

CAS Course on Optics Design

Otwock, September 2015

Information at:

http://cern.ch/Werner.Herr/CAS2015_COURSE

CAS 2015 course on optics design

■ Aims:

- From the lectures to praxis
- Design a realistic machine optics with various features

■ Not a lecture, but following a series of steps (as exercises) applying what was learned in previous lectures

■ Done by you in close collaboration with the tutors and your colleagues

■ The **MADX** program is used for this course

Procedure and basic steps

- Introduction to MAD-X (2 talks)
- Work on 8 exercises:
 - Design of periodic machine with desired properties (1-2)
 - Correction of chromaticity and orbit imperfections (3-5)
 - Design of a dispersion suppressor (6)
 - Design of a β -insertion (low and high β , for experiments, collimation etc.) (7)
 - Particle tracking to study stability of your design (8)

Available tools

- Individual computers
 - LINUX operating system
 - You have MAD-X, compilers, gnuplot ...
- Bonus material:
You get all your solutions and our suggested solutions
together with the MAD-X binaries after the school.

Available to help

Werner Herr,
Guido Sterbini,
Bernhard Holzer (week 1),
Verena Kain,
Yannis Papaphilippou

for computers: Adam Wasilewski, Jacek Szlachciak

How to get in ?

- For LINUX: see instructions or ask for help
- Common accounts:
 - login name: **user**
 - password: **xxxxxxxx**
 - (always use the same computer, files are local)
- If you want to use your own computer, download MADX from website:
 - WINDOWS: **http://cern.ch/Werner.Herr/CAS2015_COURSE/exe**
 - LINUX: **http://cern.ch/Werner.Herr/CAS2015_COURSE/bin**

Where you find all that:

You find a directory: **/home/user/COURSE** on your machine (You may have it as a .tar file, extract with e.g.: tar xvf *name*)

Documentation: **/COURSE/doc**

Your exercises in: **/COURSE/doc/problems.pdf**

Examples in: **/COURSE/examples**

Solutions at: **/COURSE/solutions**

Executable:

/COURSE/bin/madx (LINUX, may have to set PATH)
(e.g.: setenv PATH "\$PATH":"*pathname*)

Introduction to MADX

Werner Herr, CERN

For all MAD details:

(<http://cern.ch/mad>)

see also:

MADX primer

MADX - part 1 (today)

- Description of the basic concepts and the language
- Define a machine and compute optical functions
- Get the parameters you want
 - Beam dimensions
 - Tune, chromaticity

Required lectures: Recap transverse dynamics, Lattice cells

MADX - part 2 (as we progress ..)

- Machines with imperfections and corrections
 - Closed Orbit distortions and correction
- Design of insertions
 - Dispersion suppressor
 - Low β insertion
- Particle tracking

Required lectures: Recap Transverse Dynamics, Lattice cells, Insertions, Non-linear dynamics

General purpose lattice programs

- For circular machines, beam lines or linacs
 - Calculate optics parameters from machine description
 - Compute (match) desired quantities
 - Simulate and correct machine imperfections
 - Simulate beam dynamics
- ➔ Used in this course: **MADX**

What is MADX ?

- The latest version in a long line of development
(Methodical Accelerator Design)
 - Used at CERN since more than 30 years for machine design and simulation (PS, SPS, LEP, LHC, ...)
 - Existing versions:
MAD8 (obsolete), **MADX (version 5)**
 - Mainly designed for large projects (LEP, LHC^{*)} , CLIC ..)
- ^{*)} Only MADX can handle the LHC ...

Why we use MADX here ?

- Multi purpose:
 - From early design to final evaluation
 - Running on all systems, Source is free and easy to extend
- Easy to understand what is happening:
 - Well defined strategy for input language
 - No hidden or invisible actions or computations
- Other programs also used somewhere else:
 - SAD (KEK: Strategic Accelerator Design)
 - BMAD (Cornell)

Data required by all optics programs ?

- Description of the machine:
 - Definition of each machine element:
e.g. a focusing quadrupole
 - Attributes of the elements:
e.g. 3.1 m long and my gradient is k_1
 - Positions of the elements:
e.g. 2341 m from the beginning
- Description of the beam(s):
Protons at 7 TeV
- Directives (what to do ?):
Give me optical functions, match the optics

How does MAD get and use this information ?

- MAD is an "interpreter":

MAD prompt is: **X: ==>**

- Accepts and (immediately) executes statements
- Statements can be assignments, expressions or initiate complex actions (commands)
- Can be used interactively or in batch
 - ➡ Reads statements from input stream or a file (has no GUI)

- Many features of a programming language (loops, if conditions, macros, subroutines ...)

MAD input language

Strong resemblance to "C" language

- Not line oriented, all statements are terminated by ;
- Comment lines start with: // or !
- Arithmetic expressions, including functions (exp, log, sin, cosh ...)
- Immediate (=) and deferred expressions (:=) (like JAVA)
- In-built random number generators for various distributions
- Predefined constants (clight, e, π , m_p , m_e ...)

MADX conventions

- Not case sensitive
- Elements placed along the reference orbit (variable **s**)
- Horizontal (assumed bending plane) and vertical variables are x and y
- Describes a **local** coordinate system moving along **s**
 - i.e. $x = y = 0$ follows the curvilinear system (reference orbit)

Examples of expressions:

- Variables can be used in expressions:
 - ANGLE = 2*PI/NBEND;
 - AIP = ATAN(SX1/SX2);
- The assignment symbols `=` and `:=` have a very different behaviour (here random number generator)!
 - `DX = GAUSS()*1.5E-3;`
The value is computed **once** and kept in DX
 - `DX := GAUSS()*1.5E-3;`
The value is recomputed **every time** DX is used

How to use MADX ?

```
$ madx
```

```
X: ==> angle = 2*pi/1232;
```

```
X: ==> value, angle;
```

```
X: ==> dx = gauss()*2.0;
```

```
X: ==> value, dx;
```

```
X: ==> value, dx;
```

```
X: ==> dx := gauss()*2.0;
```

```
X: ==> value, dx;
```

```
X: ==> value, dx;
```

How to use MADX ?

For a large machine you may need many commands
(LHC \approx 27000)

Better: store your input in a file: e.g. `my.file`

```
$ madx
```

X: ==> `call, file=my.file;` (WINDOWS or LINUX)

alternatively, redirection from the file into the parser (LINUX)

```
$ madx < my.file (LINUX)
```

Warning !

- For WINDOWS users:
 - The input file must be a plain text (ASCII) file !
 - NOT a WORD, POWERPOINT or EXCEL file ...
- For ALL users:
 - Do NOT use colours in the commands (I use them only here for the talk !)

MAD input statements (what we need)

■ Typical assignments:

- Properties of machine elements
- Set up of the lattice
- Definition of beam properties (particle type, energy, emittance ...)

■ Typical actions:

- Compute lattice functions, match optical parameters
- Assignment of errors and imperfections
- Correct machines

Recommendation: make use of the examples !

How to define machine elements ?

- MAD-X **Keywords** used to define the type of an element.
- General format:
 - *name* : *keyword, attributes;*
- Can define single *element* or *class* of elements and give it a **name** of your choice
 - Some **keywords** are predefined, **name** can be anything (but avoid blanks in the name)
 - Some examples:

Example: Definitions of magnets

Dipole (bending) magnet:

$$k_0 = \frac{1}{p/c} B_y [\text{in } m^{-1}] \left[= \frac{1}{\rho} = \frac{\text{angle}}{l} \right] [\text{in } rad/m]$$

DIP01: SBEND, L = 10.0, ANGLE = angle, K0 = k0;
name *keyword* *attributes*

Quadrupole magnet:

$$k_1 = \frac{1}{p/c} \frac{\partial B_y}{\partial x} [\text{in } m^{-2}] \left[= \frac{1}{l \cdot f} \right]$$

MQA: QUADRUPOLE, L = 3.3, K1 = k1;

Example: Definitions of magnets

Sextupole magnet:

$$k_2 = \frac{1}{p/c} \frac{\partial^2 B_y}{\partial x^2} \left[\text{in } m^{-3} \right]$$

KLSF = k_2 ;

MSXF: SEXTUPOLE, L=1.1, K2 = KLSF;

Octupole magnet:

$$k_3 = \frac{1}{p/c} \frac{\partial^3 B_y}{\partial x^3} \left[\text{in } m^{-4} \right]$$

KLOF = k_3 ;

MOF: OCTUPOLE, L=1.1, K3 = KLOF;

Other elements:

Drift space:

DR1: DRifT, L=1.1;

Marker (has no effect, mostly used as a position reference):

MK1: Marker;

Example: definitions of elements

Define a class of Quadrupole magnets:

MQF: QUADRUPOLE, L=3.3, K1 = +1.23E-02;

MQD: QUADRUPOLE, L=3.3, K1 = -1.23E-02;

QUAD01, QUAD02, ... are instances of the class MQF etc., all with the same properties:

QUAD01: MQF;

QUAD02: MQD;

QUAD03: MQF;

QUAD04: MQD;

....

Example: LHC dipole magnet

length = 14.3;

B = 8.33;

PTOT = 7.0E12;

ANGLHC = B * clight * length/PTOT;

MBLHC: SBEND, L = Length, ANGLE = anglhc;

the same example for other codes (for comparison):

BMAD → **MBLHC: SBEND**, L = Length, ANGLE = anglhc

SAD → **SBEND MBLHC** =(L = Length ANGLE = anglhc)

Try it ..

```
$ madx
```

```
X: ==> length = 14.3;
```

```
X: ==> B = 8.33;
```

```
X: ==> PTOT = 7.0E12;
```

```
X: ==> ANGLHC = B * clight * length/PTOT;
```

```
X: ==> MBLHC: SBEND, L = Length, ANGLE = ANGLHC;
```

```
X: ==> value,mblhc->angle;
```

Thick and thin elements

- Thick elements: so far all examples were thick elements (or: lenses)
- Specify **length** and **strength** separately (except dipoles !)
 - + More precise, path lengths and fringe fields correct
 - Not symplectic in tracking
 - May need symplectic integration

- Thin elements: specified as elements of **zero** length
- Specify **field integral**, e.g.: $k_0 \cdot L, k_1 \cdot L, k_2 \cdot L, \dots$
 - + Easy to use
 - + Used for tracking
 - Path lengths not correctly described
 - Fringe fields not correctly described
 - Maybe problematic for small machines

For a proper discussion see lecture on "Tools for Non-Linear Dynamics"

Special MAD element: multipoles

Multipole: general element of zero length (**thin lens**), can be used with one or more components of any order:

multip: multipole, knl := { $k_{n0}L, k_{n1}L, k_{n2}L, k_{n3}L, \dots$ };

→ knl = $k_n \cdot L$ (normal components of n^{th} order)

Very simple to use:

mul1: multipole, knl := {0, $k_1L,0,0,\dots$ };

is equivalent to definition of quadrupole ($k_1L = \int \frac{1}{p/c} \frac{\partial B_y}{\partial x} \cdot dl$)

mul0: multipole, knl = {angle,0,0,...};

is equivalent to definition of a dipole ($k_0L = \text{angle}$)

Thick and thin elements

- For all exercises: → use thin lenses (multipoles) unless explicitly requested to use thick elements
- Easier to handle and analytic calculations are precise

E.g. for a dipole you can use:

MYD: MULTIPOLE, KNL = {angle,0,0,...};

E.g. for a quadrupole with an octupole component you can use:

MYQ: MULTIPOLE, KNL := {0,k₁L,0,k₃L,0,...};

Definitions of sequence (position)

Have to assign position to the elements.

Positions are defined within a **sequence** with a **name**:

cassps: SEQUENCE, REFER=CENTRE, L=6912;

...

here specify positions of all elements ...

...

ENDSEQUENCE;

A (relative (to some other element) or absolute) position can be defined.

General format is:

name: at = (position s in metres);

or (if an element appears several times):

name: class, at = (position s in metres);

Definitions of sequence (position)

cassps: SEQUENCE, refer=centre, l=6912;

...

...

MBL01: MBLA, at = 102.7484; ! absolute position

MBL02: MBLB, at = 112.7484;

MQ01: MQA, at = 119.3984;

BPM01: BPM, at = 1.75, from MQ01; ! relative position

COR01: at = LMCV/2 + LBPM/2, from BPM01;

MBL03: MBLA, at = 126.3484;

MBL04: MBLB, at = 136.3484;

MQ02: MQB, at = 142.9984;

BPM02: BPM, at = 1.75, from MQ02;

COR02: at = LMCV/2 + LBPM/2, from BPM02;

...

...

ENDSEQUENCE;

Complete example: SPS (thick)

```
circum = 6912;
// bending magnets as thin lenses
mbsps: multipole,knl={0.007272205};

// quadrupoles and sextupoles
kqf = 0.0146315;
kqd = -0.0146434;
qfsps: quadrupole,l=3.085,k1 := kqf;
qdsps: quadrupole,l=3.085,k1 := kqd;
lsf: sextupole,l=1.0, k2 = 1.9518486E-02;
lsd: sextupole,l=1.0, k2 = -3.7618842E-02;

// monitors and orbit correctors
bpm: monitor,l=0.1;
ch: hkicker,l=0.1;
cv: vkicker,l=0.1;

cassps: sequence, l = circum;
start_machine: marker, at = 0;
qfsps, at = 1.5425;
lsf, at = 3.6425;
ch, at = 4.2425;
```

```
bpm, at = 4.3425;  
mbsps, at = 5.0425;  
mbsps, at = 11.4425;  
mbsps, at = 23.6425;  
mbsps, at = 30.0425;  
qdsp, at = 33.5425;  
lsd, at = 35.6425;  
cv, at = 36.2425;  
bpm, at = 36.3425;  
....  
....  
qdsp, at = 6881.5425;  
lsd, at = 6883.6425;  
cv, at = 6884.2425;  
bpm, at = 6884.3425;  
mbsps, at = 6885.0425;  
mbsps, at = 6891.4425;  
mbsps, at = 6903.6425;  
mbsps, at = 6910.0425;  
end_machine: marker, at = 6912;  
endsequence;
```

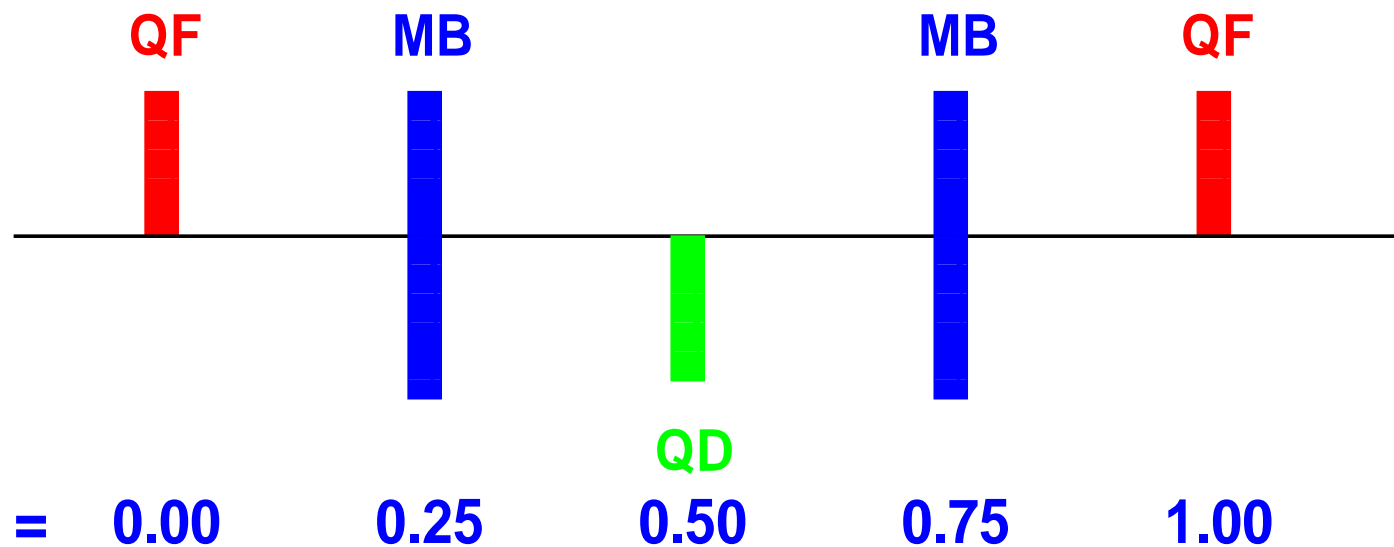


Definition of large machines ...

- For large machines with many elements:
 - Time consuming to specify every element individually (e.g. LHC more than 25000 elements needed)
 - Very inflexible (e.g. change of cell length)
- Several options:
 - Loops over elements possible
 - Elements can be combined into new objects

A very simple cell ..

cell: Length = l_{cell}



A very simple cell ..

- Positions can be defined in loops:
- Loop over number of cells (*ncell*)

```
lcell = 64;      ! cell length
ncell = 108;     ! number of cells
circum = ncell*lcell;    ! total circumference
cassps: sequence, refer=centre, l=circum;
n = 1;
while (n < ncell+1) {
    qfsps: qfsps, at=(n-1)*lcell;
    mbsps: mbsps, at=(n-1)*lcell + lcell*0.25;
    qdsps: qdsps, at=(n-1)*lcell + lcell*0.50;
    mbsps: mbsps, at=(n-1)*lcell + lcell*0.75;
    n = n + 1;
}
endsequence;
```

We have 6 lines instead of 432 (and are much more flexible !)

Nested sequences

→ Sequences can be defined and used like (new) elements:

```
cascell1: sequence, refer=centre, l=lcell;    (cascell1 is now an element)
qfsps: qfsps, at=0.0;
mbsps: mbsps, at=0.25*lcell;
qdsps: qdsps, at=0.50*lcell;
mbsps: mbsps, at=0.75*lcell;
endsequence;
```

```
allcells: sequence, refer=centre, l=nCell*lCell;
n = 1;
while (n < nCell+1) {
    cascell1, at=(n-1)*lCell;
    n = n + 1;
}
endsequence;
```

Simple MAD directives

- Define the input
- Define the beam
- Initiate computations (Twiss calculation, error assignment, orbit correction etc.)
- Output results (tables, plotting)
- Match desired parameters
- Beware: may have default values !

Input definition and selection

- Define the input:

- `call,"sps.seq";`
 - Selects a file with description of machine
 - Can be split into several files

- Activate the machine:

- `USE, sequence=cassps;`
 - Activates the sequence you want (described in `"sps.seq"`, which can contain more than one)

We still need a beam !

Some computations need to know the type of beam and its properties:

- Particle type
- Energy
- Emittance, number of particles, intensity

BEAM, PARTICLE=name, MASS=mass, NPART=Nb, CHARGE=q, ENERGY=E,.....;

Example:

BEAM, PARTICLE=proton, NPART=1.1E11, ENERGY=450,.....;

Initiate the computations

Execute an **action** (calculation of all lattice parameters around the
(circular !) machine):

`twiss;` or:

`twiss, file=output;` or:

`twiss, file=output, sequence=cassps;`

Execute an **action** (produce graphical output of β -functions):

`plot, haxis=s, vaxis=betx, bety;`

SELECT - a powerful command

Assigns parameters and options for an action, as an example **twiss**:

select,flag=twiss,column=name,s,betx,bety;

(defines output for action **twiss**: name, s, betx, bety)

Output for selected elements only, using C language regular expressions, examples:

select,flag=twiss,pattern="q.*",column=name,s,betx,bety;

shown only for elements starting with: "q"

select,flag=twiss,range="MQF01",column=name,s,betx,bety;

shown only for the element with name: "MQF01"

select,flag=twiss,range="QD[10]/QD[16]",column=name,s,betx,bety;

shown only for the elements inside the range of 10th to 16th quadrupole

Initiate the computations

Set parameters for an action with the **SELECT** command
(or defaults are used)

Calculation of Twiss parameters around the machine, store **selected** lattice functions on file **twiss.out** and plot β -functions:

```
select,flag=twiss,column=name,s,betx,bety;  
twiss, sequence=cassps, file=twiss.out;
```

```
plot, haxis=s, vaxis=betx, bety, colour=100;
```

Initiate the computations

Calculation of Twiss parameters around the machine, store and plot lattice functions for **quadrupoles only** (name starting with "q"):

```
select,flag=twiss,pattern="^q.*",column=name,s,betx,bety;
```

```
twiss, sequence=cassps, file=twiss.out;
```

```
plot, haxis=s, vaxis=betx, bety, colour=100;
```

Initiate the computations

Make a geometrical survey of the machine layout, available in a file:

```
select,flag=twiss,column=name,s,betx,bety;  
twiss, sequence=cassps, file=twiss.out;
```

```
survey, file=survey.cas;
```

Initiate the computations

Calculation of Twiss parameters around the machine, plot between 10th and 16th quadrupoles only:

```
select,flag=twiss,pattern="^q.*",column=name,s,betx,bety;  
twiss, sequence=cassps, file=twiss.out;
```

```
plot, haxis=s, vaxis=betx, bety, colour=100, range=qd[10]/qd[16];
```

Typical MAD example input:

```
// Read input file with machine description
call file="sps.seq";

// Define the beam for the machine
Beam, particle=proton, sequence=cassps, energy=450.0;

// Use the sequence with the name: cassps
use, sequence=cassps;

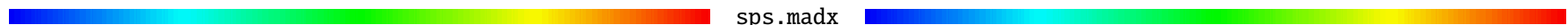
// Define the type and amount of output
select,flag=twiss,column=name,s,betx,bety;

// Execute the Twiss command to calculate the Twiss parameters
// Compute at the centre of the element and write to: twiss.out
twiss,save,centre,file=twiss.out;

// Plot the horizontal and vertical beta function between the
// 10th and 16th occurrence of a defocussing quadrupole
plot, haxis=s, vaxis=betx, bety, colour=100, range=qd[10]/qd[16];

// get the geometrical layout (survey)
survey,file=survey.cas;

stop;
```



Typical MAD example input:

```
// Read input file with machine description
call file="sps.seq";

// Define the beam for the machine
Beam, particle=proton, sequence=cassps, energy = 450.0;

// Use the sequence with the name: cassps
use, sequence=cassps;

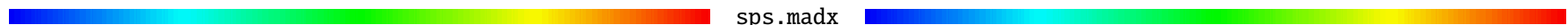
// Define the type and amount of output
select,flag=twiss,column=name,s,betx,bety;

// Execute the Twiss command to calculate the Twiss parameters
// Compute at the centre of the element and write to: twiss.out
twiss,save,centre,file=twiss.out;

// Plot the horizontal and vertical beta function between the
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plot, haxis=s, vaxis=betx, bety, colour=100, range=qd[10]/qd[16];

// get the geometrical layout (survey)
survey,file=survey.cas;

stop;
```



sps.madx

Typical MAD example input:

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use, sequence=cassps;

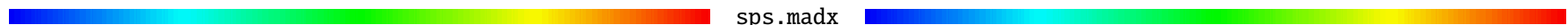
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```



Typical MAD example input:

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call file="sps.seq";

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Beam, particle=proton, sequence=cassps, energy=450.0;

// Use the sequence with the name: cassps
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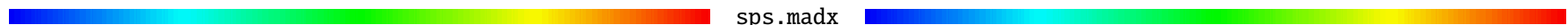
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call file="sps.seq";

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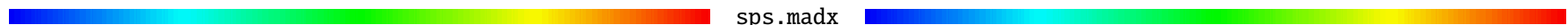
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twiss,save,centre,file=twiss.out;

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stop;
```



sps.madx

Typical MAD example input:

```
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call file="sps.seq";

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// Plot the horizontal and vertical beta function between the
// 10th and 16th occurrence of a defocussing quadrupole
plot, haxis=s, vaxis=betx, bety, colour=100, range=qd[10]/qd[16];

// get the geometrical layout (survey)
survey,file=survey.cas;

stop;
```

Typical MAD example input:

```
// Read input file with machine description
call file="sps.seq";

// Define the beam for the machine
Beam, particle=proton, sequence=cassps, energy=450.0;

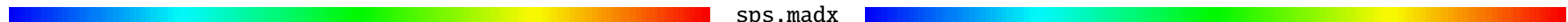
// Use the sequence with the name: cassps
use, sequence=cassps;

// Define the type and amount of output
select,flag=twiss,column=name,s,betx,bety;

// Execute the Twiss command to calculate the Twiss parameters
// Compute at the centre of the element and write to: twiss.out
twiss,save,centre,file=twiss.out;
// Plot the horizontal and vertical beta function between the \\
// 10th and 16th occurrence of a defocussing quadrupole \\
plot, haxis=s, vaxis=betx, bety, colour=100, range=qd[10]/qd[16];\\

// get the geometrical layout (survey)
survey, file=survey.cas;

stop;
```



sps.madx

Typical MAD output (summary):

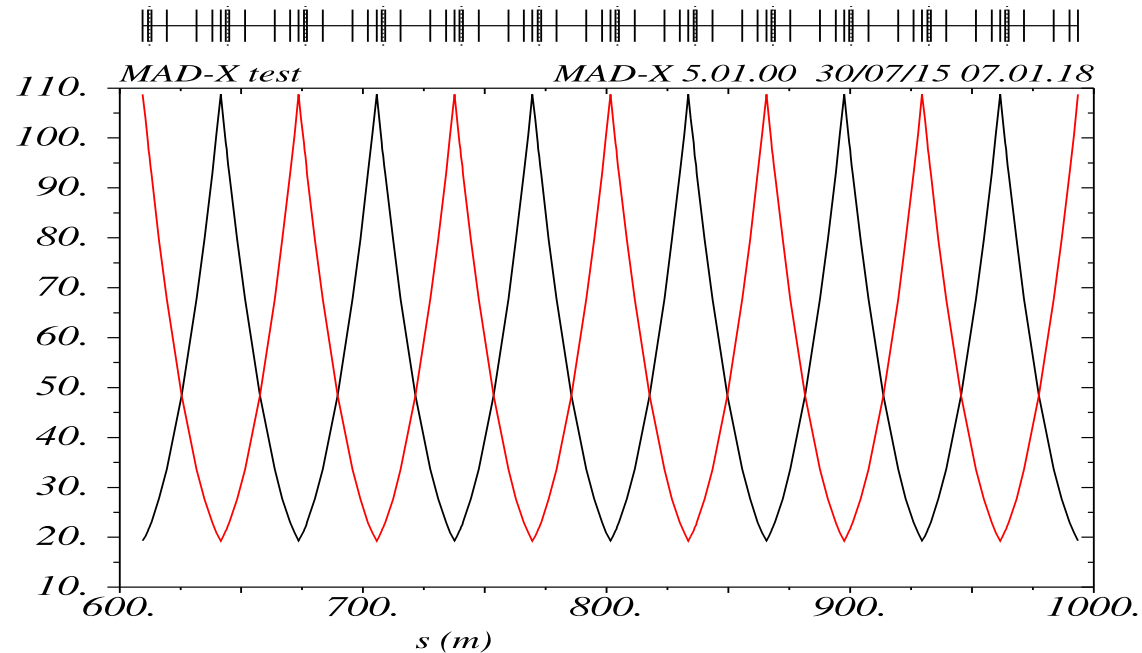
+++++ table: summ

length	orbit5	alfa	gammatr
6912	-0	0.001667526597	24.4885807
q1	dq1	betxmax	dxmax
26.57999204	-8.828683153e-09	108.7763569	2.575386926
dxrms	xcomax	xcorms	q2
1.926988371	0	0	26.62004577
dq2	betymax	dymax	dyrms
4.9186549e-08	108.7331749	0	0
ycomax	ycorms	deltap	synch_1
0	0	0	0

Typical MAD output (all elements):

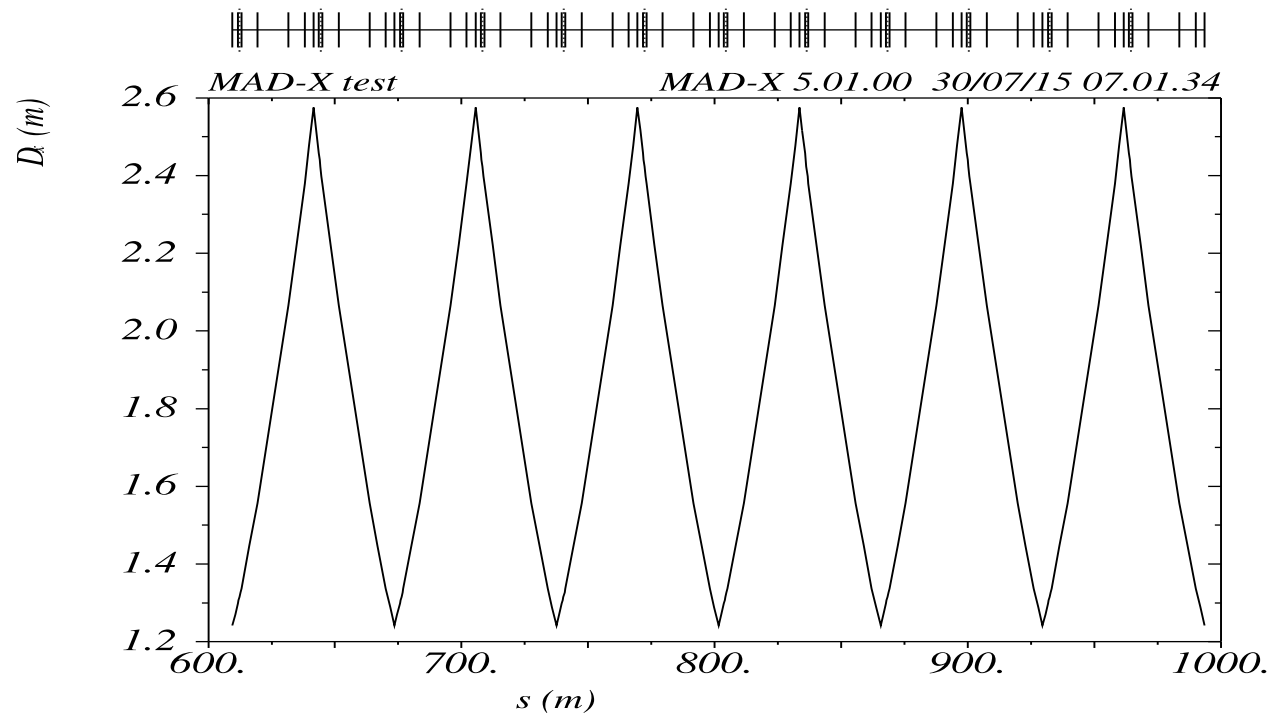
* NAME	S	BETX	BETY
\$ %s	%le	%le	%le
"CASSPS\$START"	0	101.5961579	20.70328425
"START_MACHINE"	0	101.5961579	20.70328425
"DRIFT_0"	0.77125	105.1499566	19.94571028
"QF"	1.5425	108.7763569	19.26082066
"DRIFT_1"	2.5925	103.8571423	20.21112973
"LSF"	3.6425	99.07249356	21.29615787
"DRIFT_2"	3.9424975	97.73017837	21.6309074
"CH"	4.2425	96.39882586	21.97666007
"DRIFT_3"	4.2925	96.17800362	22.03535424
"BPM"	4.3425	95.95748651	22.0943539
"DRIFT_4"	4.6925025	94.4223997	22.51590816
"MBSPS"	5.0425	92.90228648	22.95242507
"DRIFT_5"	8.2425	79.69728195	27.63752778
"MBSPS"	11.4425	67.74212222	33.5738988
"DRIFT_6"	17.5425	48.41469349	48.35614376
"MBSPS"	23.6425	33.6289371	67.68523387
"DRIFT_5"	26.8425	27.68865546	79.6433337
"MBSPS"	30.0425	22.99821861	92.85270185
"DRIFT_7"	31.7925	20.96178735	100.6058286
"QD"	33.5425	19.29915001	108.7331749
"DRIFT_1"	34.5925	20.25187715	103.8118608
.....			
.....			

Graphical output (β)



Plotted only for: range=qd[10]/qd[16]

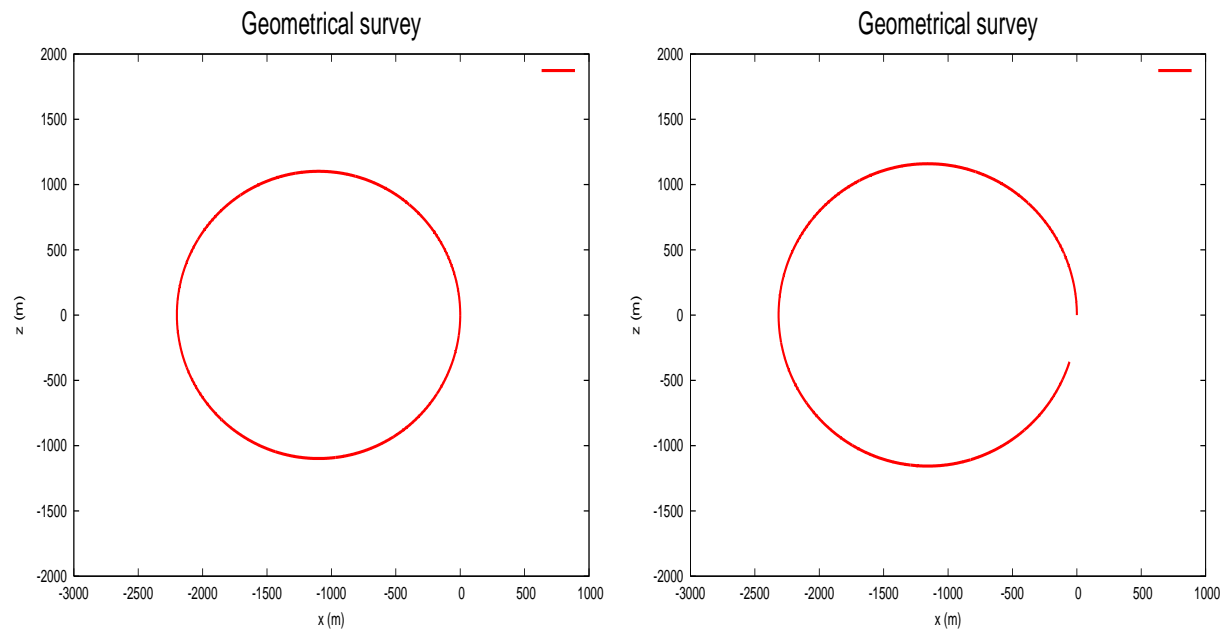
Graphical output (dispersion)



Plotted only for: range=qd[10]/qd[16]

Graphical output (geometrical survey)

- Output gives x, y, z, θ in **absolute** (terrestrial) coordinates, plotting x versus z should be a ring:



Optical matching

- To get the optical configuration you want
 - compute settings yourself or use MAD for **matching**
- Main applications:
 - Setting **global** optical parameters
(e.g. tune, chromaticity) → part 1 (following)
 - Setting **local** optical parameters
(e.g. β -function, dispersion ..) → part 2
 - Correction of imperfections → part 2

Matching global parameters

- Adjust strengths etc. to get desired properties (e.g. tune, chromaticity)
- Define the **properties** you want and the **elements** to vary
- Examples for global parameters (MAD convention):
 - Q1, Q2:(horizontal and vertical tune)
 - dQ1, dQ2:(horizontal and vertical chromaticity)

Matching global parameters - tune

!Example, match horizontal (Q1) and vertical (Q2) tunes:

!Vary the quadrupole strengths **kqf** and **kqd**

!Quadrupoles must be defined with: ..., k1:=kqf, ... etc.

```
match, sequence=cassps;
  global,sequence=cassps,Q1=26.58;    →  you want that !
  global,sequence=cassps,Q2=26.62;    →  you want that !
  vary,name=kqf, step=0.00001;    →  you vary that !
  vary,name=kqd, step=0.00001;    →  you vary that !
  Lmdif, calls=10, tolerance=1.0e-21; → Method to use ! (just copy that ...)
endmatch;
```

Matching global parameters - chromaticity

!Example, match horizontal (dQ1) and vertical (dQ2) tunes:

!Vary the sextupole strengths **ksf** and **ksd**

!Sextupoles must be defined with: ... , k2:=ksf, ... etc.

```
match, sequence=cassps;
  global,sequence=cassps,DQ1=2.0;    →  you want that !
  global,sequence=cassps,DQ2=-1.5;   →  you want that !
  vary,name=ksf, step=0.00001;    →  you vary that !
  vary,name=ksd, step=0.00001;    →  you vary that !
  Lmdif, calls=10, tolerance=1.0e-21; → Method to use ! (just copy that ...)
endmatch;
```

Changing MADX variables

- Deferred (:=) variables can be changed at any time during execution

```
use, period=cascell3;  
ksf = 0.0;  
ksd = 0.0;  
select,flag=twiss,column=name,s,betx,muy,bety,dx,dy;  
twiss,file=twiss1.out;  
  
ksf = +0.017041/20.0;  
ksd = -0.024714/20.0;  
twiss,file=twiss2.out;
```

- Useful for: closed orbit, matching, chromaticity ...etc.

(Some comments ...)

- Input language seems heavy, but:
 - Can be interfaced to data base
 - Can be interfaced to other programs (e.g. Mathematica, Python,...)
 - Programs exist to generate the input interactively
 - Allows web based applications
 - Allows to develop complex tools

Linacs and beam lines !

They are not closed machines ! They have no periodic solution !

→ There are no β -functions etc. !

Must give INITIAL optical parameters !

twiss, betx=..., bety=..., alfx=..., ;

plot, haxis=s, vaxis=betx, bety, colour=100;

WARNING: careful with **alfx, alfy** when you have thin lenses !

MADX

- END OF PART 1 -

- YOUR TURN -

MADX - part 2

■ We can:

- Design and compute a regular lattice
- Adjust machine parameters (tune, chromaticity, $\hat{\beta}$..)

■ Solutions of exercises 1 - 3 at:

<http://cern.ch/Werner.Herr/CAS2015/solutions>

■ What next:

- Machines with imperfections and corrections
- Design of a dispersion suppressor
- Design of a low β insertion

Error assignment

- MAD can assign **errors** to elements:
 - Alignment errors on all or selected elements
 - Field errors (up to high orders of multipole fields) on all or selected elements
- Errors are included in calculations (e.g. Twiss)
- Correction algorithms can be applied

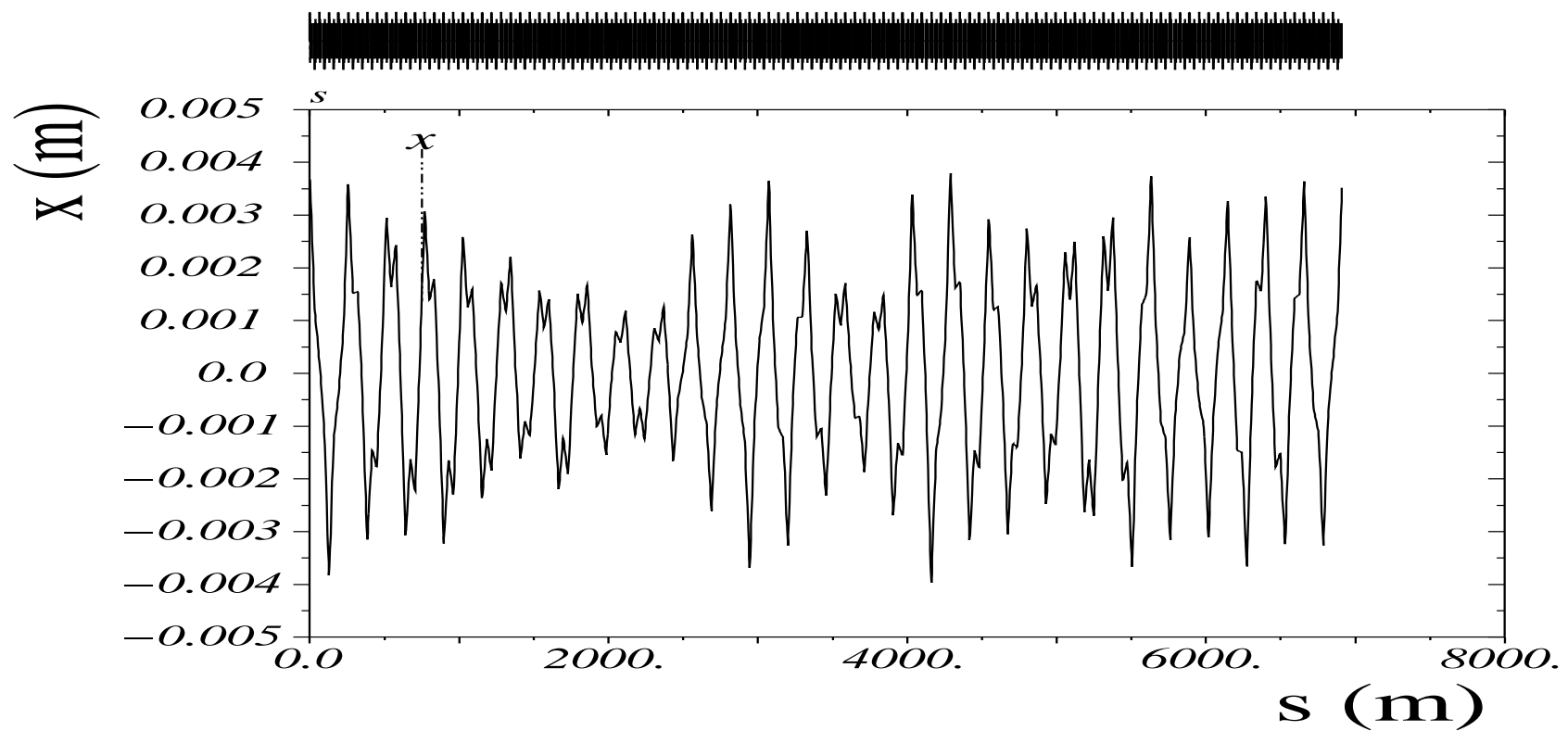
Error assignment

- Can define alignment errors (EALIGN):

```
! assign error to all elements starting with Q
select,flag=error,pattern="Q.*";
Ealign, dx:=tgauss(3.0)*1.0e-4, dy:=tgauss(3.0)*2.0e-4;
Twiss,file=orbit.out; ! compute distorted machine
plot,haxis=s,vaxis=x,y; ! plot orbits in x and y
```

- Can define field errors of any order (EFCOMP)
- Remember the `:=`!
- See MADX Primer: page 14

Orbit with alignment errors



→ Now we want to correct the orbit



How to measure an orbit ?

Needs Beam Position Monitors (keyword → MONITOR):

Gives position in one or both dimensions [*in m*]

BPMV: VMONITOR, L=0.1;

BPMV01: VMONITOR, L=0.1;

BPMV02: VMONITOR, L=0.1;

BPMV03: BPMV;

BPMH02: HMONITOR, L=0.1;

BPMHV01: MONITOR, L=0.1;

For orbit correction: consider orbit **only** at monitors ...

How to correct an orbit ?

Needs Orbit corrector magnets (keyword → HKICKER/VKICKER):

The strength of a corrector is an angle (kick) [*in rad*]

MCV: VKICKER, L=0.1;

MCV01: VKICKER, L=0.1, KICK := KCV01;

MCV02: VKICKER, L=0.1, KICK := KCV02;

MCV03: MCV, KICK := KCV03;

MCH02: HKICKER, L=0.1, KICK := KCH01;

Q: why do I use ***:=*** ?

Orbit correction algorithms in MADX

- Best kick method (MICADO) in horizontal plane:
! Selected with **MODE=MICADO**

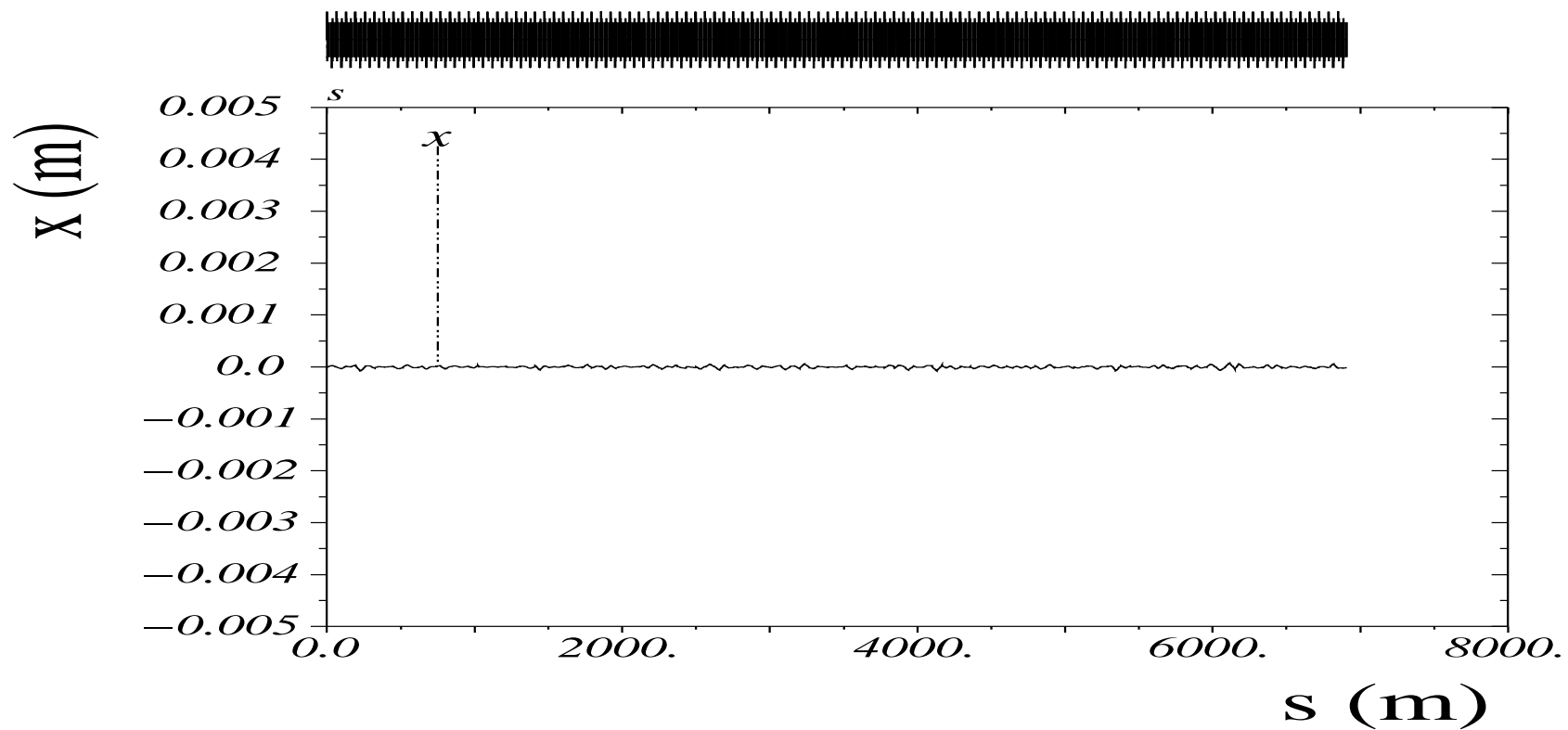
```
Correct,mode=MICADO,plane=x,  
     clist="c.tab",mlist="m.tab";
```

- Singular Value Decomposition (SVD):
! Selected with **MODE=SVD**

```
Correct,mode=SVD,plane=x,  
     clist="c.tab",mlist="m.tab";
```

- For details: see MADX Primer

Orbit after correction



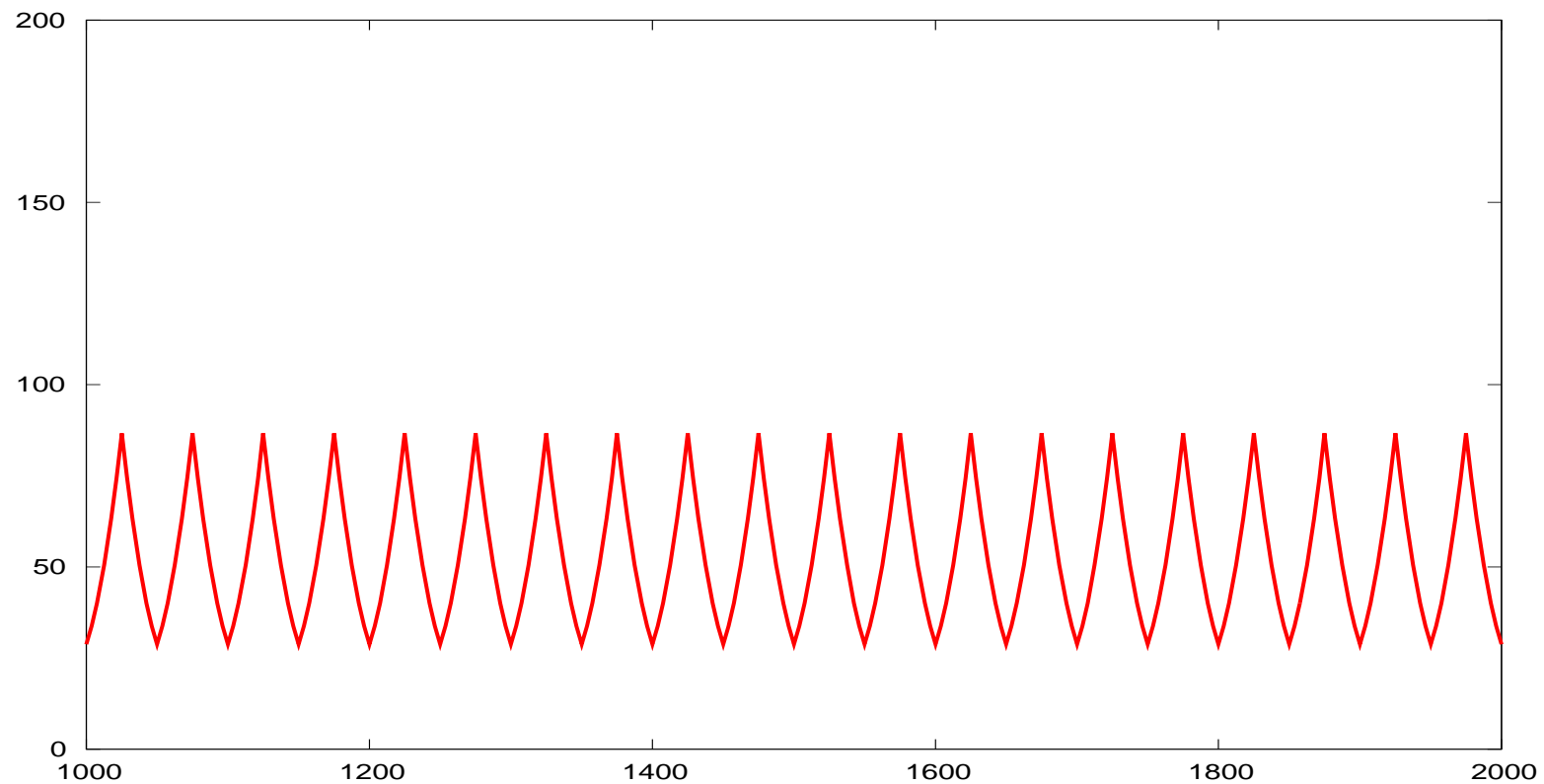
Optical matching

- To get the optical configuration you want → **matching**
- Main applications:
 - Setting **global** optical parameters (e.g. tune, chromaticity)
 - Setting **local** optical parameters (e.g. β -function, dispersion ..)
 - Correction of imperfections

Matching local parameters

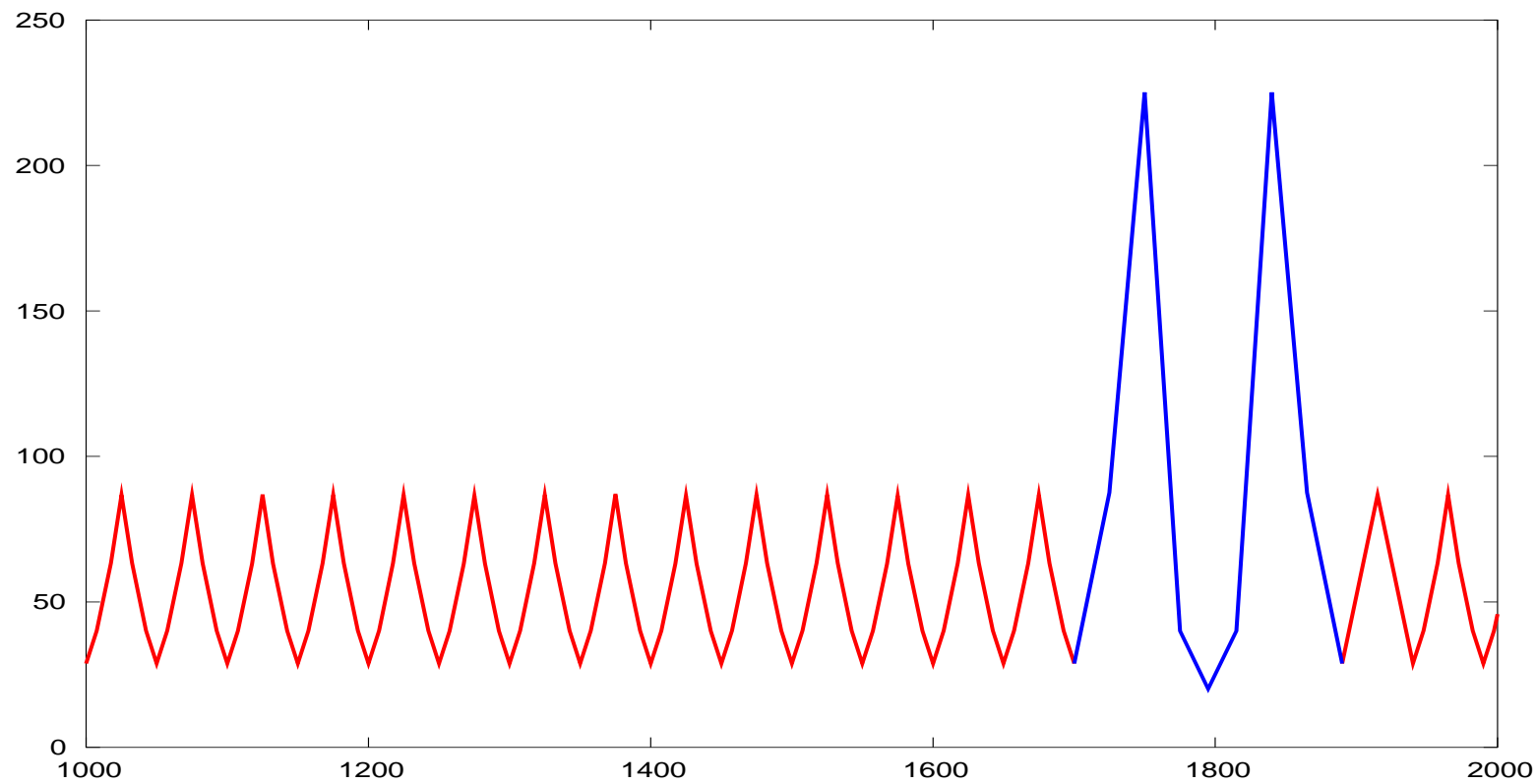
- Get local optical properties, but leave the rest of the machine unchanged
- Adjust strength of individual machine elements
- Examples for **local** matching:
 - Low (or high) β insertions
 - Dispersion suppressors

Local optical matching



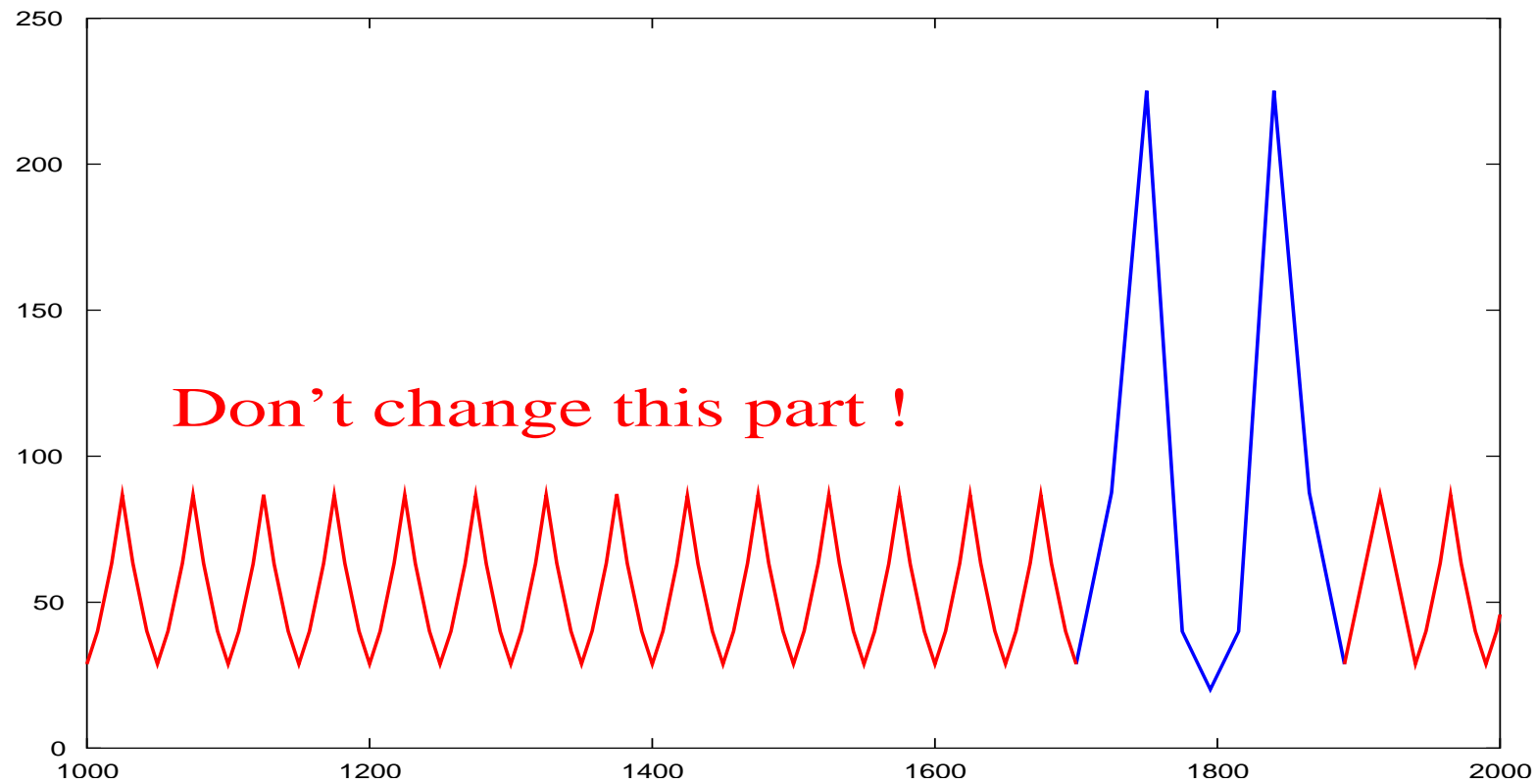
➤ What we have ...

Local optical matching



➤ What we want ...

Local optical matching



➤ What we want ...

Insertions (I)

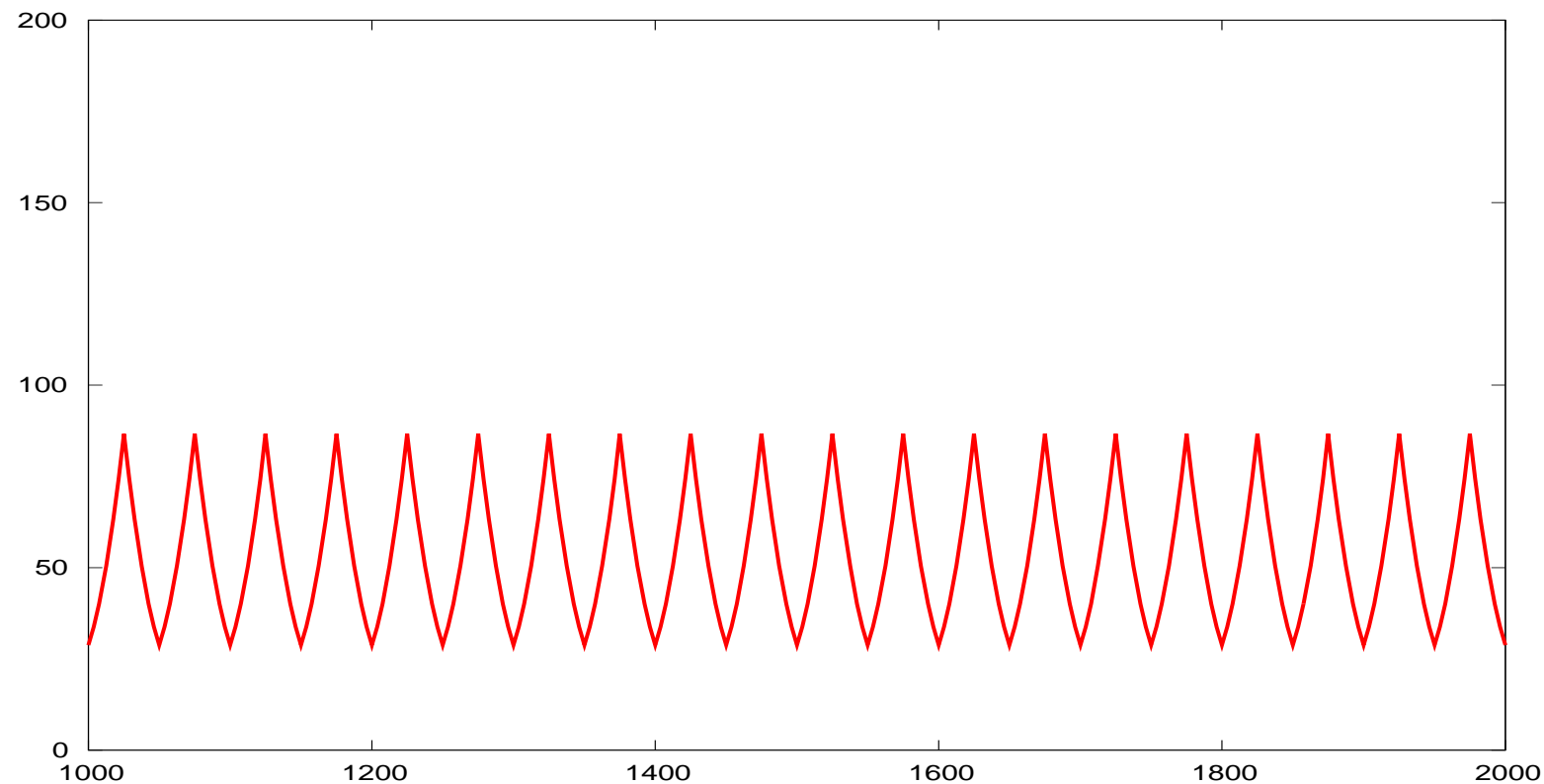
- How to add an insertion, e.g. two special cells ?
 - Start with periodic machine :

```
cassps: sequence, refer=centre, l=circum;  
start_machine: marker, at = 0;  
    n = 1;  
    while (n <= ncell) {  
        qfsps: qfsps,      at=(n-1)*lcell;  
        mbsps: mbsps,      at=(n-1)*lcell + lcell*0.25;  
        qdsps: qdsps,      at=(n-1)*lcell + lcell*0.50;  
        mbsps: mbsps,      at=(n-1)*lcell + lcell*0.75;  
        n = n + 1; }  
end_machine: marker at=circum;  
endsequence;
```

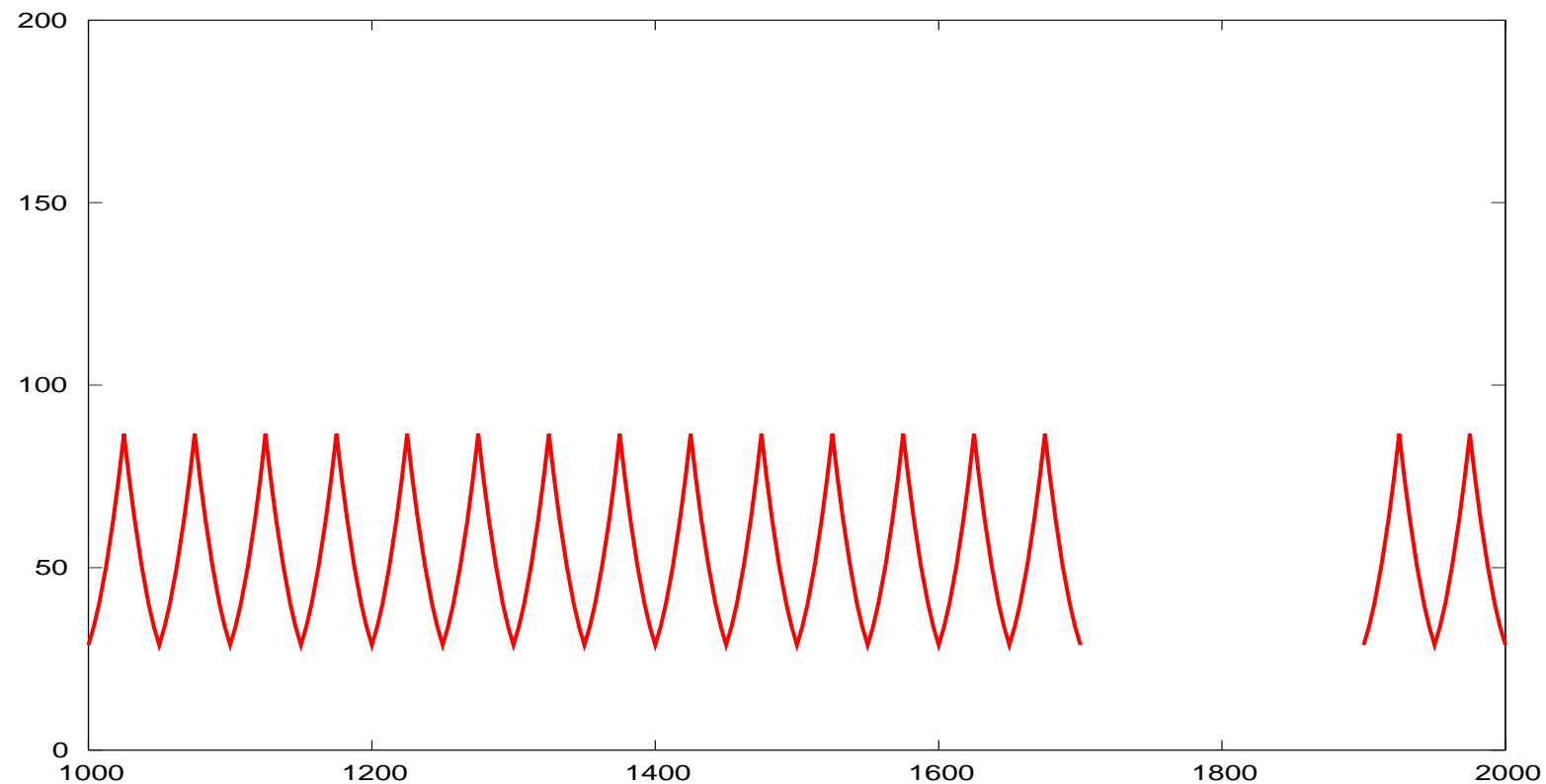
- Split it into several pieces



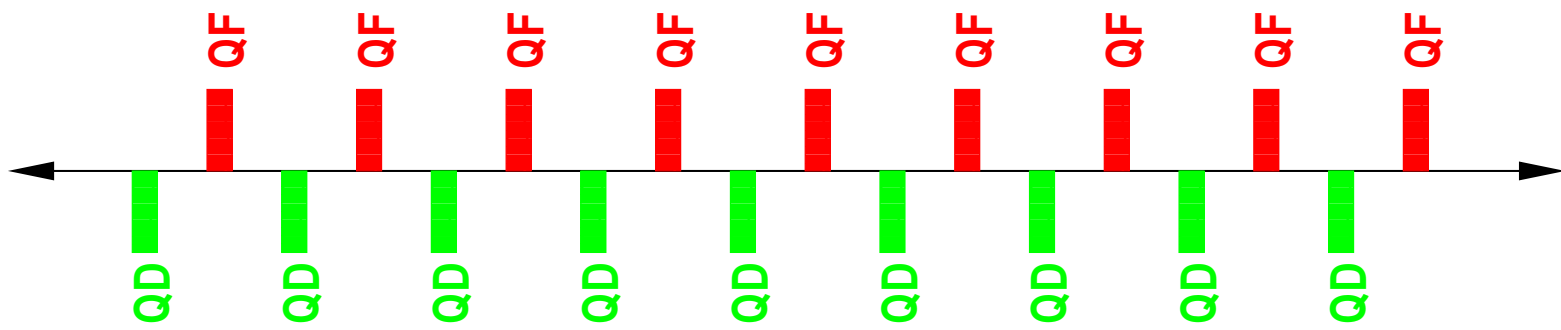
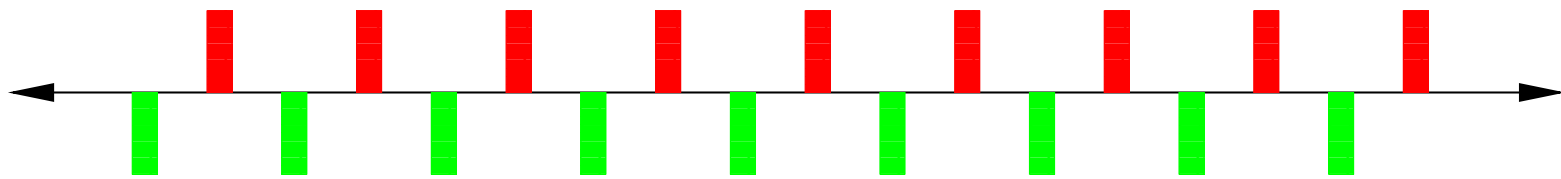
Local optical matching



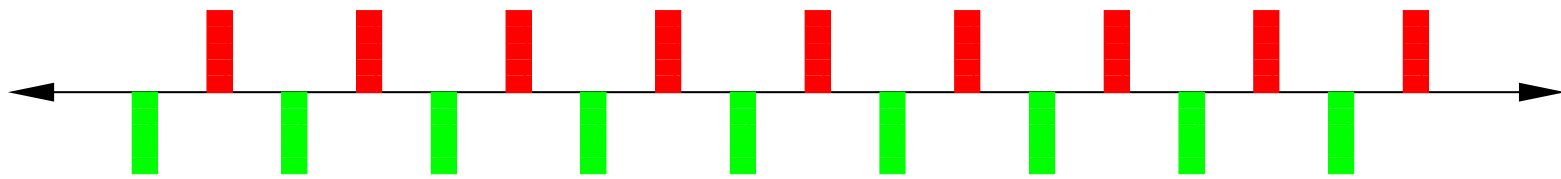
Local optical matching



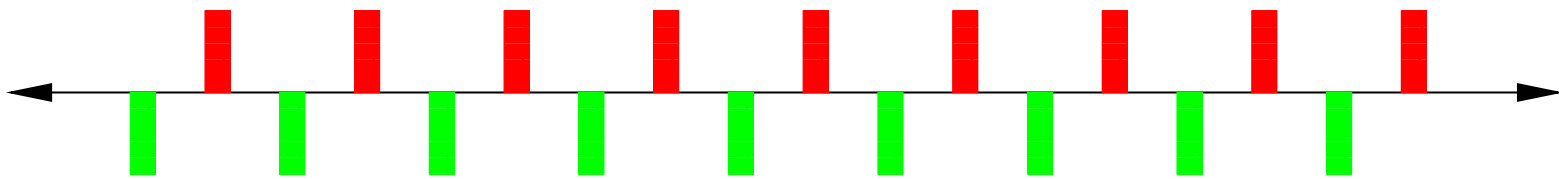
Original lattice



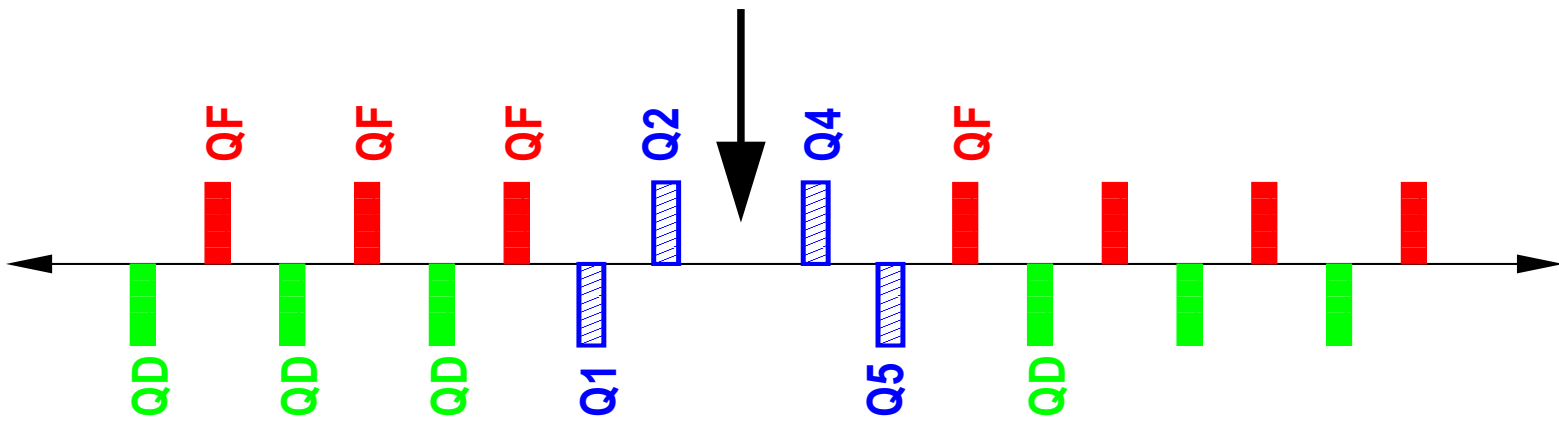
Space for insertion



Adding quadrupoles



make a symmetric drift space: $Q_2 - Q_4$



Insertions (II)

→ Split it into several pieces

```
cassps: sequence, refer=centre, l=circum;
n = 1;

while (n ≤ ncell-2) {

    qfsps: qfsps,      at=(n-1)*lcell;
    mbsps: mbsps,      at=(n-1)*lcell + lcell*0.25;
    qdssps: qdssps,    at=(n-1)*lcell + lcell*0.50;
    mbsps: mbsps,      at=(n-1)*lcell + lcell*0.75;
    n = n + 1;

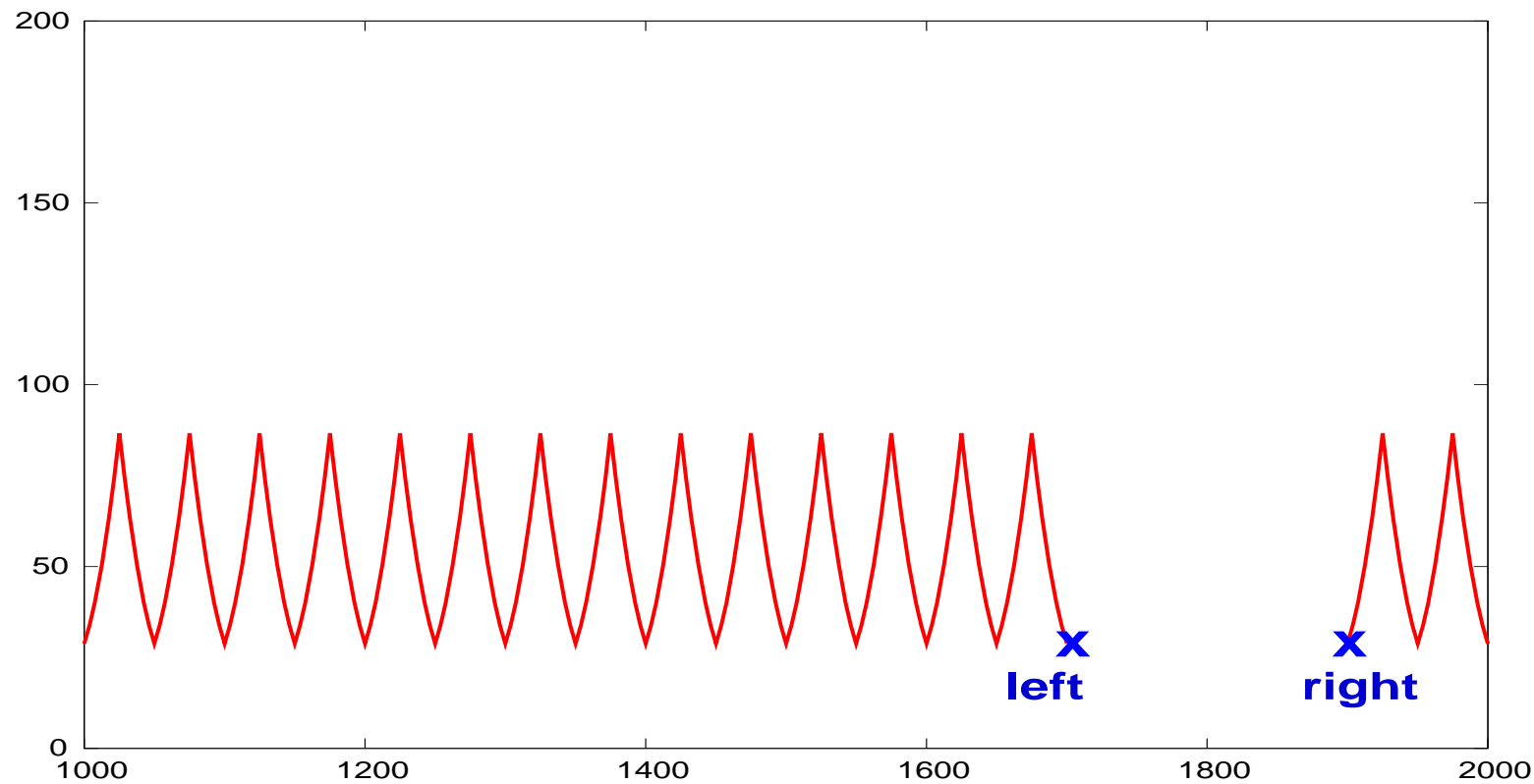
}

qf1 : qf1 ,      at=(ncell-2)*lcell;
mbsps: mbsps,      at=(ncell-2)*lcell + lcell*0.25;
qd1 : qd1 ,      at=(ncell-2)*lcell + lcell*0.50;
mbsps: mbsps,      at=(ncell-2)*lcell + lcell*0.75;

qf2 : qf2 ,      at=(ncell-1)*lcell;
mbsps: mbsps,      at=(ncell-1)*lcell + lcell*0.25;
qd2 : qd2 ,      at=(ncell-1)*lcell + lcell*0.50;
mbsps: mbsps,      at=(ncell-1)*lcell + lcell*0.75;
endsequence;
```

s1.ins.seq

Local optical matching



- Fix parameters at beginning and end of insertion

Matching techniques I(a)

- Use of markers:
 - Have no effect on the optics
 - Used to mark a position in the machine
 - Can be used as reference in matching etc.

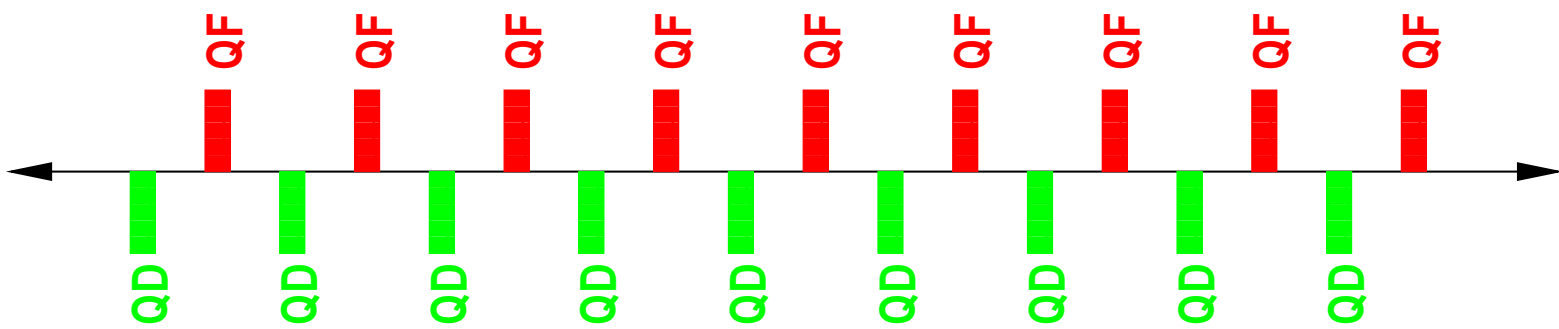
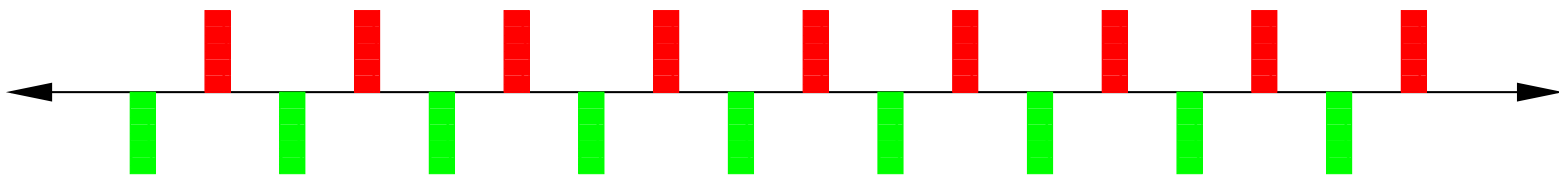
- Use:
 - left: MARKER, at=*position*;**
 - right: MARKER, at=*position*;**

Matching techniques I(b)

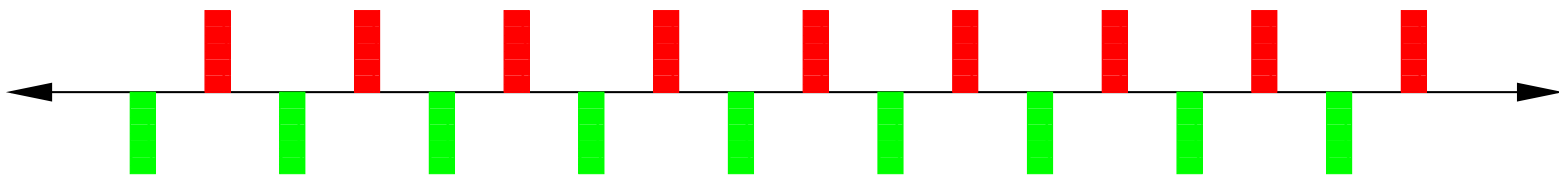
■ Markers:

- can be used with **RANGE** in **PLOT** commands:
→ **PLOT, range=left/right ...;**
- can be used with **RANGE** in **MATCH** commands:
→ **MATCH, range=left/right ...;**
- can be used with **PLACE** in **SAVEBETA** commands to store twiss functions at position of the marker
→ **SAVEBETA, label=left_beta,place=left;**

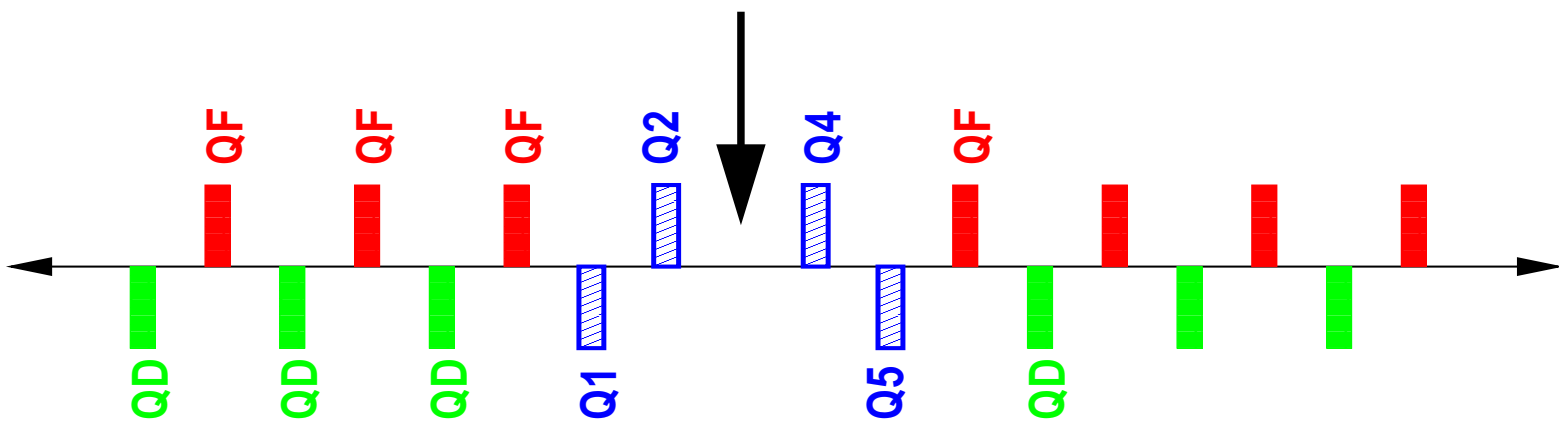
Use of MARKERS



Use of MARKERS



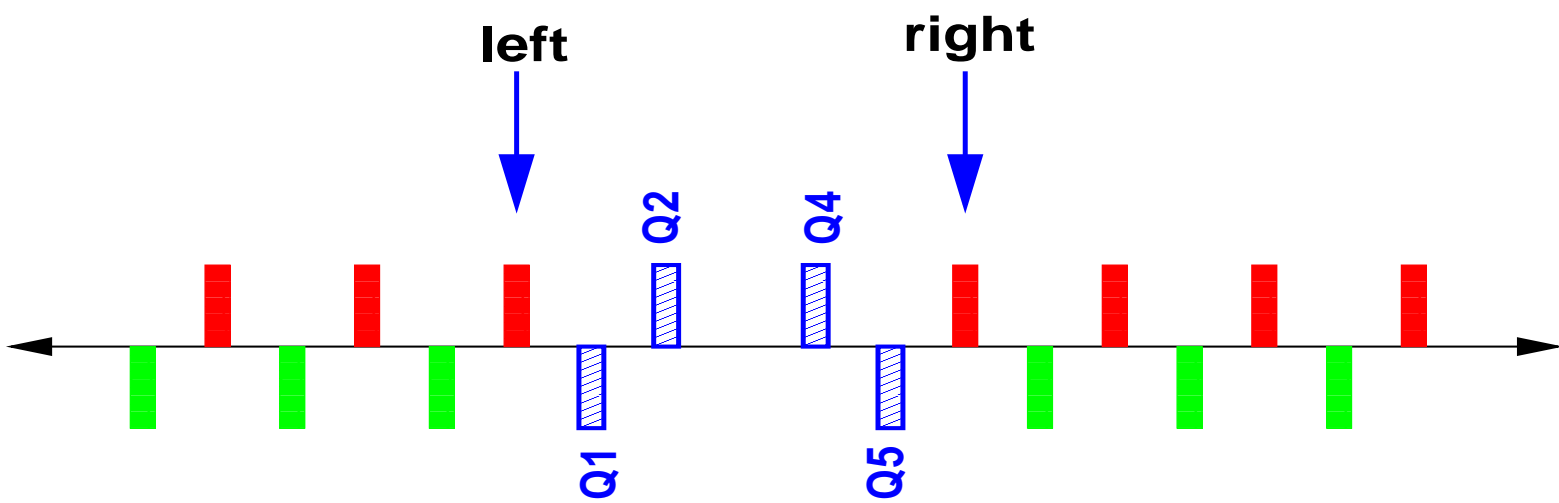
make a symmetric drift space: $Q_2 - Q_4$



Use of MARKERS



left right

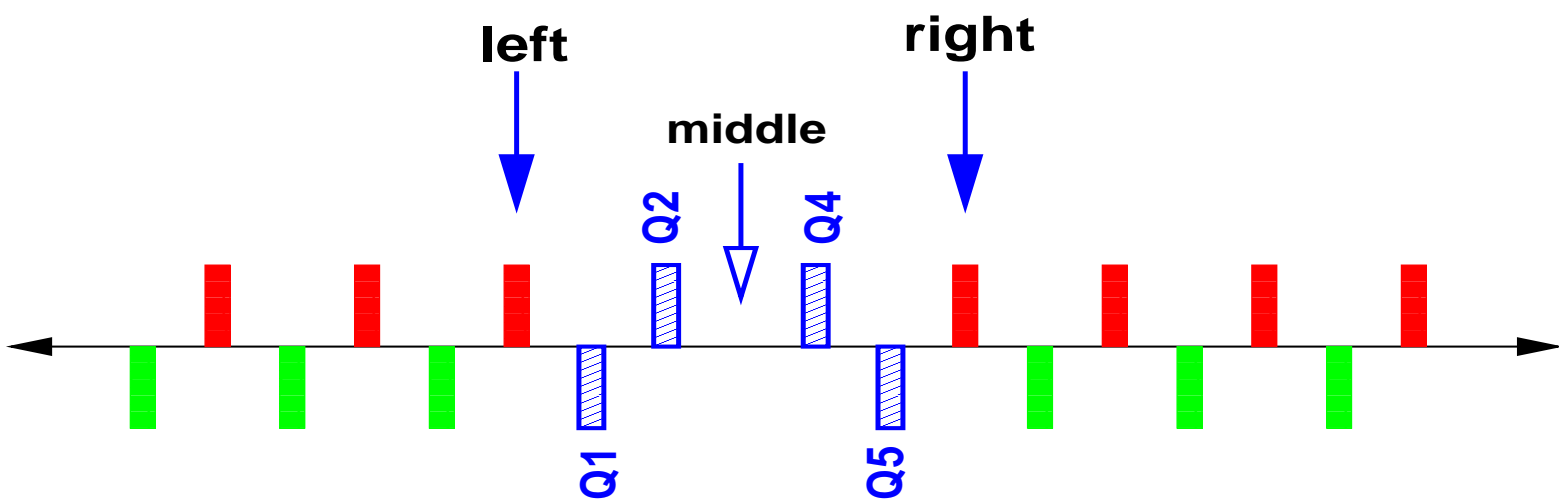


Use of MARKERS



left right

middle



Matching techniques II

- Matching is done only locally (between markers **left** and **right**), not for the whole machine, needs initial and end conditions (β_x, α_x, \dots)

```
match, range=left/right,betx=..., alfx=..., bety=...;  
vary,name=kq1.l, step=0.00001;  
vary,name=kq2.l, step=0.00001;  
! removed to become center of insertion  
// vary,name=kq3.l, step=0.00001;  
vary,name=kq4.l, step=0.00001;  
vary,name=kq5.l, step=0.00001;  
constraint,range=middle,sequence=cascell,betx=20.0,bety=50.0;  
constraint,range=right,betx=..., alfx=..., bety=..., ...;  
Lmdif, calls=100, tolerance=1.0e-21;  
endmatch;
```

Clever: Using SAVEBETA to store optical functions

```
savebeta,label=tw_left,place=left;
savebeta,label=tw_right,place=right;
twiss;
kq3.l = 0.0; ! set to 0.0 after initial computation
match, sequence=cascell,range=left/right,beta0=tw_left;
vary,name=kq1.l, step=0.00001;
vary,name=kq2.l, step=0.00001;
// vary,name=kq3.l, step=0.00001;
vary,name=kq4.l, step=0.00001;
vary,name=kq5.l, step=0.00001;
constraint,range=middle,sequence=cascell,betx=20.0,bety=50.0;
constraint,range=right,sequence=cascell,beta0=tw_right;
Lmdif, calls=100, tolerance=1.0e-21;
endmatch;
```

Matching techniques IV

- Constraints on **all quadrupoles**, using limits:

```
match, sequence=cascell;
vary, name=kqf, step=0.00001;
vary, name=kqd, step=0.00001;
constraint, pattern="^qf.*", sequence=cascell, betx < 100.0;
constraint, pattern="^qd.*", sequence=cascell, bety < 100.0;
Lmdif, calls=100, tolerance=1.0e-21;
endmatch;
```

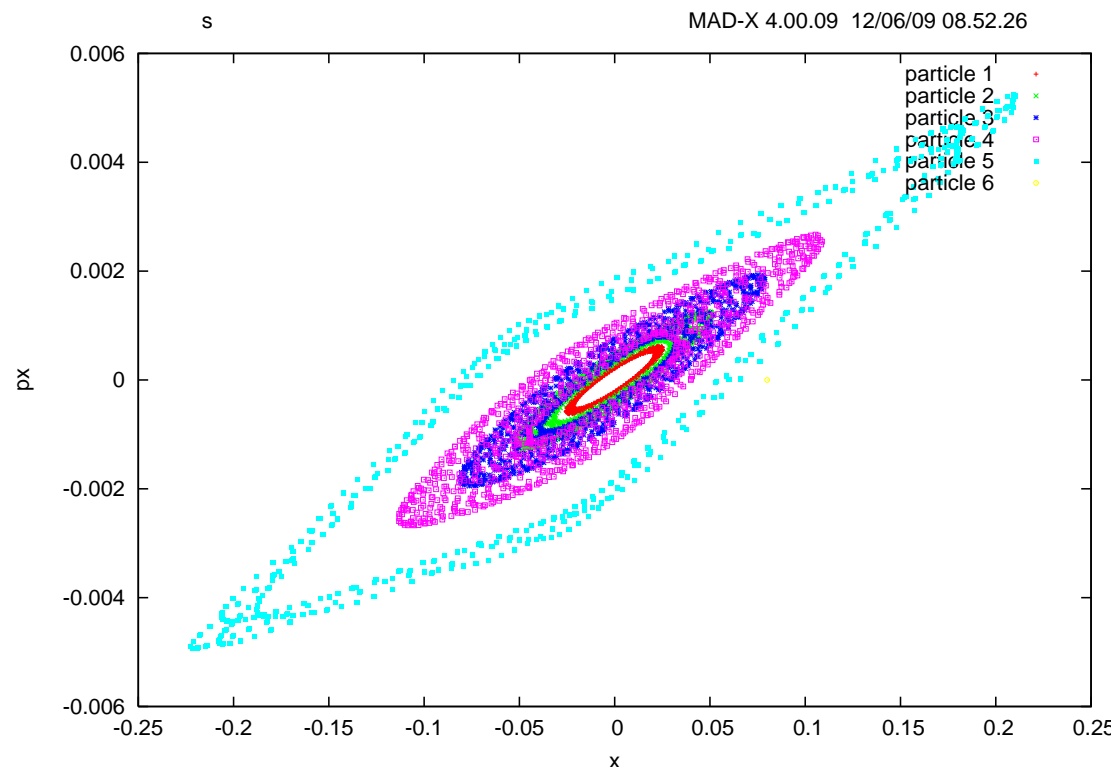
Particle tracking

- To track 4 particles for 1024 turns, add:

```
track, file=track.out, dump;  
start, x= 2e-2, px=0, y= 2e-2, py=0;  
start, x= 4e-2, px=0, y= 4e-2, py=0;  
start, x= 6e-2, px=0, y= 6e-2, py=0;  
start, x= 8e-2, px=0, y= 8e-2, py=0;  
run, turns=1024;  
endtrack;  
  
plot, file="MAD_track", table=track, haxis=x, vaxis=px,  
      particle=1,2,3,4, colour=1000, multiple, symbol=3;  
plot, file="MAD_track", table=track, haxis=y, vaxis=py,  
      particle=1,2,3,4, colour=1000, multiple, symbol=3;
```

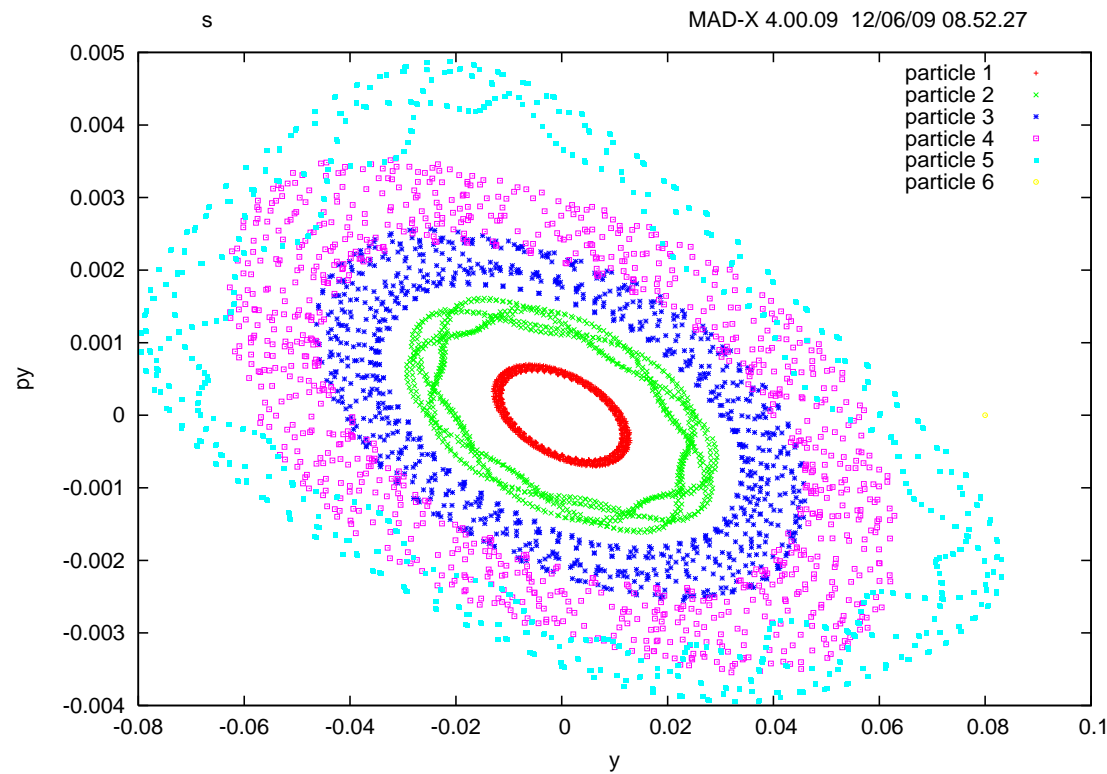
Particle tracking

- Phase space plot in horizontal coordinates:



Particle tracking

- Phase space plot in vertical coordinates:



What we do not need (here !) ...

- Higher order effects
- IBS, beam-beam elements
- Equilibrium emittance (leptons)
- RF and acceleration