```

Reflection and Repetition

An unsigned repetition count and an asterisk indicate repetition of a beam line member. A minus prefix causes reflection, i.e. all elements in the subsequence are taken in reverse order. Sub-lines of reflected lines are also reflected, but physical elements are not. If both reflection and repetition are desired, the minus sign must precede the repetition count.

Example:

```
r:     line=(g,h);
s:     line=(c,r,d);
t:     line=(2*s,2*(e,f),-s,-(a,b));
       use,period=t;
```

Attention: the repetition "2*s" will only work if "s" is itself a line. In case "s" is an element replace by "2*(s)".

Proceeding step by step, this example produces

1. Replace sub-line S:
   ```
   ((c,r,d),(c,r,d),(e,f),(e,f),(d,-r,c),(b,a))
   ```

2. replace sub-line r:
   ```
   ((c,(g,h),d),(c,(g,h),d),(e,f),(e,f),(d,(h,g),c),(b,a))
   ```
3. omit parentheses:

c, g, h, d, c, g, h, d, e, f, e, f, d, h, g, c, b, a

Note that the inner sub-line R is reflected together with the outer sub-line S.

**Replaceable Arguments**

A beam line definition may contain a formal argument list, consisting of labels separated by commas and enclosed in parentheses. Such a line can be expanded for different values of its arguments. When this line is referred to, its label must be followed by a list of actual arguments separated by commas and enclosed in parentheses. These arguments must be beam line, or element names. The number of actual arguments must agree with the number of formal arguments. All occurrences of a formal argument on the right-hand side of the line definition are replaced by the corresponding actual argument.

Example:

```plaintext
s:  line=(a, b, c);
1(x,y): line=(d, x, e, 3*y);
l4f:  line=(4*f);
lm2s: line=(-2*s);
res:  line=1(14f, lm2s);
```

Proceeding step by step, this example generates the expansion

d, f, f, f, f, e, c, b, a, c, b, a, c, b, a, c, b, a, c, b, a

Second example:

```plaintext
cel (sf, sd):  line=(qf, d, sf, d, b, d, qd, d, sd, d, b, d);
arc:           line=(cel (sf1, sd1), cel (sf2, sd2), cel (sf1, sd1));
use, period=arc;
```

This example generates the expansion

1. Replace the line CEL and its formal arguments:

   ```plaintext
   (qf, d, (sf1), d, b, d, qd, d, (sd1), d, b, d)
   (qf, d, (sf2), d, b, d, qd, d, (sd2), d, b, d)
   (qf, d, (sf1), d, b, d, qd, d, (sd1), d, b, d)
   ```

2. Omit parentheses:

   ```plaintext
   qf, d, sf1, d, b, d, qd, d, sd1, d, b, d
   qf, d, sf2, d, b, d, qd, d, sd2, d, b, d
   qf, d, sf1, d, b, d, qd, d, sd1, d, b, d
   ```

**Warning: Line Depreciation**

MADX has been developed using sequences, in fact internally the code works with sequences only. Consequently, there may exist some inconveniences when only lines are used. It is recommended to convert as soon as possible lines into sequences (by means of the save command) in a design phase and to use only sequences for a finalised machine.
Limits of Construction of Lines

Since Lines are in fact depreciated there are some limits of how they can be constructed. Please find below a running MADX run which shows an example of OK (valid) and WRONG (invalid) cases.

!---------------------------------------------------------------------
beam, PARTICLE=electron, energy=1;
qf: QUADRUPOLE, L:=1,K1:=1;
qd: QUADRUPOLE, L:=1,K1:=-1;
d: DRIFT, l=1;
m: MARKER;
rpl(a,b): LINE=(a,b);
sl: LINE=(qf,d,qd);
test0: LINE=(rpl(sl,sl)); !OK
test1: LINE=(rpl((sl),(sl)))); !OK
test2: LINE=(rpl((sl),(-sl)))); !OK
test3: LINE=(sl,-sl); !OK
test4: LINE=(rpl((3*sl),(3*sl)))); ! WRONG 
test5: LINE=(3*sl,3*sl); !OK
test6: LINE=(rpl((3*sl),(-3*sl)))); ! WRONG 
test7: LINE=(3*sl,-3*sl); !OK

use, period=test0;
twiss,BETX=1,bety=1;

use, period=test1;
twiss,BETX=1,bety=1;

use, period=test2;
twiss,BETX=1,bety=1;

use, period=test3;
twiss,BETX=1,bety=1;

use, period=test4;
twiss,BETX=1,bety=1;

use, period=test5;
twiss,BETX=1,bety=1;

use, period=test6;
twiss,BETX=1,bety=1;

use, period=test7;
twiss,BETX=1,bety=1;
!---------------------------------------------------------------------

hansg, June 17, 2002
Defining aperture in MAD-X

A new feature of MAD-X is the ability to set an aperture for a particular element, or parent of a set of elements. This removes the need of placing a collimator next to every element to do aperture tracking. The aperture of any elements can be specified (excepts drifts) by the use of the following parameters:

- **APERTYPE** This can have seven text values: CIRCLE, RECTANGLE, ELLIPSE, LHSCREEN (a superposition of a CIRCLE and a RECTANGLE), MARGUERITE (two LHSCREENS, one rotated by 90 degrees), RECTELLIPSE (a superposition of an ELLIPSE and a RECTANGLE) and RACETRACK.

- **APERTURE** This is an array of values, the number and meaning of which depends on the APERTYPE:

<table>
<thead>
<tr>
<th>APERTYPE</th>
<th># of parameters</th>
<th>meaning of parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>CIRCLE</td>
<td>1</td>
<td>radius</td>
</tr>
<tr>
<td>ELLIPSE</td>
<td>2</td>
<td>horizontal half axis, vertical half axis</td>
</tr>
<tr>
<td>RECTANGLE</td>
<td>2</td>
<td>half width and half height</td>
</tr>
<tr>
<td>LHSCREEN</td>
<td>3</td>
<td>half width, half height (of rect.) and radius (of circ.)</td>
</tr>
<tr>
<td>MARGUERITE</td>
<td>3</td>
<td>half width, half height (of rect.) and radius (of circ.)</td>
</tr>
<tr>
<td>RECTELLIPSE</td>
<td>4</td>
<td>half width, half height (of rectangle), horizontal half axis, vertical half axis (of ellipse)</td>
</tr>
<tr>
<td>RACETRACK</td>
<td>3</td>
<td>horizontal, vertical shift, radius shift</td>
</tr>
<tr>
<td>FILENAME</td>
<td>0</td>
<td>where the file contains a list of x and y coordinates outlining the shape. This option is only supported by the aperture module, see below.</td>
</tr>
</tbody>
</table>

Here is an example for setting an ELLIPTICAL aperture for the main dipoles for the LHC.

MB : SBEND, L := 1 MB, APERTYPE=ELLIPSE, APERTURE={0.02202,0.02202};

And an example for setting a FILENAME aperture for another magnet. Notice that no aperture parameters are needed.
The syntax of myfile should be like this:

\[
\begin{array}{ll}
0 & 0 \\
1 & 1 \\
& \vdots \\
\end{array}
\]

Notes concerning the use of aperture:

- There is some inconsistency in the parameter definition for the different APERTYPE. This is historical and has to be kept for backwards compatibility. Pay some attention to the parameters you introduce!
- When \texttt{MAKETHIN} is called all the thin slices inherit the aperture from their original thick lens version.
- When the SIXTRACK command is called (see the SixTrack converter module\texttt{C6T}) the apertures are ignored by default. To convert the apertures as well the APERTURE flag has to be set.
- Aperture parameters are like all parameters and are inherited by offspring. Like other parameters they can also be overridden by the offspring elements if necessary.

The APERTYPE and the APERTUREs themselves can be conveniently added to the TWISS table (see Twiss Module) by using the \texttt{SELECT} command. E.G. the command:

```
select,flag=twiss,clear;
select,flag=twiss,column-name=s,betx,alfx,mux,bety,alfy,muy,apertype,aper_1,aper_2;
```

and a subsequent TWISS command will put the aperture information together with the specified TWISS parameters into the TWISS table.

**Defining tolerances in MAD-X**

A parameter closely connected to the aperture is the sum of the mechanical and alignment tolerances. The mechanical tolerance is the maximal error margin of errors in the element body which causes a decrease of aperture, and the alignment tolerance is a misalignment of the element in the accelerator, which also causes a decrease of aperture. The tolerance is given in the transverse plane as a racetrack, like in the picture below.
A tolerance can be assigned to each element in a MAD-X sequence as a vector:

Syntax: APER_TOL = {r, g, s};

MB : SBEND, L := l.MB, APER_TOL=(1.5, 1.1, 0);

**APERTURE MODULE**

Computes the n1 values for a piece of machine. Each element is sliced into thick subelements at given intervals, and the available aperture is computed at the end of each slice. The computation is based on the last Twiss table, so it is important to run the `Twiss` and aperture commands on the same period or sequence, see the aperture example below. Also showed in the example is how n1 values can be plotted.

The minimum n1 for each element is written to the last Twiss table, to allow for matching by aperture.

- **Aperture,**

  file=filename,
  halofile=filename,
  pipefile=filename,
  range=range,
  exn=real,
  eyn=real,
  dqf=real,
  betaqfx=real,
  dp=real,
  dparx=real,
  dpary=real,
where the parameters have the following meaning:

- file: Output file with aperture table. Default = none
- halofile: Input file with halo polygon coordinates. Will suppress an eventual halo parameter. Default = none
- range: [Range given by elements. Default = #s/#e
- exn: Normalised horizontal emittance. Default = 3.75e-6
- eyn: Normalised vertical emittance. Default = 3.75e-6
- dqf: Peak linear dispersion [m]. Default = 2.086
- betaqfx: Beta x in standard qf [m]. Default = 170.25
- dp: Bucket edge at the current beam energy. Default = 0.0015
- dpdx: Fractional horizontal parasitic dispersion. Default = 0.273
- dpdx: Fractional vertical parasitic dispersion. Default = 0.273
- cor: Maximum radial closed orbit uncertainty [m]. Default = 0.004
- bbeat: Beta beating coefficient applying to beam size. Default = 1.1
- nco: Number of azimuth for radial scan. Default = 5
- halo: Halo parameters: \{n, r, h, v\}. n is the radius of the primary halo, r is the radial part of the secondary halo, h and v is the horizontal and vertical cuts in the secondary halo. Default = \{6, 8.4, 7.3, 7.3\}
- interval: Approximate length in meters between measurements. Actual value: nslice = nodelength/interval, nslice is rounded down to closest integer, interval = nodelength/nslice. Default = 1.0
- spec: Aperture spec, for plotting only. Gives the spec line in the plot. Default = 0.0
- notsimple: Use only if one or more beamscreens in the range are considered not to be a "simply connex". Since all MAD-X apertures are simply connex, this is only possible if an input file with beam screen coordinates are given. See below for a graphical example. Default = false.
- trueprofile: A file containing a list of magnets, and for each magnet a list of horizontal and vertical deviations from the ideal magnet axis. These values may come from measurements done on the magnet. See below for example. Default = none.
- offsetelem: A file containing a reference point in the machine, and a list of magnets with their offsets from this point described as a parabola. See below for example. Default = none.

**Not simply connex beam pipes**

Methodically, the algorithm for finding the largest possible halo is fairly simple. The distance from halo centre to the first apex (i = 0) in the halo is calculated (l_i), and the equation for a line going through these points is derived. This line is then compared with all lines making the pipe polygon to find their respective intersection coordinates. The distance h_i between halo centre and intersection...
are then divided by $l_i$, to find the maximal ratio of enlargement, as seen below. This procedure is then repeated for all apexes $i$ in the halo polygon, and the smallest ratio of all apexes is the maximal enlargement ratio for this halo to just touch the pipe at this particular longitudinal position.

There is one complication to this solution; polygons which are not simple connexes. (Geometrical definition of "simply connex": A figure in which any two points can be connected by a line segment, with all points on the segment inside the figure.) The figure below shows what happens when a beam pipe polygon is not a simple connex. The halo is expanded in such a way that it overlaps the external polygon in the area where the latter is dented inwards.
To make the module able to treat all kinds of polygons, *notsimple* must be activated. With this option activated, apexes are strategically added to the halo polygon wherever the beam pipe polygon might have an inward dent. This is done by drawing a line from halo centre to each apex on the pipe polygon. An apex with its coordinates on the intersection point line-halo is added to a table of halo polygon apexes. The result is that the halo polygon has a few “excessive” points on straight sections, but the algorithm used for expansion will now never miss a dent in the beam pipe. The use of the notsimple option significantly increases computation time.
Trueprofile file syntax

This file contains magnet names, and their associated displacements of the axis for an arbitrary number of S, where So is the start of the magnet and Sn the end. The interval between each S must be regular, and X and Y must be given in meters. The magnet name must be identical to how it appears in the sequence. The displacements are only valid for this particular magnet, and cannot be assigned to a family of magnets. n1 is calculated for a number of slices determinated by the number of Si.
Layout of file:

magnet.name1
So  X  Y
Si  X  Y
Sn  X  Y

magnet.name2
So  X  Y
Si  X  Y
Sn  X  Y

etc.

Example of file:

!This is the start of the file.
!Comments are made with exclamation marks.

mb.a14r1.b1
0   0.0002   0.000004
7.15 1.4e-5   0.3e-3
14.3 0.000000032 4e-6

!Further comments can of course be added

mq.22r1.b1
0   0.3e-5   1.332e-4
1.033 0.00034   0.3e-9
2.066 0   0.00e-2
3.1 4.232e-4 0.00000003

!This is the end of the file.

Offsetelem file syntax

This file contains coordinates describing how certain elements are displaced w.r.t. a given reference point in the machine. It might be used with elements in insertions, or other special-purpose elements that has a magnet axis which does not coincide with the reference trajectory. We operate with two coordinate system, s,x and s,y, where the reference point is the origin and the actual element axis is described as a parabola with coefficients A, B and C. For each element we give two sets of coefficients, one for horizontal displacement and one for vertical:

\[ X_{\text{offs}}(s) = Ax*s^2 + Bx*s + Cx \]

and

\[ Y_{\text{offs}}(s) = Ay*s^2 + By*s + Cy \]

. The coordinate systems are in meters.
Layout of file: --- FOR MADX VERSION 3.XX AND OLDER ONLY---

reference.point

magnet.name1
Ax   Bx   Cx
Ay   By   Cy

magnet.name2
Ax   Bx   Cx
Ay   By   Cy

e tc.

Example of file:

!This is the start of the file.
!First we give a reference point. The origin of the
!coordinate system will be at the START of this element.

mq.12r1.b1

!Then we give a list of elements and their displacement
!w.r.t. the reference point.

mcbxa.3l2
0   -2.56545   -3
0   -2.3443666  0

!The next nodes use the same reference point.
!Elements offset w.r.t. another point must be given in another file,
!together with the new reference point.

mcbxa.3r2
0.3323  32.443355 -0.84
0.2522  32.554363  0.0

!This is the end of the file.

Layout of file: --- FOR MADX VERSION 4.XX ONWARDS : now TFS format ---

note that variable names changes with : Ax -> DDX_OFF, Bx -> DX_OFF, Cx -> X_OFF, same for
Y The column S_IP is useless but mandatory (!). It results from a misunderstanding. Content is
ignored. In a future version, it will be suppressed (but will not induce an error if present).

@ NAME             %06s "OFFSET"
@ TYPE             %06s "OFFSET"
@ REFERENCE        %10s "mq.12r1.b1"
* NAME          S_IP       X_OFF     DX_OFF    DDX_OFF    Y_OFF    DY_OFF     DDY_OFF
"mq.12r1.b1"    0.0        -3.0      -2.56545    0.0       0.0    -2.3443666    0.0
"mcbxa.3r2"     0.0        -0.84     32.443355  0.3323     0.0    32.554363    0.2522

A python script to convert a file from the old V.3.XX format to the new V4.xx can be found at :

/afs/cern.ch/eng/lhc/optics/V6.503/aperture/convert_offsets.py

usage : convert_offsets.py filename
As an example we see in the picture below how the horizontal axes of elements m1 and m2 do not coincide with the reference trajectory.

The $X_{\text{ref}}(s)$ and $Y_{\text{ref}}(s)$ of the reference trajectory are calculated via an internal call to the Survey module. $X_{\text{offs}}(s)$ and $Y_{\text{offs}}(s)$ are derived from the coefficients given in the file. The resulting

$$X_{\text{tot}}(s) = X_{\text{ref}}(s) - X_{\text{offs}}(s)$$

and

$$Y_{\text{tot}}(s) = Y_{\text{ref}}(s) - Y_{\text{offs}}(s)$$

are taken into account in the aperture calculations.

**Aperture command example**

The aperture module needs a Twiss table to operate on. It is important not to USE another period or sequence between the Twiss and aperture module calls, else aperture looses its table. One can choose the ranges for Twiss and aperture freely, they need not be the same.

```
use, period=lhcb1;
select, flag=twiss, range=mb.a14r1.bl/mb.a17r1.bl, column=keyword,name,parent,k0l,k1l,s,betx,bety,n1;  
twiss, file=twiss.bl.data, betx=beta.ip1, bety=beta.ip1, x=x.ip1, y=y.ip1, py=py.ip1;  
plot,haxis=s,vaxis=betx,bety,colour=100;

select, flag=aperture, column=name,n1,x,dy;  
aperture, range=mb.b14r1.bl/mb.a17r1.bl, spec=5.235;  
plot,table=aperture,noline,vmin=0,vmax=10,haxis=s,vaxis=n1,spec,on_elem,style=100;
```
The `select` command can be used to choose which columns to print in the output file.

Column names: name, n1, n1x_m, n1y_m, apertype, aper_1, aper_2, aper_3, aper_4, rtol, xtol, ytol, s, betx, bety, dx, dy, x, y, on_ap, on_elem, spec

n1 is the maximum beam size in sigma, while n1x_m and n1y_m is the n1 values in si-units in the x- and y-direction.

aper_# means for all apertypes but racetrack:
aper_1 = half width rectangle
aper_2 = half height rectangle
aper_3 = half horizontal axis ellipse (or radius if circle)
aper_4 = half vertical axis ellipse

For racetrack, the aperture parameters will have the same meaning as the tolerances:
aper_1 and xtol = horizontal displacement of radial part
aper_2 and ytol = vertical displacement of radial part
aper_3 and rtol = radius
aper_4 = not used

On_elem indicates whether the node is an element or a drift, and on_ap whether it has a valid aperture. The Twiss parameters are the interpolated values used for aperture computation.

When one wants to plot the aperture, the table=aperture parameter is necessary. The normal line of hardware symbols along the top is not compatible with the aperture table, so it is best to include noline. Plot instead the column on_elem along the vaxis to have a simple picture of the hardware. Spec can be used for giving a limit value for n1, to have something to compare with on the plot. This example provides a plot,
where we see the $n1$, beta functions and the hardware symbolized by $on_{elem}$.
SixTrack: Produce input files for tracking in SixTrack

In dynamic aperture studies SixTrack is often used because of its speed and controllability. However the input files are notoriously difficult to produce by hand. This command may be used to produce SixTrack files from a sequence in MAD-X’s memory.

N.B.: The files contain all information concerning optics, field errors and misalignments. Hence these should all be set and a

TWISS, SAVE;

command should always be issued beforehand.

The generation of the SixTrack input files is then done by the command:

SIXTRACK, CAVALL,
MULT_AUTO_OFF,
MAX_MULT_ORD,
SPLIT,
APERTURE,
RADIUS = ref. radius of magnets;

The parameters are defined as:

- CAVALL - (optional flag) This puts a cavity element (SixTrack identifier 12) with Volt, Harmonic Number and Lag attributes at each location in the machine. Since for large hadron machines the cavities are typically all located at one particular spot in the machine and since many cavities slow down the tracking simulations considerably all cavities are lumped into one and located at the first appearance of a cavity. This default is enforced by omitting this flag.
- MULT_AUTO_OFF - (optional flag, default = .FALSE.) If .TRUE. this module does not process zero value multipoles. Moreover, multipoles are prepared in SixTrack (file fc.3) to be treated up to the order as specified with MAX_MULT_ORD.
- MAX_MULT_ORD - (optional parameter, default = 11) Process up to this order for mult_auto_off = .TRUE.
- SPLIT - (optional flag) OBSOLETE. This splits all the elements in two. This is for backwards compatibility only. The user should now use the MAKETHIN command instead.
- APERTURE - (optional flag) Set this to convert the apertures from MAD-X to SixTrack, so SixTrack will track with aperture.
- RADIUS - (optional, default value is 1m). This sets the reference radius for the magnets. This argument is optional but should normally be set.
- Note: the bv flag is presently ignored
- WARNING: SixTrack and c6t are presently set up for names of a maximum of 16 characters!!!!!
Therefore, it is mandatory to respect this limit for MAD-X names.

The command will then always produce the following file:

- fc.2 - contains the basic structure of the lattice.

and may produce any or all of the following files, depending on the sequence:

- fc.3 - contains the multipole mask(s).
- fc.3.aux - contains various beam parameters.
- fc.8 - contains the misalignments and tilts.
- fc.16 - contains the field errors and/or combined multipole kicks.
- fc.34 - file with various optics parameters at various locations (not needed by SixTrack but may be used as input to [SODD])

For a full description of these files see [SixTrack] and for information on running SixTrack see [Run Environment].

Mark Hayes 20.06.02
MAKETHIN: Slice a sequence into thin lenses

This module converts a sequence with thick elements into one composed entirely of thin elements as required by the default MAD-X tracking.

Slicing is done by the MAKETHIN command:

\[
\text{MAKETHIN, SEQUENCE=sequence name, STYLE=slicing style;}
\]

The parameters are defined as:

- **SEQUENCE** chooses the sequence you wish to slice.
- **STYLE** (optional) chooses the slicing style. The options are:
  - SIMPLE: this is a simplified slicing algorithm which produces any number of equal strength slices at equidistant positions with the kick in the middle of each slice.
  - COLLIM: this is the default slicing for collimators. If only one slice is chosen it is placed in the middle of the old element. If two slices are chosen they are placed at either end. Three slices or more are treated as one slice.
  - [TEAPOT] (default): this is the standard slicing as used by MAD9. N.B. This has a maximum of four slices for any one object.

By default all elements are converted to one thin element positioned at the center of the thick element. To get a greater slicing for certain elements use a standard SELECT command with FLAG=MAKETHIN and CLASS, RANGE or PATTERN:

\[
\text{SELECT, FLAG=MAKETHIN, CLASS=class, RANGE=range, SLICE=no of slices;}
\]

The created thin lens sequence has the following properties:

- The created sequence has the same name as the original. The original is therefore no longer available and has to be reloaded if it is needed again.
- The slicer also slices any inserted sequence used in the main sequence. These are also given the same names as the originals.
- Any component changed into a single thin lens has the same name as the original.
- If a component is sliced into more than one slice, the individual slices have the same name as the original component and a suffix .1, .2, etc... and a marker will be placed at the center with the original name of the component.
Hints:

- See the examples for makethin.
- Compare the optics before and after slicing with makethin. Consider to increase the number of slices and rematch after makethin to reach the required accuracy.
- Consider to replace rbend by sbend + thin quads taking into account the edge focusing before slicing with makethin.
- The selection works on the current sequence. Consider to insert a "USE,SEQUENCE=.." before SELECT

Helmut Burkhardt, September 2005
DYNAP: Tunes, Tune Footprints, Smear and Lyapunov Exponent

DYNAP can be called instead of RUN inside a TRACK command:

```
DYNAP,TURNS=real, FASTUNE=logical,LYAPUNOV=real,MAXAPER:={..,..,..,..,..,..},ORBIT=logical;
```

For each previously entered start command, DYNAP tracks two close-by particles over a selected number of turns, from which it obtains the betatron tunes with error, the action smear, and an estimate of the lyapunov exponent. Many such companion particle-pairs can be tracked at the same time, which speeds up the calculation. The smear is defined as
\[
2.0 \frac{(w_{xy_{\text{max}}} - w_{xy_{\text{min}}})}{(w_{xy_{\text{max}}} + w_{xy_{\text{min}}})},
\]
where the \( w_{xy_{\text{min,max}}} \) refer to the minimum and maximum values of the sum of the transverse betatron invariants \( w_x + w_y \) during the tracking. The tunes are computed by using an FFT and either formula (18) or formula (25) of CERN SL/95-84 (AP), depending on whether the number of turns is less-equal or larger than 64.

The arguments have the following meaning:

- **TURNS**: The number of turns to be tracked (default: 64, present maximum: 1024).
- **FASTUNE**: A logical flag. If set, the tunes are computed (default: false).
- **MAXAPER**: If the particle phase-space coordinates exceed certain maximum values, the particle is considered lost. The maximum aperture is a vector of 6 real numbers (default: \((0.1, 0.01, 0.1, 0.01, 1.0, 0.1)\)).
- **LYAPUNOV**: The launch distance between two companion particles added to the \( x \) coordinate (default: \(1.0 \times 10^{-7} \) m).
- **ORBIT**: A logical flag. If set, the flag orbit is true during the tracking and its initialization (default: true). **This flag should be set to be true, if normalized coordinates are to be entered.**

Example:

```
BEAM, PARTICLE=ELECTRON, ENERGY=50, EX=1.E-6, EY=1.E-8, ET=0.002, SIGT=1.E-2;
... USE, PERIOD=FODO;
... TRACK;
START, X=0.0010, Y=0.0017, PT=0.0003;
DYNAP, FASTUNE, TURNS=1024, LYAPUNOV=1.e-7;
ENDTRACK;
...```

The first command defines the beam parameters. It is essential that the longitudinal emittance \( ET \) is set. The command **use** selects the beam line or sequence. The **track** activates the tracking module, **start** enters the starting coordinates (more than one particle can be defined), **dynap** finally tracks two nearby particles.
with an initial distance lyapunov for each start definition over turns revolutions, and endtrack terminates the execution of the tracking module.

The results are stored in the DYNAP and DUNAPTUNE tables, and can be obtained by the commands

```
value,table(dynap,smear);
resp.
value,(dynaptune,tunx),(dynaptune,tuny),(dynaptune,dtune);
```

More generally, all results can be printed to a file, using the commands

```
write,table=dynap,file;
write,table=dynaptune,file;
```

The output file 'lyapunov.data' lists the turn number and phase distance between the two Lyapunov partners, respectively, allowing for visual inspection of chaoticity.
Fully Coupled Motion and Radiation

**EMIT: Equilibrium Emittances**

The command

```
EMIT,DELTAP=real,TOL=tolerance;
```

adjusts the RF frequencies such as to obtain the specified average energy error. More precisely, the revolution frequency $f_0$ is determined for a fictitious particle with constant momentum error

$$\text{DELTAP} = \delta_b = \delta(E) / p, c$$

which travels along the design orbit. The RF frequencies are then set to

$$f_{RF} = h f_0.$$  

If the machine contains at least one RF cavity, and if synchrotron radiation is on, the EMIT command computes the equilibrium emittances and other electron beam parameters using the method of [Chao]. In this calculation the effects of quadrupoles, sextupoles, and octupoles along the closed orbit is also considered. Thin multipoles are used only if they have a fictitious length LRAD different from zero.

If the machine contains no RF cavity, if synchrotron radiation is off or if the longitudinal motion is not stable, it only computes the parameters which are not related to radiation.

The tolerance is for the distinction static/dynamic: if for the eigenvalues of the one-turn matrix, $|e_{val}_5| < \text{tol}$ and $|e_{val}_6| < \text{tol}$, then the longitudinal motion is not considered, otherwise it is. The default for TOL is 1.000001.

In the current implementation, the BEAM values of the emittances and beam sizes are only updated for deltap = zero. Example:

```
RFC: RFCAVITY,HARMON...,VOLT=...;
    BEAM,ENERGY=100.0,RADIATE;
    EMIT,DELTAP=0.01;
```

**Remark:** This module assumes nearly constant lattice functions inside elements. This assumption works for many machines, like LEP [see example], but it fails when the lattice functions largely vary inside single elements. In the later case it is advised to slice the elements as shown in ALBA.

---

**R. Tomás**

Last updated: 148
Error Definitions

This chapter describes the commands which provide error assignment and output of errors assigned to elements. It is possible to assign alignment errors and field errors to single beam elements or to ranges or classes of beam elements.

Elements, classes or ranges of elements are selected by the SELECT command.

ATTENTION: since errors can only be assigned to machine elements, it is necessary to FLATTEN a sequence if it includes other sequences.

Errors can be specified both with a constant or random values. Error definitions consist of four types of statements listed below. They may be entered after having selected a beam line by means of a USE command.

WARNING: any further USE command will destroy the assigned errors. Use the ESAVE option to save and reload errors.

- EALIGN: Define Misalignments
- Field Errors
  - EFCOMP: Components
- EOPTION: Set Error Options
- EPRINT: List Machine Imperfections
- ESAVE: Save Machine Imperfections and read back from file

Werner Herr 18.6.2002
EALIGN: Define Misalignments

Alignment errors are defined by the EALIGN command. The misalignments refer to the local reference system for a perfectly aligned machine. Possible misalignments are displacements along the three coordinate axes, and rotations about the coordinate axes. Alignment errors can be assigned to all beam elements except drift spaces. The effect of misalignments is treated in a linear approximation. A beam position monitor can be given read errors in both horizontal and vertical planes. Monitor errors (MREX, MREY, MSCALX and MSCALY) are ignored for all other elements. Each new EALIGN statement replaces the misalignment errors for all elements in its range, unless EOPTION,ADD=TRUE has been entered.

Alignment errors are defined by the statement

SELECT,FLAG=ERROR,RANGE=range,CLASS=name,PATTERN=string;
EALIGN, DX=real,DY=real,DS=real,
    DPHI=real,DTHETA=real,DPSI=real,
    MREX=real,MREY=real,
    MSCALX=real,MSCALY=real,
    AREX=real,AREY=real;

and elements are now selected by the SELECT command. The attributes are:

DX: The misalignment in the x-direction for the entry of the beam element (default: 0 m). DX>0 displaces the element in the positive x-direction

DY: The misalignment in the y-direction for the entry of the beam element (default: 0 m). DY>0 displaces the element in the positive y-direction

DS: The misalignment in the s-direction for the entry of the beam element (default: 0 m). DS>0 displaces the element in the positive s-direction

DPHI: The rotation around the x-axis.
A positive angle gives a greater x-coordinate for the exit than for the entry (default: 0 rad).

DTHETA: The rotation around the y-axis according to the right hand rule (default: 0 rad).

DPSI: The rotation around the s-axis according to the right hand rule (default: 0 rad).

MREX: The horizontal read error for a monitor. This is ignored if the element is not a monitor
If MREX>0 the reading for x is too high (default: 0 m).

MREY: The vertical read error for a monitor. This is ignored if the element is not a monitor
If MREY>0, the reading for y is too high (default: 0 m).

AREX: The misalignment in the x-direction for the entry of an aperture limit (default: 0 m). AREX>0 displaces the element in the positive x-direction
AREY: The misalignment in the y-direction for the entry of an aperture limit (default: 0 m).
AREY>0 displaces the element in the positive y-direction

MSCALX: The relative horizontal scaling error for a monitor. This is ignored if the element is not a monitor. If MSCALX>0 the reading for x is too high (default: 0). A value of 0.5 implies the actual reading is multiplied by 1.5.

MSCALY: The relative vertical scaling error for a monitor. This is ignored if the element is not a monitor. If MSCALY>0 the reading for y is too high (default: 0). A value of -0.3 implies the actual reading is multiplied by 0.7.

Example:

```
SELECT,FLAG=ERROR,CLASS=MQ;
EALIGN,DX=0.002,DY=0.0004*RANF(),DPHI=0.0002*GAUSS();
```

Assigns alignment errors to all elements of class MQ.

```
SELECT,FLAG=ERROR,PATTERN="QF.*";
EALIGN,DX=0.001*TGAUSS(2.5),DY=0.0001*RANF();
```

Assigns alignment errors to all elements starting with "QF". TGAUSS(2.5) means a Gaussian distribution cut at 2.5 sigma.

![Figure 1: Example of Misplacement in the (x, s)-plane.](image)
Figure 2: Example of Misplacement in the (x, y)-plane.

Figure 3: Example of Misplacement in the (y, s)-plane.

Figure 4: Example of Read Errors in a monitor

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Werner Herr 18.6.2002
Field Errors

Field errors can be entered as relative or absolute errors. Different multipole components can be specified with different kinds of errors (relative or absolute). Relations between absolute and relative field errors are listed below.

In MAD8 two commands were used for that purpose: EFIELD and EFCOMP. Only EFCOMP was implemented in MAD-X since it provides the full functionality of EFIELD and there was no need for duplication.

All field errors are specified as the integrated value \( \int K ds \) of the field components along the magnet axis in m \(^{-1}\). There is no provision to specify a global relative excitation error affecting all field components in a combined function magnet. Such an error may only be entered by defining the same relative error for all field components.

Field errors can be specified for all magnetic elements by the statement

```
SELECT,FLAG=ERROR,RANGE=range,CLASS=name,PATTERN=string;
EFCOMP, ORDER:=integer,RADIUS:=real,
      DKN:={dkn(0),dkn(1),dkn(2),...},
      DKS:={dks(0),dks(1),dks(2),...},
      DKNR:={dknr(0),dknr(1),dknr(2),...},
      DKS:={dksr(0),dksr(1),dksr(2),...};
```

and elements are now selected by the SELECT command. Each new EFCOMP statement replaces the field errors for all elements in its range (s). Any old field errors present in the range are discarded or incremented depending on the setting of [EOPTION,ADD]. EFCOMP defines them in terms of relative or absolute components.

The attributes are:

ORDER: If relative errors are entered for multipoles, this defines the order of the base component to which the relative errors refer. This reference strength \( k_{\text{ref}} \) always refers to the normal component. To use a skew component as the reference the reference radius should be specified as a negative number. The default is zero.

Please note that this implies to specify \( k_0 \) to assign relative field errors to a bending magnet since \( k_0 \) is used for the normalization and NOT the ANGLE.

RADIUS: Radius \( R \) were dknr(i) or dksr(i) are specified for 0...i...20 (default 1 m). This attribute is required if dknr(i) or dksr(i) are specified. If \( R \) is negative, the skew component is used for the reference strength.

DKN(i): Absolute error for the normal multipole strength with \((2i+2)\) poles (default: 0 m \(^{-1}\)).

DKS(i): Absolute error for the skewed multipole strength with \((2i+2)\) poles (default: 0 m \(^{-1}\)).

DKNR(i): Relative error for the normal multipole strength with \((2i+2)\) poles (default: 0 m \(^{-1}\)).
DKSR(i): Relative error for the skewed multipole strength with \((2i+2)\) poles (default: 0 m\(^{-3}\)).

**Time memory effects:**
The relative errors can be corrected for possible time memory effects. A correction term is computed and added to the relative error.
The correction term is parametrized as a 3rd order polynomial in the reference strength \(k_{ref}\) according to:

\[
\Delta = \sum (c_i \cdot k_{i\ ref}) \quad i = 0...3
\]

The coefficients \(c_i\) for the polynomial must be supplied in the command.
Two additional parameters and options are required:

**HYSTER:** if it is set to 1 applies the correction term derived from the reference strength and the coefficients.

**HCOEFFN** and **HCOEFFS:** coefficients (normal and skew) for the computation of the correction term. The 4 coefficients are specified in increasing order, starting with the 0th order. Each group of four coefficients is valid for one order of the field errors. Trailing zeros can be omitted, preceding zeros must be given.

Examples:
Example 1 (assign relative errors to quadrupoles):

```plaintext
select, flag=error, pattern="q.*";
efcomp, order=1, radius=0.010,
dknr:={0,4e-1,1e-3,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0},
dksr:={0,4e-1,1e-3,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0};
```

Example 2 (add time memory effect to relative errors):

```plaintext
select, flag=error, pattern="q.*";
efcomp, order=1, radius=0.020, hyster=1,
hcoffn:={0.000,0.000,0.000,0.000, // coefficients multipole order 0
         0.001,0.000,0.000,0.000, // coefficients multipole order 1
         0.000,0.000,0.002,0.000}, // coefficients multipole order 2
dknr:={0,1e-2,2e-4,5e-5,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0},
dksr:={0,1e-2,2e-4,5e-5,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0};
```

See also: [Random values](#) and [Deferred expressions](#)
EPRINT: List Machine Imperfections

This command prints a table of errors assigned to elements. The range for these elements has to be specified. Field errors are printed as absolute errors, because all relative errors are transformed to the corresponding absolute error at definition time. An error print is requested by the statement

```
SELECT,FLAG=ERROR,RANGE=range,CLASS=name,PATTERN=string;
EPRINT;
```

and elements are now selected by the `SELECT` command. A listing for ALL elements, i.e. not only the selected, can be obtained with the command

```
EPRINT,FULL=TRUE;
```

In that case, the SELECT command has no effect.

Werner Herr 18.6.2002
ESAVE: Save Machine Imperfections and read back from file

Writing errors to a file:
This command saves a table of errors assigned to elements on a file, using a format which can be read in again to obtain the same results. This allows dumping the errors and reloading them after a new USE command. The range for these elements has to be specified. An error save is requested by the statement

```
ESAVE,FILE=string;
```

Example:

```
SELECT,FLAG=ERROR,RANGE=range,CLASS=name,PATTERN=string;
ESAVE,FILE=err.file;
```

and elements selected by the [SELECT] command are saved to the file.

To save the errors of all elements to a file, one can use:

```
SELECT,FLAG=ERROR,FULL;
ESAVE,FILE=err.file;
```

Please note: in case of field errors, the absolute errors are saved and not relative errors.

Setting errors from a table or file:
To assign errors from a file is not a priori straightforward. It may be required to re-assign existing errors after a USE command was executed (which deletes all errors attached to a sequence).

Errors stored in the form of an internal table (errtab) can be directly attached to the appropriate positions in the sequence with the command:

```
SETERR,TABLE=errtab;
```

The table errtab can be generated internally or from an external file (errfile) with the generic command READMYTABLE.

The command sequence:

```
READMYTABLE,FILE=errfile,table=errtab;
SETERR,TABLE=errtab;
```

reads the file errfile into the table errtab and the command SETERR attaches the errors to the elements in the active sequence.

The file errfile can be produced by a preceding ESAVE command or any other utility. It should follow the format of a file generated with ESAVE (see example program).

Please note:
1. To assign correctly the errors from the file to the elements in the sequence, all elements must have individual names, otherwise an identification is not possible. Elements in the file not identified in the active sequence are ignored.
2. Errors are assigned to ALL elements found in the file and the FLAG=ERROR is set. Therefore the number of elements selected corresponding to a command like:
   SELECT, FLAG=ERROR,....;
   can be different after the execution of SETERR.
The Intra-Beam Scattering Module (IBS)

As emphasized by its name, the Intra-Beam Scattering module (IBS) computes the contribution to emittance growth rates due to Coulomb scattering of particles within relativistic beams. The formalism used in this module is that derived by J.D. Bjorken and S.K. Mtingwa [Bjorken and Mtingwa] in 1982. Contrary to other IBS-routines, the Bjorken-Mtingwa formalism takes into account the variation of the lattice parameters around the machine, rather than using average values. Consequently, the knowledge of the optical functions of the machine is required. In MAD-X, this is achieved with the ‘‘twiss’’ command.

It is well known that the intra-beam scattering growth times behave like:

$$\frac{1}{T_i} = C_i \times \frac{N}{\gamma \epsilon_x \epsilon_y \epsilon_z} \ (i = x, y, z)$$

where $C_i$ accounts for some constants and the integrals for the scattering functions, $N$ is the number of particles in the bunch, $\gamma$ is the relativistic factor and $\epsilon_i$ are the normalized emittances in the horizontal, vertical and longitudinal plane respectively. It thus follows that the second required input is a description of the beam parameters, which is achieved via the ‘‘beam’’ command (see below).

Once the optical functions and the beam parameters have been defined, the evaluation of the scattering growth times follows via the ‘‘ibs’’ command. The logical follow-up of the MAD-X commands is illustrated in the two examples provided with the IBS-module.

Input of the beam parameters

This section briefly describes the parameters which have to be present in the ‘‘beam’’ command in order to run the IBS-module:

Type of particle

The parameter ‘‘particle=’’ is mandatory. It can take one of the following three values: proton, electron or ion. For proton and electron, the parameter ‘‘particle’’ is the only one to be defined. In case ion is used, two additional parameters have to be defined, namely ‘‘mass=’’, which is typically the number of nucleons for the corresponding ion
multiplied by \( \text{nmass} \) the unified atomic mass unit [0.931494013 GeV/(c**2)] , and "charge=" for the number of charges.

**The energy**

The definition of the energy (total, kinetic, total energy of the ions or energy per nucleon) is a difficult one. In the present approach, the energy is the total energy of the particle. For ions, the expected input is the proton equivalent energy, i.e. the total energy a proton would have when circulating in the defined machine. As an illustration, in the LHC, protons will be injected with an energy of 450 GeV. Consequently, to evaluate the growth times for Lead ions at injection in the LHC, one has to input \( \text{energy}=450\times\text{charge} \). Therefore the above example of Lead at the LHC injection energy may look as follows in the MAD-X input language:

\[
\text{nucleon}=208; \text{charge}=82; \\
\text{beam,particle}=\text{ion,charge}=\text{charge,energy}=450\times\text{charge,mass}=\text{nucleon}\times\text{nmass};
\]

An important check for the correctness of the input is the printed value of the relativistic factor \( \gamma \). The latter should correspond to:

\[
\gamma_{\text{ion}} = \gamma_{\text{proton}} \times \frac{\text{charge}}{\text{nucleon}}
\]

**The number of particles**

The number of particles (or number of ions) is defined with the parameter "npart=".

**Beam sizes - Emittances**

This part of the input is used to define the normalized emittances (horizontal, vertical and longitudinal). The required parameters are the physical transverse emittances (ex= and ey= [m]), the bunch length (sigt= [m]) and the relative energy spread (sige=).

**File Attribute**

If FILE="file_name" appears MAD-X produces a table and writes on a file for each element of the machine: ELEMENT NAME, Position S [m], DELS [m] (Length Difference of consecutive Elements in the Table), TLI (Longitudinal growth time), TXI (Horizontal growth time), TYI (Vertical growth time).
Examples

The two examples provided for the module Intra-Beam Scattering illustrate the commands required to run the module. The two examples have been selected such as to highlight the differences between a computation for protons and that for ions. Both examples compute the IBS growth times at injection into the LHC. The examples are located at http://frs.home.cern.ch/frs/Xdoc/mad-X.html.

Frank Schmidt 2003-05-23
Matching Module

Before a match operation at least one sequence must be selected by means of a USE command. Matching is then initiated by the MATCH command. The matching module can act on more than one sequence simultaneously by specifying more than one sequence when INITIATING the matching mode. From this command to the corresponding ENDMATCH command MAD accepts the matching commands listed below. For a mathematical description of the minimisation procedures see [James]. In particular one may do the following:

- Define the sequence(s) the matching module will work on
- Set initial conditions for transfer line matching
- Define constraints
- Define the parameters to be varied
- Match by different methods.

The matching commands are described in detail below. Some other commands can also be issued during matching.

- **Enter and Leave Matching Mode**
  - MATCH: Initiating the Matching Mode
  - ENDMATCH: Leave Matching Mode

- **Define Variable Parameter**
  - VARY: Set Parameter to Vary

- **Constraint**
  - CONSTRAINT: Simple Constraint
  - CONSTRAINT: User Defined Variables
  - WEIGHT: Matching Weights
  - GLOBAL: Global Constraints
  - GWEIGHT: Weights for Global Constraints

- **Matching Methods**
  - LMDIF: Fast Gradient Minimisation
  - MIGRAD: Gradient Minimisation
  - SIMPLEX: Simplex Minimisation
  - JACOBIAN: Newton Minimisation

- **Expression Matching with USE_MACRO**
Oliver Brüning June, 2002; Riccardo de Maria February, 2006.
Before matching at least one \texttt{SEQUENCE} must be selected by means of a \texttt{USE} command. The matching module can act on more than one sequence simultaneously by specifying more than one sequence when \texttt{INITIATING} the matching mode:

### Initiating the Matching Module

The ’match’ command can be either used for matching a periodic cell or for matching an insertion to another part of the machine. Both matching modes are initiated by the MATCH command.

- **Cell matching:**

  In the first mode the matching routine is initiated only with one attribute specifying the sequence(s) the matching module will work on. In this matching mode the periodicity of the optics functions is enforced at the beginning and end of the selected range.

  \texttt{MATCH, SEQUENCE=’name1’, ’name2’,…,nema-n’;}

- **Insertion matching:**

  In the second mode, called insertion matching, the matching routine is initiated with two attributes: one specifying the sequence(s) the matching module will work on and one specifying the initial conditions of the optic functions for each sequence. In this case the initial values are assumed as exact.

  - **Specification of Initial Values:** The initial values of the optical functions for the insertion matching can either be specified in form of a \texttt{SAVEBETA} command or by explicitly stating the individual optic function values after the ’MATCH’ command. The two options can be entered as

    \texttt{MATCH,sequence=’name1’,’name2’,…,’name-n’,BETA0=’beta01’,’beta02’,…,’beta0n’;}

    or

    \texttt{MATCH,SEQUENCE=’sequence-name’, BETX=real,ALFX=real,MUX=real, BETY=real,ALFY=real,MUY=real, X=real,FX=real,Y=real,PY=real, DX=real,DY=real,DPX=real,DPY=real, DELTAP=real;}

> **Examples:**
- **Example 1:**

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This matches the sequence 'INSERT' with initial conditions to a new phase advance. The initial conditions are given by the periodic solution for the sequence CELL1.

- Example 2:

This matches the beam line 'INSERT' with periodic boundary conditions to a new phase advance.

The initial conditions can also be transmitted by a combination of a `SAVEBETA` command and explicit optic function specifications:

```
USE,CELL1;
SAVEBETA, LABEL=bini, PLACE=#E;
TWISS, SEQUENCE=CELL1;
USE,PERIOD=LINE1;
MATCH, SEQUENCE=LINE1, BETA0=bini, MUX=1.234, MUY=4.567;
```

This example transmits all values of the SAVEBETA array 'bini' as initial values to the MATCH command and overrides the initial phase values by the given values.

An additional [CONSTRAINT] may be imposed in other places, i.e. intermediate or end values of the optics functions at the transition point.

- More than one active sequence:

The matching module can act on more than one sequence simultaneously by specifying more than one sequence after the MATCH command:

```
MATCH, SEQUENCE=LINE1, CELL1, BETA0=bini1, bini2;
```

This example initiates the matching mode for the 'LINE1' and the 'CELL1' sequence. The [Twiss module] function of the two sequences are calculated with fixed initial conditions. The SAVEBETA array 'bini1' is used for calculating the optics functions of sequence 'LINE1' and the SAVEBETA array 'bini2' for calculating the optics functions of sequence 'CELL1'. Without the initial conditions the matching module will work in the [CELL] mode.

- Special speed flag:

The "slow" attribute enforces the old and slow matching procedure. Recently a number of parameter, like "RE56", have been added to list of matchable parameters. Nevertheless, some parameters might only be available when using the "slow" attribute.
Further attributes of the TWISS statements are:

- **RMATRIX**: If this flag is used the one-turn map at the location of every element is calculated and prepared for storage in the TWISS table. Target values for the matrix elements at certain positions in the sequence can be specified with the help of the `CONSTRAINT` command and the keywords: `RE, RE11...RE16...RE61...RE66`, where `REij` refers to the “ij” matrix component.

  > Examples:

  ```bash
  Example 1:
  MATCH,RMATRIX,SEQUENCE='name',BETA0='beta-block-name';
  CONSTRAINT,SEQUENCE=insert,RANGE=#e,RE11=-2.808058321,re22=2.748111197;
  VARY,NAME=kqf,STEP=1.0e-6;
  VARY,NAME=kqd,STEP=1.0e-6;
  
  This matches the sequence 'name' with initial conditions to new values for the matrix elements 'RE11' and 'RE22' by varying the strength of the main quadrupole circuits.
  ```

- **CHROM**: If this flag is used the chromatic functions at the location of every element are calculated and prepared for storage in the TWISS table. Target values for the chromatic functions at certain positions in the sequence can be specified with the help of the `CONSTRAINT` command and the keywords `WX, PHIX, WY, PHIY...`.

**Leave Matching Mode**

The `ENDMATCH` command terminates the matching section and deletes all tables related to the matching run.

```bash
ENDMATCH;
```

---

Oliver Brüning, October, 2003; Riccardo de Maria, January, 2008.
References


7. P. Bramham and H. Henke. private communication and LEP Note LEP-70/107, CERN.


F. James. *MINUIT, A package of programs to minimise a function of n variables, compute the covariance matrix, and find the true errors*. program library code D507, CERN, 1978.


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MAD Home Page. [MAD-8 User Guide](#)

[fc] *January 23, 1997*
Define Variable Parameter

**VARY: Define Variable Parameter**

A parameter to be varied is specified by the command

VARY, NAME=variable, STEP=real, LOWER=real, UPPER=real;

It has four attributes:

- **NAME**: The name of the parameter or attribute to be varied.
- **STEP**: The approximate initial step size for varying the parameter. If the step is not entered, MAD tries to find a reasonable step, but this may not always work.
- **LOWER**: Lower limit for the parameter (optional).
- **UPPER**: Upper limit for the parameter (optional).
- **SLOPE**: allowed change rate (optional, available only using JACOBIAN routine). Limit the parameter to increase (SLOPE=1) decrease (SLOPE=-1) only.
- **OPT**: optimal value for the parameter (optional, available only using JACOBIAN routine).

Examples:

VARY, NAME=PAR1, STEP=1.0E-4; ! vary global parameter PAR1
VARY, NAME=QL11->K1, STEP=1.0E-6; ! vary attribute K1 of the QL11
VARY, NAME=Q15->K1, STEP=0.0001, LOWER=0.0, UPPER=0.08; ! vary with limits

If the upper limit is smaller than the lower limit, the two limits are interchanged. If the current value is outside the range defined by the limits, it is brought back to range. If a parameter comes outside the limits during the matching process the matching module resets the parameter to a value inside the limits and informs the user with a message. If such a 'rescaling' occurs more than 20 times the matching process terminates. The user should either eliminate the corresponding parameters from the list of varied parameters or change the corresponding upper and lower limits before restarting the matching process. After a matching operation all varied attributes retain their value after the last successful matching iteration. Using JACOBIAN routine, STRATEGY=3, in case the number of parameters is greater the the number of constraint, if a parameter comes outside the limits, it is excluded automatically from the set of variables and a new solution is searched.

Constraints

CONSTRAINT: Simple Constraint

Simple constraints are imposed by the CONSTRAINT command. The CONSTRAINT command has three attributes:

- the SEQUENCE entry specifies the sequence for which the constraint applies.
- the RANGE entry specifies the position where the constraint must be satisfied. The RANGE can either be the name of a single element in the sequence or a range between two elements. In the later case the two element names must be separated by a ‘/’: RANGE=nam1/name2
- the optics functions to be constrained

The optic functions can be constraint in four different ways:

- lower limits: 'BETX > value' -> type1
- upper limits: 'BETX < value' -> type2
- lower and upper limits: 'BETX < value1,BETX > value2' -> type3
- target value: BETX=value -> type 4

In case one element is affected my more than one constraint command the last CONSTRAINT will be chosen. For example, one can specify the maximum acceptable beta function over a range of the sequence and specify the target beta function for one element that lies inside this range. In this case one must first specify the constraint that affects the whole range and then the constraint for the single element. This way the constraint of the target value overrides the previous constraint on the upper limit for the selected element. For example, the following constraint statements limit the maximum horizontal beta function between ‘marker1’ and ‘marker2’ to 200 meter and require a horizontal beta function of 180 meter at element ‘name1’:

```
CONSTRAINT,SEQUENCE=sequence-name,RANGE='marker1'/'marker2',BETX<200.0;
CONSTRAINT,SEQUENCE=sequence-name,RANGE='name1'/'marker2',BETX=100.0;
```

When the two constraint statements are interchanged the horizontal beta function at element ‘name1’ will only be limited to less than 200 meter and NOT constrained to 100 meter!

The CONSTRAINTS can either be specified with explicit values for the constraints of the optic functions or via a pre-calculated SAVEBETA module. The first options has the form:

```
CONSTRAINT,SEQUENCE=sequence-name,RANGE=position,BETX=real,ALFX=real,MUX=real,
BETY=real,ALFY=real,MUY=real,
X=real,FX=real,Y=real,FY=real,
DX=real,DY=real,DPX=real,DPY=real;
```
Here all **linear lattice functions** (BETX, BETY, ALFX, ALFY, MUX, MUY, DX, DY, DPX, DPY) or **chromatic lattice functions** (WX, XY, PHIX, PHIY, DMUX, DUMY, DDX, DDY, DDPX, DDPY) are constrained at the selected range to the corresponding values.

The second form of the CONSTRAINT command has the form

```
CONSTRAINT, SEQUENCE=sequence-name, RANGE=position, BETA0=beta0-name, MUX=real, MUY=real
```

Here all of (BETX, BETY, ALFX, ALFY, MUX, MUY, DX, DY, DPX, DPY) are constrained in the selected points to the corresponding values of a pre-calculated SAVEBETA module. In the above example the phases (MUX, MUY) are overridden by the numerical values specified via 'MUX=real' and 'MUY=real'. Normally ‘RANGE’ refers to a single position.

**User Defined Matching Constraints**

In addition to the nominal TWISS variables the user can define a limited set of 'user-defined' variables in the constraint statement. This allows, for example, the matching of the normalized dispersion or the mechanical aperture. The MATCH module allows four user defined variables called: mvar1, mvar2, mvar3 and mvar4. The variables can be defined according to the general variable declaration rules of deferred expressions. For example, in order to match the normalized dispersion at a certain location in the sequence one would first define a variable:

```
mvar1 := table(twiss,dx)/sqrt(table(twiss,betx));
```

After that the user has to select the variable for output in the TWISS statement (see TWISS module and SELECT for more details on the TWISS module and SELECTION statements):

```
select, flag=twiss, clear;
select, flag=twiss, column=keyword,, name, s, betx, dx, mvar1;
twiss, sequence=’sequence-name’, file=twiss.file;
```

The variable can now be referenced like any other TWISS variable in the constraint command:

```
constraint, sequence=’sequence-name’, range=’location’, mvar1=’target-value’;
```

**Matching Weights**

The matching procedures try to fulfil the constraints in a least square sense. A penalty function is constructed which is the sum of the squares of all errors, each multiplied by the specified weight. The larger the weight, the more important a constraint becomes. The weights are taken from a table of current values. These are initially set to weight default values and may be reset at any time to different values. Values set in this way remain valid until changed again. The command

```
WEIGHT, BETX=real, ALFX=real, MUX=real,
       BETY=real, ALFY=real, MUY=real,
       X=real, PX=real, Y=real, PY=real,
       DX=real, DPX=real, DY=real, DPY=real;
```

changes the weights for subsequent constraints. The weights are entered with the same name as the linear lattice functions or orbit coordinate to which the weight applies. Frequently the matching weights serve to
select a restricted set of functions to be matched.

### Default Matching Weights

<table>
<thead>
<tr>
<th>name</th>
<th>weight</th>
<th>name</th>
<th>weight</th>
<th>name</th>
<th>weight</th>
<th>name</th>
<th>weight</th>
<th>name</th>
<th>weight</th>
<th>name</th>
<th>weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>BETX</td>
<td>1.0</td>
<td>ALFX</td>
<td>10.0</td>
<td>MUX</td>
<td>10.0</td>
<td>BETY</td>
<td>1.0</td>
<td>ALFY</td>
<td>10.0</td>
<td>MUY</td>
<td>10.0</td>
</tr>
<tr>
<td>X</td>
<td>10.0</td>
<td>PX</td>
<td>100.0</td>
<td>Y</td>
<td>10.0</td>
<td>PY</td>
<td>100.0</td>
<td>T</td>
<td>0.0</td>
<td>PT</td>
<td>0.0</td>
</tr>
<tr>
<td>DX</td>
<td>10.0</td>
<td>DPX</td>
<td>100.0</td>
<td>DY</td>
<td>10.0</td>
<td>DPY</td>
<td>100.0</td>
<td></td>
<td></td>
<td></td>
<td>-</td>
</tr>
<tr>
<td>WX</td>
<td>10.0</td>
<td>PHIX</td>
<td>10.0</td>
<td>DMUX</td>
<td>100.0</td>
<td>WY</td>
<td>10.0</td>
<td>PHIY</td>
<td>10.0</td>
<td>DMUY</td>
<td>100.0</td>
</tr>
<tr>
<td>DDX</td>
<td>10.0</td>
<td>DDPX</td>
<td>100.0</td>
<td>DDY</td>
<td>10.0</td>
<td>DDPY</td>
<td>100.0</td>
<td></td>
<td></td>
<td></td>
<td>-</td>
</tr>
<tr>
<td>MVAR1</td>
<td>10.0</td>
<td>MVAR2</td>
<td>10.0</td>
<td>MVAR3</td>
<td>10.0</td>
<td>MVAR4</td>
<td>10.0</td>
<td></td>
<td></td>
<td></td>
<td>-</td>
</tr>
</tbody>
</table>

### GLOBAL: Global Matching Constraints

In addition to conventional matching constraints that specify the optics functions at a certain position in the sequence the user can also constrain global optics parameters such as, for example, the overall machine tune and the machine chromaticity. Such global optics parameters can be constrain via the GLOBAL command, having the following syntax:

```
GLOBAL,SEQUENCE=sequence-name,Q1=real,Q2=real,dQ1=real,dQ2=real,&
    ddQ1=real,ddQ2=real;
```

Global matching weights can be (re)set by the new GWEIGHT command, having attributes identical to those of GLOBAL. All the attributes are optional and have the following meaning:

- **Q1, Q2, dQ1, dQ2**
  - enable a correction of tunes and chromaticities in presence of magnetic imperfections or misalignments.

- **ddQ1, ddQ2**
  - enable a correction of nonlinear chromaticities

*Oliver Brüning*  *June, 2002*
Matching Methods

MADX currently supports four different matching algorithms:

- **LMDIF: Fast Gradient Minimisation**
  
The LMDIF command minimises the sum of squares of the constraint functions using their numerical derivatives:

  \[
  \text{LMDIF, CALLS=integer, TOLERANCE=real;}
  \]

  It is the fastest minimisation method available in MAD. The command has two attributes:
  - **CALLS**: The maximum number of calls to the penalty function (default: 1000).
  - **TOLERANCE**: The desired tolerance for the minimum (default: \(10^{-6}\)).

  Example:

  \[
  \text{LMDIF, CALLS=2000, TOLERANCE=1.0E-8;}
  \]

- **MIGRAD: Gradient Minimisation**
  
The MIGRAD command minimises the penalty function using the numerical derivatives of the sum of squares:

  \[
  \text{MIGRAD, CALLS=integer, TOLERANCE=real, STRATEGY=1;}
  \]

  The command has three attributes:
  - **CALLS**: The maximum number of calls to the penalty function (default: 1000).
  - **TOLERANCE**: The desired tolerance for the minimum (default: \(10^{-6}\)).
  - **STRATEGY**: A code for the strategy to be used (default: 1). Details are given in [James].

  Example:

  \[
  \text{MIGRAD, CALLS=2000, TOLERANCE=1.0E-8;}
  \]
**SIMPLEX: Simplex Minimisation**

The SIMPLEX command minimises the penalty function by the simplex method:

```
SIMPLEX,CALLS=integer,TOLERANCE=real;
```

Details are given in [James]. The command has two attributes:
- CALLS: The maximum number of calls to the penalty function (default: 1000).
- TOLERANCE: The desired tolerance for the minimum (default: 10**(-6)).

Example:
```
SIMPLEX,CALLS=2000,TOLERANCE=1.0E-8;
```

**JACOBIAN: Newton Minimisation**

The JACOBIAN command minimises the penalty function calculating the Jacobian and solving the linear problem. A QR or LQ decomposition is performed when the system is over or under-determined. Before starting the matching routine two optional transformations (COOL and RANDOM) are performed.

```
JACOBIAN,CALLS=integer,TOLERANCE=real,REPEAT=integer,STRATEGY=integer,COOL=real,BALANCE=real, random=real;
```

The command has the attributes:
- CALLS: The maximum number of calls to the penalty function (default: 30).
- TOLERANCE: The desired tolerance for the minimum (default: 10**(-6)).
- REPEAT: The number of call of the JACOBIAN routine (default: 1).
- BISEC: Selects the maximum number of iteration used to determin the step length which reduces the penalty function during the main iteration. A large number (i.e. 6) reduce the probability to diverge from the solution, but increase the one for being trapped in a local minum.
- STRATEGY: A code for the strategy to be used (default: 3). If STRATEGY=1 the routine resets the values of the variables which exceeds the limits. If STRATEGY=2 the routine print the Jacobian and exit without matching. If STRATEGY=3 the routine disables the variables which exceeds the limits keeping however the number of variables greater or equal to the number of the constraints.
- COOL, BALANCE: The factors which specify the following transformation:

```
if "balance" >=0
    newval=(1-cool)*oldval+cool* ( (1-balance)*maxval+balance*minval )
else
    newval=(1-cool)*oldval+cool* optval
```

where newval is the new value after the transformation, oldval is the previous value, maxval, minval, optval are the maximum value, minimum value, optimal value of the variable specified in the VARY command.
- RANDOM: The factors which specify the following transformation:
newval = (1 + random * rand()) * oldval

where newval is the new value after the transformation, oldval is the previous value, 
rand() is a stochastic variable with a uniform (-0.5,0.5) distribution.

Example:

JACOBIAN, CALLS=20, TOLERANCE=1.0E-8, STRATEGY=3, COOL=0.1, BALANCE=0.5, RANDOM=0.01;

Introduction

It is possible to match user defined expressions with the USE_MACRO keyword. The general input structure for a match command is the following:

MATCH, USE_MACRO;
... VARY statements ...
USE_MACRO, NAME=macro1;
or
macro1: MACRO={ ... madx statements};
CONSTRAINT, expr="lhs1 < | = | > rhs1";
CONSTRAINT, expr="lhs2 < | = | > rhs2";
... CONSTRAINT statements ...
MACRO 2 definition
... CONSTRAINT statements ...
MACRO n definition
... CONSTRAINT statements ...
... METHODS statements ...
ENDMATCH;

The algorithm for evaluating the penalty function is the following:

- execute the first macro,
- evaluate and compute the difference between the lhs and the rhs the first set of expressions,
- in case of other macros, evaluates in order the macro and the expressions
- the set of differences are minimized by the selected method using the variables defined in the VARY statements.

Initiating the Matching Module with USE_MACRO

With:

MATCH, USE_MACRO;

the 'match' command can be used for matching any expression which can be defined through expression. It requires a slightly different syntax.

VARY statements

In the USE_MACRO mode the vary statement follows the same rules of the other modes explained in the section Define Variable Parameter.
Macro definitions

The macro to be used in the matching routine can be defined in two ways:

- using USE_MACRO statement:

  USE_MACRO, NAME=macro1;

  defining a new macro on the fly using the usual syntax for macros.

After a macro definition is necessary to define a set of constraints exclusively with the following syntax:

```plaintext
CONSTRAINT, expr=  "lhs = rhs";
```

or

```plaintext
CONSTRAINT, expr=  "lhs < rhs";
```

or

```plaintext
CONSTRAINT, expr=  "lhs > rhs";
```

where "lhs" and "rhs" are well defined MadX expressions. Other set of macro and constraints can be defined afterwards.

Examples

The following example the USE_MACRO mode can emulate a matching script which uses the normal syntax.

Normal syntax:

```plaintext
MATCH,SEQUENCE=LHCB1,LHCB2;
VARY, NAME=KSF.B1, STEP=0.00001;
VARY, NAME=KSD.B1, STEP=0.00001;
VARY, NAME=KSF.B2, STEP=0.00001;
VARY, NAME=KSD.B2, STEP=0.00001;
GLOBAL,SEQUENCE=LHCB1,DQ1=QPRIME;
GLOBAL,SEQUENCE=LHCB1,DQ2=QPRIME;
GLOBAL,SEQUENCE=LHCB2,DQ1=QPRIME;
GLOBAL,SEQUENCE=LHCB2,DQ2=QPRIME;
LMDIF, CALLS=10, TOLERANCE=1.0E-21;
ENDMATCH;
```

USE_MACRO syntax:

```plaintext
MATCH,USE_MACRO;
VARY, NAME=KSF.B1, STEP=0.00001;
VARY, NAME=KSD.B1, STEP=0.00001;
VARY, NAME=KSF.B2, STEP=0.00001;
VARY, NAME=KSD.B2, STEP=0.00001;
M1: MACRO={ TWISS,SEQUENCE=LHCB1; };
CONSTRAINT, EXPR= "TABLE(SUMM,DQ1)=QPRIME";
CONSTRAINT, EXPR= "TABLE(SUMM,DQ2)=QPRIME";
M2: MACRO={ TWISS,SEQUENCE=LHCB2; };
```
CONSTRAINT, EXPR= "TABLE(SUMM,DQ1)=QPRIME";
CONSTRAINT, EXPR= "TABLE(SUMM,DQ2)=QPRIME";
LMDIF, CALLS=10, TOLERANCE=1.0E-21;
ENDMATCH;

Oliver Brüning  October, 2003; Riccardo de Maria  February, 2006.
Matching Examples

All matching examples and the related files for executing the MADX sample jobs can be found on the 'afs' directory under:

/afs/cern.ch/group/si/slap/share/mad-X/test_suite/match/V3.02.03.

•

Simple Periodic Cell
Match a simple cell to given phase advances:

FIVE-CELL

•

Simple Periodic Cell
Match the matrix elements of the linear transfer matrix at the end of a sequence 5 periodic cells:

RMATRIX

•

Transfer line with initial conditions
Match a sequence of 5 periodic cells with initial conditions to given beta-functions at the end of the sequence:

Transfer line

•

Global tune matching in a sequence of 5 periodic cells
Match the global tune of a sequence of 5 periodic cells:

Global tune
Global tune matching for the LHC
Match the global tune for beam1 of the LHC:

Global chromaticity matching for the LHC
Match the global chromaticity for beam1 of the LHC:

Global chromaticity matching for both beams of the LHC
Match the global chromaticity for beam1 and beam2 of the LHC:

IR8 insertion matching for beam1 of the LHC
Match the insertion IR8 with initial conditions to given values of the optics functions at the IP and the end of the insertion:

IR8 insertion matching for beam1 of the LHC with upper limits on the optics functions
Match the insertion IR8 with initial conditions to given values of the optics functions at the IP and the end of the insertion while limiting the maximum acceptable beta functions over the whole insertion:

Simultaneous orbit matching at IP8 for beam1 and beam2 of the LHC
Match simultaneously the orbit of beam1 and beam of the LHC at IP8 with initial conditions to the same given values at the IP:
Orbit matching at IP8 for beam1 and beam2 of the LHC

- **IR8 beta squeeze for beam1 using JACOBIAN matching routine**
  
  Try to find a beta squeeze for IR8 starting from 10 meters.

- **Matching first and second order chromaticity of the LHC using USE_MACRO option.**
  
  Match simultaneously the first and second order chromaticity by defining macros which compute them using the TWISS command or PTC.

- **Matching s position using VLENGTH flag.**
  
  match the positions of elements and the total sequence length for a simple sample sequence.

- **Matching s position using USE_MACRO.**
  
  match the positions of elements and the total sequence length for a simple sample sequence using USE_MACRO.

*Oliver Brüning* June, 2002; *Riccardo de Maria* August, 2007.
Orbit Correction

This chapter describes the commands which can be used to correct the closed orbit or a trajectory. The distorted orbit is taken from an internal or external TFS table.

Purpose of this Module:
The purpose of this orbit module is to provide some basic tools to assess the performance of an orbit correction system of a machine in the design phase.
Although some interface is available, it cannot and does not provide the full functionality expected from a dedicated online orbit correction and steering program.

- **CORRECT: Correction commands and parameters**
- **Activate/Deactivate correctors and monitors**
- **READ/WRITE corrector settings**
- **COPTION: Global Correction Options**

*Werner Herr 22.10.2008*
CORRECT: Orbit Correction

The CORRECT statement makes a complete closed orbit or trajectory correction using the computed values at the monitors from the Twiss table. The CORRECT command has the following format (not all possible options included, some options are valid only for special algorithms):

```plaintext
CORRECT, ORBIT=myorbit,MODEL=mymodel,TARGET=mytarget,
   FLAG=ring,MODE=lsq,
   MONERROR=integer,MONON=real,MONSCALE=real,
   PLANE=x,COND=integer,RESOUT=integer,
   CLIST=file1,MLIST=file2;
```

The command CORRECT is set up with defaults which should allow a reasonable correction for most cases with a minimum of required options (see Example 1 below). The orbit correction must always be preceded by TWISS commands which generate Twiss tables. The most recent Twiss table is assumed to contain the optical parameters and the distorted orbits. The options used in the CORRECT command are:

- **FLAG**: FLAG can be "ring" or "line", either a circular machine or a trajectory is corrected. Default flag is "ring".
- **MODE**: MODE defines the method to be used for corrections. Available modes are LSQ, MICADO and SVD. The first performs a least squares minimization using all available correctors. The mode SVD uses a Singular Value Decomposition to compute a correction using all available correctors. The latter can also be used to condition the response matrix for the modes LSQ or MICADO (using COND=1). It is highly recommended to precede a LSQ correction by a SVD conditioning (set COND=1). The mode MICADO is a "best kick" algorithm. Naive use or using it with a large number of correctors (see option NCORR) can give unexpected results. To avoid the creation of local bumps, it is recommended to precede a MICADO correction by a SVD conditioning (set COND=1). Default mode is MICADO.
- **PLANE**: If this attribute is x, only the horizontal correction is made; if it is y, only the vertical correction is made. (This differs from the MAD8 implementation). Default plane is horizontal.
- **COND**: When COND is 1, a Singular Value Decomposition is performed and the response matrix CONDITIONed to avoid linearly dependent correctors. This can be used to avoid creation of artificial bumps during a LSQ or MICADO correction (requires some computing time). Please note: this option is not robust since it depends on parameters which control the determination of singular values and redundant correctors. These can be set with the commands SNGVAL and SNGCUT. Both parameters depend on the machine and may need adjustment. Default values are adjusted to large machines and "reasonable" performance for smaller machines.
- **NCORR**: Only used by the MICADO algorithm. Defines the number of correctors to be used, unless set to 0 in which case all available correctors are used. Default is 0 (all available correctors).
- **SNGVAL**: Used to set the threshold for finding singular values with the COND command. (Hint: smaller number finds fewer singular values). Use with care!
Default is 2.0
- **SNGCUT**: Used to set the threshold for finding redundant correctors with the COND command. (Hint: larger number finds fewer redundant correctors).
  Use with extreme care!
  Default is 50.0

- **MONERROR**: When MONERROR is 1, the alignment errors on monitors assigned by [EALIGN] MREX and MREY are taken into account, otherwise they are ignored.
  Default is 0.

- **MONSCALE**: When MONSCALE is 1, the scaling errors on monitors assigned by [EALIGN] MSCALX and MSCALY are taken into account, otherwise they are ignored.
  Default is 0.

- **MONON**: MONON takes a real number between 0.0 and 1.0. It determines the number of available monitors. If the command is given, each monitor is considered valid with a probability MONON. In the average a fraction (1.0 - MONON) of the monitors will be disabled for the correction, i.e. they are considered not existing. This allows to study the effect of missing monitors.
  Default is 1.0 (100%).

- **CORRLIM**: A limit on the maximum corrector strength can be given and a WARNING is issued if it is exceeded by one or more correctors. Please note: the strengths computed by the correction algorithms are NOT limited, only a warning is printed!
  Default is 1.0 mrad.

Normally the last active table provides the orbit to be corrected and the model for the correction. This can be overwritten by the appropriate options. Optionally, these tables can be given names like in: TWISS, TABLE=name; (see documentation on TWISS command). To use these named tables, one of the following optional parameters must be used:

- **ORBIT**: When this parameter is given, the orbit to be corrected is taken from a named table. The default is the last (named or unnamed) Twiss table.

- **MODEL**: When this parameter is given, the model for the correction is taken from a named Twiss table. The default is the last (named or unnamed) Twiss table.

- **TARGET**: When this parameter is given, the correction is made to a named target orbit, pre-computed with a TWISS command. Default is correction to the zero orbit.
  Please note: the target orbit is normally computed by MAD within the same job.

Two attributes affect the printing of tables and results:

- **CLIST=** file: Corrector settings (in units of rad) before and after correction printed to file
- **MLIST=** file: Monitor readings (in units of m) before and after correction printed to file
- **RESOUT**: This command outputs the results for all monitors and all correctors in a computer readable format if its integer argument is larger than 0. The argument is added to the output. Useful to analyze runs with loops to produce large statistics.

  **ATTENTION**: May produce gigantic outputs for large machines.

- **TWISSUM**: If the argument of twissum is larger than 0, it prints maximum orbit and r.m.s. for both planes taken from the Twiss summary table in computer readable form. Allows to analyze orbits etc. at elements that are not monitors or correctors. The argument is added to the output. Only for output: no correction is made, all other commands are ignored.
Obsolete commands or options:

ITERATE, ITERMAX /* Done with loop feature in MAD commands */
THREADER, THRTOL, WROBIT /* Not part of orbit correction module */
MLIST, M2LIST /* Replaced by MLIST */
C1LIST, C2LIST /* Replaced by CLIST */
GETORBIT, PUTORBIT /* Replaced by generic TFS access */
GETKICK, PUTKICK /* Replaced by generic TFS access */

EXAMPLES (for complete MAD input files see section on examples):
Example 1 (correct orbit in horizontal plane, taken from most recent Twiss table, using default algorithm (MICADO)):
CORRECT,PLANE=x;
Example 2 (no correction, only output of Twiss summary):
CORRECT,TWISSUM=1;
Example 3 (correct orbit in horizontal plane, corrector and monitor output on table):
CORRECT,PLANE=x,MODE=lsq,CLIST=corr.out,MLIST=mon.out;
Example 4 (correct orbit in horizontal plane, use alignment and scaling errors, 15% of orbit correctors faulty): CORRECT,PLANE=x,MONERROR=1,MONSCALE=1,MONON=0.85;

Last updated: 22.10.2008
Werner Herr 14.06.2006
Activate/Deactivate Correctors or Monitors

To provide more flexibility with orbit correction two commands are provided:

**USEMONITOR**, STATUS=flag,
[,SEQUENCE=sequence][,RANGE=range][,CLASS=class][,PATTERN=regex]

**USEKICK**, STATUS=flag,
[,SEQUENCE=sequence][,RANGE=range][,CLASS=class][,PATTERN=regex]

The purpose of the two commands is:

- **USEMONITOR**: Activates or deactivates a selection of beam position monitors. This command affects elements of types MONITOR, HMONITOR, or VMONITOR.
- **USEKICK**: Activates or deactivates a selection of orbit correctors. This command affects elements of types KICKER, HKICKER, or VKICKER.

Both commands have the same attributes:

- **STATUS**: If this flag is true (on), the selected elements are activated. Active orbit monitor readings will be considered, and active correctors can change their strengths in subsequent correction commands. Inactive elements will be ignored subsequently.
- **SEQUENCE**: The sequence can be specified, otherwise the current sequence is used for this operation.
- **RANGE, CLASS, PATTERN**: The usual selection commands are used to identify the elements for this operation.

Example:

USE,...                    ! set working beam line
...                        ! define imperfections
USEKICK,RANGE=..., OFF;    ! deactivate selected correctors
USEMONITOR,RANGE=..., OFF; ! deactivate selected monitors
CORRECT,NCORR=32;          ! uses different set of correctors
CORRECT,NCORR=..., OFF;    ! uses different set of correctors

*Werner Herr* 18.6.2002
This page is under construction, options presently only available in MADX development version.

**CSAVE:** Write orbit correctior settings to file

**SETCORR:** Set orbit correctior settings

[Werner Herr] 18.6.2002
**OPTION: Set Orbit Correction Options**

The random generator for MAD is taken from [[Knuth]]
In the orbit program monitors can be randomly disabled and
the correct option command specifies different seeds for random values:

**OPTION,SEED=integer,PRINT=2**

- **SEED**: Selects a particular sequence of random values.
  A SEED value is an integer in the range [0...999999999] (default: 123456789).
  SEED alone continues with the current sequence
  See also:[Random values]
  SEED may be an expression.
- **PRINT**: This flag can take integer values and controls the printout.
  In general: the higher its value the more printout is produced.
  For PRINT=0 no output is produced.
  The default value is 1 (Correction summary is given).

Example:

**OPTION,SEED=987456321,PRINT=2;**

[Werner Herr] 18.6.2002
Values contained in MAD-X tables can be plotted in the form column versus column, with up to four differently scaled vertical axes; furthermore, if the horizontal axis is the position "s" of the elements in a sequence, then the symbolic machine can be plotted above the curves as well. In certain conditions True interpolation inside the element is available (through calls to the Twiss module for each slice). The "environment" (interpolation, line thickness, annotation size, PostScript format) can be set with the `setplot` command.

```
plot, vaxis=vname1,vname2,...,vnamen,
vaxis1=vname1,vname2,...,vnamen, vaxis2=vname1,vname2,...,vnamen,
vaxis3=vname1,vname2,...,vnamen, vaxis4=vname1,vname2,...,vnamen,
haxis=vname, hmin=real, hmax=real, vmin=real, vmax=real, bars=integer,
style=integer, colour=integer, symbol=integer, noversion=logical,
interpolate=logical, noline=logical, notitle=logical, marker_plot=logical,
range_plot=logical, table=table_name, particle=particle1,particle2,...,particlen,
multiple=logical, title=string, range=range, file=file_name_start,
ptc=logical, ptc_table=table_name, trackfile=table_name;
```

where the parameters have the following meaning:
- **vaxis**: one or several variables from the table to be plotted against the (only) vertical axis.
- **vaxis1**: one or several variables from the table to be plotted against the vertical axis number 1 (out of 4 possible ones).
- **vaxis2**: one or several variables from the table to be plotted against the vertical axis number 2 (out of 4 possible ones).
- **vaxis3**: one or several variables from the table to be plotted against the vertical axis number 3 (out of 4 possible ones).
- **vaxis4**: one or several variables from the table to be plotted against the vertical axis number 4 (out of 4 possible ones).
- **Important**: vaxis and vaxis1 are **exclusive in their application**!
- **haxis**: name of the horizontal variable
- **hmin**: lower horizontal edge
- **hmax**: upper horizontal edge; to be used, both hmin and hmax must be given.
- **vmin**: lower edges of vertical axes, up to four numbers
- **vmax**: upper edges of vertical axes, up to four numbers; both vmin and vmax must be given for an axis to be effective.
- **bars**: 0 (default) or 1 - in the latter case, all curve points coming from the table are connected
with the horizontal axis by vertical bars.

- **Style**: 1 (default), 2, 3, or 4: curve style, being solid, dashed, dotted, and dot-dashed; a value of 100 makes MAD-X use these four styles in turn for successive curves in the same plot. If style is 0 no curve is printed between points. N.B. If symbol and style are null at the same time, style is forced to its default value (= 1).
- **Colour**: 1 (default), 2, 3, or 5: colour, being black, red, green, blue, and magenta; a value of 100 makes MAD-X use these five colours in turn for successive curves.
- **Symbol**: 0 (default), 1, 2, 3, 4, or 5: none, dot, "+", "*", circle, and "x". These symbols are potted at all curve points; there size may have to be adapted (see below).
- **NoVersion**: logical, default=false. If set true, the information concerning the madx version and the date are suppressed from the title. This option frees more space for the user’s title.
- **Interpolate**: logical, default=false. Normally the curve points from the table are connected by straight lines; if "interpolate" is requested, then on-momentum Twiss parameters such as beta, alfa, and dispersion are interpolated with calls to the Twiss module inside each element, for all other variables splines are used to smooth the curves.
- **NoLine**: logical, default=false. If s is the horizontal variable, then the machine will be plotted in symbolic form above the curve plot (except for tables having been read back into MAD-X). This may result in a thick black block if the horizontal scale is too large. "noline" allows the user to suppress the machine plotting.
- **NoTitle**: logical, default=false. If true, suppresses the title line.
- **MarkerPlot**: logical, default=false. If true, plotting is done also at the location of marker elements. This is only useful for the plotting of non-continuous functions like the "N1" from the aperture module. Beware that the PS file might became very large if this flag is invoked.
- **RangePlot**: logical, default=false. Needed to allow to specify a plotting range also for user defined horizontal axis.
- **Table**: name of the table to be plotted from (default: twiss). If it is track, the data to be plotted are taken from the tracking files generated for each required particle as defined by the attribute particle. The name of this file has the following format: file name as defined by the attribute trackfile, the observation point fixed to 1 and the particle number, e.g. testtrack.obs0001.p0003. If the required file has not been generated by the previous MAD-X command track, no plot is done for that particle. The plot is obtained through the gnuplot package. N.B. the previous track command should contain the attribute dump. The tracking plots appends the plots to an existing file specified via filename appended by .ps. The user should make sure that this file does not exist before starting a MAD-X run!
- **Particle**: one or several numbers associated to the tracked particles for which the specified plot has to be displayed.
- **Multiple**: logical, default=false. If true all the curves generated for each tracked particle are put on one plot. Otherwise there will be one plot for each particle.
- **Title**: plot title string; if absent, the last overall title is used; if no such overall title as well, the sequence name is used.
- **Range**: horizontal plot range given by elements.
- **Filename**: start of the file name for the Postscript file(s). Only the first occurrence of such a name will be used. Default is "madx" or "madx_track" if the table attribute is track. Depending on the format (.ps or .eps, see below) the plots will either all be written into one file filename.ps, or one per plot into file_name01.eps, file_name02.eps, etc.
- **Ptc**: logical, default=false. If set true, the data to be plotted are taken from the table defined by
the attribute \textit{ptc\_table} which is expected to be generated previously by the ptc package. The data belong to the column identified by one of the names set in the definition of the ptc twiss table. Interpolation is not available and the attribute \textit{interpolate} has no effect.

- \textit{ptc\_table}: name of the ptc twiss table to be plotted from (default: \textit{ptc\_twiss})
- \textit{trackfile}: first part of the name of the files containing tracking data for each particle (default: \textit{track})

\section*{SETPLOT}

\begin{verbatim}
setplot, post=integer, font=integer, lwidth=real, xsize=real, ysize=real,
ascale=real, lscale=real, sscale=real, rscale=real;
\end{verbatim}

where the parameters have the following meaning:

- \textit{post}: default = 1. If =1, makes one PostScript file (.ps) with all plots; if =2, makes one Encapsulated PostScript file (.eps) per plot.
- \textit{font}: there are two defaults: 1 for screen plotting: this uses characters made from polygons; -1 for PostScript files; this is Times-Italic. There are various fonts available for positive and negative integers, best to be tried out, since they will look different on different systems anyway. GhostView will show strange vertical axis annotations, but the printed versions are normally OK.
- \textit{lwidth}: default = 1. Allows the user to set the curve line width. Depends on the system as well, so to be tried out.
- \textit{xsize}: bounding box size for PostScript, default=27 cm.
- \textit{ysize}: bounding box size for PostScript, default=19 cm.
- \textit{ascale}: annotation character height scale factor, default=1.
- \textit{lscale}: axis label character height scale factor, default=1.
- \textit{sscale}: curve symbol (see above) scale factor, default=1.
- \textit{rscale}: axis text character height scale factor, default=1.

\section*{RESPLOT}

\begin{verbatim}
resplot;
\end{verbatim}

resets all defaults for the setplot command.

This command will execute the Second Order Detuning and Distortion as described in the paper of J. Bengtsson and J. Irwin "Analytical Calculation of Smear and Tune Shift" (SSC-232, February 1990), on the beam line defined by the last USE command followed by a TWISS command. It is based on the stand-alone program written by Frank Schmidt in November 1998 - January 1999 who also extended the analytical computation to the second order distortion (cfr. Beam Physics Note 60 F. Schmidt "SODD: A physics Guide"). It consists of three parts:

**Subroutine detune (launched by the attribute detune)**

It calculates the detuning function terms in first and second order in the strength of the multipoles. If the attribute print_at_end has been set, the following two files (and the corresponding madx tables) are created:

*detune_1_end* containing five columns:

1) 'multipole order', 2) '(hor., ver. plane => (1/2)', 3) 'hor. or ver. detuning', 4) 'order of horizontal invariant', 5) 'order of vertical invariant'.

*detune_2_end* containing five columns:

1) 'first multipole order', 2) 'second multipole order', 3) 'horizontal detuning', 4) 'order of horizontal invariant', 5)'order of vertical invariant'.

If the attribute print_all has been set, the following two files (and the corresponding madx tables) are created:

*detune_1_all* containing five columns:

1) 'multipole order', 2) '(hor., ver. plane => (1/2)', 3) 'hor. or ver. detuning', 4) 'order of horizontal invariant', 5)'order of vertical invariant'.

*detune_2_all* containing five columns:

1) 'first multipole order', 2) 'second multipole order', 3) 'horizontal detuning', 4) 'order of horizontal invariant', 5) 'order of vertical invariant'.
Subroutine distort1 (launched by the attribute distort1)

It calculates the distortion function and the Hamiltonian terms in first order in the strength of the multipoles. If the attribute print_at_end has been set, the two files (and the corresponding madx tables) are created:

\textit{distort\_1\_F\_end} containing eight columns:

1) 'multipole order', 2) 'cosine part of distortion', 3) 'sine part of distortion', 4) 'amplitude of distortion', 5) 'j', 6) 'k', 7) 'l', 8) 'm'.

\textit{distort\_1\_H\_end} containing eight columns:

1) 'multipole order', 2) 'cosine part of Hamiltonian', 3) 'sine part of Hamiltonian', 4) 'amplitude of Hamiltonian', 5) 'j', 6) 'k', 7) 'l', 8) 'm'.

If the attribute print_all has been set, the following two files (and the corresponding madx tables) are created:

\textit{distort\_1\_F\_all} containing eleven columns:

1) 'multipole order', 2) 'appearance number in position range', 3) 'number of resonance', 4) 'position', 5) 'cosine part of distortion', 6) 'sine part of distortion', 7) 'amplitude of distortion', 8) 'j', 9) 'k', 10) 'l', 11) 'm'.

\textit{distort\_1\_H\_all} containing eleven columns:

1) 'multipole order', 2) 'appearance number in position range', 3) 'number of resonance', 4) 'position', 5) 'cosine part of Hamiltonian', 6) 'sine part of Hamiltonian', 7) 'amplitude of Hamiltonian', 8) 'j', 9) 'k', 10) 'l', 11) 'm'.

Subroutine distort2 (launched by the attribute distort2)

It calculates the distortion function and Hamiltonian terms in second order in the strength of the multipoles. If the attribute print_at_end has been set, the following two files (and the corresponding madx tables) are created:

\textit{distort\_2\_F\_end} containing nine columns:

1) 'first multipole order', 2) 'second multipole order', 3) 'cosine part of distortion', 4) 'sine part of distortion', 5) 'amplitude of distortion', 6) 'j', 7) 'k', 8) 'l', 9) 'm'.

\textit{distort\_2\_H\_end} containing nine columns:

1) 'first multipole order', 2) 'second multipole order', 3) 'cosine part of Hamiltonian', 4) 'sine part of Hamiltonian', 5) 'amplitude of Hamiltonian', 6) 'j', 7) 'k', 8) 'l', 9) 'm'.
N. B. The first row of every file is a header containing the names of the columns. This row is absent in the internal tables.

- **SODD**

  sodd,
detune=logical,
distort1=logical,
distort2=logical,
start_stop = start, stop
multipole_order_range = first, last
noprint = logical
print_all = logical
print_at_end = logical
nosixtrack = logical

where the parameters have the following meaning:
- detune : logical, default=false. If true, the detune subroutine is executed.
- distort1 : logical, default=false. If true, the distort1 subroutine is executed.
- distort2 : logical, default=false. If true, the distort2 subroutine is executed.
- start_stop : longitudinal interval of the beam line (in m). start and stop should be given as real numbers.
- multipole_order_range : the lowest and the largest multipole order which will be taken in account. first and last should be given as integers.
- noprint : logical, default=false. If true, no file or internal table will be created to keep the results. In this case the attributes print_all or print_at_end have no effect.
- print_all : logical, default=false. If true, the files and internal tables containing results at each multipole will be generated.
- print_at_end : logical, default=false. If true, the files and internal tables containing results at the end of the position range will be generated.
- nosixtrack : logical, default=false. If true, the input file fc.34 will not be generated internally by invoking the conversion routine of sixtrack and the user should provide it before the execution of the sodd command.
- A more detailed description can be found in [AB-note-2004-069](#)
[damico] September 10, 2004
GEOMETRIC LAYOUT

The SURVEY command computes the coordinates of all machine elements in a global reference system. These coordinates can be used for installation. In order to produce coordinates in a particular system, the initial coordinates and angles can be specified. The computation results are written on an internal table (survey) and can be written on an external file. Each line contains the coordinates at the end of the element. The last "USEd" sequence is used except if another one is specified.

-----------------------------------------------------------------------------------------
WARNING : in the case a machine geometry is constructed with thick lenses, the circumference will change if the structure is converted into thin lenses (via the makethin command). This is an unavoidable feature. ONLY the structure with thick lenses must be used for practical purposes.
INFORMATION : The skew dipole component of a MULTIPOLE element (MULTIPOLE, KSL={FLOAT}) is NOT taken into account in the survey calculation. You should use a tilted normal MULTIPOLE or BEND instead.
-----------------------------------------------------------------------------------------

The survey calculation is launched by a single command line with the following syntax :
SURVEY, x0=double, y0=double, z0=double, theta0=double, phi0=double, psi0=double,
file=string, table=string, sequence=string;

<table>
<thead>
<tr>
<th>parameter</th>
<th>meaning</th>
<th>default value</th>
</tr>
</thead>
<tbody>
<tr>
<td>x0</td>
<td>initial horizontal transverse coordinate</td>
<td>0.0</td>
</tr>
<tr>
<td>y0</td>
<td>initial vertical transverse coordinate</td>
<td>0.0</td>
</tr>
<tr>
<td>z0</td>
<td>initial longitudinal coordinate</td>
<td>0.0</td>
</tr>
<tr>
<td>theta0</td>
<td>initial horizontal angle</td>
<td>0.0</td>
</tr>
<tr>
<td>phi0</td>
<td>initial vertical angle</td>
<td>0.0</td>
</tr>
<tr>
<td>psi0</td>
<td>initial transverse tilt</td>
<td>0.0</td>
</tr>
<tr>
<td>file</td>
<td>name of external file</td>
<td>null (default name survey)</td>
</tr>
<tr>
<td>table</td>
<td>name of internal table</td>
<td>null (default name survey)</td>
</tr>
<tr>
<td>sequence</td>
<td>name of sequence to be surveyed</td>
<td>last used sequence</td>
</tr>
</tbody>
</table>
Example : average LHC ring with CERN coordinates.

REAL CONST R0 = 1.0;                   ! to obtain the average ring
OPTION, -echo, -info;
CALL, file="V6.4.seq.070602";         ! follow this link for the file
OPTION, echo;
BEAM, particle=proton, energy=450, sequence=lhcb1;
USE, period=lhcb1;
! SELECT, flag=survey.clear;            ! uncomment if the optional select below is used
! optional SELECT to specify a class and the output columns
! SELECT, flag=survey, class=marker, column=name,s,psi;

SURVEY, x0=-2202.21027, z0=2710.63882, y0=2359.00656, theta0=-4.315508007, 
phi0=0.0124279564, psi0=-0.0065309236, file=survey.lhcb1;
WRITE, table=survey;                     ! to display the results immediately
STOP;
!***********      The external file "survey.lhcb1" can now be read ***********
F.Tecker, March 2006
SXF file input and output

The command

```
SXFWRITE, FILE=filename;
```

writes the currently (i.e. last) USEd sequence with all alignment and field errors in SXF format onto the file specified. This then represents one "instance" of the sequence, where all parameters are given by numbers rather than expressions; the file can be read by other programs to get a complete picture of the sequence.

The command

```
SXFREAD, FILE=filename;
```

reads a file in SXF format, stores the sequence away and USEs it(!) in order to keep the existing errors. The following does therefore work:

Example:

```
job 1:
! define sequence MYSEQ
use,mysequ;
! add alignment errors and field errors
sxfwrite, file=file;
stop;

job 2:
sxfread, file=file;
twiss;
stop;
```

_hansg_ January 24, 1997
TFS File Format

[TFS] files (Table File System) have been used in the LEP control system. The MAD program knows only coded TFS files. The TFS format has been chosen for all table output of MAD-X. TFS formatted tables can be read back into MAD-X, and may then be further processed.

- Descriptor Lines
- Column Formats
- TFS file example

hansg  June 17, 2002
Descriptor Lines

MAD-X writes the following descriptors on all tables:

- **COMMENT**: The current title string from the most recent TITLE command.
- **ORIGIN**: The version of MAD-X used.
- **DATE**: The date of the MAD-X run.
- **TIME**: The wall clock time of the MAD-X run.
- **TYPE**: The type of the table: e.g. TWISS

Additional descriptors exist in the Twiss table as well as the Track tables.

*hansg*  *June 17, 2002*
Twiss TFS file header

The format of the twiss table is best illustrated with an TFS file example

It should be mentioned that MAD-X allows to access parameters from twiss and other tables using the table access function.

hansg June 17, 2002
Column Formats

The column formats used are listed in the TFS columns table.

<table>
<thead>
<tr>
<th>C format</th>
<th>Meaning</th>
<th>C format</th>
</tr>
</thead>
<tbody>
<tr>
<td>%hd</td>
<td>Short integer</td>
<td>(%8d)</td>
</tr>
<tr>
<td>%le</td>
<td>Long float</td>
<td>(%-18.10g)</td>
</tr>
<tr>
<td>%ks</td>
<td>String of length k</td>
<td>(&quot;&quot;-18s&quot;&quot;)</td>
</tr>
</tbody>
</table>

Control lines begin with the TFS control character, followed by a blank. Data lines begin with two blanks. Columns are also separated by one blank character. The column width is chosen such as to accommodate the large of the column name and the data values of the column.

hansg June 17, 2002
TOUSCHEK: Touschek Lifetime and Scattering Rates

The TOUSCHEK module computes the Touschek lifetime and the scattering rates around a lepton or hadron storage ring, based on the formalism of Piwinski [A. Piwinski, "The Touschek Effect in Strong Focusing Storage Rings," DESY-98-179; see also Piwinski’s article on Touschek lifetime in the Handbook of Accelerator Physics and Engineering (A. Chao, M. Tigner, eds.), World Scientific, 1999].

The syntax of the TOUSCHEK command is:

```
TOUSCHEK, FILE;
```

TOUSCHEK should be called after a TWISS command. One or several cavities with rf voltages should be defined prior to calling TWISS and TOUSCHEK. [Warning: Calling EMIT between the TWISS and TOUSCHEK commands leads to TOUSCHEK using wrong beam parameters, even if the BEAM command is reiterated.]

The momentum acceptance is taken from the bucket size taking into account the energy loss per turn \( U_0 \) from synchrotron radiation. The value of \( U_0 \) is computed from the second synchrotron radiation integral \( \text{synch}_2 \) in the TWISS summ table (\( \text{synch}_2 \) is calculated only when the TWISS option 'chrom' is invoked), using Eq. (3.61) in Matt Sands’ report SLAC-121, which was generalized to the case of several harmonic rf systems. If \( \text{synch}_2=0 \), not defined, or not calculated, zero energy loss is assumed.

In the case of several rf systems with nonzero voltages, it is assumed that the lowest frequency system defines the phase of the outer point on the separatrix when calculating the momentum acceptance, and that all higher-harmonic systems are either in phase or in anti-phase to the lowest frequency system. (Note: if a storage rings really uses a different rf scheme, one would need to change the acceptance function in the routine \text{cavtousch} for that ring.)

The arguments have the following meaning:

- **FILE**: The name of the output file (default: 'touschek')

Example:

```
BEAM, PARTICLE=PROTON, ENERGY=450, NPART=1.15e11, EX=7.82E-9, EY=7.82E-9, ET=5.302e-5, SIGE=7.164e-4, SIGT=0.1124, RADIATE=TRUE;
...
USE, PERIOD=FODO;
```
The first command defines the beam parameters. It is essential that the longitudinal emittances and bunch length are set. The command \textit{use} selects the beam line or sequence. The next command assigns a value to the cavity rf voltage \textit{vrf} (example name). The \textit{select} clears previous assignments to the \textit{twiss} module, \textit{twiss} calculates and saves the values of all twiss parameters for all elements in the ring; the \textit{touschek} command computes the Touschek lifetime and writes it to the file 'touschek' (default name).

The results are stored in the \textit{TOUSCHEK} tables, and can be written to a file (with the default name 'touschek' in the example above), or values can be extracted from the table using the value command as follows:

\begin{verbatim}
value,table(touschek,name),table(touschek,s),table(touschek,tl),table(touschek,tlw),table(touschek,tlitot);
\end{verbatim}

where 'name' denotes the name of a beamline element, \textit{s} the position of the center of the element, \textit{tl} the instantaneous Touschek loss rate within the element, and \textit{tlw} the instantaneous rate weighted by the length of the element divided by the circumference (its contribution to the total loss rate), and \textit{tlitot} the accumulated loss rate adding the rates over all beamline elements through the present position. The value of \textit{tlitot} at the end of the beamline is the inverse of the Touschek lifetime in units of 1/s.

Also, all results can be printed to a file using the command

\begin{verbatim}
write,table=touschek,file;
\end{verbatim}

The MADX Touschek module was developed by Catia Milardi and Frank Zimmermann.

\begin{flushright}
\texttt{frank} 11.03.2008
\end{flushright}
Twiss Module

The TWISS command causes computation of the linear lattice functions and optionally of the chromatic functions. The coupled functions are calculated in the sense of Edwards and Teng. For the uncoupled cases they reduce to the C and S functions. It operates on the working beam line defined in the latest USE command. One can also specify either a SEQUENCE="sequence_name" or a LINE="line_name" on the TWISS command. Moreover, one can restrict the TWISS calculation to a desired RANGE.

The relative energy error DELTAP may be entered in one of the 2 forms

DELTAP=real[, real] DELTAP=initial:final:step

The first form lists several numbers, which may be general expressions, separated by commas. The second form specifies an initial value, a final value, and a step, which must be constant expressions, separated by colons.

Examples:

DELTAP=0.001                    ! a single value
DELTAP=0.001,0.005              ! two values
DELTAP=0.001:0.007:0.002        ! four values

If DELTAP is missing, MAD-X uses the value 0.0.

Further attributes of the TWISS statements are:

- CHROM: A logical flag. If set, MAD-X also computes the chromatic functions. Please note that this option is needed for a proper calculation of the chromaticities in the presence of coupling!
- FILE: If FILE="file_name" appears MAD-X writes a full TFS Twiss table on the disk file "file_name". FILE alone is equivalent to FILE="twiss":
- TABLE (overrides SAVE): MAD-X creates a full Twiss table in memory and gives it the name TWISS, unless TABLE="table_name" appears on the command, then it is called table_name. This table includes linear lattice functions as well as the chromatic functions for all positions. An important new feature of MAD-X is the possibility to access entries of tables and in particular the twiss table (see table access).
- CENTRE: This flag enforces the calculation of the linear lattice functions at the center of the element instead of the end of it. Mind you that since this is inside the element the closed orbit includes the misalignment of the element.
- RMATRIX: If this flag is used the the one-turn map at the location of every element is calculated and prepared for storage in the TWISS table. Using the SELECT command and using the column RE,
RE11...RE16...RE61...RE66 these components will be added to the TWISS table, i.e. with "column, RE" and "column, REij" one gets all or the component "ij" respectively.

- SECTORMAP: This flag initiates the calculation of a sector map as described at SECTORMAP.
- SECTORFILE: Used to write SECTORMAPs to the file SECTORFILE="file_name", if missing the output of SECTORMAP will go to the file "sectormap" with the format as found in SECTORMAP.
- KEEPORBIT: The keeporbit attribute (with an optional name, keeporbit="name") stores the orbit under this name at the start, and at all monitors.
- USEORBIT: The useorbit attribute (with an optional name, useorbit="name") uses the start value provided for the closed orbit search; the values at the monitors are used by the threader.
- COUPLE (obsolete): This MAD8 option can no longer be set since TWISS in MAD-X is always calculated in coupled mode. MAD-X computes the coupled functions in the sense of Edwards and Teng. For the uncoupled cases they reduce to the C and S functions.
- Twiss calculation is 4D only!: The Twiss command will calculate an approximate 6D closed orbit when the accelerator structure includes an active cavity. However, the calculation of the Twiss parameters are 4D only. This may result in apparently non-closure of the beta values in the plane with non-zero dispersion. The full 6D Twiss parameters can be calculated with the ptc_twiss command.
- RIPKEN: This flags will calculate the Ripken-Mais TWISS parameters (beta11, beta12, beta21, beta22, alfa11, alfa12, alfa21, alfa22, gamal1, gamma12, gamma21 and gamma22) using betx, bety, alfx, alfy, gamax, gamay, R11, r12, R21 and R22 as input.

The tables are suitable for plot. After a successful TWISS run MAD-X creates an implicit table of summary parameters named "summ" which includes tunes, chromaticities etc (Please note that the chrom option is needed for a proper calculation of the chromaticities in the presence of coupling!) versus the selected values of DELTAP. Notice that in MAD-X DELTAP is converted in PT, which is used as longitudinal variable. Dispersive and chromatic functions are hence derivatives with respects to PT see table. These summary parameters can later be accessed via the table access function using the aforementioned implicit table named "summ". There is no way to change the name of this summary table.

**Twiss Parameters for a Period**

The simplest form of the TWISS command is

```
TWISS, DELTAP=real{,value},CHROM, TABLE=table_name;
```

It computes the periodic solution for the specified beam line for all values of DELTAP entered (or for DELTAP = 0, if none is entered).

Example:

```
USE,period=OCT;TWISS,DELTAP=0.001,CHROM;
```

This example computes the periodic solution for the linear lattice and chromatic functions for the beam line OCT. The DELTAP value used is 0.001. Apart from saving computing time, it is equivalent to the command sequence.
RING: LINE=(\text{4*(OCT,-OCT)}); USE,period=RING; TWISS,DELTAP=0.001,CHROM;

Initial Values from a Periodic Line

It is possible to track the lattice functions starting with the periodic solution for another beam line. If this is desired the TWISS command takes the form

\text{TWISS, DELTAP=real\{,value\},LINE=beam-line, MUX=real,MUY=real, \text{TABLE=table\_name};}

No other attributes should appear in the command. For each value of DELTAP MAD-X first searches for the periodic solution for the beam line mentioned in LINE=beam-line: The result is used as an initial condition for the lattice function tracking.

Example:

CELL: LINE=(\ldots);INSERT: LINE=(\ldots); USE,period=INSERT; TWISS,LINE=CELL,DELTAP=0.0:0.003:0.001,CHROM,FILE;

For four values of DELTAP the following happens: First MAD-X finds the periodic solution for the beam line CELL: Then it uses this solution as initial conditions for tracking the lattice functions of the beam line CELL: Output is also written on the file TWISS:

If any of the beam lines was defined with formal arguments, actual arguments must be provided:

\text{CELL(SF,SD): LINE=(\ldots);INSERT(X): LINE=(\ldots); USE,period=INSERT; TWISS,LINE=CELL(SF1,SD1);}

Given Initial Values

Initial values for linear lattice functions and chromatic functions may also be numerical. Initial values can be specified on the TWISS command:

\text{TWISS, BETX=real,ALFX=real,MUX=real, BETY=real,ALFY=real,MUY=real, \text{DX=real,DPX=real, DY=real,DPY=real, \text{\ldots  !coupling matrix, \text{TABLE=table\_name, \text{TOLERANCE=real, \text{DELTAP=real:real:real;}}}}}

All initial values for linear lattice functions and chromatic functions are permitted, but BETX and BETY are required. Moreover, a beta0 block can be added as filled by the \text{savebeta} command or see below. The lattice parameters are taken from this block, but will be overwritten by explicitly stated lattice parameters. As entered, the initial conditions cannot depend on DELTAP, and can thus be correct only for one such value.

Tolerance

This value defines the maximum closed orbit error of all six orbit components during the closed orbit search. The default value is 1.e-6. The value is only valid for the current twiss command; a permanent value can be entered via the \text{COGUESS} command.

\text{SAVEBETA: Save Lattice Parameters}

Initial lattice parameters can be transferred for later commands, in particular for twiss or the \text{match module}, by using the \text{savebeta} command sequence.
It should be mentioned that parameters can be also accessed from tables using the \texttt{table access} function.

\texttt{USE,period=\ldots;SAVEBETA,LABEL=name,PLACE=place,SEQUENCE=s\_name;TWISS,\ldots;}

When reaching the \texttt{place} in the sequence "s\_name" during execution of TWISS, MAD-X will save a \texttt{beta0} block with the \texttt{label} name: This block is filled with the values of all lattice parameters in place. Example 1:

\texttt{USE,period=CELL;SAVEBETA,LABEL=END,PLACE=#E,SEQUENCE=CELL;TWISS;USE,period=INSERT;TWISS,BETA0=END;}

This first example calculates the \texttt{periodic solution} of the line CELL, and then track lattice parameters through INSERT, using all end conditions (including orbit) in CELL to start.

Example 2:

\texttt{USE,period=CELL;SAVEBETA,LABEL=END,PLACE=#E,SEQUENCE=CELL;TWISS;USE,period=INSERT;TWISS,BETX=END->BETY,BETY=END->BETX;}

This is similar to the first example, but the beta functions are interchanged (overwritten).

\texttt{\textcopyright \, 06-Apr-2003. Revised in February 2007.}
PTC Set-up Parameters

The E. Forest's Polymorphic Tracking Code (PTC) is a kick code, allowing a symplectic integration through all accelerator elements giving the user full control over the precision (number of steps and integration type) and exactness (full or extended Hamiltonian) of the results. The degree of exactness is determined by the user and the speed of his computer. The main advantage is that the code is inherently based on the map formalism and provides users with all associated tools.

The PTC code is actually a library that can be used in many different ways to create an actual module that calculates some property of interest. Several modules using the PTC code have been presently implemented in MAD-X. These MADX-PTC modules are executed by the following commands: ptc_twiss, ptc_normal, ptc_track, ptc_track_line. To perform calculations with these MADX-PTC commands, the PTC environment must be initialized, handled and turned off by the special commands within the MAD-X input script.

Synopsis

```
PTC_CREATE_UNIVERSE, sector_nmul_max=integer, sector_nmul=integer,
ntpsa=logical, symprint=logical;
PTC_CREATE_LAYOUT, time=logical, model=integer, method=integer,
nst=integer, exact=logical, offset_deltap=double,
errors_out=logical, magnet_name=string, resplit=logical,
thin=double, xbend=double, even=logical;
..........................
PTC_READ_ERRORS, overwrite=logical;
..........................
PTC_MOVE_TO_LAYOUT, index=integer;
..........................
PTC_ALIGN;
..........................
PTC_END;
```

Commands

```
PTC_CREATE_UNIVERSE;
sector_nmul_max=integer, sector_nmul=integer
```

Description

The "PTC_CREATE_UNIVERSE" command is needed to set-up the PTC environment.
### Options

<table>
<thead>
<tr>
<th>Option</th>
<th>Meaning</th>
<th>Default Value</th>
<th>Value Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>SECTOR_NMUL_MAX</td>
<td>Global variable in PTC needed for exact sector bends defining up to which order Maxwell’s equation are solved [a, page 76-77]. The value of SECTOR_NMUL_MAX must not be smaller than SECTOR_NMUL otherwise MAD-X stops with an error.</td>
<td>10</td>
<td>integer</td>
</tr>
<tr>
<td>SECTOR_NMUL</td>
<td>Global variable in PTC needed for exact sector bends defining up to which order the multipole are included in solving Maxwell’s equation up to order SECTOR_NMUL_MAX. Multipoles of order N with N &gt; SECTOR_NMUL and N ≤ SECTOR_NMUL_MAX are treated a la SixTrack.</td>
<td>10</td>
<td>integer</td>
</tr>
<tr>
<td>NTPSA</td>
<td>This attribute invokes the second DA package written in C++ and kindly provided by Lingyun Yang <a href="mailto:lyyang@lbl.gov">lyyang@lbl.gov</a>. Etienne Forest has written the wrapper to allow both the use of the legendary DA packages written in Fortran by Martin Berz (default) or this new DA package. It is expected that this DA package will allow for the efficient calculation of a large number of DA parameters.</td>
<td>.FALSE.</td>
<td>logical</td>
</tr>
<tr>
<td>SYMPRINT</td>
<td>This flag allows the supression of the printing of the check of symplecticity. It is recommended to leave this flag set to TRUE.</td>
<td>.TRUE.</td>
<td>logical</td>
</tr>
</tbody>
</table>

### PTC_CREATE_LAYOUT.

```plaintext
* time=logical, model=integer, method=integer,
* nst=integer, exact=logical, offset_deltap=double, errors_out=logical, magnet_name=string,
* resplit=logical, thin=double, xbend=double, even=logical;
```

**Description**

The "PTC_CREATE_LAYOUT" command creates the PTC-layout according to the specified integration method and fills it with the current MAD-X sequence defined in the latest USE command.

The logical input variable time controls the coordinate system that is being used.

**Options**
Remarks

**TIME:** at small energy \((\beta_0 \ll 1\)\), momentum-dependent variables like dispersion will depend strongly on the choice of the logical input variable "time". In fact, the derivative \((\partial \delta / \partial \beta_p)\) and \((\partial \delta / \partial p)\) are different by the factor \(\beta_0\). One would therefore typically choose the option "time=false", which sets the fifth variable to the relative momentum deviation \(\delta_p\).

**MODEL:** 1 for "Drift-Kick-Drift"; 2 for "Matrix-Kick-Matrix"; 3 for "Delta-Matrix-Kick-Matrix" (SixTrack-code model).

**NST:** sets the same value for all "thick" elements \((l > 0)\) of a beam-line. Please note, that each individual element may have its own NST value **(see below)**.

**PTC_READ_ERRORS**, overwrite=logical;

**Description**

The "PTC_READ-ERRORS" command let’s you read any numbers of "errors_read" tables **READMYTABLE**

**Options**

<table>
<thead>
<tr>
<th>Option</th>
<th>Meaning</th>
<th>Default Value</th>
<th>Value Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>OVERWRITE</td>
<td>Flag to either OVERWRITE the read-in errors (on request by using this flag) or by DEFAULT just add them to multipole components already present.</td>
<td>.FALSE.</td>
<td>logical</td>
</tr>
</tbody>
</table>

**Remarks**
**PTC_MOVE_TO_LAYOUT,**
index=integer;

*Description*
Several PTC layouts can be created within a one PTC-"universe". The layouts are automatically numbered with sequential integers by the MAD-X code. The "PTC_MOVE_TO_LAYOUT" is used for an activation of a requested layout and the next PTC commands will be applied to this active PTC layout until a new PTC layout will be created or activated.

**Option**

<table>
<thead>
<tr>
<th>Option</th>
<th>Meaning</th>
<th>Default Value</th>
<th>Value Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>INDEX</td>
<td>Number of the PTC layout to be activated.</td>
<td>1</td>
<td>integer</td>
</tr>
</tbody>
</table>

**PTC_ALIGN;**

*Description*
The "PTC_ALIGN" command is used to apply the MAD-X alignment errors to the current PTC layout.

**PTC_END;**

*Description*
The "PTC_END" command is turning off the PTC environment, which releases all memory back to the MAD-X world proper;

**Additional Options for Physical Elements**

[SBEND | RBEND | QUADRUPOLE | SEXTUPOLE | OCTUPOLE | SOLENOID ],
l=double, ......,tilt=double, ...., nst=integer, ...., knl:={0, double, double,..}, ksl:={0, double, double,..};

*Description*
1. The full range of normal and skew multipole components on the bench can be specified for the following physical elements: sbend, rbend, quadrupole, sextupole, octupole and solenoid. Multipole coefficients are specified as the integrated value $\int K ds$ of the field components along the magnet axis (see the table below). These multipole components in PTC are spread over a whole element, if $l > 0$. This is a considerable advantage of PTC input compare to MAD-X which allows only thin multipoles.

2. To preserve the reference orbit of straight elements, dipole components for those elements are ignored, knl(0)=0, ksl(0)=0.
3. Individual NST values for a particular "thick" element ($l > 0$) can be specified. For example, in MAD-X any RF cavity is represented by a single kick, while PTC splits the RF cavity into (global) NST segments. In this way, PTC considers properly transit-time effects of the cavity. In case, one wants to reproduce the approximate results of MAD-X, one can use NST=1 for RF cavity in PTC.

**Multipole on Bench (PTC only)**
<table>
<thead>
<tr>
<th>Option</th>
<th>Meaning</th>
<th>Default Value</th>
<th>Value Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>KNL</td>
<td>The normal multipole coefficient</td>
<td>0 [m(^{-1})]</td>
<td>double array</td>
</tr>
<tr>
<td>KSL</td>
<td>The skew multipole coefficient</td>
<td>0 [m(^{-1})]</td>
<td>double array</td>
</tr>
</tbody>
</table>

Remarks

Length l: Bending magnets (shend, rbend) are treated as "markers", if l = 0.
Additional Field Errors: A full range of multipole field errors can be additionally specified with the EFCOMP command. Errors are added to the above multipole fields on the bench.

Caution

A user has to understand that PTC exists inside of MAD-X as a library. MAD-X offers the interface to PTC, i.e. the MAD-X input file is used as input for PTC. Internally, both PTC and MAD-X have their own independent databases which are linked via the interface.

With the "PTC_CREATE_LAYOUT" command, only numerical numbers are transferred from the MAD-X database to the PTC database.
Any modification to the MAD-X database is ignored in PTC until the next call to "PTC_CREATE_LAYOUT".

For example, a deferred expression of MAD-X after a "PTC_CREATE_LAYOUT" command is ignored within PTC.

Examples

Examples for any MADX-PTC module contain the above PTC set-up commands.

References


See Also

ptc_twiss, ptc_normal, ptc_track, ptc_track_line

V. Kapin (ITEP) and F. Schmidt, March 2006
Overview of MAD-X Tracking Modules

A number of particles with given initial conditions can be tracked through a beam-line or a ring. The particles can be tracked either for a single passage or for many turns.

While MAD-X [a] is keeping most of the functionality of its predecessor [MAD-8], the trajectory tracking in MAD-X is considerably modified comparing to MAD-8. The reason is that in MAD-8 the thick lens tracking is inherently not symplectic, which implies that the phase space volume is not preserved during the tracking, i.e. contrary to the real particle the tracked particle amplitude is either growing or decreasing.

The non-symplectic tracking as in MAD-8 has been completely excluded from MAD-X by taking out the thick lens part from the tracking modules. Instead two types of tracking modules (both symplectic) are implemented into MAD-X.

The first part of this design decision is the thin-lens tracking module (thintrack) which tracks symplecticly through drifts and kicks and by replacing the end effects by their symplectic part in form of an additional kick on either end of the element. This method demands a preliminary conversion of a sequence with thick elements into one composed entirely of thin elements (see the MAKETHIN command). The details of its usage are given on the page "thintrack".

The second part of this design decision is to produce a thick lens tracking module based on the PTC code [b] that allows a symplectic treatment of all accelerator elements giving the user full control over the precision (number of steps and integration type) and exactness (full or extended Hamiltonian) of the results.

The first PTC thick-lens tracking module is named ptc_track. It has the same features as the thin-lens tracking code (thintrack) except it treats thick-lenses in a symplectic manner.

There is a second PTC tracking module called the line tracking module (ptc_track_line). It is meant for tracking particles in CLIC in fact it treats beam-lines containing traveling-wave cavities and includes a beam acceleration.

References

a) F. Schmidt, ”MAD-X PTC Integration”, Proc. of the 2005 PAC Conference in Knoxville, USA, pp.1272.


See Also

PTC Set-up Parameters
Thin-Lens Tracking Module (thintrack)

The thin-lens tracking module of MAD-X performs element per element tracking of (one to many) particle trajectories in the last used sequence. Only thin elements are allowed (apart from the element drift), which guarantees the symplecticity of the coordinate transformation. Any lattice can be converted into a "thin element" lattice by invoking the makethin command.

Synopsis

```
TRACK, onepass, deltap= double, dump;  
  START, x= double, px= double, y= double, py= double, t= double, pt= double;  
  RUN, turns= integer;  
ENDTRACK;
```

Commands

```
TRACK, deltap= double, onepass, dump, onetable, file= string;   (MAD-X version 1)
TRACK, deltap= double, onepass, damp, quantum, dump, aperture, onetable, file= string;  
(MAD-X version 2)
TRACK, deltap= double, onepass, damp, quantum, dump, aperture, onetable, recloss, file= string;  
(MAD-X version 3)
[commands];  
ENDTRACK;
```

Description

The TRACK command initiates trajectory tracking by entering the thin-lens tracking module. Several options can be specified, the most important being dump, deltap and aperture.

Inside the block TRACK-ENDTRACK a series of initial trajectory coordinates can be specified by the START command (as many commands as trajectories). This will be usually done in a while-loop. Note that the coordinates are either canonical coordinates or action-angle variables!

- For usual tracking (single/multi-turn), all coordinates are specified with respect to the actual closed orbit (possibly off-momentum, with magnet errors) and NOT with respect to the reference orbit.
- If the option onepass is used, the coordinates are specified with respect to the reference orbit. The name "onepass" might be misleading: Still tracking can be single- or multi-turn!

The tracking is actually started with the RUN command, where the option turns defines for how many turns the particles will be tracked in the given sequence.

If the option dump is used, the particle coordinates are written to files at each turn. The
output files are named automatically. The name given by the user is followed by .obsnnnn(observation point), followed by .pnnnn(particle number). Hence filenames look like track.obs0001.p0001.

Tracking is terminated by the command \textit{ENDTRACK}.

Options

<table>
<thead>
<tr>
<th>Option</th>
<th>Meaning</th>
<th>Default Value</th>
<th>Value Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>DELTAP</td>
<td>relative momentum offset for reference closed orbit (switched off for onepass)</td>
<td>0.0</td>
<td>double</td>
</tr>
<tr>
<td>ONEPASS</td>
<td>the sequence is treated as transfer line (no stability test, ie. no closed-orbit search)</td>
<td>.FALSE. = closed-orbit search</td>
<td>logical</td>
</tr>
<tr>
<td>DAMP</td>
<td>introduce synchrotron damping (needs RF cavity, RADIATE in BEAM)</td>
<td>.FALSE. = no damping</td>
<td>logical</td>
</tr>
<tr>
<td>QUANTUM</td>
<td>introduce quantum excitation via random number generator and tables for photon emission</td>
<td>.FALSE. = no excitation</td>
<td>logical</td>
</tr>
<tr>
<td>DUMP</td>
<td>write the particle coordinates in files (names generated automatically)</td>
<td>.FALSE. = no file generated</td>
<td>logical</td>
</tr>
<tr>
<td>APERTURE</td>
<td>particle is lost if its trajectory is outside the aperture of the current element.</td>
<td>.FALSE. = no aperture check</td>
<td>logical</td>
</tr>
<tr>
<td>ONETABLE</td>
<td>write all particle coordinates in a single file</td>
<td>.FALSE. = one file per particle</td>
<td>logical</td>
</tr>
<tr>
<td>RECLOSE</td>
<td>create a table named &quot;trackloss&quot; in memory with lost particles' coordinates</td>
<td>.FALSE. = no table</td>
<td>logical</td>
</tr>
<tr>
<td>FILE</td>
<td>name for the track table</td>
<td>&quot;track&quot;, &quot;trackone&quot;</td>
<td>string</td>
</tr>
<tr>
<td>UPDATE</td>
<td>parameter update per turn</td>
<td>.FALSE. = no update</td>
<td>string</td>
</tr>
</tbody>
</table>

Remarks

\textit{IMPORTANT:} If an RF cavity has a no zero voltage, synchrotron oscillations are automatically included. If tracking with constant momentum is desired, then the voltage of the RF cavities has to be set to zero. If an RF cavity has a no zero voltage and DELTAP is non zero, tracking is done with synchrotron oscillations around an off-momentum closed orbit.

DELTAP

Defining a non-zero \textit{deltap} results in a change of the beam momentum/energy without changing the magnetic properties in the sequence. This leads to a new closed orbit, the off-momentum closed orbit. Particle coordinates are then given with respect to this new closed orbit, unless the option \textit{onepass} is used!
ONEPASS

If the option *onPASS* is used, no closed orbit is searched, which also means that no stability test is done. Use this option if you want to get the particles' coordinates with respect to the reference orbit rather than the closed orbit. Unfortunately the name is misleading, but for backwards compatibility it is kept. "onPASS" does **NOT** restrict the tracking to one turn only!

APERTURE

- If the *aperture* option is applied, the *apertype* and *aperture* information of each element in the sequence is used to check whether the particle is lost or not. For further information on the definition of apertures and different aperture types, see the documentation of the [APERTURE](#) module.

- In case no aperture information was specified for an element, the following procedure will currently take place:
  → No aperture definition for element → Default apertype/aperture assigned (currently this is *apertype= circle, aperture = {0}*)
  → If tracking with *aperture* is used and an element with *apertype= circle* AND *aperture= {0}* is encountered, then the first value of the *maxaper* vector is assigned as the circle’s radius (no permanent assignment!). See option [maxaper](#) for the default values.
  ⇒ Hence even if no aperture information is specified by the user for certain elements, default values will be used!

RECLOSS

Traditionally, when a particle is lost on the aperture, this information is written to stdout. To allow more flexible tracking studies, the lost particles’ coordinates and further information can also be saved in a table in memory. Usually one would save this table to a file using the WRITE command after the tracking run has finished. The following information is available in the TFS table "trackloss":
- Particle ID (number)
- Turn number
- Particle coordinates (x,px,y,py,t,pt)
- Longitudinal position in the machine (s)
- Beam energy
- Element name, where the particle is lost

UPDATE

Changed behaviour for time variation in tracking. Use track command option ’update’ (e.g.: ’track, onepass, update;’) to use the following additions:
- Introduced special variable (’tr$turn’) that can be used in expressions like ’KICK:= sin(tr$turn)’ and is updated at each turn during tracking.
- Introduced special macro (’tr$macro’) that can be user-defined (’tr$macro(turn): macro = {whatever depending on turnnumber;};’) and is executed/updated at each turn during tracking. (Macro is necessary e.g. for table access.)
START, x= double, px= double, y= double, py= double, t= double, pt= double;
START, fx= double, phix= double, fy= double, phiy= double, ft= double, phit= double;

Description
After the TRACK command, a series of initial trajectory coordinates has to be given by means of a START command (as many commands as trajectories). The coordinates can be either canonical coordinates,
START, X= double, PX= double, Y= double, PY= double, T= double, PT= double;
or action-angle coordinates,
START, FX= double, PHIX= double, FY= double, PHIY= double, FT= double, PHIT= double;
For this case the normalised amplitudes are expressed in number of r.m.s. beam size \( F_X \), \( F_Y \), \( F_T \) (the actions being computed with the emittances in the BEAM command) in each mode plane. The phases are \( \Phi_X \), \( \Phi_Y \) and \( \Phi_T \) expressed in radian. In the uncoupled case, we have in the plane mode labelled \( z \),
\[
Z = F_z \sqrt{E_z} \cos(\Phi_z), \quad P_z = F_z \sqrt{E_z} \sin(\Phi_z),
\]
where \( E_z \) is the r.m.s. emittance in the plane \( Z \).

Options

<table>
<thead>
<tr>
<th>Option</th>
<th>Meaning</th>
<th>Default Value</th>
<th>Value Type</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>X, PX, Y, PY, T, PT</td>
<td>canonical coordinates</td>
<td>0.0</td>
<td>double</td>
<td>m</td>
</tr>
<tr>
<td>FX, PHIX, FY, PHIY, FT, PHIT</td>
<td>action-angle coordinates</td>
<td>0.0</td>
<td>double</td>
<td>rad</td>
</tr>
</tbody>
</table>

Remarks
- For usual tracking (single/multi-turn), all coordinates are specified with respect to the actual closed orbit (possibly off-momentum, with magnet errors) and NOT with respect to the reference orbit.
- If the option onepass is used, the coordinates are specified with respect to the reference orbit. The name "onepass" might be misleading: Still tracking can be single- or multi-turn!

OBSERVE, place= string;

Description
Coordinates can be recorded at places that have names. Such observation points are specified by the command OBSERVE (as many commands as places). The output files are named automatically. The name given by the user is followed by .obsnnnn(observation point), followed by .pnnnn(particle number). Hence filenames look like track.obs0001.p0001.

Options

<table>
<thead>
<tr>
<th>Option</th>
<th>Meaning</th>
<th>Default Value</th>
<th>Value Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>PLACE</td>
<td>name of the observation point</td>
<td></td>
<td>string</td>
</tr>
</tbody>
</table>
Remarks
If no OBSERVE command is given, but the dump option in the TRACK command is used, the particles trajectory coordinates are still recorded. The observation point is then the starting point of the sequence.

RUN, maxaper= double array, turns= integer, ffile= integer;
Description
The actual tracking itself is launched by the RUN command. Via the option turns the user can specify how many turns will be tracked.

Options

<table>
<thead>
<tr>
<th>Option</th>
<th>Meaning</th>
<th>Default Value</th>
<th>Value Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAXAPER</td>
<td>upper limits for the six coordinates</td>
<td>{0.1, 0.01, 0.1, 0.01, 1.0, 0.1}</td>
<td>double array</td>
</tr>
<tr>
<td>TURNS</td>
<td>number of turns</td>
<td>1</td>
<td>integer</td>
</tr>
<tr>
<td>FFILE</td>
<td>periodicity for printing coordinates</td>
<td>1</td>
<td>integer</td>
</tr>
</tbody>
</table>

Remarks
The limits defined by the maxaper option are only being taken into account if the aperture option of the TRACK command is used.

Remarks
- Plotting is possible in MAD-X, however it can also be done externally by using the files created by TRACK.
- The following internal tables are created while tracking: tracksumm, trackloss, and trackone or track.obs$$$.p$$$. (depending on option onetable).
  These internal tables can be accessed via the table-access functions.

See Also
APERTURE, MAKETHIN

A. Koschik February 2007
Thick-Lens Tracking Module
(PTC-TRACK Module)

The PTC-TRACK module \[a\] is the symplectic thick-lens tracking facility in MAD-X \[b\]. It is based on PTC library written by E.Forest \[c\]. The commands of this module are described below, optional parameters are denoted by square brackets ([ ]). Prior to using this module the active beam line must be selected by means of a \[USE\] command. The general \[PTC environment\] must also be initialized.

Synopsis

```
PTC_CREATE_UNIVERSE;
PTC_CREATE_LAYOUT, model=integer, method=integer, nst=integer, [exact];

..........................
PTC_START, .....;

..........................
PTC_OBSERVE,.....;

..........................
PTC_TRACK, .....;

..........................
PTC_TRACK_END;

..........................
PTC_END;
```

Commands

**PTC_START**,

```
x=double, px=double, y=double, py=double, t=double, pt=double,
fx=double, phix=double, fy=double, phiy=double, ft=double, phit=double ;
```

**Description**

To start particle tracking, a series of initial trajectory coordinates has to be given by means of **PTC_START** command (as many commands as trajectories). It must be done before the **PTC_TRACK** command. The coordinates can be either **canonical coordinates** \((x, px, y, py, t, pt)\) or action-angle coordinates \((fx, phix, fy, phiy, ft, phit)\), which are expressed by the normalized amplitude, \(F_z\) and the phase, \(\Phi_z\) for the \(z\)-th mode plane \((z=x,y,t)\). The actions are computed with the values of the emittances, \(F_z\), which must be specified in the preceding **BEAM** command. \(F_z\) are expressed in number of r.m.s. beam sizes and \(\Phi_z\) are expressed in radians.
Options

<table>
<thead>
<tr>
<th>Option</th>
<th>Meaning</th>
<th>Default Value</th>
<th>Value Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>X, PX, Y, PY, T, PT</td>
<td>canonical coordinates</td>
<td>0.0</td>
<td>double</td>
</tr>
<tr>
<td>FX, PHIX, FY, PHIY, FT, PHIT</td>
<td>action-angle coordinates</td>
<td>0.0</td>
<td>double</td>
</tr>
</tbody>
</table>

Remarks

1. If the option `closed_orbit` in the `PTC_TRACK` command is active (see below), all coordinates are specified with respect to the actual closed orbit (possibly off-momentum with magnet errors) and NOT with respect to the reference orbit. If the option `closed_orbit` is absent, then coordinates are specified with respect to the reference orbit.

2. In the uncoupled case, the canonical and the action-angle variables are related with equations

\[
\begin{align*}
    z &= F_z(E_z)^{1/2} \cos(\Phi_z), \\
    p_z &= F_z(E_z)^{1/2} \sin(\Phi_z).
\end{align*}
\]

3. The use of the action-angle coordinates requires the option `closed_orbit` in the `PTC_TRACK` command.

4. If both the canonical and the action-angle coordinates are given in the `PTC_START` command, they are summed after conversion of the action-angle coordinates to the canonical ones.

**PTC_OBSERVE,**

place=string;

Description

Besides of the beginning of the beam-line, one can define an additional observation points along the machine. Subsequent `PTC_TRACK` command will then record the tracking data on all these observation points.

Option

<table>
<thead>
<tr>
<th>Option</th>
<th>Meaning</th>
<th>Value Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>PLACE</td>
<td>name of observation point (markers are very much preferred)</td>
<td>string</td>
</tr>
</tbody>
</table>

Remarks

1. The first observation point at the beginning of the beam-line is marked as "start".
2. It is recommended to use labels of markers in order to avoid usage observations at the ends of thick elements.
3. The data at the observation points other than at "start" can be produced by two different means:
   a) traditional `MADX` element-by-element tracking (use option `element_by_element`);
b) coordinate transformation from "start" to the respective observation point using high-order PTC transfer maps (required option closed_orbit; turned off options radiation and element_by_element).

PTC_TRACK,
deltap=double, icase=integer, closed_orbit, element_by_element, turns=integer,
dump, onetable, maxaper=double array, norm=integer, norm_out,
file=[string], extension=string, ffile=integer,
radiation, radiation_model1, radiation_energy_loss, radiation_quadr,
beam_envelope, space_charge;

Description
The PTC_TRACK command initiates trajectory tracking by entering the thick-lens tracking module. Several options can be specified, the most important are presented in table "Basic Options". There are also switches to use special modules for particular tasks. They are presented in the table "Special Switches". The tracking can be done element-by-element using the option element_by_element or "turn-by-turn" (default) with coordinate transformations over the whole turn. Tracking is done in parallel, i.e. the coordinates of all particles are transformed through each beam element (option element_by_element) or over full turns. The particle is lost if its trajectory is outside the boundaries as specified by maxaper option. In PTC, there is a continuous check, if the particle trajectories stays within the aperture limits. The Normal Form calculations (required option closed_orbit) is controlled by norm_no and norm_out are used.

Basic Options
ICASE

DELTAP

CLOSED_ORBIT

ELEMENT_BY_ELEMENT

TURNS

DUMP

ONETABLE

MAXAPER

NORM_NO

NORM_OUT

FILE

EXTENSION

FFILE

Remarks

ICASE: has a highest priority over other options:
  a) RF cavity with non-zero voltage will be ignored for icase=4, 5;
  b) A non-zero deltap will be ignored for icase=4, 6.

However, if RF cavity has the voltage set to zero and for icase=6, the code sets icase=4.

DELTAP: is ignored for icase=6, but the option offset_deltap of the command
PTC_CREATE_LAYOUT may be used, if

the reference particle should have an momentum off-set as specified by offset_deltap

CLOSED_ORBIT: It must be used for closed rings only. This option allows to switch ON
the Normal Form analysis, if required. If CLOSED_ORBIT is off, the sequence is treated
as a transfer line.

NORM_NO=1: makes the Normal Form linear (always true for MAD8/X).

FILE: The output file endings are: .obsnnnn(observation point), followed by .pnnnn
(particle number),
if the `onetable` option is not used.

### Special Switches

<table>
<thead>
<tr>
<th>Option</th>
<th>Meaning</th>
<th>Default Value</th>
<th>Value Type</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>RADIATION</strong></td>
<td>turn on the synchrotron radiation calculated by an internal procedure of PTC</td>
<td>.FALSE. logical</td>
<td></td>
</tr>
<tr>
<td><strong>RADIATION_MODEL1</strong></td>
<td>switch to turn on the radiation according to the method given in the Ref. [c]</td>
<td>.FALSE. logical</td>
<td></td>
</tr>
<tr>
<td><strong>RADIATION_ENERGY_LOSS</strong></td>
<td>adds the energy loss for radiation_model1</td>
<td>.FALSE. logical</td>
<td></td>
</tr>
<tr>
<td><strong>RADIATION_QUADR</strong></td>
<td>adds the radiation in quadrupoles. It supplements either radiation or radiation_model1</td>
<td>.FALSE. logical</td>
<td></td>
</tr>
<tr>
<td><strong>BEAM_ENVELOPE</strong></td>
<td>turn on the calculations of the beam envelope with PTC</td>
<td>.FALSE. logical</td>
<td></td>
</tr>
<tr>
<td><strong>SPACE_CHARGE</strong></td>
<td>turn on the simulations of the space charge forces between particles.</td>
<td>.FALSE. logical</td>
<td></td>
</tr>
</tbody>
</table>

#### Remarks

1. **RADIATION**: Has precedence over radiation_model1.
2. **RADIATION_MODEL1**: Additional module by F. Zimmermann. The model simulates quantum excitation via a random number generator and tables for photon emission. It can be used only with the element-by-element tracking (option `element-by-element`).
3. **RADIATION_ENERGY_LOSS**: Of use for radiation_model1.
4. **BEAM_ENVELOPE**: It requires the options `radiation` and `icase=6`.
5. **SPACE_CHARGE**: This option is under construction and is reserved for future use.
PTC_TRACK_END;

Description
The PTC_TRACK_END command terminate the command lines related to the
PTC_TRACK module.

TRACKSUMM table

The starting and final canonical coordinates are collected in the internal table "tracksumm"
(printed to the file with WRITE command).

Examples
Several examples are found on the here

The typical tasks

The following table facilitates the choice of the correct options for a number of tasks.

<table>
<thead>
<tr>
<th>CLOSED_ORBIT</th>
</tr>
</thead>
<tbody>
<tr>
<td>ELEMENT_BY_ELEMENT</td>
</tr>
<tr>
<td>PTC_START, X, PX, ...</td>
</tr>
<tr>
<td>PTC_START, FX, PHIX,</td>
</tr>
<tr>
<td>NORM_NO</td>
</tr>
<tr>
<td>NORM_OUT</td>
</tr>
<tr>
<td>PTC_OBSERVE</td>
</tr>
<tr>
<td>RADIATION</td>
</tr>
<tr>
<td>RADIATION_MODEL1</td>
</tr>
<tr>
<td>RADIATION_ENERGY_LOSS</td>
</tr>
<tr>
<td>RADIATION_QUAD</td>
</tr>
<tr>
<td>BEAM_ENVELOPE</td>
</tr>
<tr>
<td>SPACE_CHARGE</td>
</tr>
</tbody>
</table>

1) The tracking of a beam-line with default parameters.
2) As 1), but with element-by-element tracking and an output at observation points.
3) Tracking in a closed ring with closed orbit search and the Normal Forms calculations.
   Both canonical and action-angle input/output coordinates are possible. Output at observation points is
   produced via PTC maps.
4) Similar to "3)" except that output at observation points is created by element-by-element tracking.
5) The with PTC radiation.

References for PTC-TRACK
a) V. Kapin and F. Schmidt, PTC modules for MAD-X code, to be published as CERN internal note by the end of 2006


c) E. Forest, F. Schmidt and E. McIntosh, Introduction to the Polymorphic Tracking Code, KEK report 2002-3, July 2002


See Also

<table>
<thead>
<tr>
<th>Overview of MAD-X Tracking Modules</th>
<th>PTC Set-up Parameters</th>
<th>thintrack</th>
<th>PTC-TRACK</th>
</tr>
</thead>
<tbody>
<tr>
<td>Examples</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

V. Kapin (ITEP) and F. Schmidt, July 2005; revised in April, 2006
PTC_TRACKLINE

PERFORMS A PARTICLE TRAJECTORY TRACKING
WITH ACCELERATION USING PTC

USER MANUAL

SYNOPSIS

PTC.Trackline,
turns                 [integer, 1, 0 ],
onetable             [logical, false, true ],
everystep           [logical, false, true],
tableallsteps        [logical, false, true],
gcs                  [logical, false, true],
file                  [string, "track", "track" ],
rootntuple           [logical, false, true],
extension            [string, ",", "] ;
<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Type</th>
<th>Default value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>turns</td>
<td>integer</td>
<td>1</td>
<td>Number of turns</td>
</tr>
<tr>
<td>onetable</td>
<td>logical</td>
<td>false</td>
<td>Present, but value not specified</td>
</tr>
<tr>
<td></td>
<td></td>
<td>true</td>
<td>If false, tracking data are written to a single table for each track for each observation point. Table names follow the naming <code>filename.obsMMMM.pNNNN</code>, where <code>filename</code> is settable prefix with <code>file</code> parameter (see below), MMMM is observation point number and NNNN is track number. If true, all data are written to single table called onetable.</td>
</tr>
<tr>
<td>file</td>
<td>string</td>
<td>&quot;track&quot;</td>
<td>Name of file where track parameters are written, see description of onetable switch above</td>
</tr>
<tr>
<td>rootntuple</td>
<td>logical</td>
<td>false</td>
<td>true</td>
</tr>
<tr>
<td>everystep</td>
<td>logical</td>
<td>false</td>
<td>true</td>
</tr>
<tr>
<td>gcs</td>
<td>logical</td>
<td>false</td>
<td>true</td>
</tr>
</tbody>
</table>
Description

This MAD-X command performs ray tracking that takes into account acceleration in traveling wave cavities. It must be invoked in the scope of correctly initialized PTC environment, i.e. after PTC_CREATE_UNIVERSE and PTC_CREATE_LAYOUT commands and before corresponding PTC_STOP. All tracks that are spawned with PTC_START commands beforehand PTC_TRACKLINE command is issued are tracked. Track parameters are dumped at every defined observation point (see PTC_OBSERVE command). Please note that MAD-X always creates observation point at the end of a sequence. Depending on value of onetable switch, all output information is stored in one table (and also file), or in one table per track per observation point is written if the switch is false. The user must note that track parameters plotting (see PLOT command) is only possible if onetable switch is set to false (status as for Feb. 2006). This unfortunate solution is the legacy of the regular MAD-X track command, that is designed for circular machines where the user usually tracks a few particles for many turns rather than many particles for one turn each.

Tracks that do not fit in aperture are immediately stopped.

Behavior of PTC calculations can be adapted with PTC_SETSWITCH command and with appropriate switches of PTC_CREATE_LAYOUT command.

Command parameters and switches

turns

integer, default value 1, no default value if value explicitly not specified

Number of turns around sequence. If layout is not closed then its value is enforced to 1.

onetable

boolean, default value false, if value explicitly not specified then true

If true then only one table is created and one file is written to disk. If false one file per track per observation point is written. File format is filename.obsNNNN.pMMMM, where NNNN and MMMM are numbers of observation point and track, respectively. Filename is defined by the switch described below.

file

character string, default is "track"

name of file where track parameters are written, see description of onetable switch above.

PROGRAMMERS MANUAL

The routine PTC_TRACKLINE is implemented in file madx_ptc_trackcavs.f90 Its single parameter is the number of observation points.

The call sequence from MAD-X interpreter is the following:
exec_command in madxp.c;
pro_ptc_trackline in madxn.c; This routine creates appropriate tables where the track parameters are
stored, and after execution of the Fortran routine dumps filled table(s) to files.

w_ptc_trackline_ in wrap.f90; Just interface to the appropriate Fortran module

ptc_trackline in madx_ptc_trackline.f90

The key routine that enables appropriate calculation of beam and track parameters in the presence of
traveling wave cavities is setcavities.

Firstly, the ptc_trackline routine finds out which are the observation points. For this purpose array of
integers observedelements is allocated. Its length is equal to the number of elements in the sequence. All
elements are zero by default. If an element with an index n is an observation point then
observedelements[n] is equal to 1. This solution enables fast checking if track parameters should be sent to
a table after a given element.

Further setcavities subroutine is called if it was not executed yet before.

PTC_TRACKLINE reads the track initial parameters from the table with the help of gettrack function
(implemented in C in file madxn.c). For the performance reasons gettrack creates a two dimensional array
and buffers there all the initial track parameters upon first call. The array is destroyed with a call of
deletetrackstrings positions function that is performed at the very end of ptc_trackline subroutine.

Tracking itself is implemented in a doubly nested loop. The external one goes over all initiated tracks, and
the internal one performs tracking of a given track element by element. The key PTC routine is called
TRACK. It propagates a track described by an array of 6 real numbers, denoted as X in equations below.
The important issue is that they are the canonical variables. In order to follow the standard MAD-X
representation the values that are written to tables and files are scaled appropriately to the reference
momentum for a given element. In the general case it changes along the line if traveling wave cavities are
present. Hence, momenta xp, yp and zp are

\[
\begin{align*}
zp &= \sqrt{(1+x(5))^2 - x(2)^2 - x(4)^2} \\
xp &= x(2)/zp \\
yp &= x(4)/zp
\end{align*}
\]

where array x containing 6 elements is the track position in the PTC representation, i.e. x(1) is horizontal
spacial coordinate, x(2) - horizontal momentum, x(3) - vertical spacial coordinate, x(4) - vertical
momentum, x(5) - \(\frac{\Delta p}{p_0c}\), x(6) - longitudinal coordinate (caution, the exact meaning depends on the PTC
settings, see [PTC_SETSWITCH command].
PTC_SETSWITCH

routine that sets the internal PTC switches

USER MANUAL

SYNOPSIS

PTC_SETSWITCH,
  debuglevel = [i,0],
  maxacceleration = [l, true, true],
  exact_mis = [l, false, true],
  totalpath = [l, false, true],
  radiation = [l false, true],
  fringe = [l, false, true],
  time = [l, true, true];

Description

Using this command the user can set switches of PTC and the MAD-X-PTC interface, adapting this way the program behavior to his needs.

Command parameters and switches

debuglevel

  integer, default value 1, no default value if value explicitly not specified

  Sets the level of debugging printout 0 none, 4 everything

maxacceleration

  logical, default true, if value explicitly not specified then true

  Switch saying to set cavities phases so the reference orbit is always on the crest, i.e. gains max energy

exact_mis

  logical, default false, true if value explicitly specified

  Switch ensures exact misalignment treatment.

totalpath

  logical, default false, true if value explicitly specified
If true, the 6th variable of PTC, i.e. 5th of MAD-X is the total path. If false it is deviation from the reference particle, which is normally the closed orbit for closed layouts.

**radiation**

*logical, default false, if value explicitly not specified then true:*

Sets the radiation switch/internal state of PTC.

**fringe**

*logical, default false, true if value explicitly specified:*

Sets the fringe switch/internal state of PTC. If true the influence of the fringe fields is evaluated for all the elements.

Please note that currently fringe fields are always taken to the account for some elements (f.g. traveling wave cavities) even if this flag is set to false. The more detailed list of the elements will be provided later, when the situation in this matter will be definitely settled.

**time**

*logical, default true, if value explicitly not specified then true :*

If true, Selects time of flight rather than path length. (*cT* to be precise) as the 6th variable of PTC, i.e. 5th of MAD-X.

**PROGRAMMERS MANUAL**

Values of the switches are stored in Fortran 90 module mad_ptc_intstate (mad_ptc_intstate.f90). The command is processed by pro_ptc_setswitch C function in file madxn.c It call an appropriate routines of the Fortran module to set each of the switches:

- ptc_setdebuglevel
- ptc_setaccel_method
- ptc_setexactmis
- ptc_setexactsim
- ptc_setfradiation
- ptc_settotalpath
- ptc_settime
- ptc_setfringe
PTC_SettCavities

f90 routine that adjusts cavities and sets appropriate reference momenta for a layout containing traveling wave cavities

PROGRAMMERS MANUAL

CAUTION For the time being cavities MUST not be placed one after another, and at least a marker must be inserted between two neighboring accelerating structures. Otherwise, program will stop with the error message.

Description

This routine sets up the properties of a layout and traveling wave cavities. The main goal is to update reference beam energy for the elements that follow a traveling wave cavity. It traces the synchronous particle, i.e. one that has all its parameters set to zero at the beginning of the layout under study. At the point it arrives to a cavity, the parameters of the latter one are adjusted according to the switches defined by the user. There are 2 cases

1. Leaves all parameters untouched
2. Phase of cavity is adjusted so it gives the maximum acceleration Afterwards to the calculated phase the lag is added. This setting is acquired using set_switch command, setting maxaccel parameter to true.

Afterwards, the synchronous particle is tracked through traveling wave cavity and its energy gain is known. This energy becomes the reference one for all the elements downstream of the cavity. The particle is tracked further to the next cavity, for which the procedure described above is repeated.

Parameters of the cavities are dumped to the file named twcavsettings.txt.

At the end patches at the ends of the cavities are set, so the parameters after them are calculated taking to the account reference energy increase.

The exact program behavior depends on the PTC switches settings.

Please note that in PTC phase velocity of a cavities wave is always equal to speed of light. Hence, if PTC internal state TIME is TRUE, what is the most correct setting, then voltage seen by a particle is varying along the structure. If TIME is FALSE, track is assumed to fly with speed of light and in such case a particle moves together with the wave front.
PTC_TWISS Module
(Ripken Optics Parameters)

The PTC_TWISS module of MAD-X is based on the PTC code. It is a supplementary to the TWISS module. The Twiss parameters are calculated in Ripken’s style (invented by G. Ripken in 1970 [31] and most accessible in Ref. [b]). These parameters were available in MAD8 using the TWISS3 command. This module is a typical example of the advantages when using PTC and its Normal Form technique (and of course the object-oriented Fortran90 coding): once the rather modest programming has been performed the Twiss calculation will always be automatically correct for all machine conditions like closed orbit, coupling or after a new element has been introduced into the code. In a traditional coding like in MAD8 this depends on reprogramming and modifying the code at various places which is inherently error-prone.

The PTC_TWISS tracks a special representation of the beam in three degrees of freedom. It works on the coupled lattice functions defined in Ref.[b], which are essentially the projections of the lattice functions for the eigen-modes on the three planes. The PTC_TWISS lists the projections of the ellipses of motion onto the three planes \((x, p_x), (y, p_y), (\ell, p_\ell)\) expressed them via the Ripken’s parameters \(b_{k j}, a_{k j}, g_{k j}\) along with the phase advances \(m_j\) in selected positions, where index \(k=1...3\) refers to the plane \((x,y,...)\), and the index \(j=1...3\) denotes the eigen-mode. The PTC_TWISS also calculates the dispersion values \(D_1,...,D_4\). In the MAD-X commands and tables, these parameters are denoted as \(\beta_{11},...\beta_{33}, \alpha_{11},...\alpha_{33}, \gamma_{11},...\gamma_{33}, \mu_1,...,\mu_3, \text{disp}_{1},...,\text{disp}_{4}\), respectively.

The Ripken parametrization can be transformed into the Edwards-Teng parametrization (used in twiss proper) using the formulae of Ref. [d]. The parameters are noted as \(\beta_x, \beta_y, \alpha_x, \alpha_y\) and the coupling matrix: \(R_{11}, R_{12}, R_{21}\) and \(R_{22}\). In absence of coupling it holds: \(\beta_x=\beta_{11}, \beta_y=\beta_{22}, \alpha_x=\alpha_{11}\) and \(\alpha_y=\alpha_{22}\).

PTC_TWISS can also compute the \(\delta p/p\)-dependency of the Twiss parameters. The column names \(\beta_{11p},...\beta_{33p}, \alpha_{11p},...\alpha_{33p}, \gamma_{11p},...\gamma_{33p}\) denote the derivatives of the optics parameters w.r.t. \(\delta p/p\). If one is interested in evaluating \(\delta p/p\)-dependency of the Twiss parameters, one must ensure that the order (no) of the map is set to 2 at least. The derivatives of the dispersion w.r.t. \(\delta p/p\) have column names: \(\text{disp}_{1p},...\text{disp}_{4p}\). Second and third order derivatives have respective column names: \(\text{disp}_{1p^2},...\text{disp}_{4p^2}\) for the second order, and \(\text{disp}_{1p^3},...\text{disp}_{4p^3}\) for the third order.

In addition, we compute momentum compaction factor \(a_c\) up to 1st order (for icase=5) or 3rd order (for icase=56). The values appear in the header of the ptc_twiss output file (zero means the value has not been computed). This feature is currently only available in the development version.
For clarification: in the 4-D case, there is the following correspondence between MAD-X and the Ripken’s notations: $\beta_{11} \cdot b_x, \beta_{12} \cdot b_x, \beta_{21} \cdot b_y, \beta_{22} \cdot b_y$, while in the uncoupled 4-D case $\beta_{11}$ is the same as the classical $b_x (\beta_x)$ and $\beta_{22}$ is $b_y (\beta_y)$, while $\beta_{12}$ and $\beta_{21}$ are zero. When there is coupling all $\beta_{NN}$ are non-zero and $\beta_{11}, \beta_{22}$ are distinctively different from $b_x, b_y$, respectively.

PTC_TWISS also tracks the eigenvectors and prints them to Twiss table according to the SELECT command (flag=ptc_twiss). Either all 36 components or particular components of the eigenvectors can be selected with eign or eignij, respectively ($j =$ number of eigenvector, $i =$ number of coordinate $\{x, p_x, y, p_y, t, p_t\}$).

For ring lattices, PTC_TWISS computes momentum compaction, transition energy, as well as other one-turn characteristics such as the tunes $Q_1, Q_2$ and if icase=6 with cavity $Q$s and chromaticities (for $no>=2$).

Synopsis

```
PTC_CREATE_UNIVERSE;
PTC_CREATE_LAYOUT, model=integer, method=integer, nst=integer, [exact];
..............................
SELECT, flag=ptc_twiss, clear;
SELECT, flag=ptc_twiss, column=name, s, beta11,...,beta33,alfa11,..., alfa33,gama11,...,gama33,
beta11p,...,beta33p,alfa11p,...,alfa33p,gama11p,...,gama33p,
mu1,...,mu3,
disp1,...,disp4,
disp1p,...,disp4p,
disp1p2,...,disp4p2,
disp1p3,...,disp4p3,
[eign], eign11, ...,eign16,...,eign61,...,eign66;
..............................
PTC_TWISS;
..............................
PTC_END;
```

Commands

```
PTC_TWISS,

  icase=integer, deltap=double, closed_orbit, slice_magnets,
  range=string, file=[string], table=[string],
  initial_matrix_table, initial_matrix_manual, initial_map_manual, beta0=string,
  betx=double, alfx=double, mux=double,
  bety=double, alfy=double, muy=double,
  dx=double, dpx=double, dy=double, dpy=double,
  x=double, px=double, y=double, py=double, t=double, pt=double,
  re11=double, re12=double, ... ,re16=double,
```

................................................................
236
re61=double, re62=double, ... ,re66=double;

Description

The PTC_TWISS command causes computation of the Twiss parameters in the Ripken’s style. It operates on the working beam line defined in the latest USE command. Several options can be specified, the most important being icase, deltap, closed_orbit, slice_magnets, no, and file, table. (see the table below). Other options should be specified for particular tasks. Applications for the PTC_TWISS command are similar to the TWISS command. The PTC_TWISS can be applied to two basic tasks. It can calculate either a periodical solution or a solution with initial conditions.

Options

<table>
<thead>
<tr>
<th>Option</th>
</tr>
</thead>
<tbody>
<tr>
<td>ICASE</td>
</tr>
<tr>
<td>NO</td>
</tr>
<tr>
<td>DELTAP</td>
</tr>
<tr>
<td>CLOSED_ORBIT</td>
</tr>
<tr>
<td>DELTAP_DEPENDENCY</td>
</tr>
<tr>
<td>SLICE_MAGNETS</td>
</tr>
<tr>
<td>CENTER_MAGNETS</td>
</tr>
<tr>
<td>FILE</td>
</tr>
<tr>
<td>TABLE</td>
</tr>
<tr>
<td>SUMMARY_FILE</td>
</tr>
<tr>
<td>SUMMARY_TABLE</td>
</tr>
<tr>
<td>RANGE</td>
</tr>
<tr>
<td>INITIAL_MATRIX_TABLE</td>
</tr>
<tr>
<td>INITIAL_MATRIX_MANUAL</td>
</tr>
<tr>
<td>INITIAL_MAP_MANUAL</td>
</tr>
<tr>
<td>RE11,..., RE66</td>
</tr>
<tr>
<td>BETA0</td>
</tr>
<tr>
<td>betx, alfx, mux,orthy, alfy, muy, dx, dpx, dy, dpy</td>
</tr>
<tr>
<td>x, px, y, py, t, pt</td>
</tr>
</tbody>
</table>

Remarks

ICASE: It can be internally corrected by the code. For example, if RF cavity has the voltage set to zero and for icase=6, the code sets icase=4.
Periodical Solution

**PTC_TWISS**, 
`icase=integer, deltap=double, closed_orbit, range=string, file[=string], table[=string]`;

Description

This is the simplest form of the *PTC_TWISS* command, which computes the periodic solution for a specified beam line. It may accept all basic options described in the above table.

Evaluation of Twiss parameters inside magnets

**PTC_TWISS**, 
`icase=integer, deltap=double, closed_orbit, slice_magnets range=string, file[=string], table[=string]`;

Description

This computes the periodic solution for a specified beam line and evaluates the Twiss parameters at each thin-slice (a.k.a "integration-node") inside magnets. The number of such integration-nodes is given by the number of steps (nst) selected when creating the PTC layout. All other basic options described in the above table may be selected.

Example

An example is found in the [PTC_TWISS Examples] repository.

Solution with Initial Conditions

**Code Logic:**

```plaintext
IF ("initial_matrix_table"=ON .AND. 
& {the map-table exists}) THEN
    (from a Map-Table)
ELSEIF("initial_map_manual"=ON) THEN
    (from a Given Map File)
ELSEIF("initial_matrix_manual"=ON) THEN
    (from a Given Matrix)
ELSEIF(BETA0 block =ON) THEN
    (from Twiss Parameters via BETA0-block)
ELSE
    (from Given Twiss Parameters)
ENDIF
```

**Initial Values from a Map-Table**

*(obtainable by a preceding *PTC_NORMAL* command):*

**PTC_TWISS**, 
`icase=integer, deltap=double, closed_orbit, range=string, file[=string], table[=string], initial_matrix_table;`
**Description**

*PTC_TWISS* calculates a solution with initial conditions given as a map-table of preceding ring or beam-line. It requires the input option `initial_matrix_table` and an existence of the map-table in memory, which was generated by a preceding *PTC_NORMAL* command.

**Example**

An example is found in the *PTC_TWISS Examples* in the folder "Example3".

---

**Initial Values from a Map-File**

(\textit{obtainable by a preceding} *PTC_NORMAL* \textit{command}):

*PTC_TWISS*,

\texttt{icase=integer, deltap=double, closed\_orbit,}
\texttt{range=string, file[=string], table[=string],}
\texttt{initial\_map\_manual;}

**Description**

*PTC_TWISS* calculates a solution with initial conditions given as a map-file fort.18 obtained from a preceding ring or beam-line. It requires the input option `initial_map_manual` and an existence of the map-file named a fort.18 file, which was generated by a preceding *PTC_NORMAL* command.

**Example**

An example is found in the *PTC_TWISS Examples* in the folder "Example3".

---

**Initial Values from a Given Matrix**:

*PTC_TWISS*,

\texttt{icase=integer, deltap=double, closed\_orbit,}
\texttt{range=string, file[=string], table[=string],}
\texttt{initial\_matrix\_manual,}
\texttt{re11=double, re12=double, ... ,re16=double,}
\texttt{.................................}
\texttt{re61=double, re62=double, ... ,re66=double;}

**Description**

*PTC_TWISS* calculates a solution with initial conditions given by the matrix, which is "manually" entered on the command-line. It requires the option `initial_matrix_manual`. MAD-X expects a symplectic 6x6 transfer matrix as input.

**Example**

An example is found in the *PTC_TWISS Examples* in the folder "Example4".

---

**Initial Values from Twiss Parameters via BETA0-block**:

*PTC_TWISS*,

\texttt{icase=integer, deltap=double, closed\_orbit,}
\texttt{range=string, file[=string], table[=string],}
\texttt{beta0=string;}

**Description**

*PTC_TWISS* calculates a solution with initial conditions given by Twiss parameters, which are transferred from the \textit{BETA0}-block. The data in the \textit{BETA0}-block have to be filled by a combination of the \texttt{SAVEBETA} and \texttt{TWISS} commands of a preceding ring or beam-line. Note, that this case is limited to uncoupled motion of the preceding machine.

**Example**

An example is found in the *PTC_TWISS Examples* in the folder "Example1".
**Initial Values from the Given Twiss Parameters:**

PTC_TWISS,

\[
  icase=\text{integer}, \ \text{deltap}=\text{double}, \ \text{closed\_orbit}, \\
  \text{range}=\text{string}, \ \text{file}=[\text{string}], \ \text{table}=[\text{string}], \\
  \text{betx}=\text{double}, \ \text{alfx}=\text{double}, \ \text{mux}=\text{double}, \\
  \text{bety}=\text{double}, \ \text{alfy}=\text{double}, \ \text{muy}=\text{double}, \\
  \text{dx}=\text{double}, \ \text{dpx}=\text{double}, \ \text{dy}=\text{double}, \ \text{dpy}=\text{double}, \\
  \text{x}=\text{double}, \ \text{px}=\text{double}, \ \text{y}=\text{double}, \ \text{py}=\text{double}, \\
  \text{t}=\text{double}, \ \text{pt}=\text{double};
\]

**Description**

PTC_TWISS calculates a solution with initial conditions given by the Twiss parameters, which are explicitly typed on the command line. Note, that this case is also limited to uncoupled motion of the preceding ring or beam-line.

**Example**

An example is found in the PTC_TWISS Examples in the folder Example2.

**References for PTC_TWISS**


c) K. Zhang, "PTC twiss with initial TWISS parameters", MAD-X Meeting 13 (04.07.2005), slides in ppt.


See Also

TWISS PTC_TWISS Examples

V. Kapin (ITEP) and F. Schmidt, March 2006
PTC_NORMAL Module
(Non-Linear Machine Parameters)

The PTC_NORMAL module of MAD-X is based on PTC code. This module takes full advantage of the PTC Normal Form analysis which is a considerable upgrade of what was available with the Lie Algebra technique used in MAD8. It allows to calculate dispersions, chromaticities, anharmonicities and Hamiltonian terms to very high order. In fact, the order is only limited by the RAM memory of your computer and your patience to wait for the results.

The number of terms per order increases with some power law. The internal MAD-X tables are not adequate to keep such large amounts of data. On the other hand, only a reduced set of this data is actually needed by the user. Thus a much easier and flexible solution is to gather the users requirements with a series of special MAD-X command called SELECT_PTCNORMAL. A special MAD-X table is dynamically built using just those commands and it will be filled by the next call to the PTC_NORMAL-command.

Another essential advantage of this table is the fact that it is structured to facilitate exchange of Normal Form (including Hamiltonian terms of high order) between MAD-X modules. The immediate goal is to use this table to allow non-linear matching inside the present MAD-X MATCHING module.

Synopsis

```plaintext
PTC_CREATE_UNIVERSE;
PTC_CREATE_LAYOUT, model=integer, method=integer, nst=integer,
[exact];

.........................
SELECT_PTC_NORMAL, dx,..., gnfu;

.........................
PTC_NORMAL;
WRITE, table=normal_results, file=normal_results;

.........................
PTC_END;
```

Commands

```plaintext
SELECT_PTC_NORMAL,
dx=integer, dpx=integer, dy=integer, dpy=integer,
q1=0, dq1=integer, q2=0, dq2=integer,
anhx=integer array, anhy=integer array,
gnfu=integer,0,0, haml=integer,0,0,
eign=integer, integer;
```
Description

The `SELECT_PTC_NORMAL` command selects parameters to be calculated by the next `PTC_NORMAL` command. The dispersion and tune parameters are defined by a name and an integer number specifying their order. For example, the notation "dx=2" means the horizontal dispersion to second order $D_x^{(2)} = \partial^{(2)} x_{co} / \partial \delta_p^{(2)}$, where "co" is abbreviation of "closed orbit". The anharmonisities are defined by a name and three integer numbers: the first is the order of $\varepsilon_1$, the second is the order of $\varepsilon_2$, the third is the order of $\delta_p$. For example, the notation "anhx=2,0,0" means second order in $\varepsilon_1$; $\partial^{(2)} q_1 / \partial \varepsilon_1^{(2)}$.

Components of the eigenvectors at the end of the structure can be specified by two integers: the first integer defines the eigenvector number, the second integer defines the coordinate $\{ x, p_x, y, p_y, t, p_t \}$.

The Generating Function can be specified by $\{ n, 0, 0 \}$. The positive and negative values of $n$ define the order of upright or skew resonances, respectively. The integers $n_2$ and $n_3$ are reserved for a future upgrade. For example, "gnfu=-5, 0, 0" will calculate all Generating Function terms for skew decapoles. In the output table, one finds the cosine, sine and amplitude coefficients as denoted by "GNFC", "GNFS", and "GNFA", respectively. Similarly, the Hamiltonian terms can be specified by $\{ n, 0, 0 \}$. The positive and negative values of $n$ define the order of upright or skew resonances, respectively. For example, "haml=3, 0, 0" will calculate all Hamiltonian terms for upright sextupoles. In the output table, one finds the cosine, sine and amplitude coefficients as denoted by "HAMC", "HAMS", and "HAMA", respectively.

Parameters

<table>
<thead>
<tr>
<th>Notation</th>
<th>Meaning</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>DX, DPX, DY, DPY</td>
<td>dispersions, $D_x^{(n)}$, $D_{px}^{(n)}$, $D_y^{(n)}$, $D_{py}^{(n)}$</td>
<td>$n$</td>
</tr>
<tr>
<td>Q1, Q2</td>
<td>horizontal and vertical tunes $q_1^{(0)}$, $q_2^{(0)}$</td>
<td>0</td>
</tr>
<tr>
<td>DQ1, DQ2</td>
<td>derivatives of horizontal and vertical tunes $\partial^{(n)} q_1 / \partial \varepsilon_1^{(n)}$, $\partial^{(n)} q_2 / \partial \delta_p^{(n)}$</td>
<td>$n$</td>
</tr>
<tr>
<td>ANHX, ANHY</td>
<td>Anharmonicities</td>
<td>$n(\varepsilon_1)$, $n(\varepsilon_2)$, $n(\delta_p)$</td>
</tr>
<tr>
<td>GNFU</td>
<td>Generating Function</td>
<td>$n$, $0$, $0$</td>
</tr>
<tr>
<td>HAML</td>
<td>Hamiltonian</td>
<td>$n$, $0$, $0$</td>
</tr>
<tr>
<td>EIGN</td>
<td>Eigenvector (the $n_2$-th component of the $n_1$-th eigenvector)</td>
<td>$n_1$, $n_2$</td>
</tr>
</tbody>
</table>
**PTC_NORMAL**, 
nice=integer, normal, closed_orbit,  
nno=integer, map_table, deltap=double;

**Description**

The calculation of the parameters specified by the preceding `SELECT_PTC_NORMAL` commands is initiated by the `PTC_NORMAL` command, which operates on the working beam line defined in the latest `USE` command. The options for `PTC_NORMAL` command are described in the table below.

**Options**

<table>
<thead>
<tr>
<th>Option</th>
<th>Meaning</th>
<th>Default Value</th>
<th>Value Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>ICASE</td>
<td>the user-defined dimensionality of the phase-space (4, 5 or 6)</td>
<td>4</td>
<td>integer</td>
</tr>
<tr>
<td>NO</td>
<td>the order of the map.</td>
<td>1</td>
<td>integer</td>
</tr>
<tr>
<td>CLOSED_ORBIT</td>
<td>the switch to turn on the closed orbit calculation.</td>
<td>.FALSE.</td>
<td>logical</td>
</tr>
<tr>
<td>DELTAP</td>
<td>relative momentum offset for reference closed orbit</td>
<td>0.0</td>
<td>double</td>
</tr>
<tr>
<td>MAPTABLE</td>
<td>turn on the map-table in memory</td>
<td>.FALSE.</td>
<td>logical</td>
</tr>
<tr>
<td>NORMAL</td>
<td>turn on the calculation of the Normal Form</td>
<td>.FALSE.</td>
<td>logical</td>
</tr>
</tbody>
</table>

**Remarks**

**MAPTABLE**: (requires \( \text{no}=1 \)) creates the one-turn matrix which can be used by the next `PTC_TWISS` command.

**Example**

The simple example is located on the Web-page for the `PTC_NORMAL` example.
References for PTC_NORMAL


See Also

PTC_NORMAL example PTC Set-up Parameters

V. Kapin (ITEP) and F. Schmidt, March 2006
MAD-X-PTC interface documentation - Auxiliaries

Available documents

- PTC_Knob.html
- PTC_SetKnobValue.html
- PTC_PrintParametric.html
- PTC_PrintFrames.html
- PTC_Select.html
- PTC_SelectMoment.html
- PTC_SetSwitch.html
- PTC_DumpMaps.html
- PTC_SetCavities.html
- PTC_EPlacement.html
- ptc_general.html
- ptc_track.html
- ptc_track_line.html
- ptc_twiss.html
- ptc_normal.html
- Under Construction Match_WithPTCKnobs.html
- Under Construction PTC_Moments.html
- this document
PTC_KNOB

USER MANUAL

SYNOPSIS

PTC_KNOB,
  elementname = [s, none],
  kn    = [i, {-1}],
  ks    = [i, {-1}],
  exactmatch = [l, true, true];

Description

Sets knobs in PTC calculations (currently only in PTC_TWISS, PTC_NORMAL will follow). Knobs appear as the additional parameters of the phase space. Twiss functions are then obtained as functions of these parameters (Taylor series). Also map elements might be stored as functions of knobs, see \texttt{ptc_select} command description to learn how to request given element to be stored as a Taylor series. The parametric results can be further

1. written to a file with \texttt{ptc_printparametric};
2. plotted and studied using rviewer command (rplot plugin);
3. used to obtain very quickly approximate values of lattice functions for given values of knobs \texttt{ptc_setknobvalue}. This feature is the foundation of a fast matching algorithm with PTC.

Example

\texttt{dog leg chicane} Dipolar components of both rbends and dipolar and quadrupolar components of the focusing quads set as knobs. Some first and second order map coefficients set to be stored as parametric results. \texttt{ptc_twiss} command is performed and the parametric results are written to files in two formats. \texttt{dog leg chicane} Knobs values are matched to get requested lattice functions.

Command parameters and switches

\texttt{elementname}

\textit{string in range format,}
Specifies name of the element containing the knob(s) to be set.

kn,ks

*list of integers*,

Defines which order

exactmatch

*logical, default true, if value explicitly not specified then true*

Normally a knob is a property of a single element in a layout. The specified name must match 1:1 to an element name. This is the case when exactmatch is true.

Knobs might also be set to all family of elements. In such case the exactmatch switch must be false. A given order field component of all the elements that name starts with the name specified by the user become a single knob.

initial

—
PTC_SETKNOBVALUE
USER MANUAL

SYNOPSIS

PTC_SETKNOBVALUE,
  elementname = [s, none] ,
  kn    = [i, {-1}],
  ks    = [i, {-1}],
  value = [r] ;

Description

With this command the user set a given knob value. In its effect all the values in

- the twiss table used by the last ptc_twiss command
- the columns specified with ptc_select parametric=true;

are reevaluated using the buffered parametric results.

The parameters of the command basically contains the fields that allow to identify uniquely the knob and
the value to be set.

Example

dog leg chicane: strength of dipol field component in quads is matched to obtain required R56 value.

Command parameters and switches

  elementname
    string in range format,

    Specifies name of the element containing the knob to be set.

  kn,ks
    list of integers,

    Defines the knob
**value**

*real, default 0, if value explicitly not specified then 0*

Specifies the value the knob is set to.
PTC_KNOB

USER MANUAL

SYNOPSIS

PTC_KNOB,
  elementname = [s, none],
  kn    = [i, {-1}],
  ks    = [i, {-1}],
  exactmatch = [l, true, true];

Description

Sets knobs in PTC calculations (currently only in PTC_TWISS, PTC_NORMAL will follow). Knobs appear as the additional parameters of the phase space. Twiss functions are then obtained as functions of these parameters (taylor series). Also map elements might

Example

Not yet ready: postion of quads is matched to obtain required R566 value.

Command parameters and switches

   elementname string in range format,

   Specifies name of the element containing the knob(s) to be set.

   kn, ks list of integers,

   Defines which order

   exactmatch logical, default true, if value explicitly not specified then true

   Normally a knob is a property of a single element in a layout. The specified name must match 1:1 to an element name. This is the case when exactmatch is true.
Knobs might be alos set to all family of elements. In such case the exactmatch switch must be false. Filed components of all the elements that name starts with the name specified by the user become a single knob.
PTC_PRINTFRAMES

USER MANUAL

SYNOPSIS

PTC_PRINTFRAMES,
file = [s, none] ,
format = [s, text] ;

Description

Print to a specified file PTC geometry of a layout.

Example

Dog leg chicane with some elements displaced with help of ptc_eplacement:

Command parameters and switches

file
string,
Specifies name of the file.

format
string, default "text"

Format of geometry. Currently two formats are accepted:
text
Prints a simple text file.
rootmacro
Creates root macro that produces 3D display of the geometry.
PTC_SELECT
USER MANUAL

SYNOPSIS

PTC_SELECT,
table = [s, none, none],
column = [s, none, none],
polynomial = [i, none],
monomial = [s, none],
parametric = [l, false, true],
quantity = [s, none] ; "

Description

Selects map elements to be:

a) Stored in a user specified table and column. Table and column must be specified than, and such table with such column must exists.

b) Stored as a function (taylor series) of knobs if any is defined. Than, parametric should be set to true. Both a) and b) can be joined in one command.

Examples

dog leg chicane: strength of quads is matched to obtain required T112 value.

dog leg chicane: position of quads is matched to obtain required T566 value.

dog leg chicane: dipols and quads strengths are matched with the help of knobs to obtain required momentum compaction and Twiss functions.

Command parameters and switches

table
    string,
Specifies name of the table where values should be stored.

column
string,

Specifies name of the table where values should be stored.
polynomial
integer,

Specifies row of the map.

monomial
string composed of digits

Defines monomial of the polynomial in PTC nomenclature. Its length should be equal to number of
variables. Each of digits corresponds to the exponent of a variable. Monomial 'ijklmn' defines
$x^i p_x^j y^k p_y^l \Delta T^m (\Delta p/p)^n$. For example, element=2 and monomial=1000000 defines coefficient of
the second polynomial (that defines $p_x$) close to $x$, in the other words it is R21.

parametric
logical, default false, if value explicitly not specified then true

If it is true, and any knobs are defined the map element is stored as the parametric result.

PROGRAMMERS MANUAL

The command is implemented pro_ptc_select function in madxn.c and by subroutine addpush in
dadx_ptc_knobs.f90, that is part of madx_ptc_knobs_module

On the very beginning the existence of the table and within column is checked. In the case of failure, error
message is printed and the function is abandoned.

The command parameters are passed as the arguments of addpush Fortran routine. A selection is stored in
a type called tablepush_poly defined madx_ptc_knobs.inc. A newly created object is added to array named
pushes.

More then one element might be stored in a single table, so the module must assure that each of tables is
augmented only ones for each magnet (or integration slice). For that purpose array of tables to be
augmented (named tables) is stored separately and we assure that a table is listed here only ones. This is
simply done by checking if a table name is not already listed before adding a new element to the array.

In case the user requested an element to be stored in the parametric format, and column in the array of
parametric results is reserved and the index of the column is remembered in index field of tablepush_poly
type is filled. In the other case this field is equal to zero.

The routine ptc_twiss (defined in file madx_ptc_twiss.f90), after tracking each of magnets in the
sequence, calls putusertable routine. This routine loops over selected elemeents defined in the pushes table.
For each of them it extracts the requested element from the map using .sub. operator of PTC and stores it
in the defined table and column. If index field is not zero and any knob is defined, it extracts the
polynomial using .par. operator, and stores it in the 2D array called results, in the row corresponding to
the number of the magnet (or integration step) and column defined by the index field.
PICT_SELECT_MOMENT

USER MANUAL

SYNOPSIS

PTC_SELECT_MOMENT,

```
table     = [s, none, none],
column    = [s, none, none],
moments   = [s, none],
moments   = [i, {0}],
parametric = [l, false, true],
```

Description

Selects a moment to be:

a) Stored in a user specified table and column.

b) Stored as a function (taylor series) of knobs if any is defined. Then, parametric switch should be set to true.

Both a) and b) can be joined in one command.

Examples

ATF2

Command parameters and switches

```
moment_s
```

*list of coma separated strings composed of up to 6 digits*

Defines moment of the polynomial in PTC nomenclature. String 'ijklmn' (i,j,k,l,m,n are digits) defines \(x^i p_x^j y^k p_y^l \Delta T^m (\Delta p/p)^n\). For example, moments=1000000 defines \(x^1\).

Note that for input we always use MAD-X notation where dp/p is always the 6th coordinate. (Internaly PTC dp/p is the 5th coordinate. We perform automatic conversion that is transparent for the user.) As the consequence RMS in dp/p is always defined as 000002, even in 5D case.
This notation allows to define more than one moment with one command. In this case, the corresponding column names are as the passed strings with "mu" prefix. However, they are always extended to 6 digits, i.e. the trailing 0 are automatically added. For example, if specified moments=2, the column name is mu200000.

This method does not allow to pass bigger numbers than 9. If you need to define such a moment, use the command switch below.

**moment**

*list of up to 6 coma separated integers,*

Defines a moment. For example: moment=2 defines \( <x^2>\), moment=0,0,2 : \( <y^2>\), moment=0,14,0,2 : \( <px^{14}py^2>\), etc.

**table**

*string, default "moments"*

Specifies name of the table where the calculated moments should be stored.

**column**

*string*

Ignored if *moments* is specified. Defines name of the column where values should be stored. If not specified then it is automatically generated from moment the definition \( <x^1p_x^1y^kpy^l\Delta T^m(\Delta p/n)> \)

\( \Rightarrow \) mu_i-j_k-l_m_n (numbers separated with underscores).

**parametric**

*logical, default false, if value explicitly not specified then true*

If it is true, and any knobs are defined the map element is stored as the parametric result.
PTC_DUMPMAPS

USER MANUAL

SYNOPSIS

PTC_DUMPMAPS,
file = [s, ptcmaps, ptcmaps];

implemented by subroutine ptc_dumpmaps() in madx_ptc_module.f90

Description

PTC_DUMPMAPS dumps linear part of the map for each element of the layout into specified file.

Command parameters and switches

file

string, default value "ptcmaps", default value if value explicitly not specified is "ptcmaps"

Specifies name of the file to which the matrices are dumped to.

PROGRAMMERS MANUAL

The command is implemented by subroutine ptc_dumpmaps() in madx_ptc_module.f90. The matrix for a single element is obtained by tracking identity map through an element, that is initialized for each element by adding identity map to the reference particle. For the elements that change reference momentum (i.e. traveling wave cavity) it is tracked to the end of the following marker, that has updated reference momentum. Hence, each cavity must be followed by a marker. If it is not, setcavities subroutine detects error and stops the program.
PTC_EPLACEMENT

USER MANUAL

SYNOPSIS

PTC_EPLACEMENT,
range = [s, none],
x = [r, 0], y = [r, 0], z = [r, 0],
phi = [r, 0],
theta = [r, 0],
onlyposition = [l, false, true],
onlyorientation = [l, false, true],
autoplacedownstream = [l, false, true],
refframe = [s, gcs];

Description

Places a given element at required position and orientation. All rotations are made around the front face of the element.

Example

Dog leg chicane position of quads is matched to obtain required R566 value.

Command parameters and switches

range

string in range format,

Specifies name of the element to be moved.

x,y,z

real,

Coordinate of the front face of the magnet.

phi, theta

real,
polar (in xz plane, around z axis) and azimuthal (around x axis) angles, respectively.

**refframe**

*string, default gcs*

Defines the coordinate system with respect to which coordinates and angles are specified. Possible values are:

- **gcs**
  - global coordinate system
- **current**
  - current position
- **previouselement**
  - end face of the previous element

**onlyposition**

*logical, default false, if value explicitly not specified then true*

If true, only translation are performed and orientation of the element is not changed.

**onlyorientation**

*logical, default false, if value explicitly not specified then true*

If true, only rotations are performed and position of the element is not changed.

**autoplacedownstream**

*logical, default true, *

if true all the elements downstream are placed at default positions in respect to the moved element, if false the rest of the layout stays untouched.

**surveyall**

*logical, default true, if value explicitly not specified then true*

If true, survey of all the line is performed after element placement at new position and orientation. It is implemented mainly for the software debugging purposes. If patching was performed correctly, the global survey should not change anything.

---

**PROGRAMMERS MANUAL**

The command is implemented pro_ptc_eplacement function in madxn.c and by subroutine ptc_eplacement() in madx_p tc_eplacement.f90.

Sopecified range is resolved with help of get_range command. Number of the element in the current sequence is resolved and passed as the parameter to the fortran routine. It allows to resolve uniquely the corresponding element in the PTC layout.

TRANSLATE_Fibre and ROTATE_Fibre routines of ptc are employed to place and orient an element in space. These commands adds rotation and translation from the current position. Hence, if the specified reference frame is other then "current", the element firstly needs to be placed at the center of the reference frame and then it is moved about the user specified coordinates.
After element placement at new position and orientation patch needs to be recomputed. If autoplacedownstream is false then patch to the next element is also recomputed. Otherwise, the layout is surveyed from the next element on, what places all the elements downstream with default position with respect to the moved element.

At the end all the layout is surveyed, if surveyall flag is true, what normally should always take place.
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Matching with PTC knobs

USER MANUAL

This matching procedure takes advantage of the parametric results that are accessible with PTC. Namely, parameters occurring in the matching constrains are obtained as functions (polynomials) of the matching variables. In other words, each variable is a knob in PTC calculation. Evaluation of the polynomials is relatively fast comparing to the regular PTC calculation what makes finding the minimum with the parametrized constraints very fast.

However, the algorithm is not faster in a general case:

1. The calculation time dramatically increases with number of parameters and at some point penalty rising from this overcomes the gain we get from the fast polynomial evaluation.
2. A parametric result is an approximation that is valid only around the nominal parameter values.

The algorithm:

1. Buffer the key commands (ptc_varyknob, constraint, ptc_setswitch, ptc_twiss or ptc_normal, etc) appearing between match, useptcknobs=true; and any of matching actions calls (migrad,lmdif,jacobian, etc)
2. When matching action appears,
   a) set "The Current Variables Values" (TCVV) to zero
   b) perform THE LOOP, i.e. points 3-17
3. Prepare PTC environment (ptc_createuniverse, ptc_createlayout)
4. Set the user defined knobs (with ptc_knob).
5. Set TCVV using ptc_setfieldcomp command
6. Run a PTC command (twiss or normal)
7. Run a runtime created script that performs a standard matching; all the user defined knobs are variables of this matching.
8. Evaluate constraints expressions to get the matching function vector (I)
9. Add the matched values to TCVV
10. End PTC session (run ptc_end)
11. If the matched values are not close enough to zeroes then goto 3
12. Prepare PTC environment (ptc_createuniverse, ptc_createlayout)
13. Set TCVV using ptc_setfieldcomp command
   ( --- please note that knobs are not set in this case --- )
14. Run a PTC command (twiss or normal)
15. Evaluate constraints expressions to get the matching function vector (II)
16. Evaluate a penalty function that compares matching function vectors (I) and (II) See points 7 and 14
17. If the matching function vectors are not similar to each other within requested precision then goto 3
18. Print TCVV, which are the matched values.

SYNOPSIS

MATCH, use_ptcknob=true;

PTC_VARYKNOB:
    initial = [s, none],
    element = [s, none],
    kn     = [i, -1],
    ks     = [i, -1],
    exactmatch = [l, true, true],
    trustrange = [r, 0.1],
    step    = [r, 0.0],
    lower   = [r, -1.e20],
    upper   = [r, 1.e20];

END_MATCH;

For the user convenience the limits are specified in the MAD-X units (k1,k2, etc). This also applies to
dipolar field where the user must specify limits of k0=angle/path_length. This guarantees consistency in
treatment of normal and skew dipol components.

Important: Note that inside the code skew magnets are represented only by normal component and tilt, so
the nominal skew component is always zero. Inside PTC tilt can not become a knob, while skew
component can. Remember about this fact when setting the limits of skew components in the matching.
When the final results are exported back to MAD-X, they are converted back to the "normal" state, so the
nominal skew component is zero and tilt and normal component are modified accordingly.

trustrange - defines the range the expansion is trusted

Description

Example

[dog leg chicane]
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PTC_MOMENTS
USER MANUAL

SYNOPSIS

PTC_MOMENTS,
no = [1, 1],
xdistr = [s, gauss, gauss],
ydistr = [s, gauss, gauss],
zdistr = [s, gauss, gauss],

Description

Calculates moments previously selected with the \texttt{ptc\_select\_moment} command. It uses maps saved by the \texttt{ptc\_twiss} command, hence, the savemaps switch of \texttt{ptc\_twiss} must be set to true (default) to be able to calculate moments.

Examples

\begin{verbatim}
ATF2
\end{verbatim}

Command parameters and switches

\begin{itemize}
\item \textit{no} \hspace{1cm} \textit{integer}
  \begin{itemize}
  \item order of the calculation, maximally twice the order of the last \texttt{twiss}
  \end{itemize}
\item \textit{xdistr, ydistr, zdistr} \hspace{1cm} \textit{string defining type of distribution for x, y, z dimension, respectively,}
  \begin{enumerate}
  \item \texttt{gauss} - Gaussian
  \item \texttt{flat5} - flat distribution in the first of variables (dp over p) of a given dimension and Delta Dirac in the second one (T)
  \item \texttt{flat56} - flat rectangular distribution
  \end{enumerate}
\end{itemize}
Known Differences to Other Programs

Definitions

MAD uses full 6-by-6 matrices to allow coupling effects to be treated, and the canonical variable set \((x, p_x / p_0), (y, p_y / p_0), (-ct, \text{delta}(E) / p_0 c)\), as opposed to other programs most of which use the set \((x, x'), (y, y'), (-\text{delta}(s), \text{delta}(p)/p_0)\). Like [Dragt], MAD uses the relative energy error \(p_y / p_0\), which is equal the relative momentum error \(\text{delta} = \text{delta}(p)/p_0\) multiplied by \(\beta = v/c\).

As from Version 8.13, MAD8 uses an additional constant momentum error \(\text{delta}_x\) in all optical calculations. The transfer maps contain the exact dependence upon this value; therefore the tunes for large deviations can be computed with high accuracy as opposed to previous versions.

The choice of canonical variables in MAD still leads to slightly different definitions of the lattice functions. In MAD the Courant-Snyder invariants in [Courant and Snyder] take the form

\[
W_x = \gamma_x x^2 - 2 \alpha_x x p_x + \beta_x p_x^2
\]

Comparison to the original form

\[
W_x = \gamma_x x^2 - 2 \alpha_x x' x + \beta_x x'^2
\]

shows that the orbit functions cannot be the same. A more detailed analysis, using

\[
x' = p_x / (1 + \text{delta})
\]

shows that all formulas can be made consistent by defining the MAD orbit functions as

\[
\beta_{xM} = \beta_{xC} * (1 + \text{delta}), \alpha_{xM} = \alpha_{xC}, \gamma_{xM} = \gamma_{xC} / (1 + \text{delta})
\]

For constant \(\text{delta}_x\) along the beam line and \(\text{delta} = 0\), the lattice functions are the same. In a machine where \(\text{delta}\) varies along the circumference, e.g. in a linear accelerator or in an electron-positron storage ring, the definition of the Courant-Snyder invariants must be generalised. The MAD invariants have the advantage that they remain invariants along the beam line even for variable \(\text{delta}\).

With the new method this problem occurs in Twiss module only for non-constant \(\text{delta}\).
Treatment of Energy Error in TWISS

It has been noted in [Milutinovic and Ruggiero] that MAD returned tunes which are too low for non-zero delta. The difference was found to be quadratic in delta with a negative coefficient. This problem has been eliminated thanks to the new treatment of momentum errors from MAD8 Version 8.13 onwards.

[hanse] January 24, 1997
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- **YROTATION**: Rotation About the Vertical Axis
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Effect of the bv flag in MAD-X

When reversing the direction ("V") of a particle in a magnetic field ("B") while keeping its charge constant, the resulting force $V \times B$ changes sign. This is equivalent to flipping the field, but not the direction.

For practical reasons the properties of all elements of the LHC are defined in the MADX input as if they apply to a clockwise proton beam ("LHC beam 1"). This allows a single definition for elements traversed by both beams. Their effects on a beam with identical particle charge but running in the opposite direction ("LHC beam 2") must then be reversed inside the program.

In MADX this may be taken into account by setting the value of the BV attribute in the Beam commands. In the case of LHC beam 1 (clockwise) and beam 2 (counter-clockwise), treated in MADX both as clockwise proton beams, the Beam commands must look as follows:

```
Beam, sequence=lhcb1, particle=proton, pc=450, bv=+1;
Beam, sequence=lhcb2, particle=proton, pc=450, bv=-1;
```
MAD-X News

This is a loose collection of new features, new sample jobs, and other possibly interesting remarks concerning MAD-X.

Work log between releases madX-4_00_09 and madX-4_00_19

Log report started Thu May 7 20:01:28 2009, ended Thu May 7 20:02:54 2009

elaface

Fixed argument of wrong type passed to the mtlmdl Fortran subroutine
warning fixed

Now the check for OSTYPE works with darwin9.0

frs

1) New TPSA package by lingyun.yang@gmail.com 2) General clean-up

Preliminary fix of memory crash - courtesy JBJ

Some rearrangements by Etienne and final clean-up of PTC

Workaround for a array out-of-bound problem due to LDA=16000 being too small. 1) LDAMAX set to 110’000 & lda_used =100000 2) Program stops when an out-of-bound array access is attempted. 3) Real fix needed to make LDA dynamic

Missing general "public" statements which creates pseudo bugs

More clean-up for the TPSA upgrade

Fortran Clean-up: indenting, remove potentially uninitialized variable and also remove unused variables

Minor clean-up concerning: character strings, possibly uninitialized variables and unused variables

277
ONLINE needs: libmdblib.a libmdbmth.a librpnlib.a libSDDS1.a libSDDS1c.a libz.a SLC5 needed: libX11.a

MAD-X production version 4.00.19
version madX-4_00_18_dev
Version 4.00.17: add LRAD to kickers
Version 4.00.16
Version 4.00.15
MAD-X version 4.00.14
MAD-X version madX-4_00_12_dev
version madX-4_00_12_dev
version madX-4_00_11_dev
version 4.00.10

version

Add attribute "range_plot" to allow range also for user defined horizontal axis (courtesy HG)

Allowing plotting at markers using the "marker_plot" attribute. Courtesy HG

Add ntpsa flag (if present the new C++ TPSA package by lingyun.yang@gmail.com in invoked) and the symprint flag pronts the symplecticity flag by default.

Remove "harmless" occurrence "dipole_bv" by the more obvious one "other_bv"

remove tabs

Clean-up unused variables

Take out: Unused external reference RESULT_FROM_NORMAL found with latest: NAG Fortran Compiler Release 5.2(668)

Minor clean-up

Get the compiler directives like "ifdef __WIN32_DLL working in F90 files using the Lahey compiler. Special Fujitsu flags for lf95 invoked e.g. lf95 -c -o1 -tp -ife ".-Cpp" -ife ".D_WIN32_DLL" %FPP%\c_tpsa_interface.F90 -winconsole -ml msvc These compiler flags can be found at: http://www.lahey.com/docs/fujitsu%20compiler%20option%20list.pdf
To wrap up for Windows

Modification for c_tpsa_interface.F90

Added new tpsa package

Fixing the passing of a double array instead of an integer array.

Etienne’s clean-up of AF

Fix C/C++ nonstandard features fix pseudo-bug in c_tpsa_interface.F90 found by NAG f95. Integer shall not be defined as an array of dimension 1.

Latest cleanup of Lingyun’s TPSA including tpsa.dll needed for Windows

Clean-up

- gfortran broken in gcc4.4 - Therefore back to g95 however the LIBX flags must be fixed according to gcc being used

Fix the exclusion of f90 & F90 files with and without NTPSA

Yet another upgrade for MAC using g95

Clean-up for MAC

Generalization for f90/F90

Compiler preprocessing for c_tpsa_interface to drop DLL for LINUX

Add new tpsa

missing -fno-range-check flag for gfortran in particular for MAC

Back to standard "LF95"

Further adjustments to safeguard running with "gfortran" which can be steered with the new "SLC4" flag.

Fix further gfortran for SLC4

Default compiler oh lxplus: lf95 gfortran with -Wall -pedantic DEBUG flags even for Fortran

gfortran explanation for SLC4

Darwin fixes

no ONLINE as standard

More small fixes: - o4 off for lf95 - proper libraries for ONLINE

Fix gfortran Home link use proper LINK options for MAC

hbu
using semi automatic object file list

Makefile mac compatible

jbj

made more robust the capture of the Twiss data of the drift preceeding the current element - solving the ‘1st slice wrong n1’ supressed the memory crash when aperture called twice with the same offsetlem in tfs format

nougaret

further prevent output mixup through forced flush when crossing the border in the other direction (i.e. Fortran calling C, which is more rare than the reverse)

handle flushing unit 6 on Intel ifort compiler

invoke ’call flush(6)’ as ’flush(6)’ with Intel compiler

suppress compilation warnings

skowron

Added filling of track summ table for ptc_trackline. Now the user can check what were the final coordinates of tracking

    If plugin support: link dynamically; if debug: do not put -O4 optimization;
g95 option: add proper debug flags

Work log between releases madX-4_00_09 and madX-4_00_19

Log report started Tue May 5 20:01:26 2009, ended Tue May 5 20:02:49 2009

elafce

Fixed argument of wrong type passed to the mtlmdl Fortran subroutine

warning fixed

Now the check for OSTYPE works with darwin9.0

frs

1) New TPSA package by lingyun.yang@gmail.com 2) General clean-up

Preliminary fix of memory crash - courtesy JBJ

Some rearrangements by Etienne and final clean-up of PTC

GET_C_J routine no longer needed

Missing general "public" statements which creates pseudo bugs
More clean-up for the TPSA upgrade

Fortran Clean-up: indenting, remove potentially uninitialized variable and also remove unused variables

Minor clean-up concerning: character strings, possibly uninitialized variables and unused variables

ONLINE needs: libmdblib.a libmdbmth.a librpnlib.a libSDDS1.a libSDDS1c.a libz.a SLC5 needed: libX11.a

MAD-X production version 4.00.19

version madX-4_00_18_dev

Version 4.00.17: add LRAD to kickers

Version 4.00.16

Version 4.00.15

MAD-X version 4.00.14

MAD-X version madX-4_00_12_dev

version madX-4_00_12_dev

version madX-4_00_11_dev

version 4.00.10

version

Add attribute "range_plot" to allow range also for user defined horizontal axis (courtesy HG)

Allowing plotting at markers using the "marker_plot" attribute. Courtesy HG

Add ntpsa flag (if present the new C++ TPSA package by lingyun.yang@gmail.com is invoked) and the symprint flag prints the symplecticity flag by default.

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remove tabs

Clean-up unused variables

Take out: Unused external reference RESULT_FROM_NORMAL found with latest: NAG Fortran Compiler Release 5.2(668)

Minor clean-up
Get the compiler directives like "ifdef _WIN32_DLL working in F90 files using the Lahey compiler. Special Fujitsu flags for lf95 invoked e.g. lf95 -c -o1 -tp -ife "-Cpp" -ife "-D_WIN32_DLL" %FPP%\c_tpsa_interface.F90 -winconsole -ml msvc These compiler flags can be found at: http://www.lahey.com/docs/fujitsu%20compiler%20option%20list.pdf

To wrap up for Windows

Modification for c_tpsa_interface.F90

Added new tpsa package

Fixing the passing of a double array instead of an integer array.

Etienne’s clean-up of AF

Fix C/C++ nonstandard features fix pseudo-bug in c_tpsa_interface.F90 found by NAG f95. Integer shall not be defined as an array of dimension 1.

Latest cleanup of Lingyun’s TPSA including tpsa.dll needed for Windows

Clean-up

- gfortran broken in gcc4.4 - Therefore back to g95 however the LIBX flags must be fixed according to gcc being used

Fix the exclusion of f90 & F90 files with and without NTPSA

Yet another upgrade for MAC using g95

Clean-up for MAC

Generalization for f90/F90

Compiler preprocessing for c_tpsa_interface to drop DLL for LINUX

Add new tpsa

missing -fno-range-check flag for gfortran in particular for MAC

Back to standard "LF95"

Further adjustments to safeguard running with "gfortran" which can be steered with the new "SLC4" flag.

Fix further gfortran for SLC4

Default compiler oh lxplus: lf95 gfortran with -Wall -pedantic DEBUG flags even for Fortran

gfortran explanation for SLC4

Darwin fixes
no ONLINE as standard

More small fixes: - o4 off for lf95 - proper libraries for ONLINE
Fix gfortran Home link use proper LINK options for MAC
hbu
using semi automatic object file list
Makefile mac compatible

jbj
made more robust the capture of the Twiss data of the drift preceeding the current element - solving the '1st slice wrong n1’ supressed the memory crash when aperture called twice with the same offset elem in tfs format
nougaret

further prevent output mixup through forced flush when crossing the border in the other direction (i.e. Fortran calling C, which is more rare than the reverse)

handle flushing unit 6 on Intel ifort compiler

invoke 'call flush(6)' as 'flush(6)' with Intel compiler
suppress compilation warnings
skowron

Added filling of track summ table for ptc_trackline. Now the user can check what were the final coordinates of tracking

If plugin support: link dynamically; if debug: do not put -O4 optimization;
g95 option: add proper debug flags

Work log between releases madX-4_00_09 and madX-4_00_19

Log report started Mon May 4 20:02:18 2009, ended Mon May 4 20:03:41 2009

elaface
Fixed argument of wrong type passed to the mtlmdl Fortran subroutine
warning fixed
Now the check for OSTYPE works with darwin9.0
frs

1) New TPSA package by lingyun.yang@gmail.com 2) General clean-up
Preliminary fix of memory crash - courtesy JBJ

Some rearrangements by Etienne and final clean-up of PTC

GET_C_J routine no longer needed

Missing general "public" statements which creates pseudo bugs

More clean-up for the TPSA upgrade

Fortran Clean-up: indenting, remove potentially uninitialized variable and also remove unused variables

Minor clean-up concerning: character strings, possibly uninitialized variables and unused variables

ONLINE needs: libmdblib.a libmdbmth.a librpnlib.a libSDDS1.a libSDDS1c.a libz.a SLC5 needed: libX11.a

version madX-4_00_18_dev

Version 4.00.17: add LRAD to kickers

Version 4.00.16

Version 4.00.15

MAD-X version 4.00.14

MAD-X version madX-4_00_12_dev

version madX-4_00_12_dev

version madX-4_00_11_dev

version 4.00.10

version

Add attribute "range_plot" to allow range also for user defined horizontal axis (courtesy HG)

Allowing plotting at markers using the "marker_plot" attribute. Courtesy HG

Add ntpsa flag (if present the new C++ TPSA package by lingyun.yang@gmail.com in invoked) and the symprint flag pronts the symplecticity flag by default.

Remove "harmless" occurrence "dipole_bv" by the more obvious one "other_bv"

remove tabs
Clean-up unused variables

Take out: Unused external reference RESULT_FROM_NORMAL found with latest: NAG Fortran Compiler Release 5.2(668)

Minor clean-up

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handle flushing unit 6 on Intel ifort compiler

invoke 'call flush(6)' as 'flush(6)' with Intel compiler

supress compilation warnings

skowron

Added filling of track summ table for ptc_trackline. Now the user can check what were the final coordinates of tracking

If plugin support: link dynamically; if debug: do not put -O4 optimization; g95 option: add proper debug flags

Work log between releases madX-4_00_07 and madX-4_00_09

Log report started Fri Mar 27 20:02:08 2009, ended Fri Mar 27 20:03:03 2009
frs

New files needed for MAD-X Version 4

Clean-up

Version 4.00.09

version 4.00.08

No lonfer needed for MAD-X Version 4

Clean wipes out fortran wrapper stuff

New Makefiles for Linux/Mac & Windows include all previous features

Work log between releases madX-4_00_00 and madX-4_00_07


frs

total_da_size set to very large (courtesy Piotr & Etienne)

madX-4_00_07_dev & first candidate for the production version

madX-4_00_06_dev

Fix of the faulty REPLACE command (coutesy HG)

madX-4_00_05_dev

madX-4_00_04_dev

Fix of the memory crash due to the USE command in a while loop (found by EB - fixed courtesy HG)

MAD-X version: 4.00.02

Version 4.00.01; fixing the crash due to conflict with markers (courtesy HG)

Return NULL; needed to continue if the "no_fatal_stop" flag enforces program continuation

Harmon no longer ignored (courtesy HG)

Fix lethal bug in DELETE command (courtesy HG)

suppress forbidden TAB character

reorganize twbtin more logically
The chromatic functions wx, phix, wy, phiy could not be initialized properly for lines.

simplifying length fix

proper fix of doubling of the length of the machine with chrom & centre

suppress double length in summ table for chrom & currpos option

preparation for SLC5 in 32 & 64 bits

nougaret

Compute derivatives of the dispersion w.r.t. deltap
to scale from 32 to 64 bits platforms, obtain pointers size with sizeof(uintptr_t)
one-turn parameters such as the tune should now depend on deltap

merged Frank and Piotr’s modifications

skowron

Removal one line from generated ROOT macros for plotting since the ROOT command disappeared in the new versions

protection against seg fault in case the command is executed before ptc_create_universe

Bug in knobs corrected

Twiss table was extended, definitions for new new columns were added in ptc_madx_knobs.inc, but the code was not modified appropriately and uninitialized univ. taylors were left in the results array

Swapped putusertable with puttwisstable so we values from the user table are copied properly to the twiss table if requested. See ptc_secordmatch example in the testsuite.

With modern versions of ROOT libraries were split to more files. Updated the code to lead all that are needed in ROOT version 2.21

yisun

Remove two useless messages.

zwe

use beam_bv flag to change sign for beam two in simultaneous orbit correction
Modifications to adapt to beam1/beam2 conventions for two beam orbit correction.

Resolve problem with RESOUT option

Work log between releases madX-3_04_72 and madX-4_00_00

Log report started Mon Feb 16 07:02:11 2009, ended Mon Feb 16 07:03:06 2009
frs

readtable stops with fatal_error when file with table does not exist (courtesy HG)

production version: madX-4_00_00
version: madX-3_04_77
Version: madX-3_04_76_dev
version: madX-3_04_75_dev

Fix the matching problem and the faulty set-up of the twiss_chrom flag (courtesy HG)

Intermediate version with improvement concerning chromaticity including coupling. More work concerning a hiccup in the matching of the LHC still under way. (courtesy HG)

Initializing q1_val_p & q2_val_p clean-up to avoid compiler warning

Solenoid becomes a marker if the integrated strength is zero otherwise a fake very short is used.

Take out doogy format for strlen - trivial clean-up

The flag chrom can now be set on demand. (courtesy HG)

Fixing small bug concerning undefined deltap. (courtesy YS)

hbu

keep new parameter mech_sep in slicing

jbj

/* BJ 13.02.2009. - added check |x-e| < dist_limit - removed useless calculations of sqrt - made consistent use of dist_limit and min_double */ /*
BJ 13.02.2009 check if point x = (xm,ym) is in the segment [s,e] with s =
(startx,starty) and e = (endx,endy) by computing cosfi = (x-s).(x-e) /
|x-s| |x-e|. cosfi = -1 : x is in first check if |x-s| and |x-e| are not too small.If yes for one of them : in if OK , the zero divide check must be superfluous. But keep it anyway. */

Work log between releases madX-3_04_68 and madX-3_04_72

Log report started Sun Feb 8 07:01:42 2009, ended Sun Feb 8 07:03:01 2009
1) Fix lethal error (no consequence for LHC) 2) Put back "etall" into DA vector headers (both courtesy EF)

Fixing the Twiss chromaticity problem in presence of coupling by running Twiss twice with different deltap (which is set to 1e-9) and calculate it numerically. Good workaround! (courtesy HG)

MAD-X version: madX-3_04_71_dev
MAD-X version: madX-3_04_70_dev
Version: madX-3_04_69_dev

Fixing an inconsistent "TARGET" declaration found by NAG. (courtesy EF)

Putting back incorrect chromaticity calculation

1) Fix proper disp & ddisp definition for ring with respect to lines 2)
Suppress chromaticity calculation in case of coupling since it is plain wrong.

First MAD-X LHC commissioning version 3.04: 09.07.2007

1) Allowing reading and writing of SDDS data sets for communication with the LHC control system.
2) A large amount of medium size and smaller code changes and bug fixes.
3) Full kinase development for CLIC purposes.
4) New Jacobian matching method.
5) Non-linear and parametric matching.

All changes for each file:

====================================================================
MAD-X proper
---------
C Files:
--------
- minor C inconsistencies and some clean-up
- Uninitialized variables
- aperture.c
  - moved aperture code from maden.c to new file aperture.c
- corrections:
  - more apex in halo polygon

====================================================================
- corrected the construction of recelipces in the general case
- secured potentially dangerous division by zero

- The brute force quick and dirty fix of frs has been reverted for a proper
  fixing at the source of the problem, i.e. assigning different names for
different multipoles. It has been shown that SixTrack gives identical
results for the "brute force" and the "proper solution".
  Courtesy Hans Grote (honorable ABP group member)
- Fixing the problem of using the same name for very different multipoles.
  This is a quick fix and a more rigorous solution is needed.
- ISO/IEEE9890 were treated as thick elements in the single element list
  but later treated as thin elements leading to screwed up linear optics
  in SixTrack. They are kept as distinct elements and are not joined with
  surrounding drifts.

gxx11c,c gxx11psc
- fix the month number in the ps files
madxc,c madxp,c madxe,c
- New aperture code from madan.c to new file aperture.c
- corrections
- corrected the construction of recelipces in the general case
- secured potentially dangerous division by zero
- Avoid divisions by zero in the aperture module.
- Avoid C concatenation and some clean-up
- mad-X_3_66 bugs found by EK
- Array overflow reported from valgrind removed
- Special not-existing End marker has been dropped. courtesy HG
- macro mode compiling, three functions moved from matchc2.c to madan.c
- Correct table_list initialized to 0
- Add warning when user macro option is used with too many variables
- Introducing aptol_(1,2,3) "rtol", "atol", "ytol" to be available in MAD-X
  input and e.g. to be added to a TWISS table. Courtesy HG
- Now table(x,y) or stable(a,b,c) or things like that will not be modified.
  - First step for node input tracking is ptc_trackline
- Fix a bug when you use a macro which defines another macro for matching
  - maex-0.8.6: "Final" fix of the TILT saga! Tilt is calculated exclusively
  in twiss.c following the strategy:
  1) Some changed constants, next,oct,etc, separated into
     but NOT dipole or multipole
  2) TILT input in the external tilt: \( \text{+k} \rightarrow \text{+ks for tilt < 0} \)
  3) & k & ks represent an internal tilt
  4) at each element the total tilt & sqrt(k^2 + ks^2) is calculated
     including field errors, i.e. the correct way which might cause
     differences with MAD8
  5) TILT input has been adjusted appropriate
  6) Frankie effects on "error" and "wait" will still be tested
  7) Many Thanks for HG for his help!
  - Bug corrected, replaced with fabs
  - Updated match with macros
  - Constraints for ranges with match_use_macro implemented.
  - Debugging
  a) madan: Wrong sign of TILT when calculated from k & ks
  b) madan_ptc_module: Total rewrites of TILT stuff!
  - TILT clean-up courtesy HG
  - Moments calculation fully implemented, map buffering in ptc_twiss
  - New ptc_twiss, so A_ is tracked. This makes possible tracking of moments
    (to be computed).
  - Introducing aptol_(1,2,3) "rtol", "atol", "ytol" to be available in MAD-X
    input and e.g. to be added to a TWISS table. Courtesy HG
  - Adding gxx option following madp_ptc_script_module

  Problems:
  1) not displayed in table.
  2) "delay out in ptc_twiss header unless a twist command was done before
     ptc_twiss".

  Solution:
  1) Set up p properly
  2) Put "delay" into header of the ptc_twiss table permanently
  - Fix routine read Gotham by exchanging with and with:
  - Fixing the plot: the header is not read if it does not exist
    (courtesy HG):
    1) Stefan Sorge new module keeper!!!!
  2) Second order detuning with proper "hor" and "ver" names. In fact,
     vertical terms were missing.
  3) Bad memory bug found by RM. Piotr found using valgrind the solution:
     in ptc_twiss.c, internal use of the "TILT" routine is essential
     because the character string has a \#["TILT at the end.
  4) In plot there is still the crash when reading the header - temporary
     solution applies:
  - Fix the readtable routine which stumbled over a blank line. (courtesy HG)
  - First step for node input tracking is ptc_trackline
  - Adding a c routine to read headers of TFS tables (courtesy HG)
  - Taking delay from table header both for TWISS and ptc_twiss (but
    NOT in case of a SUMM table!) and place on the plot.
  - Move the net_variable routine from madan.c to madan.c. Needed to compile
    madxe.c
  - New setvar command, fixed readtable bug
  - Removed redundant "delay" printout.
  - Exact same matching implemented, now passing name with x
  - Function assigned, the same as name of node in MAD but with capital letters
  - Updated match with macros
  - Constraints for ranges with match_use_macro implemented
  - Adding "e1", "e2", "h1", "h2", "hgap", "fint", "fintx" to the twiss table
  - Removed charge setting to the my_ring layout to make ptc_twiss running,
    redundant printouts removed
  - Moment calculation fully implemented, map buffering in ptc_twiss
  - Few bugs corrected (e.g. map initialized to null instead of spare when
    initial twiss provided). Moments seem to work (to be tested yet)
For further details see the comments at the top of matchptcknobs.c file.

- Implemented:
  - Redundant printf removed
  - change in the print out of match summary when USE_MACRO
  - Corrected bug in match use_macro in match2_evaluate_expressions
  - Increased number of constraints
  - New jacobian routine with svd. Option COND added for controlling the SVD.
  - add warning when use_macro option is used with too many variables
  - fix compilation warnings
  - Constraints for ranges with match,use_macro; implemented
  - Updated match with knobs
  - mpar made compiling, three functions moved from matchc2.c to madxu.c
  - Printouts redirected toprt_file
  - Windows "CL".
  - Comment corrected
  - Experimental knob file generation
  - Not enough memory for buffers ==> FODO: LINE={34*(CELL)}; failed when cell was a line itself
  - cleanup of gino
  - Adding gino command following make ptc_script_module
  - PTC knobs (pol_blocks) almost completely interfaced to MAD-X.
  - New ptc_normal accepted in matching with ptcknob: bug corrections
  - Changing the conflicting "ksl" for the integrated solenoid strength to "ksi".
    This name is reserved for the vector of the integrated skew "ksi" (not "ksl")
  - Correct bug in read_table: can read long integers now
  - Solution: courtesy Hans Grote
  - "string function" tabstring count start at 1 courtesy Hans Grote
  - Printout cleanup
  - changing table entries to more logical names
  - closing unit 34 to allow multiple SODD runs
  - Sceleton for knobs and arbitrary element placement implemented. Lattice
    visualization via ROOT macro. Printing detailed lattice geometry in PTC.
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  - Sceleton for knobs and arbitrary element placement implemented. Lattice
    visualization via ROOT macro. Printing detailed lattice geometry in PTC.
1. Minor corrections and protections against segmentation violation.
   - Error flag is monitored in main so if an error occurs during macro execution it is handled appropriately.

matchptcknobs.c
- Functions defined elsewhere should be defined with extern so linker does not complain about multiple definitions
- Algorithm made more stable
- Typo corrected
- Introduced correct treatment of magnet families
- Protection against deletion of NOLL pointer added
- Parameter matching of initial conditions works now, final tests and debugging to be done
- Updated match with knobs
  - New ptc_twiss, so A_ is tracked. This makes possible tracking of moments (to be completed).
  - Corrected bug in match_use_kmacro in match2_evaluate_expressions
  - New ptc_normal accepted in matching with ptcknobs, bug corrections
  - Implemented:
    1. ptc_setfieldcomp that set any order field strength to requested value. It enables matching of higher order field components.
    2. Special matching mode use_ptcknob. It implements kind of macro that emplys parametric FTC calculations to perform matching in a faster manner.
    3. New ptc corrections and protections against segmentation violation.

rplot.c
- In some systems it is needed to load manually all needed ROOT libraries
- Bug correction
- Now also ptc_normal accepted in matching with ptcknobs, bug corrections
- Code counting
- Turn number added in rplot
- rviewer plugin interface now on rplot is a plugin instead of compiled in optional code
- New bug in knobs corrected; attempt to delete not properly associated taylors in case no knobs are set by the user

C Header Files:
- mod.h
  - Add parameter (units) for orbit correction
  - Added: Knowledge type for matching mode, protection against multiple inclusion.

modl.h
- Definitions for tables used by Slice Tracking with PTC
- Momentum compaction "Alpha" included into TWD table for matching.
- Add "polarity" parameter to the twins table
- Updated match with knobs
  - Adding "a1", "a2", "a1", "a2", "b1", "b2", "f1", "f2" to the twins table
  - Supressing "maxa" in favor of "null" - request by Thys Kessels
  - Adding node value "maxa" (maximum K value) and "maxb" (maximum current value)
- New tracking feature by Andres Gomez Alonso:
  - Using flag "prio" in the tracking command creates a table called "tracklist", which keeps a record of lost particles. It can be saved using the *"write, table=tracklist"* command.
  - Added madx data types , mainly used in 2003 module
  - New function "prio" courtesy Hans Grote
  - Odd table names using small letters only
  - Closing unit 34 to allow multiple SODD runs

madx.h
- Experimental knob file generation
  - Added option that tells to free memory at the end of the program execution.
  - For option to printout tracked table added that switches on/off track parameters storage in memory for every slice
  - Fix traditional matching of alf
  - First step for node layout tracking in ptc_trackline
  - Added ptc names of twins functions to constraint, hence one can set constraint in a range
  - New setvar command, fixed readtable bug
  - Add "polarity" parameter to the twins table
  - Clean-up
  - winp.f90
- Fixing the crash for dew > max = multipoles larger than 10. This set-up requires to solve Maxwell's equation up to SECTOR_NMUL_MAX. The default is set to 10 to avoid excessive computing time. This is now safeguarded in code.
- To this end the parameters SECTOR_NMUL and SECTOR_NMUL_MAX are transferred from "ptc_create_layout" to "ptc_create_universe" such that these global parameters can be set early enough. Internally in PTC the parameter "lda_used" is increased where needed from 1500 to 3000 and set back.
- Moreover Kriste has done the following modifications to make this possible:
  - The modification I made in the new PTC I sent you are as follows:
2. Special matching mode use_ptcknob. It implements kind of macro
that emplys parametric PTC calculations to perform matching in a faster manner.
For further details see the comments at the top of matchptcknobs.c file.
3. Minor corrections and protections against segmentation vilation.
   - PTC knobs (pol_blocks) almost completely interfaced to MAD-X.
   - User sets a knob with ptc_knob command.
   - Twiss parameters and user specified (with ptc_select) map components are
     buffered in memory after every element in form of taylor series.
     They can be dumped to text file in two formats with ptc_printparametric command.
     They can be also visualized and further studied with rviewer from rplot plugin.
   - Further, user can set numeric values of knobs with ptc_setknobvalue what
     updates all numeric values of the parameters in the tables.
     This way knobs can be used in matching.
   - New tracking feature by Andres Gomez Alonso:
     Using flag "recloss" in the tracking command creates a table called
     "trackloss", which keeps a record of lost particles. It can be saved
     using the "write, table= trackloss" command.
   - change in beambeam command: usage of scattering beam
     with different radial shapes is possible:
     parameters:
     - knobs implemented with PTC with pol_blocks; command to dump parametric
       results to file or stdout; content of ptc_madx_tablepush.f90 moved to
       ptc_madx_knobs.f90, the former one removed
     - Element placement options added
       - Commands for 3D read and write
       - Skeleton for knobs and arbitrary element placement implemented. Lattice
         visualization via MAD-X macros; printing detailed lattice geometry in PTC.
         Several small bug corrections and mere code cosmetics.
         - introducing FORTRAN and MAD-X in PTC. To this end 2 flags have been
           introduced in the MAD-X dictionary madxdict.h:
           1) ptcblock: if true it use a PTC type REND
           2) truerbend: if true it use THWRENDB; if false it uses WEDGRBEND
           - option for ptc_trackline position given in global coordinate system added.
           - changing the conflicting "ksi" for the integrated solenoid strength to
             "ksi": this name is reserved for the vector of the integrated skew
             multipoles "ksl=[]". Thin solenoids can now have normal "ksi" and skew
             "ksi" multiple errors in PTC, ignored in madx proper. Thin solenoids
             are presently not considered in PTC.
           - PTC_Enforce6D implemented
           - defaultlevel default has been 0 now 1 as planned originally
matchptcknobs.h
   - Updated match with knobs
   - Now also ptc_normal accepted in matching with ptc_knobs; bug corrections
   - Implemented:
     1. pc_setfieldcomp that set any order field strengh
        to requested value. It enables matching of higher order field components.
   2. Special matching mode use_ptcknob. It implements kind of macro
      that emplys parametric PTC calculations to perform matching in a faster manner.
      For further details see the comments at the top of matchptcknobs.c file.
   3. Minor corrections and protections against segmentation vilation.
pilot.h
   - Code cosmatics
   - Turn number added in plot
   - rviewer plugin interfaced, from now on rplot is a plugin instead of compiled in
     optional code
   - Sever bug in knobs corrected: attempt to delete not properly assocoated taylors
     in case no
knobs are set by the user
sdds.h
   - New files for MAD-X On-Line Modeling Version
FORTRAN Files:
-------------
Changes:
--------
emit.F and all Fortran Files
   - Clean-up of unused variables
   - Fortran Clean-Up
match.F
   - tentative new output for matching var
matchdc.F
   - better scaling Jacobian
   - Penalty Function printed with twice larger precision
   - New Jacobian routine with opt. Option COND added for controlling the SV.
   - Increased number of constraints
   - change in calls behavior for JACOBIAN
matchdf1.F
1) Just the routines needed => 5 times smaller; 15% => 10%
2) Fortran90 Clean-up
   - Changes needed to compile routine daniol without optimization in extra file
matchdf2.F. Otherwise code gets stuck in matching procedures. In all Linux
Makefiles matchdf2.F is compiled when using pg7. For the Fortran90 compilers
gfortran (95, 96, 95.000), and gfortran as optimised routine is used as provided by
carlos tresoldi. (Note: For use of gfortran the option -fno-automatic, -fno-calling-seq.
-if "-O0.txt" was needed) special undocumented Fujitsu compile flag of
the compiler (1995) to compile this special Fortran90 version of daniol.
   - Replacing the DIAMIC by an improved Fortran90 use as proposed by
Andres Gomez Alonso.
matchdf3.F
   - remove Makefile map offending code
   - New Jacobian routine with opt. Option COND added for controlling the SV.
   - Increased number of constraints
matchdf4.F
   - Needed to compile routine daniol without optimization. Otherwise code
     gets stuck in matching procedures.
1) TILT input is the external tilt (+k == +ks for tilt < 0)
   but NOT dipole or multipole.
2) These changes concern quad, sext, oct, elec separator;
   in twiss.F following the strategy:
   - Add "polarity" parameter to the twiss table
   - Momentum compaction "alfa" included into TWISS table for matching.
   subroutines tmbb_flattop and tmbb_hollowparabolic in twiss.F
   flattop and hollow parabolic radial density profile, i.e. in the
   - Closed orbit implemented in the maps for the beambeam element with
   twiss.F are presently not considered in PTC.
   "ksl" multipole errors in PTC, ignored in madx proper. Thin solenoids
   - Changing the conflicting "ksl" for the integrated solenoid strength to
   for bbshape=3: fractional width of the parabolic part
   width:   for bbshape=2: fractional width of edge region
   3 hollow-parabolic
   bbshape: 1 (default) Gaussian, standard as before
   parameters:
   - change in beambeam command: usage of scattering beam
   - "trackloss", which keeps a record of lost particles. It can be saved
   using flag "recloss" in the tracking command creates a table called
   - New tracking feature by Andres Gomez Alonso:
   make -f Makefile_nag. Fixed.
   - Inconsistent variable declarations of z0 encountered by Piotr when using
   - Some unnecessary changes taken out again by Andres Gomez Alonso.
   - PTC knobs (pol_blocks) almost completely interfaced to MAD-X.
   User sets a knob with ptc_knob command.
   Tokens parameters and user specified with ptc_select map components are
   in memory after every element in form of Taylor series.
   They can be changed to text file in two formats with ptc_parametriccommand.
   They can be also visualized and further studied with viewer from split plugin.
   Further, user can set numeric values of knobs with ptc_setknobvalue what
   updates all numeric values of the parameters in the tables.
   This way knobs can be used in matching.
   - knobs implemented with PTC with pol_blocks command to dump parametric results
to file or stdout; content of ptc_map_table in ptc제도 move to ptc_mds_table.F. in
   the former one removed
   knobs implemented with PTC with pol_blocks; command to dump parametric results
   to file or stdout; content of ptc_mds_table in ptc제도 move to ptc_mds_table.F. in
   the former one removed
   - Solution for knobs and arbitrary element placement implemented. Lattice
   visualization via ROOT macros. Printing detailed lattice geometry in PTC. Several
   small bug corrections and some code cosmetics.
   - PTC_Enforce6D implemented
   small bug corrections and some code cosmetics.
   - Skeleton for knobs and arbitrary element placement implemented. Lattice
   visualization via ROOT macros. Printing detailed lattice geometry in PTC. Several
   small bug corrections and some code cosmetics.
   - PTC_Enforce6D implemented
   - New ptc_twiss, so A_ is tracked. This makes possible tracking of moments
   (not to be completed).
   - User sets a knob with ptc_knob command.
   - Implemented:
   1. ptc_parametric command that set any order field strength
   to requested value. It enables matching of higher order field components.
   2. Special matching code used inside. It implements kind of macros
   that employs parametric PTC calculations to perform matching in a faster manner.
   For further details see the comments at the top of ptc_parametric command.
   - PTC knobs (pol_blocks) almost completely interfaced to MAD-X.
   User sets a knob with ptc_knob command.
   Token parameters and user specified with ptc_select map components are
   buffered in memory after every element in form of Taylor series.
   They can be changed to text file in two formats with ptc_parametric command.
   They can be also visualized and further studied with viewer from split plugin.
   Further, user can set numeric values of knobs with ptc_setknobvalue which
   updates all numeric values of the parameters in the tables.
   This way knobs can be used in matching.
   - knobs implemented with PTC with pol_blocks command to dump parametric results
to file or stdout; content of ptc_mds_table in ptc제도 move to ptc_mds_table.F. in
   the former one removed
   - Solution for knobs and arbitrary element placement implemented. Lattice
   visualization via ROOT macros. Printing detailed lattice geometry in PTC. Several
   small bug corrections and some code cosmetics.
   - PTC_Enforce6D implemented
   - Suppression of excessive printing (courtesy SS)
   1) Stefan Sorge new module keeper!!!!
   2) Second order detuning with proper "hor" and "ver" names. In fact,
   vertical terms were missing.
   - Close unit 34 even on error output
   - odd.F
   - Argument *ax* was removed from argument list of subroutine trnum called
   here (line 532 in trnum.F), because it did not coincide with the argument
   of the real subroutine trnum in trnum.F.
   - Bug found by Stefan Sorge in trnum/trinumid: Variables "bsol" and "bsolid" were
   undefined, leading to erroneous results. Corrected.
   - Fixing: Unwanted changes commented.
   - The aperture of the collimators was checked before undergoing the (possible)
   rotation at the entry of the element. Corrected now: If collimator has a roll
   angle ('tilt'), coordinates are transformed to the rotated local coordinate
   system and only after that apertures are checked.
   - Add MY flag to the modules
   - Some unnecessary changes taken out again by Andres Gomez Alonso.
   inconsistencies in variable declaration of s0 encountered by Piotr when using
   make -f Makefile_nag. Fixed.
   - New tracking feature by Andres Gomez Alonso:
   - Preliminary fix of the crash when the aperture table is used
   - Some unnecessary changes taken out again by Andres Gomez Alonso.
   - Inconsistent variable declaration of s0 encountered by Piotr when using
   make -f Makefile_nag. Fixed.
   - New tracking feature by Andres Gomez Alonso:
   - Preliminary fix of the crash when the aperture table is used
   - Some unnecessary changes taken out again by Andres Gomez Alonso.
   - Inconsistent variable declaration of s0 encountered by Piotr when using
   make -f Makefile_nag. Fixed.
   - New tracking feature by Andres Gomez Alonso:
   - Preliminary fix of the crash when the aperture table is used
   - Some unnecessary changes taken out again by Andres Gomez Alonso.
   - Inconsistent variable declaration of s0 encountered by Piotr when using
   make -f Makefile_nag. Fixed.
   - New tracking feature by Andres Gomez Alonso:
   - Preliminary fix of the crash when the aperture table is used
   - Some unnecessary changes taken out again by Andres Gomez Alonso.
   - Inconsistent variable declaration of s0 encountered by Piotr when using
   make -f Makefile_nag. Fixed.
   - New tracking feature by Andres Gomez Alonso:
   - Preliminary fix of the crash when the aperture table is used
   - Some unnecessary changes taken out again by Andres Gomez Alonso.
   - Inconsistent variable declaration of s0 encountered by Piotr when using
   make -f Makefile_nag. Fixed.
   - New tracking feature by Andres Gomez Alonso:
   - Preliminary fix of the crash when the aperture table is used
   - Some unnecessary changes taken out again by Andres Gomez Alonso.
   - Inconsistent variable declaration of s0 encountered by Piotr when using
   make -f Makefile_nag. Fixed.
   - New tracking feature by Andres Gomez Alonso:
2) & ks represent an internal tilt 
3) at each element the total tilt k sqrt(k**2 + ks**2) is calculated 
   including field errors, i.e. the correct way which might cause 
   differences with MAD8
4) PTC has been adjusted appropriate 
5) Possible effects in "survey" and "emit" will be tested 
   - safeguard atan2 against both arguments equal to zero
   - Fixing TILT in multipole kick and make TILT proper in thick octupole.
   - Change in beambeam command: usage of scattering beam 
     with different radial shapes is possible:
     parameters: 
     bbshape: 1 (default) Gaussian, standard as before 
     2 flattop (or trapezoidal) 
     3 hollow-parabolic 
     width: for(bbshape)=2: fractional width of edge region 
     for(bbshape)=3: fractional width of the parabolic part 
   - Change in "ksl" for the integrated solenoid strength to 
     "ksi". This name is reserved for the vector of the integrated skew 
     multipoles "kx,ky;". This misnomer can now have normal "ksi" and skew 
     "ksl" multiple errors in PTC, ignored in madx proper. Thin solenoids 
     are presently not considered in PTC.
   - Add BV flag to the solenoids 
   - Suppressing "imax" in favor of "calib" - request by Thys Risselada 
     - Adding node value "kmax" (maximum K value) and "imax" (maximum Current value) 
     - Add BV flag to the solenoids 
     - Adding node value "kmax" (maximum K value) and "imax" (maximum Current value) 
     - Add BV flag to the solenoids 
     - Correction of an error occurring in subroutine tmbb_flattop for ftrk=.false.
     - Unused dummy variable IMASS in subr. photon is removed
     - Added seterrorflag routine that sets the error flag in c part if an error 
       occurred 
     - Matplotlib files: 
       - Clean-up of fi files 
       - Momentum compaction "zmax" included into TWISS table for matching. 
       - Add "polarity" parameter to the twiss table 
     - New ptc_twiss, so A_ is tracked. This makes possible tracking of moments 
       (to be completed). 
PTC MODULES 
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Changes: -------
- Bug Corrected: Parametric twiss results where not scaled with energy 
  - Moments calculation fully implemented; map buffering in ptc_twiss 
  - New ptc_twiss, so A_ is tracked. This makes possible tracking of moments 
    (to be completed).
  - Implemented: 
    1. ptc_setfieldcomp that set any order field strengh 
       to requested value. It enables matching of higher order field components. 
    2. Special matching mode use_ptcknob. It implements kind of macros 
       that emplys parametric PTC calculations to perform matching in a faster manner. 
       For further details see the comments at the top of matchingptcknobs.c file. 
  - Added header lines to ROOT macro that display layout geometry. This makes 
    possible to compile a macro, what is a must in the case of lengthy machines. 
  - Removed charge setting to the my_ring layout to make ptc_twiss running, 
    redundant printouts removed 
  - Few bugs corrected (f.g. map initialized to nd2 instead of npara when 
    initial twiss provided). Moments seem to work (to be tested yet) 
  - New ptc_twiss, so A_ is tracked. This makes possible tracking of moments 
    (to be completed).
  - add (lp) to logical function; indenting
- Enforce 6d implemented
- PTC_Enforce6D implemented
- defaultlevel default has been 3 now i as planned originally

madx_ptc_knobs.f90
- Bug Corrected: Parametric twist results where not scaled with energy
- Exact name matching implemented, now passing name with 
  - knobs for Initial parameters
  - New ptc_twiss, so A_ is tracked. This makes possible tracking of moments
    (to be completed).
- PTC first changes stay October 2004 thussless, cutting
  - Updated to the new nomenclature (beta12->beta22,...)
  - Bug corrected in the treatment of 4D and 5D cases; cosmetics;
  - Universal taylor nullified at the initialization level
  - Implemented:
    1. ptc_twissdump that set any order field strength
       to requested value. It enables matching of higher order field components.
      2. Special matching mode use_ptcknob: It implements kind of magic
         that empies parametric PTC calculations to perform matching in a faster manner.
      3. Minor corrections and protections against segmentation violation.
        - don't print to file trailing blanks in the buffer

- PTC knobs (pol_blocks) almost completely interfaced to MAD-X.
  - user sets a knob with ptc_knob command.
  - Twists parameters and user specified (with ptc_select) map components are
    buffered in memory after every element in form of taylor series.
  - They can be dumped to text file in two formats with ptc_prinptcparametric command.
    Can be also visualized and further studied with viewer from plot plugin.
  - Run can be set to stop at first set of knobs with ptc_setparameter what
    updates all numeric values of the parameters in the database.
  - This can be done in shell in red:
  - Inverse bug in knobs corrected: attempt to delete not properly associated
    knobs in case no knobs are set by the user.
  - knobs implemented with PTC with pol_blocks; command to dump parametric
    results to file or screen; content of ptc_madx_tablepush.f90 moved to
    ptc_madx_knobs.f90, the former one removed
  - debug info printed only at appropriate debuglevel
  - skeleton for knobs and arbitrary element placement implemented.
  - lattice visualization via ROOT macro. Printing detailed lattice geometry in PTC.
  - Several small bug corrections and some code cosmetics.

madx_ptc_module.f90
  - in many treated as TN a knn now
  1) madx_ptc_module without the ptc_normal stuff
  2) fixing the "password" bug
     - Fix to restricted ptc output format
  3) be_mstar: preliminary fix of uninitialized variable NADVERTISE_MER
  4) madx_ptc_module: Fix of ptc_normal by fixing the string comparison
  5) madx_ptc_module: madx_ptc_rais: write a read traditional %A map format

PTC version including spin
- madx-3.01.Rr: "Final" fix of the TILT maps! Tilt is calculated exclusively
  in twiss.f following the strategy:
  1) These changes concern quad, next, oct, elec separator;
     but NOT dipole or multipole
  2) TILT input is the external tilt (+k ==+ks for tilt t 0)
  3) s & k be represent an internal tilt
  4) at each element the total tilt & sqrt(k**2 + ks**2) is calculated
     - the correct way which might cause differences with MAD-X
  5) If ptc has been adjusted appropriate
  6) Possible effects on "survey" and "emit" will be tested
  7) Many Thanks for HG for his help!
  - Reduced debug info available only in high level debug mode
  - VORNAME assigned, the same as name of node in MAD-X but with capital letters
  - Too long line split into 3 lines
  - useless print statements
  - Fixing the crash for madx + exact + multiples larger than 10. This set-up
    requires to solve Maxwell's equation up to SECTOR_NMUL_MAX. The default is
    set to 10 to avoid excessive computing time. This is now safeguarded in
    madx. To this end the parameters SECTOR_NMUL and SECTOR_NMUL_MAX are
    transferred from "ptc_create_layout" to "ptc_create_universe" such that
    these global parameters can be set early enough. Internally in PTC the
    parameter "lda_used" is incremented where needed from 1500 to 3000 and set
    back. Moreover Etienne has done the following modifications to make this
    possible:
    The modification I made in the new PTC I sent you are as follows:
    1) PTC_Enforce6D implemented to order SECTOR_NMUL_MAX. For all
       multiples <= SECTOR_NMUL then Maxwell's is solved to order SECTOR_NMUL_MAX.
       For multiples above SECTOR_NMUL, they are treated as a straight
       So if you have written to order 10, you may work with Maxwell's only to order SECTOR_NMUL and
       Maxwell's 10 as far as Maxwell's is concerned.
     2) Special matching mode use_ptcknob. It implements kind of magic
       that empies parametric PTC calculations to perform matching in a faster manner.
       3) minor corrections and protections against segmentation violation.
      - updated match with knobs
      - Fix unused madx definition
      - Missing declarations of dpt and dpt added
      - Debugging
        - madx:
          Wrong sign of TILT when calculated from k & ks
        - madx_ptc_module: Total exercise of TILT stuff
          - Corrected code for lab
        - In 6D, before setting in internal state only,6,D we remove delta, otherwise
          delta stays
          - replacing "maxn" by "*max2" thereby fixing the sign for quad, next, oct
          - set my_fetchargs="" [preliminary fix]
          - removed change setting to the my_fetch interface to make ptc_madx running,
            otherwise it crashes.
          - Moments calculation fully implemented, map buffering in ptc_twiss
          - Initialize moments and charge of MAD-X before set_madx
        - new ptc_twiss, so A_ is tracked. This makes possible tracking of moments
          (to be completed).
        - Initial orbit NOT closed orbit for initial betas
        - In case of instability in normal form, the code sets the global error
          flag and returns to the main command loop instead of fatal
        - fulfilled formalistic request for a change of the definition of nomenclature
          of the ptc_twiss variables: beta, alpha and gamma
      - Making unstable behavior in Normal form a fatal error
      - All PTC track commands and NormalForm executions are checked for unstable behavior

- 5) Possible effects on "survey" and "emit" will be tested
- 4) PTC has been adjusted appropriate
- 3) madx_ptc_module & madx_ptc_twiss: write & read traditional DA map format
- 2) madx_ptc_module: Fix of "ptc_normal" by fixing the string comparison
- 1) Se_status: preliminary fix of uninitialized variable RADIATION_NEW

madx_ptc_knobs.f90
- defaultlevel default has been 0 now 1 as planned originally
- PTC_Enforce6D implemented
- Updated to the new nomenclature (beta12->beta22,...)
- Bug Corrected in the treatment of 4D and 5D cases; cosmetics;
  - Implemented:
    1. ptc_twissdump that set any order field strength
       requested value. It enables matching of higher order field components.
First fill user tables and at the end TWISS table.
- Implemented:
  1. ptc_set avaliação that set any order field strength to requested value. It enables setting of higher order field components.
  2. Special matching node use point. It implements kind of syncs that employ parametric FTC calculations to perform matching in a faster manner.
- For further details see the comments at the top of madxptc_knobs.c file.
- Minor corrections and protections against segmentation violation.
- FTC books (pol_blocks) almost completely interfaced to MAD-X.
User sets a knob with ptc_knob command.

Twice parameters and user specified (with ptc_select) map components are buffered in memory after every element in form of taylor series. They can be dumped to text file in two formats with ptc_printparametric command.
They can be also visualized and further studied with viewer from rplot plugin.
Further, user can set numeric values of knobs with ptc_setknobvalue what updates all numeric values of the parameters in the tables.

This way knobs can be used in matching.
- Several bugs in knobs corrected: attempt to delete not properly associated taylor in case no knobs are set by the user
- Knobs implemented with FTC with pol_blocks command to dump parametric results to file or stdout; context of ptc_madx_tablepush.f90 moved to ptc_madx_knobs.f90, the former one removed.
- Corrected taylor with parameters
- Selection for knobs and arbitrary element placement implemented.
- Lattice visualization via ROOT macro. Printing detailed lattice geometry in FTC.
- Several small bug corrections and some code cosmetics.
- Introduced HIERARCHY and HIERARCHY in FTC. To this end 2 flags have been introduced in the MAD-X dictionary madxdict.h:
  1) ptcrbend: if true it uses HIERARCHY, if false it uses HIERARCHY
  2) treedist: if true it uses HIERARCHY, if false it uses HIERARCHY
- logical lp -> 4 and vice versa so NAG does not cry
- problem causing compiler warning removed
- open and close of unit 21 only for "ptcknob" now
- Error flag implemented that signals that error code occurred
- Changing the conflicting "ksi" for the integrated solenoid strength to "ksi". This name is reserved for the vector of the integrated skew multipolar "ksi'". This solenoids can have normal "ksi" and skew "ksi" multiple errors in FTC, ignored in madx proper. This solenoids are presently not considered in FTC.
- Check of initial conditions provided by the user on input.
- FTC_defdefault implemented. If 6D TWISS calculation is performed with initial conditions (beta0 block) then non-zero beta is required
- Fix writing 5/5 components of closed_orbit to twiss table
- Bug corrected: writing maps should not only in debug mode

madx_pcnormal.f90
- Parsed out the stuff for the "pc_normal" module since "madx_pcnorm" is already very large.

madx_pcs3rcscript.f90
- Adding gino command following madx_pcs_script_module

madx_pcs_options.f90
- Cleaned code so NAG warnings are minimized now: mainly unused variables
- RF cavity treated as WN cavity now, bug correction
- temporary fix of non-existing "ksi" member
- RF cavity treated as WN cavity now
- Error flag implemented that signals that error code occurred

madx_pcs_tablepush.f90
- Bugs implemented with FTC with pol_blocks command to dump parametric results to file or stdout; context of ptc_madx_tablepush.f90 moved to ptc_madx_knobs.f90, the former one removed.
- Selection for knobs and arbitrary element placement implemented.
- Lattice visualization via ROOT macro. Printing detailed lattice geometry in FTC.
- Several small bug corrections and some code cosmetics.
- Proper handling of 6D: 5th column and row are swapped with 6th ones.
- Bug corrected: writing maps should not only in debug mode

madx_pcs_track_run.f90
- Unused dummy variable in madx photons is removed
- Unused variables detected with Makefile_log are removed
- The re-initialization of the buffer
- IHARM6 created a fatal bug, fixed by setting IHARM6=5.
- IHARM other than 6, 1, 5, 15, 51, 4 throws a fatal error
- For safety a "Forgotten nose" statement plugged into every subroutine and also every function, independently if strictly needed or not!
- FTC.Track tables: The last element in MAD (not PTC)!
- Mat-tracks by TRACK of FTC are blocked and tracking is terminated
- Serious crash in ptc_track for unstable particles due to uninitialized FTC aperture check variable.
- Corrected from windows to unix encoding
- Bug in madx_pcs_track_run.f90 for IHARM=5 is fixed (num. of Ieigen >= 4)
- Debug output only at Debuglevel=4

madx_pcs_tracktabs.f90
- Removed bug in node tracking making impossible tracking for closed layouts
- First step for node layout tracking in ptc_tracktabs
- New ptc_tabs, so A, is tracked. This makes possible tracking of moments (to be completed).
- Track global coordinates corrected -> position is given with respect to fiber not magnet
- Task number added in gino
- Added switch to ptc_tracktabs command so user can choose if tracks should start from a single. The feature is only accessible if gino plugin is installed and madx is compiled with plugin support.
- No error for knobs and arbitrary element placement implemented.
- Lattice visualization via ROOT macro. Printing detailed lattice geometry in FTC.
- Several small bug corrections and some code cosmetics.
- Change "logical(lmp)" to the correct "logical(lmp)"
- Error flag implemented that signals that error code occurred, option for track position given is global coordinate system added.
- Debug level 9 removed completely

madx_pcs_twiss.f90
- madx_pcs_twiss.f90: bugs found by RH
- Problem:
1) gt not displayed in table.
2) Delrap act in ptc_twiss unless a twiss command was done before ptc_twiss!
Solution:
1) Set up p properly.
2) Put "&output" into header of the ptc_twiss table permanently.
   - Fix routine readmatrix by exchanging &it and &acti
   - Adding delrap to ptc_twiss in case of initial conditions.
1) make_ptc_module without the ptc_normal stuff
2) fixing the "substring" bug
   - Don't display output in debug 0 mode
1) p0c: preliminary fix of uninitialized variable RADIATION_HM
2) make_ptc_module: Fix of "ptc_normal" by fixing the string comparison
3) make_ptc_module & make_ptc_twiss: write a read traditional 3a map format
   - Tips twiss becomes "public" to overcome problem with q95. Probably okay in g95 but may be overly picky.
   - Bug corrected q95 was written to twiss table instead of energy.
   - Bug corrected: Parameter twiss results where not scaled with energy
   - Few protections against seg faults added. Redundant debug info available only in high level debug mode.
   - knobs for Initial parameters
   - Moments initialization for 5D in twiss, moments not available for SECTOR_NMUL_MAX = 10.
   - Removed charge setting to the sey_ring layout to make ptc_twiss running, redundant parameters removed.
   - Moments calculation fully implemented, map differing in ptc_twiss
   - New ptc_twiss to SECTOR_NMUL_MAX = 10 instead of errors when initial values provided. Moments seem to work (to be tested yet)
   - New ptc_twiss to A_ is tracked. This makes possible tracking of moments (to be completed).

wrap.f90
   - needed changes to accept new "make_ptc_normal_module".
   - First step for node layout tracking in ptc_trackline
   - Clean-up.
   - Added feature that allows to set values of several knobs and only at the end recalculates in tables.
   - Normally all tables are recalculated after setting a new value.
   - However, it does show parametric matching.
   - New command ptc_refreshtable
   - Updated match with knobs.
   - In addmom, t and delta swapped so the MADX input corresponds to the MADX nomenclature.
   - Moments calculation fully implemented, map differing in ptc_twiss
   - New ptc_twiss, A_ is tracked. This makes possible tracking of moments (to be completed).
   - Etienne's own stuff
   - Adding gin command following make_ptc_script_module.
   - Implemented.
1. ptc_setfieldcomp that set any order field strength to requested value. It makes matching of higher order field components.
2. Special matching mode use_gt95k. It implements kind of match that employs parametric PTC calculations to perform matching in a faster manner.
   - For further details see the comments at the top of matchptcknobs.c file.
3. Major corrections and protections against segmentation violation.
   - PTC knobs (pol_blocks) almost completely interfaced to MAD-X.
   - User sets a knob with ptc_knob command.
   - Twiss parameters and user specified (with ptc_select) map components are buffered in memory after every element in form of Taylor series.
   - They can be dumped to text file in two formats with ptc_printparametric command. They can be also visualized and further studied with viewer from gptk plugin.
   - Further, user can set numeric values of knobs with ptc_setknobvalue what updates all numeric values of the parameters in the tables.
   - This way knobs can be used in matching.
   - knobs implemented with PTC with pol_blocks; command to dump parametric results to file or stdout; content of ptc_madx_tablepush.f90 moved to ptc_madx_knobs.f90, knobs implemented with PTC with pol_blocks; command to dump parametric results to file or stdout; content of ptc_madx_tablepush.f90 moved to ptc_madx_knobs.f90, the former one removed
   - Skeleton for knobs and arbitrary element placement implemented. Lattice visualization via ROOT macro. Printing detailed lattice geometry in PTC. Several small bug corrections and some code cosmetics.
   - PTC_Enforce6D implemented

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Changes:

- PTC May 2007
  - VORMAKE assigned, the same as some of node in MAD but with capital letters
  - Fixing the crash for den = n mult = 10. This set-up requires to solve Maxwell's equation up to SECTOR_NMUL_MAX. The default is set to 10 to avoid excessive computing time. This is now safeguarded in madx. To this end the parameters SECTOR_NMUL and SECTOR_NMUL_MAX are transferred from "ptc_create_layout" to "ptc_create_universe" such that the former one is fixed and the numbers shown in ptc create universe are meaningful. The parameter "lmmax" is incremented where needed from 150 to 3000 and set back. Moreover Etienne has done the following modifications to make this possible:
    - The modification I made in the new PTC I sent you are as follows:
      - You select SECTOR_NMUL and SECTOR_NMUL_MAX. For all multiples <= SECTOR_NMUL, then maxwell's is solved to order SECTOR_NMUL_MAX.
      - For multiples above SECTOR_NMUL, they are treated a la sixtrack.
      - As for example, if you have errors to order 10, you may bother with maxwell's only to order 10 but real_max=5 as far as Maxwell's is concerned.
      - Multipole higher will be sixtrack multipole.
- New PTC
- PTC with crash security.

PTC
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Files:

- make_ptc_module: Fix of "ptc_normal" by fixing the string comparison
- make_ptc_module & make_ptc_twiss: write a read traditional 3a map format
   - Tips twiss becomes "public" to overcome problem with g95. Probably okay in g95 but may be overly picky.
   - Bug corrected q95 was written to twiss table instead of energy.
   - Bug corrected: Parameter twiss results where not scaled with energy
   - Few protections against seg faults added. Redundant debug info available only in high level debug mode.
   - knobs for Initial parameters
   - Moments initialization for 5D in twiss, moments not available for SECTOR_NMUL_MAX = 10.
   - Removed charge setting to the sey_ring layout to make ptc_twiss running, redundant parameters removed.
   - Moments calculation fully implemented, map differing in ptc_twiss
   - New ptc_twiss to SECTOR_NMUL_MAX = 10 instead of errors when initial values provided. Moments seem to work (to be tested yet)
   - New ptc_twiss to A_ is tracked. This makes possible tracking of moments (to be completed).

wrap.f90
   - needed changes to accept new "make_ptc_normal_module".
   - First step for node layout tracking in ptc_trackline
   - Clean-up.
   - Added feature that allows to set values of several knobs and only at the end recalculates in tables.
   - Normally all tables are recalculated after setting a new value.
   - However, it does show parametric matching.
   - New command ptc_refreshtable
   - Updated match with knobs.
   - In addmom, t and delta swapped so the MADX input corresponds to the MADX nomenclature.
   - Moments calculation fully implemented, map differing in ptc_twiss
   - New ptc_twiss, A_ is tracked. This makes possible tracking of moments (to be completed).
   - Etienne's own stuff
   - Adding gin command following make_ptc_script_module.
   - Implemented.
1. ptc_setfieldcomp that set any order field strength to requested value. It makes matching of higher order field components.
2. Special matching mode use_gt95k. It implements kind of match that employs parametric PTC calculations to perform matching in a faster manner.
   - For further details see the comments at the top of matchptcknobs.c file.
3. Major corrections and protections against segmentation violation.
   - PTC knobs (pol_blocks) almost completely interfaced to MAD-X.
   - User sets a knob with ptc_knob command.
   - Twiss parameters and user specified (with ptc_select) map components are buffered in memory after every element in form of Taylor series.
   - They can be dumped to text file in two formats with ptc_printparametric command. They can be also visualized and further studied with viewer from gptk plugin.
   - Further, user can set numeric values of knobs with ptc_setknobvalue what updates all numeric values of the parameters in the tables.
   - This way knobs can be used in matching.
   - knobs implemented with PTC with pol_blocks; command to dump parametric results to file or stdout; content of ptc_madx_tablepush.f90 moved to ptc_madx_knobs.f90, knobs implemented with PTC with pol_blocks; command to dump parametric results to file or stdout; content of ptc_madx_tablepush.f90 moved to ptc_madx_knobs.f90, the former one removed
   - Skeleton for knobs and arbitrary element placement implemented. Lattice visualization via ROOT macro. Printing detailed lattice geometry in PTC. Several small bug corrections and some code cosmetics.
   - PTC_Enforce6D implemented

--------
Changes:

- PTC May 2007
  - VORMAKE assigned, the same as some of node in MAD but with capital letters
  - Fixing the crash for den = n mult = 10. This set-up requires to solve Maxwell's equation up to SECTOR_NMUL_MAX. The default is set to 10 to avoid excessive computing time. This is now safeguarded in madx. To this end the parameters SECTOR_NMUL and SECTOR_NMUL_MAX are transferred from "ptc_create_layout" to "ptc_create_universe" such that the former one is fixed and the numbers shown in ptc create universe are meaningful. The parameter "lmmax" is incremented where needed from 150 to 3000 and set back. Moreover Etienne has done the following modifications to make this possible:
    - The modification I made in the new PTC I sent you are as follows:
      - You select SECTOR_NMUL and SECTOR_NMUL_MAX. For all multiples <= SECTOR_NMUL, then maxwell's is solved to order SECTOR_NMUL_MAX.
      - For multiples above SECTOR_NMUL, they are treated a la sixtrack.
      - As for example, if you have errors to order 10, you may bother with maxwell's only to order 10 but real_max=5 as far as Maxwell's is concerned.
      - Multipole higher will be sixtrack multipole.
- New PTC
- PTC with crash security.
- First BB
- PTC version including spin
- 0-tone Etienne:
  Therefore I included this Zip file which contains the newest PTC. There are
  a few minor bugs related to patches in the present CVS version of PTC. This
could affect the CHANGEREF command of Frank. It is fixed in this new PTC. In
addition I included some routines in pointers.f90 and the script file for the
example.
  - Etienne's cleanup
  - Change faulty print out of Totalpath

1) _se_status: preliminary fix of uninitialized variable RADIATION_NEW
2) madx_ptc_module: Fix of "ptc_normal" by fixing the string comparison
3) madx_ptc_module & madx_ptc_twiss: write & read traditional DA map format

- PTC first changes stay October 2006 thinlens, cutting

Introducing THINLENS in WAVEMAP is PTC. To this end 2 flags have been
introduced in the MAD-X dictionary madxdict.h:

1) ptcrend: if true it uses a PTC type WAVEMAP
2) f3rend: if true it uses WAVEMAP; if false it uses WAVEMAPR

- Etienne O-tone:
  I fixed a bug in the exit patches part of the backward survey. This bug was
noticed while doing the Daphne backward ring. Patching was done correctly
but the survey command was moving the layout. This reflects a bug in either
patching or survey: dangerous.

- The definition is:
integer, pointer:: CAVITY_TOTALPATH ! REAL PILL BOX =1 , FAKE =0  default

- Big bug in GETMAT7R and GETMAT7d DH wrong: "basically model 2 method 4 is
messed up"

- New PTC Etienne end of visit a) thin lense b) BB

Spb_fake_gino_sub.f90
- New file needed for "Gino Version"

pointers.f90
- reason had tempo comment
- Etienne's Gino stuff
  - Adding gino command following madx_ptc_script_module
    - logical lp -> 4 and vice versa so NAG does not cry

Spc_pointers.f90, st_pointers.f90,
- Replaced by st_pointers.f90
- New 2007 PTC: this file replaces pointers.f90

Sq_outpt_gtc.f90
- New routines needed by PTC including spin

Sqa_beam_beam_ptc.f90, Sqb_accel_ptc.f90, Sr_spin.f90
- New routines needed by PTC including spin

=============================================================================

FPF
---
Files:
------
a_def_all_kind.inc, a_def_element_fibre_layout.inc, a_def_frame_patch_chart.inc,
a_def_sagan.inc, a_def_worm.inc, a_scratch_size.f90, b_da_arrays_all.f90,
c_dabnew.f90, d_lielib.f90, h_definition.f90, i_tpsa.f90, j_tpsalie.f90,
k_tpsalie_analysis.f90, l_complex_taylor.f90, m_real_polymorph.f90,
n_complex_polymorph.f90, o_tree_element.f90

Changes:
--------
- Throw out unused variables
  - Fix single/double precision definition that crashed NAG F95
  - logical=>logical(lp) needed for NAG f95
- Bug corrected - only first 10 elements of an array was zeroed instead of whole
- Check of initial conditions provided by the user on imput.
- pointers initialized to null in universal_taylor

m_define_newda.f90, f_newda.f90, g_newLielib.f90
- Experimental NEWDA no longer for this PTC version

=============================================================================

Makefiles
---------
Files:
------
Makefile, Makefile.bat, Makefile.prof, Makefile_develop, Makefile_g95,
Makefile_gdb, Makefile_fortran, Makefileจิต, Makeonline

Changes:
--------
- moved aperture code from madan.c to new file aperture.c
  - more apex in halo polygon
  - corrected the construction of rectangles in the general case
  - secured potentially dangerous division by zero
  - cleaned code on NAG warnings are minimized now: mainly unused variables

- PTC May 2007
  - "madx_ptc_normal_module"

- PTC versions including spin
  - Changes needed to compile routine dianal without optimization in extra file
  - included a file that gets stuck in matching procedures. In all Linux
    Makefiles matchlib2.F is compiled when using g77. For the Fortran90 compilers
    g95, f95(NAG) and gfortran an optimized routine is used as provided by
    Andy Vaught, the "g95" maintainer. For Windows the special compile flag
    -D_G95 was needed (special undocumented function compile flag)
  - the Laby 1756 to compile this special Fortran90 version of dianal.

=============================================================================
as a comment.
  - change in the print out of match summary when USE_MACRO
    - Implemented:
      1. ptc_setfieldcomp that set any order field strength
         to requested value. It enables matching of higher order field
         components.
      2. Special matching mode use_ptcknob. It implements kind of_macros
         that emplys parametric PTC calculations to perform matching in a faster manner.
         For further details see the comments at the top of matchingptcknobs.f file.
      3. Minor corrections and protections against segmentation violation.
        - Linker option added to export main program symbols so the function can be
          used from plugins
        - plugin support switched off by default
        - optional plugin support added, that requires dynamic linking. Switched-off
          by default.
        - corrected error
          - knobs implemented with PTC with pol_blocks; command to dump parametric
            resuls to file or stdout; content of ptc_madx_tablepush.f90 moved to
            ptc_madx_knobs.f90, the former one removed
          - skeleton for knobs and arbitrary element placement implemented. Lattice
            visualization via ROOT macros. Printing detailed lattice geometry in PTC.
            Several small bug corrections and some code cosmetics.
        - Missing dependencies added.
        - PTC with crash security!

First BB
Makefile.bat
  - JMJ: Added line to compile fetchlib2.f and modified lines to link it into
    cmde.exe and madxp.exe.

Makefile
  - Makefile using gfortran - not working yet for PTC

Makefile
  - New files for MAD-X On-Line Modeling Version

=============================================================================
MAD-X production version 3.03: 04.05.2006

1) Documentation of standard PTC modules
2) New PTC module "ptc_track_line", i.e. lines with acceleration
3) Thin lens tracking in agreement with Ripken theory and PTC
4) Non-linear matching via encapsulated ptc_normal commands

All changes for each file:

Makefile
  - replacing macros by madx
  - combining to the end on request of Etienne
  - adapting Makefiles for non-linear matching and PTC upgrade
  - PTC upgrade: Proper Thin Lens Lattice

=============================================================================
Makefile.bat
  - replacing macros by madx
  - passing to the end on request of Etienne
  - Take out old Sb_1 and Sb_1.obj files
  - Update Madx classes
  - combine to the end on request of Etienne
  - non-linear matching and PTC upgrade
  - PTC upgrade: Proper Thin Lens Lattice

=============================================================================
madX/Makefile.prof
  - replacing macros by madx
  - passing to the end on request of Etienne
  - Makefile for profiling courtesy PS

=============================================================================
Makefile_develop
  - replacing macros by madx
  - passing to the end on request of Etienne
  - adapting Makefiles for non-linear matching and PTC upgrade
  - PTC upgrade: Proper Thin Lens Lattice

=============================================================================
Makefile_g95
  - replacing macros by madx
  - passing to the end on request of Etienne
  - adapting Makefiles for non-linear matching and PTC upgrade
  - PTC upgrade: Proper Thin Lens Lattice
  - Merged with version MAD-X 3.02.29

=============================================================================
Makefile_nag
  - Makefile for the NAG compiler -- for the moment not operational

=============================================================================
Sa_extend_poly.f90
  - PTC with dvds implemented in the travelling wave cavity. The voltage is
    given by: V=V0-dvds*z
  - Updated to           energybefore = nfen%energy*1000.

=============================================================================
Sb_1_pol_template.f90
  - No longer needed after PTC upgrade 25.04.2005

=============================================================================
Sb_2_pol_template.f90
  - No longer needed after PTC upgrade 25.04.2005

=============================================================================
Sd_frame.f90
  - Updated to           energybefore = nfen%energy*1000.

=============================================================================
Se_status.f90
  - Drop the printing of "NO=10" in curvabend
  - Adapting Makefiles for non-linear matching and PTC upgrade
  - PTC upgrade: Proper Thin Lens Lattice

=============================================================================
Makefile.mac
  - Makefile for the MAC requires -- for the moment not operational

fa_madx_poly.f90
  - PTC with dvds implemented in the travelling wave cavity. The voltage is
    given by: V=V0-dvds*z
    Updated to            energybefore = nfen%energy*1000.

=============================================================================
fa_madx_poly_template.f90
  - No longer needed after PTC upgrade 23.04.2005

fa_madx_poly_template.f90
  - No longer needed after PTC upgrade 23.04.2005

=============================================================================
fa_top.f90
  - Updated to            energybefore = nfen%energy*1000.

=============================================================================
fa_bottom.f90
  - Drop the processing of "PNT" if wanted
  - adapting Makefiles for non-linear matching and PTC upgrade
  - PTC upgrade: Proper Thin Lens Lattice
- Updated to madX-3_02_16; bug corrected in madx_ptc_setcavs.f90

1) Remove residual left-over definition of double precision numbers. Should all be in a_scratch_size.f90.
2) The logical needs to be defined as "logical(l)". Several instances found. Should all be in a_scratch_size.f90.

Adapting Makefiles for non-linear matching and PTC upgrade

- PTC upgrade: Proper Thin Lens Lattice
- Domain specific code:

2) The logical needs to be defined as "logical(lp)". Several instances found. Should all be in a_scratch_size.f90.

1) Remove residual left-over definition of double precision numbers. Should all be in a_scratch_size.f90.

Adaptation of the Makefiles for non-linear matching and PTC upgrade

- PTC upgrade: Proper Thin Lens Lattice
- Domain specific code:

1) Remove residual left-over definition of double precision numbers. Should all be in a_scratch_size.f90.
2) The logical needs to be defined as "logical(l)". Several instances found. Should all be in a_scratch_size.f90.

Adapting Makefiles for non-linear matching and PTC upgrade

- PTC upgrade: Proper Thin Lens Lattice
- Domain specific code:

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1) Remove residual left-over definition of double precision numbers. Should all be in a_scratch_size.f90.

Adapting Makefiles for non-linear matching and PTC upgrade

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Adapting Makefiles for non-linear matching and PTC upgrade

- PTC upgrade: Proper Thin Lens Lattice
- Domain specific code:

2) The logical needs to be defined as "logical(lp)". Several instances found. Should all be in a_scratch_size.f90.

1) Remove residual left-over definition of double precision numbers. Should all be in a_scratch_size.f90.
2) Add CT variable print-out in 5D
- text of print-out is corrected
at Closed_ORBIT=OFF.
- calculation of CO is removed when ELEMENT_BY_ELEMENT is forced to ON

Fixing the closed orbit at the observation points for ELEMENT_BY_ELEMENT
madx_ptc_track_run.f90
=============================================================================
-Needed for PTC upgrade 25.04.2005

Fix write statement that was changed by automatic clean-up, IE replacing put pointers to the end on request of Etienne

pointers.f90

Updated to madX-3_02_16; bug corrected in madx_ptc_setcavs.f90

o_tree_element.f90

Revision

matchlib.F

Revision

matchjc.F

Revision

matchc2.c

Revision

madxu.c

Updated to madX-3_02_16; bug corrected in madx_ptc_setcavs.f90

2) Grow table "normal_results" if needed
1) Adding "eign" to ptc_normal table "normal_results"

-eigenvector calculation in ptc_twiss

-The eigenvalue keyword is consistent set to the 2 characters "eign"

-New matching with macros that enables fitting of non-linear parameters with PTC

-Clean-up

madxn.c

Updated to madX-3_02_16; bug corrected in madx_ptc_setcavs.f90

2) Grow table "normal_results" if needed
1) Adding "eign" to ptc_normal table "normal_results"

-eigenvector calculation in ptc_twiss

-The eigenvalue keyword is consistent set to the 2 characters "eign"

-New matching with macros that enables fitting of non-linear parameters with PTC

-Clean-up

madxdict.h

-Reverting to old warning routine

madxp.c

Updated to madX-3_02_16; bug corrected in madx_ptc_setcavs.f90

2) Grow table "normal_results" if needed
1) Adding "eign" to ptc_normal table "normal_results"

-eigenvector calculation in ptc_twiss

-The eigenvalue keyword is consistent set to the 2 characters "eign"

-New matching with macros that enables fitting of non-linear parameters with PTC

-Clean-up

abploc with "strict" version of madxdev

-version 3.02.25 => PTC examples (_track, _normal, _twiss) are checked on

-Merge with version 3.02.29

2) Grow table "normal_results" if needed
1) Adding "eign" to ptc_normal table "normal_results"

-eigenvector calculation in ptc_twiss

-The eigenvalue keyword is consistent set to the 2 characters "eign"

-New matching with macros that enables fitting of non-linear parameters with PTC

-Clean-up

madx_ptc_trackcavs.f90

Updated to madX-3_02_16; bug corrected in madx_ptc_setcavs.f90

2) Grow table "normal_results" if needed
1) Adding "eign" to ptc_normal table "normal_results"

-eigenvector calculation in ptc_twiss

-The eigenvalue keyword is consistent set to the 2 characters "eign"

-New matching with macros that enables fitting of non-linear parameters with PTC

-Clean-up
Only radiation part of code still contains delta.
Pull DC potentials (Sniper) code.
Closed orbit still computed by twiss.
Merged with version MAD-X 3.02.29

trrun.F
- Change the definition of the kicker. The acting on px/py now instead of x’/y’.
- AK/FS
- Cleaned version of thintrack. No delta_p dependence internally any more.
- Only radiation part of code still contains delta.
- Full 6D equations (Ripken) used.
- Some further improvements.
- Closed orbit still computed by twiss.
Merged with version MAD-X 3.02.29

user2_photon.f90
1) Remove debug printing of signs
- remove bug (division by zero) due to dipole errors

wrapp.f
- Change the definition of the kicker. The acting on px/py now instead of x’/y’.
- AK/FS
- Cleaned version of thintrack. No delta_p dependence internally any more.
- Only radiation part of code still contains delta.
- Full 6D equations (Ripken) used.
- Some further improvements.
- Closed orbit still computed by twiss.
Merged with version MAD-X 3.02.29

MAD-X version 3.02.14: 12.04.2006
1) PTC modules have been cleaned up and are all documented by now
2) PTC.Track.Intermediate has been improved including a new user oriented
   interface and maintained by Piotr Skowronski
3) Numerous bug fixes and clean-ups

All changes for each file:

MAD-X version 3.02.05: 22.03.2006
1) Merge PTC module. All are now documented by now
2) New match mode
3) Fix priority order between ICASE and DELTAP and cavities
4) Proper conversion between variables for 5D
5) Clean-Up
6) Replace "double precision" by generic "real(dp)
7) Merged newmatch-060411 with recent HEAD developments
All changes for each file:

1) Plug in "public" in each module
2) Partial unitialized array in "FIND_ORBIT_LAYOUT_noda" in "So_fitting.f90".
3) A couple of missing "implicit none" statements as described before.
4) check_iteration, check_interpolate_x, check_interpolate_y where
MAD-X version 3.02.01: 08.02.2006

All changes for each file:

- run_madx.f90
  - Modified to allow more flexible PTC modeling using madx_main.f90 and
  - The link option 'static' does not yet work for 'madxdev' using g95.

Makefile_g95

- Adding new PTC file Sg_1_fitted.f90
- Bug fix of dependence of Sg_1_fitted.f90

run_madx.f90

- Modified to allow more flexible PTC modeling using madx_main.f90 and
- Default location for checked-out files changed, comment added.

Makefile.bat

- Missing dependence on madxreg.c
- Adding new PTC file Sg_1_fitted.f90

All changes for each file:

- MAD-X Source
  - 3) Worldwide CVS read access from:
    - 4) PTC upgrade
  - 2) "Jacobian" matching
  - 1) Many bug fixes

MAD-X version 3.02.01: 08.02.2006
The normalization of $\mu$ to $2\pi$ has been done in "twiss" and "embedded twiss".

The deltap variable was wrongly defined in the match section, this lead to ...

Field error components are added as well.

Several warnings during a MAD-X run.

Adverse effect when trying to read in 2 previously "save"d sequences.

Hans Grote non-dynamic buffer fix.
MAD-X version 3.00.01: 07.09.2005

Overview:

All changes for each file:
MAD-X version 2.13.09: 09.03.2005

Overview:

All changes for each file:

MAD-X version 2.13:  FS 23.11.2004
MAD-X version 2.12: FS 29.09.2004

MAD-X version 2.11: FS 02.06.2004

MAD-X version 2.10: FS 27.03.2004

MAD-X version 2.00: FS 24.11.2003
MAD-X version 1.12:  FS 04.07.2003


MAD-X version 1.10:  HG & FS 20.01.2003

MAD-X version 1.09:  FS 09.12.2002

MAD-X version 1.08:  FS 18.11.2002

MAD-X version 1.06:  FS 16.10.2002

MAD-X version 1.05:  HG 25.9.2002

New sample job to create footprints for LHC: HG 18.9.2002
RPlot

Introduction
RPlot is a MAD-X plug-in that provides additional functionality using [ROOT]. It contains several tools

RVIEWER
plotting tool that handles the results in parametric form

What makes it different from the standard PLOT module of MAD-X is that it is also able to deal with the parametric results. RPlot provides graphical user interface that allows to choose which functions shall be drawn, set its ranges and adjust all the details of the plot formatting. Of course, the result is immediately visible on the screen, in contrary to the standard plot tool that is able to work solely in the batch mode. The user can choose several formats to save his plot, including postscript, gif, pdf, root macro and many others.

RVIEWER is able to draw the lattice functions
1. along the layout
2. at given position in function of one or two knobs
It provides a convenient way to set the knob values. As the value is set, the plotted functions are immediately drawn for the new value.

In order to run RVIEWER simply issue "rviewer;" command

RTRACKSTORE
enables storage of the tracking data in ROOT NTuple/Tree format

Ntuple and its modern extension called Tree are formats designed for storing particle tracking data. It is proven to provide the fastest data writing and reading thanks to column wise I/O operations. It is commonly used for data storage by HEP experiments. Additionally, ROOT provides automatical ZIP data compression that is transparent for the user algorithms. Moreover, ROOT provides wide set of very comfortable tools for advanced analysis and plotting of the data stored in Trees.

Additionally, we plan to extend RVIEWER functionality that would provide intuitive graphical user interface to most commonly used features in particle tracking in accelerators. Thanks to that, the user is not forced to learn how to use the ROOT package.

Currently the feature is enabled only for tracking using the ptc_trackline command, however, it will be extended to other tracking modes.
Download
The newest version is available here

Installation
Prerequisite: ROOT must be installed beforehand compilation and whenever the user wants to use the plug-in. See explanations on [ROOT webpage](#).

To install RPLOT
1. Unpack the archive, it will create directory rplot
   
   ```
   tar xzvf rplot-X.XX.tgz
   ```

2. Change to rplot directory
   
   ```
   cd rplot
   ```

3. Type
   
   ```
   make install
   ```

Examples

SYNOPSIS

RVIEWER;

PROGRAMMERS MANUAL

To be continued...
References

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P. Bramham and H. Henke. private communication and LEP Note LEP-70/107, CERN.

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[*Ir*] May 01, 2003