20.4 Solutions of Exercises of Chapter 9: Weak Focusing Synchrotron

6511 **9.1**

6512 Construct Saturne I. Spin Resonances

A photo of Saturne I synchrotron can be found in Fig. 9.3. A schematic layout of the ring and 90 deg cell is given in Fig. 9.23. This figure as well as Tab. 9.2 which lists the parameters of the synchrotron, will be referred to in building the Saturne I ring in the following.

6517 (a) A model of Saturne I synchrotron.

⁶⁵¹⁸ DIPOLE is used to simulate the 90° cell dipole. It is necessary to have Fig. 20.55 at hand (in addition to the users' guide), when filling up the data list under DIPOLE. Some comments regarding these data:

• DIPOLE is defined in a cylindrical coordinate system.

• The bending sector is 90 degrees, however the field region extent AT has to encompass the fringe fields, at both ends of the 90 deg sector. A large 5 deg extension is taken, for a total AT=100 deg which ensures absence of truncation of the fringe fields at the AT sector boundaries, over the all radial excursion of the beam.

• RM is given the curvature radius value, $RM = B\rho/B = 0.274426548_{[Tm]}/0.03259493_{[T]} = 8.4193 m$, this makes magnet positioning and closed orbit checks easier (see below). Note that the field and reference rigidity are, as for the simulation input data, for 3.6 MeV, the injection energy, this is an arbitrary choice.

• ACENT=50 deg is the reference azimuth, for the positioning of the entrance and exit effective field boundaries (EFB). It is taken in the middle of the AT range, an arbitrary choice.

• The entrance radius in the AT sector is $RE = RM/\cos(AT - \omega^+) = RM/\cos(5^\circ)$, with $\omega^+ = 45$ deg the positioning of the entrance EFB with respect to ACENT. And similarly for the positioning of the exit reference frame, $RS = RM/\cos(AT - (ACENT - \omega^-)) = RM/\cos(5^\circ)$ with $\omega^- = -45$ deg the positioning of the exit EFB. Note that $\omega^+ - \omega^- = 90^\circ$, the value of the bend angle.

• The entrance angle TE identifies with the extension to the 90 deg sector, namely, ⁶⁵³⁹ TE=5 deg. And similarly for the positioning of exit frame, 5 deg downstream of ⁶⁵⁴¹ the exit EFB, thus TS=5 deg.

In order to build the cell it is a good idea to proceed by steps:

(i) first build a 90 deg deviation sector in the hard edge model (Tab. 20.46). FAISCEAU located next to DIPOLE indicates that a trajectory entering DIPOLE at radius R=RM, normally to the EFB (thus, $Y_0 = 0$ and $T_0 = 0$ in OBJET) exits with Y=0 and T=0. Data validation at this stage can be performed by comparing DIPOLE's transport matrix computed with MATRIX, and the theoretical expectation (after Eq. ??, Sec. 19.5, numerical values truncated to 4th decimal)



Fig. 20.55 This figure gives a representation of the data that define a dipole magnet, using DIPOLE [1]



MATRIX computation outcomes from raytracing can be found for comparison in Tab. 20.47.

It is leisurable, at this point, to choose to add fringe fields. This is the case in Exercise 9.2, however the hard edge model will be carried on with, here.

(ii) next, assemble a cell including this dipole with a pair of half-drifts at each ends, 2 m each (Fig. 9.23 and Tab. 20.48).

Table 20.46 Simulation input data file: a 45 degree sector bend in the hard edge model. The magnet is spilt into two identical halves, this is in order to allow access to particle coordinates at the center of the dipole, via FAISTORE or (here) FAISCEAU. The reference trajectory has equal entrance and exit position, and opposite sign angles. It coincides with the arc R=RM. MATRIX computes the transport matrix of the dipole (bottom of this Table), for comparison with the fringe field model, and possible comparison with matrix codes outcomes. This input data file is named SatI_DIP.inc and defines the Saturne I cell sequence segment S_SatI_DIP to E_SatI_DIP, for use with INCLUDE in subsequent exercises

```
File name: SatL_DIP.inc
Saturne I. Hard edge dipole model. Transport matrix.
'08JET'
0.274426548e3
                                                            ! Reference Brho: 3.6 MeV proton.
! Create an 11 particle set, proper for MATRIX computation.
1
'MARKER' S Satt DIP
                                                                                                            ! Cell dipole begins here.

      'DIPOLE' upstream_half
      ! Analytical modeling of a dipole magnet.

      0
      ! II=2 here, to log trajectory coordinates in zgoubi.plt. at integration steps.

      45. 841.93
      ! Field region angle=90; reference radius set to curvature radius value.

      22.5 0.3259493638 -0.6 0. 0.
      ! Reference angle ACENT set to AT/2; Bo field at RM; indices, all zero.

      0
      0.
      ! EFB 1 with fringe field extent.

      22.5 0. 1.266 -1.266 1.266
      1.266
      ! Angle to ACENT; face angle; face is straight.

 .0 0
 4 .1455 2.2670 -.6395 1.1558 0.0.0.
-22.5 0. 1.E6 -1.E6 1.E6 1.E6
0. 0.
0 0.
                                                                                                                          ! EFB 3. Unused.
.0 0.
4 .1455 2.2670 -.6395 1.1558 0. 0. 0.
22.5 0. 1.E6 -1.E6 1.E6 1.E6
                                                                                                                     ! Enge coefficients
                                                                               ! Angle to ACENT; face angle; face is straight.
1 EFB 2.
! EFB 3. Unused.
                                                               ! Degree of interpolation polynomial; flying grid sizing.
! Integration step size. It can be large in uniform field.
! Positioning of entrance and exit frames.
2 841.93 0. 841.93 0.
'MARKER' E_SatI_DIP
                                                                                                               ! Cell dipole ends here.
'FAISCEAU'
                                                                 ! Local particle coordinates.
! Compute transport matrix, from trajectory coordinates.
 'MATRIX
1 0
'END'
```

Table 20.47 Outcomes of the simulation file of Tab. 20.46

An excerpt from zgoubi.res. Coordinates of the first particle (the reference trajectory) and its path length under FAISCEAU, at OBJET on the left hand side below, locally on the right hand side:

	3	Keyword,	label(s)	: I	FAISCEAU									IPASS= 1
						TRACE	DU FAISCEAU							
						(follows	element #	2)						
						11	TRAJECTOIRES							
					OBJET						FAISCE	EAU		
		D	Y(cm)	T(r	nr) Z(cm)) P(m	r) S(cm)		D-1	Y(cm)	T(mr)	Z(cm)	P(mr)	S(cm)
0	1	1.0000	0.000	0.0	0.000	0.00	00 0.0000	0	.0000	0.000	0.000	0.000	0.000	1.322501E+03

Transport matrix of Saturne I 90 degree sector bend, in the hard edge model, two difference cases of integration step size, namely, 4 cm and 1 m (an excerpt of MATRIX computation, from zgoubi.res). It can be checked against matrix transport expectations. The "first order symplectic conditions" are very small in the 4 cm step size case, which is an indication of accurate numerical integration of the trajectories across DIPOLE; the reference trajectory (first one) exits better aligned (reference coordinates, before change of frame for MATRIX computation, are closer to zero):

```
- Case of 4 cm step size:
```

4 Keyword, label(s) : MATRIX IPASS= 1
 Reference, before
 change of frame
 (particle # 1
 D-1,Y,T,Z,S,time)

 0.00000000E+00
 4.53054326E-07
 6.27843350E-07
 0.00000000E+00
 6
 0.00000000E+00 1.32250055E+03 4.41138700E-02 TRANSFER MATRIX ORDRE 1 (MKSA units) 0.545795 11.1544 0.00000 0.00000 0.00000 9.56022 0.00000 0.00000 0.346711 -8.629576E-02 0.00000 0.00000 0.545795 -6.294423E-02 0.00000 0.00000 1.32487 0.00000 11.1544 0.545795 0.00000 0.00000 9.56022 0.00000 0.00000 0.00000 10.1951 0.346711 0.00000 0.00000 9.56022 1.32487 0.00000 0.00000 5.17640 1.00000 0.00000 0.00000 0.00000 1.00000 DetY-1 = 0.0000000278, R12=0 at -20.44 m, DetZ-1 = R34=0 at 0.000000045 -29.41 m
 R12=0 at
 -20.44
 m,
 R34=0 at
 -29.41
 r

 First order symplectic conditions (expected values = 0):
 2.7767E-08
 4.4762E-09
 0.000
 0.000
 0.000 0.000 - Case of 1 m step size: 4 Keyword, label(s) : MATRIX IPASS= 1 Reference, before change of frame (particle # 1 - D-1,Y,T,Z,s,time) : 0.00000000E+00 -7.54923113E-03 -1.08904867E-02 0.00000000E+00 0.0000000E+00 1.32249873E+03 4.41138091E-02
 TRANSFER
 MATRIX
 ORDRE
 1
 (MKSA units)

 0.545757
 11.1567
 0.00000
 < 0.00000 0.00000 0.00000 0.00000 0.00000 1.32517 10.1954 0.00000 0.346750 1.32486 9.56148 0.00000 0.00000 1.00000 5.17692 0.00000 0.00000 0.00000 0.00000 1.00000 DetY-1 = 0.0003978566, DetZ-1 = 0.0000685588 R12=0 at -20.43 m, R34=0 at -29.40 rst order symplectic conditions (expected values = 0) : 3.9786E-04 6.8559E-05 0.000 0.000 m First 0.000 0.000

Table 20.48 Simulation input data file: Saturne I cell, obtained by assembling DIPOLE taken from Tab. 20.46 together with two half-drifts. This input data file is named SatI_cell.inc and defines the Saturne I cell sequence segment S_SatI_cell to E_SatI_cell, for possible use in INCLUDE statements in subsequent exercises

File name: SatI_cell.inc.
! Saturme I, one cell of the 4-period ring.
'MARKER' SatICellMATRIX_S ! Just for edition purposes.
'OBJET'
0.27426548e3 ! Reference Brho: 3.6 MeV proton.
0.01.001.001.001 ! Create an 11 particle set, proper for MATRIX computation.
.001.01.001.001.001 ! Create an 11 particle set, proper for MATRIX computation.
.001.01.001.001 ! Create an 11 particle set, proper for MATRIX computation.
.001.01.001.001 ! Create an 11 particle set, proper for MATRIX computation.
.001.01.001.001 ! Create an 11 particle set, proper for MATRIX computation.
.001.01.001.001 ! Coordinate sampling.
0.0.0.0.0.1. ! Reference trajectory: all initial coordinates nul, relative rigidity D=1.
'MARKER' S_SatI_cell
'NARKER' S_SatI_cell
'NARKER' L_SatI_DIP:E_SatI_DIP!
'AMARKER' E_SatI_cell
'MARKER' L_SatI_cell
'MARKER' S_AICCELMATRIX_E ! Local particle coordinates.
'FMISG'
'MARKER' S_AICCELMATRIX_E ! Just for edition purposes.
'END'

The TWISS command down the sequence (Tab. 20.48) produces the periodic beam matrix results shown in Tab. 20.49

 Table 20.49
 Results obtained running the simulation input data file of Tab. 20.48, Saturne I cell

 - an excerpt from zgoubi.res

6557

The TWISS command also produces a zgoubi.TWISS.out file which details the optical functions along the sequence (at the downstream end of the optical elements. The header of that file details the optical parameters of the structure (Tab. 20.50).

⁶⁵⁵⁵ Lattice parameters

Table 20.50 An excerpt of zgoubi.TWISS.out file resulting from the execution of the Saturne I cell simulation input data file of Tab. 20.48. Note that the ring (4-period) wave numbers are 4 times the cell values Q1, Q2 displayed here. Optical functions (betatron function and derivative, orbit, phase advance, etc.) along the optical sequence are listed as part of zgoubi.TWISS.out following th header. The top part and last line of that listing are given below

C	LENGTH	%le	17.22500552			
C	ALFA	%le	1.919448707			
C	ORBIT5	%le	-0			
C	GAMMATR	%le	0.7217914685			
C	Q1	%le	0.1810314404	[fractional]		
C	Q2	%le	0.2221459901	[fractional]		
C	DQ1	%le ·	-0.6022172911			
C	DQ2	%le	0.3800544183			
C	DXMAX	%le	2.10586311E+01	@ DXMIN	%le	2.10482503E+01
C	DYMAX	%le	0.0000000E+00	@ DYMIN	%le	0.0000000E+00
C	XCOMAX	%le	2.10528899E-01	@ XCOMIN	%le	0.0000000E+00
C	YCOMAX	%le	0.0000000E+00	@ YCOMIN	%le	0.0000000E+00
C	BETXMAX	%le	1.57006971E+01	<pre>@ BETXMIN</pre>	%le	1.44132839E+01
¢	BETYMAX	%le	1.30884296E+01	@ BETYMIN	%le	1.14110171E+01
C	XCORMS	%le	6.05227342E-04			
¢	YCORMS	%le	 not compute 	d		
C	DXRMS	%le	2.98427468E-03			
¢	DYRMS	%le	0.0000000E+00			

Top and bottom four lines (truncated) of zgoubi.TWISS.out optical functions listing, including the periodic β_x , β_y (β_Y , β_Z in zgoubi notations) and D_x (η_Y in zgoubi notations) values at cell ends:

# alfx	btx	alfy	bty	alfl	btl	Dx	Dxp
3.1789380E-009	1.4426805E+001	-8.1670406E-009	1.1411067E+001	0.0000000E+000	0.0000000E+000	2.1058631E+001	1.1261478E-003
1.3369744E-008	1.4426805E+001	-1.7451004E-009	1.1411067E+001	0.0000000E+000	0.0000000E+000	2.1048250E+001	1.6725930E-009
1.3369744E-008	1.4426805E+001	-1.7451004E-009	1.1411067E+001	0.0000000E+000	0.0000000E+000	2.1048250E+001	1.6725930E-009
-1.3863082E-001	1.4704066E+001	-1.7526845E-001	1.1761604E+001	0.0000000E+000	0.0000000E+000	2.1048250E+001	1.6725930E-009
1.3919473E-001	1.4692541E+001	1.7526999E-001	1.1761559E+001	0.0000000E+000	0.0000000E+000	2.1048250E+001	1.6725925E-009
1.3919473E-001	1.4692541E+001	1.7526999E-001	1.1761559E+001	0.0000000E+000	0.0000000E+000	2.1048250E+001	1.6725925E-009
4.3382070E-004	1.4413284E+001	7.7156422E-007	1.1411017E+001	0.0000000E+000	0.0000000E+000	2.1048250E+001	1.6725925E-009
4.3382070E-004	1.4413284E+001	7.7156422E-007	1.1411017E+001	0.0000000E+000	0.0000000E+000	2.1048250E+001	1.6725925E-009

6561 Tune scan

A simulation is given in Tab. 20.51, derived from Tab. 20.48: TWISS has been replaced by MATRIX, a REBELOTE do loop repeatedly changes n. A plot of the scan is given in Fig. 20.56, some detailed values are given in Tab. 20.52.

Table 20.51 Simulation input data file: tune scan, using REBELOTE to repeatedly change *n*. Beam matrix and wave numbers are computed by MATRIX, from the coordinates of the 11 particle sample generated by OBJET[KOBJ=5]

Saturne I, tune scan. 'MARKER' SatI_Qscan_S ! Just for edition purposes. 'OBJET 0.274426548e3 ! Reference Hroh: 3.6 MeV proton. ! Create an 11 particle set, proper for MATRIX computation. . 001 .01 .001 .001 .001 .001 .001 ! Coordinate sampling 0. 0. 0. 0. 0. 1. ! Reference trajectory: all initial coordinates nul, relative rigidity D=1. 1 ! Reference Brho: 3.6 MeV proton. 'MARKER' S_SatI_cell 'DRIFT' half_drift 200. 'INCLUDE' 1 ./SatI_DIP.inc[S_SatI_DIP:E_SatI_DIP] 'DRIFT' half drift 200. 'MARKER' E_SatI_cell 'FAISCEAU' ! Local particle coordinates. MATRIX 1 11 PRINT ! Comoute a 10+4 period transport matrix, and tunes. Save outcomes to zgoubi.MATRIX.out. 'REBELOTE' ! A do loop: repeat the section above commencing at the top of the file, 10 1.1 0 1 10 times DIPOLE 6 -0.757:-0.5 ! Change the value of parameter 30 (namely, n) in DIPOLE (prior to repeating). ! in any DIPOLE in the sequence. 'SYSTEM' gnuplot <./gnuplot_MATRIX_Qxy.gnu
'MARKER' SatI_Qscan_E</pre> ! Plot tunes vs index.
! Just for edition purposes. 'END'

gnuplot script to obtain Fig. 20.56:

Ulplot Script to Obtain Fig. 20.55.
./gnuplot_MATRIX_Qsy.gnu
set xlabel "index m'sst yllabel "{/Symbol n}_x, {{/Symbol n}_x^2+{/Symbol n}_y^2}{1/2}"
set y2label "{/Symbol n}_y"; set xtics; set ytics nomirror; set y2tics nomirror; ncell=4
set key t 1; set key maxrow 2; set yrange [:1.3]; set y2range [:1.66]
n1 = -0.75; dmc(.757-5)/01; R=10.9658; rho=8.4193
plot \
"zgoubi.MATRIX.out" u (n1+(\$61-1)*dn): \
(\$61517 \$56 *ncell :1/0) wp pt 5 lt 1 lw .5 lc rgb "red" tit "{/Symbol n}_x " ,\
"zgoubi.MATRIX.out" u (n1+(\$61-1)*dn): \
(\$61517 \$56 *ncell :1/0) wp pt 5 lt 1 lw .5 lc rgb "red" tit "{/Symbol n}_x " ,\
"zgoubi.MATRIX.out" u (n1+(\$61-1)*dn): \
(\$61517 \$57 *ncell :1/0) wp pt 5 lt 1 lw .5 lc rgb "red" tit "{/Symbol n}_y " ,\
"zgoubi.MATRIX.out" u (n1+(\$61-1)*dn): \
(\$61517 \$57 *ncell :1/0) axes xj2 w pt 6 lt 3 lw .5 lc rgb "blue" tit "{/Symbol n}_y " ,\
"zgoubi.MATRIX.out" u (n1+(\$61-1)*dn): \
(\$61517 \$57 *ncell :1/0) axes xj2 w pt 6 lt 3 lw .5 lc rgb "blue" tit "{/Symbol n}_y " ,\
"zgoubi.MATRIX.out" u (n1+(\$61-1)*dn): \
(\$61517 \$557*ncell :1/0) axes xj2 w pt 6 lt 3 lw .5 lc rgb "blue" tit "{/Symbol n}_y " ,\
"zgoubi.MATRIX.out" u (n1+(\$61-1)*dn): \
(\$61517 \$557*ncell :1/0) axes xj2 w pt 6 lt 3 lw .5 lc rgb "blue" tit "theor. " ,\
"zgoubi.MATRIX.out" u (n1+(\$61-1)*dn): \
(\$61517 \$557*ncell :1/0) axes xj2 w pt 7 lt 1 lc rgb "blue" tit "theor. " ,\
"zgoubi.MATRIX.out" u (n1+(\$61-1)*dn): \
"zgoubi.MATRIX.ou



Table 20.52 Dependence of wave numbers on index n, from numerical raytracing (columns denoted "ray-tr.") and from theory

	۱	'Y	ν_Z				
n	ray-tr.	$\sqrt{(1-n)rac{R}{ ho_0}}$	ray-tr. $\sqrt{n\frac{R}{\rho_0}}$				
0.5	0.810353	0.806987	0.810353 0.806987				
0.6	0.724125	0.721791	0.888583 0.884010				
0.7	0.626561	0.625089	0.960806 0.954840				
0.757	0.563635	0.562580	0.999804 0.992955				

(b) Betatron functions of Saturne I cell.

Among the various ways to produce the betatron functions along the sequence (and throughout the DIPOLEs), here are two possibilities, based on the storage of particle coordinates in zgoubi.plt during stepwise raytracing:

 a direct way consists in using OBJET[KOBJ=5.1] and transport the 11-particle set so obtained across the sequence. Then, betaFromPlt from zgoubi toolbox [2] can be used to compute the transport matrix, step by step across the sequence, from the coordinate values logged in zgoubi.plt during the stepwise integration;

2. an indirect way consists in launching a few particles on a common invariant (horizontal and/or vertical) and subsequently plot the s-dependent quantities $\hat{Y}^2(s)/\epsilon_Y$ and/or $\hat{Z}^2(s)/\epsilon_Z$. The maximum value of the latter, a function of the distance s, is the betatron function along the sequence, $\beta_{Y,Z}(s)$.

The second method is used here (this is an arbitrary choice. Exercises may be found in the various Chapters, that use the first method and may be referred to, if desired).

The input data file to derive the betatron function following method (2) above is given in Tab. 20.53. The initial ellipse parameters (under OBJET) are the periodic values, namely, $\alpha_Y = \alpha_Z = 0$, $\beta_Y = 14.426$ m, $\beta_Z = 11.411$ m, they are a sub-product of the TWISS procedure performed in (a), to be found in zgoubi.TWISS.out (Tab. 20.50). The resulting envelopes and their squared value are shown in Fig. 20.57. Note that this raytracing also provides the coordinates of the 60 particles on their common upright invariant

$$x^2/\beta_x + \beta_x x'^2 = \epsilon_x/\pi$$

at start and at the end of the cell (with x standing for either Y or Z, and $\epsilon_{Y,Z}/\pi$ =

 10^{-4} , here). This allows checking that the initial ellipse parameters (under OBJET,

Tab. 20.53) are effectively periodic values, and that the raytracing went correctly,

namely by observing that the initial and final ellipses do superimpose (Fig. 20.58).

 Table 20.53
 Simulation input data file: raytrace 60 particles across Saturne I cell to generate beam envelopes. Store particle data in zgoubi.plt, along DRIFTs and DIPOLEs. The INCLUDE file and segments are defined in Tab. 20.48



Fig. 20.57 Left: horizontal and vertical envelopes as generated by plotting the coordinates Y(s) (greater excursion, red, along the drifts and at dipole center) or Z(s) (smaller excursion, blue) across the Saturne I cell, of 60 particles evenly distributed on a common $10^{-4} \pi \mu m$ invariant, either horizontal or vertical (while the other invariant is zero). Right: a plot of $Y^2(s)/\epsilon_Y$ and $Y^2(s)/\epsilon_Y$: the extrema identify with $\beta_Y(s)$ and $\beta_Z(s)$, respectively. The extrema extremorum values are $\hat{\beta}_Y = 14.4 \text{ m}$ and $\hat{\beta}_Z = 15.7 \text{ m}$, respectively. These plots are obtained using zpop, which reads stepwise particle data from zgoubi.plt



Fig. 20.58 Sixty particles evenly distributed on a common periodic invariant (of value either $\epsilon_Y = 10^{-4}\pi\mu$ m and $\epsilon_Z = 0$, left plot, or the reverse, right plot) have been tracked from start to end of the cell. These periodic invariants are defined assuming the periodic ellipse parameters determined from prior TWISS, given in Tab. 20.50; values resulting from an *rms* match of the coordinates are given in the figure, and do agree with those TWISS data. The figure shows the good superposition of the start and end invariants (the start and end *rms* match ellipse parameters show negligible difference), which confirms the correct value of the periodic ellipse parameters, namely, left graph: horizontal phase space at start (crosses) and end (dots) of the cell; right graph: vertical phase space at start (crosses) and end (dots) of the cell

6584 Dispersion function

Raytracing off-momentum particles on their chromatic closed orbit provides the periodic dispersion function. In order to do so, the input data file of Tab. 20.53 can be used, it just requires changing OBJET to the following:

 6588
 'OBJET'

 6589
 0.274426548e3
 ! Reference Brho: 3.6 MeV proton.

 6590
 2
 ! Create particles individually.

 6591
 3
 ! Three particles

 6592
 +.21056 0.0.0.0.1.0001 'p' ! Chromatic orbit coordinates Y0 and T0 for D=1.001 relative rigidity.

 6593
 0.0.0.0.0.0.0.09999 'm' ! Chromatic orbit coordinates Y0 and T0 for D=0.999 relative rigidity.

 6594
 -.21056 0.0.0.0.0.0.9999 'm' ! Chromatic orbit coordinates Y0 and T0 for D=0.999 relative rigidity.

The position and angle of the chromatic particles, which are offset by $\Delta p/p = \pm 10^{-4}$, are drawn from the value of the periodic dispersion $\eta_Y = 21.05$ m and its derivative $\eta'_Y \approx 0$ (Tab. 20.50), namely, $Y_0 = \eta_Y \Delta p/p = \pm 0.2105$ cm and $T_0 = \eta_Y \Delta p/p = 0$.

Running Tab. 20.53 simulation file with this new OBJET produces the following coordinates at FAISCEAU, located at the end of the sequence (an excerpt from zgoubi.res):

6603		13	Keyword,	label(s)	: FA	ISCEAU								IPASS= 1	
6604							TRACE DU	J FAISCEAU							
6605							(follows el	.ement #	12)						
6606							3 TF	AJECTOIRES							
6607						OBJET					FAISC	EAU			
6608			D	Y(cm)	T(mr) Z(cm)	P(mr)	S(cm)	D-1	Y(cm)	T(mr)	Z(cm)	P(mr)	S(cm)	
6609	р	1	1.0001	0.211	0.00	0.000	0.000	0.0000	0.0001	0.211	0.000	0.000	0.000	1.722831E+03	1
6610	ο	1	1.0000	0.000	0.00	0.000	0.000	0.0000	0.0000	0.000	0.000	0.000	0.000	1.722501E+03	2
6611	m	1	0.9999	-0.211	0.00	0.000	0.000	0.0000	-0.0001	-0.210	0.000	0.000	0.000	1.722170E+03	3

The local coordinates Y, T (under FAISCEAU, right hand side) and initial coordinates Y_0 , T_0 (under OBJET, left hand side) are identical (to better than 5 μ m, 0.5 μ rad accuracy, respectively - zgoubi.fai can be consulted for greater precision on these values), confirming the periodicity of these chromatic trajectories. Figure 20.59 shows the particle trajectories through the cell DIPOLE, they appear to be at constant radius as expected.

Fig. 20.59 A plot of the radial excursion, across DIPOLE body (namely, AT=90° extent,(Tab. 20.48), of an onmomentum particle (its radial position along the dipole body is $R_0 \approx 8.4193$ m, corresponding to Y=0 in this graph) and two particles at respectively $dp/p = \pm 10^{-3}$. A graph obtained using zpop, which reads stepwise particle data from zgoubi.plt: menu 7; 1/1 to open zgoubi.plt; 2/[6,2] to select Y versus distance; 7 to plot



6618 (c) Some verifications regarding the model.

The field along large excursion orbits can be logged in zgoubi.plt, using option IL=2 (or 20, or 200, etc. for printout every 10, or 100, etc. integration step) under DIPOLE.

The simulation file of Tab. 20.53 is used to raytrace five particles, with OBJET changed to the following:

0024	OBJEI	
625	0.274426548e3	! Reference Brho: 3.6 MeV proton.
626	2	! Create particles individually,
627	5 1	! five particles.
628	+0.21056 0. 0. 0. 0. 1.01 'p'	! Chromatic orbit coordinates for D=1.01 relative rigidity.
629	0. 0. 0. 0. 0. 1. '0'	! On-momentum closed orbit.
630	-0.21056 0. 0. 0. 0. 0.99 'm'	! Chromatic orbit coordinates for D=0.99 relative rigidity.
631	0. 0. 5. 0. 0. 1. 'm'	! Initial vertial excursion is Z0= 5 cm off-mid-plane.
632	0. 0. 20. 0. 0. 1. 'm'	! Initial vertial excursion is Z0=20 cm off-mid-plane.
633	1 1 1 1 1	

Apart from the on-momentum particle (2nd in the list) this OBJET defines two particles on $\Delta p/p = \pm 1\%$ chromatic orbit (1st and 3rd in the list), this is an excursion of a few tens of centimeters, large as requested, as $\Delta x \approx 38 \times dp/p$. OBJET also defines 2 particles launched into the cell at respectively $Z_0 = 5$ cm and $Z_0 = 20$ cm.

The magnetic field as a function of the azimuthal angle in DIPOLE frame, along these trajectories across the upstream DIPOLE of the cell, is shown in Fig. 20.60. The field curves for the first four trajectories essentially superimpose except for the fringe field regions (Fig. 20.60), due to the wedge angles. This behaves as expected. Detail inspection is possible, from the detailed particle coordinate and field data in zgoubi.plt - this is out of the scope of the present question.

The field along the 5th particle trajectory features overshoots (Fig. 20.60), this is due to the very large vertical excursion ($Z \approx 20$ cm in the entrance fringe field region). It looks reasonable, however it may be an artifact in the case that the high order derivatives of the field in that region are large, resulting from the truncated Taylor series method used for off mid-plane field extrapolation [1, Sec. 1.3.3].

-8.3

Zgoubi Zpop



(T)

angle (rad

Fig. 20.60 Magnetic field along 5 different trajectories across the upstream DIPOLE, including four large horizontal and vertical excursion cases 359

(d) Sinusoidal approximation of the betatron motion.

The approximation

$$y(\theta) = A \cos(v_Z \theta + \phi)$$

is checked here considering the vertical motion (considering the horizontal motion
 leads to similar conclusions). The value of the various parameters in that expression
 are determined as follows:

- the particle raytraced for comparison is launched with an initial excursion $Z_0(\theta = 0) = 5 \text{ cm}$ (4th particle in OBJET, above). At the launch point (middle of the drift) the beam ellipse is upright (Fig. 20.58), whereas phase space motion is clockwise, thus take

A = 5 cm and
$$\phi = \pi/2$$

- the vertical betatron of the 4-cell ring tune is (Tab. 20.50)

 $v_{\rm Z} = 4 \times 0.222146 = 0.888284$

 $-\theta = s/R$ and $R = \oint ds/2\pi$ with (Tab. 20.50)

 $2\pi R$ = circumference = $2\pi \times 10.9658 = 68.9 \text{ m}$

The comparison with a trajectory obtained from raytracing is given in Fig. 20.61 and confirms the validity of the sinusoidal approximation.





(e) An acceleration cycle. Symplecticity checks. 6655

Eleven particles are launched for a 30,000 turn tracking at a rate of

$$\Delta W = q\hat{V} \cos \phi_s = 200 \times \sin 150^0 = 100 \text{ keV/turn}$$

 $(E: 3.6 \rightarrow 3.0036 \text{ GeV})$, all evenly distributed on the same initial vertical invariant 6656

$$Z^2/\beta_Z + \beta_Z Z'^2 = \epsilon_Z/\pi \tag{20.14}$$

6657

with $\epsilon_Z/\pi = 10^{-4}$ m, or, normalized, $\beta \gamma \epsilon_Z/\pi = 0.08768 \times 10^{-4}$ m. The simulation file is given in Tab. 20.54. CAVITE[IOPT=3] is used, it provides an RF phase independent boost

$$\Delta W = q\hat{V}\sin\phi_s$$

as including synchrotron motion is not necessary here, even better, this ensures 6658 constant depolarizing resonance crossing speed, so precluding any possibility of 6659 multiple crossing (it can be referred to [3] regarding that effect). 6660

Table 20.54 Simulation input data file: track 11 particles launched on the same vertical invariant, with quasi-zero horizontal invariant. The INCLUDE adds the Saturne I cell four times, the latter is defined in Tab. 20.48 and Fig. 9.23

Saturne I ring. Polarization landscape. 'MARKER' SatIPolarLand_S ! Just for edition purposes. '08JET' ! Reference Brho: 3.6 MeV proton. 8 ! Create a set of 60 particles evenly distributed on the same invariant; 111 1 ! case of 60 particles on a vertical invariant; use 60 1 1 instead for horizontal invariant. 0. 0. 0. 0. 0. 1. 1. 14.426 le-4 ! Periodic optical functions and invariant value, horizontal and 0. 11.411 le-4 ! Veriodic optical functions and invariant value, momentum spread. ! Commented. ! Reference Brho: 50 MeV proton. ! Create an 11 particle set, proper for MATRIX computation. !'MCOBJET' !1.03527036749193e3 !3 .3 !200 1200 12 2 2 2 2 2 2 10. 0. 0. 0. 0. 1. 10. 14.426 25e-6 3 10. 11.411 10e-6 3 10. 1. 1.e-8 3 1123456 234567 345678 ! Periodic alpha_Y, beta_Y, and invariant value; ! Periodic alpha_Z, beta_Z, and invariant value. 'PARTICUL' ! Necessary data in order to allow (i) spin trackingand, and (ii) acceleration. ! Switch on spin tracking, ! all initial spins vertical. PROTON 'SPNTRK' 3 'FAISCEAU' 'FAISTORE' b_polarLand.fai ! Log particle data in b_polarLand.fai, turn-by-turn; "b_" imposes ! binary write, which results in faster i/o. 'SCALING' 1 1 DIPOLE ! Causes field increase in DIPOLE, in correlation to particle ! rigidity increase by CAVITE. -1 1. 1 ! 4 cells follow. 'INCLUDE' 1 4* ./SatI_cell.inc[S_SatI_cell:E_SatI_cell] 'CAVITE' 3 0 0 200e3 0.523598775598 ! 20e3 0.523598775598 ! Acceleration rate is 200*0.5=100keV/turn. ! Commented: an acceleration rate of 20*0.5=10keV/turn. 'REBELOTE' 30000 0.3 99 ! 30000 0.3 99 ! Case of 100 keV/turn: ~30,000 turns from 3.6 MeV to 3 GeV. ! Commented: case of 10 keV/turn: ~300,000 turns from 3.6 MeV to 3 GeV. 'FAISCEAU' 'MARKER' SatIPolarLand_E ! Just for edition purposes. 'SPNPRT' 'END'

6661 Betatron damping

Figure 20.62 shows the damped vertical motion of the individual particles, over the acceleration range, together with the initial and final distributions of the 11 particles on elliptical invariants. Departure from the matching ellipse at the end of the acceleration cycle, 3 GeV (Eq. 20.14 with $\epsilon_Z/\pi = 1.0745 \times 10^{-6}$ m), is marginal.



Fig. 20.62 Left: damped vertical motion, from 3.6 MeV to 3.004 GeV in 30,000 turns. Right: the initial coordinates of the 11 particles (squares) are taken on a common invariant $\epsilon_Z(0) = 10^{-4} \pi m$ (at 3.6 MeV, $\beta \gamma = 0.0877$, thus $\beta \gamma \epsilon_Z(0) = 8.77 \times 10^{-6} \pi m$); the final coordinates after 30,000 turns (crosses) appear to still be (with negligible departure) on a common invariant of value $\epsilon_Z(\text{final}) = 2.149 \times 10^{-6} \pi m$ (at 3.004 GeV, $\beta \gamma = 4.08045$) thus $\beta \gamma \epsilon_Z(\text{final}) = 8.77 \times 10^{-6} \pi m$, equal to the initial value

66666 Degree of non-symplecticity of the numerical integration

The degree of non-symplecticity as a function of integration step size is illustrated in Fig. 20.63. The initial motion is taken paraxial, vertical motion is considered as it resorts to off-mid plane Taylor expansion of fields [1, DIPOLE Sec.], a stringent test as the latter is expected to deteriorate further the non-simplecticity inherent to the Lorentz equation integration method (a truncated Taylor series method [1, Eq. 1.2.4]).



Fig. 20.63 Turn-by-turn evolution of the normalized invariant, $\beta\gamma\epsilon_Z(\text{turn})/\beta\gamma\epsilon_Z(0)$ (initial $\epsilon_Z(0)$ taken paraxial), for four different integration step size values: 1, 2 and 4 cm

6673 Evolution of the wave numbers

The Fortran tool tunesFromFai_iterate can be used to computes tunes as a function 6674 of turn number or energy, it reads turn by turn particle data from zgoubi.fai and 6675 computes a discrete Fourier transform over so many turns (a few tens, 100 here 6676 for instance), every so many turns (300, here) [4]. Typical results are displayed in 6677 Fig. 20.64, tunes have the expected values: $v_Y = 0.7241$, $v_Z = 0.8885$. In acceleration 6678 rate of 100 keV/turn has been taken (namely, $\hat{V} = 200 \text{ kV}$ and still $\phi_s = 150^0$), to 6679 save on computing time. SCALING with option NTIM=-1 causes the magnet field 6680 to strictly follow the momentum boost by CAVITE. 6681



Fig. 20.64 Horizontal ring tune (left vertical axis), $v_Y \approx$ 0.7241, and vertical ring tune (right vertical axis), $v_Z \approx 0.8885$, as a function of turn number, over 30,000 turns ($E : 0.0036 \rightarrow 3 \text{ GeV}$ at a rate of 100 keV/turn)

(f) Spin tracking. Bunch polarization.

Spin depolarizing resonances in Saturne I synchrotron are located at (Figs. 20.65, 20.66)

 $G\gamma_R = k \pm v_Z = k \pm 0.888284 \equiv 4 - 0.888284, 4 + 0.888284, 8 - 0.888284$

where v_Z has been taken from Tab. 20.50, or from Fig. 20.64. $G\gamma_R$ is bounded by $G\gamma(3 \text{ GeV}) = 7.525238 < 8 + v_Z$

The simulation data file to track through these resonances is the same as in question (e), Tab. 20.54, except for the following substitutions:

- substitute MCOBJET (to be uncommented) to OBJET (to be commented),

- under CAVITE substitute a peak voltage V = 20 kV to V = 200 kV,

- under REBELOTE, request a 300,000 turn cycle rather than 30,000.

MCOBJET creates a 200 particle bunch with Gaussian transverse and longitudinal densities, with the following *rms* values at 3.6 MeV:

$$\epsilon_Y/\pi = 25 \ \mu m, \quad \epsilon_Z/\pi = 10 \ \mu m, \quad \frac{dp}{p} = 10^{-4}$$

⁶⁶⁹⁰ CAVITE accelerates that bunch from 3.6 MeV to 3 GeV at a rate of $q\hat{V}\sin(\phi_s) =$ ⁶⁶⁹¹ 10 keV/turn ($\hat{V} = 20 kV, \phi_s = 30^o$), in 300,000 turns.

Figure 20.65 shows sample S_Z spin components of a few particles taken among the 200 tracked. Figure 20.66 displays $\langle S_Z \rangle$, the vertical polarization component of the 200 particle set. A gnuplot script is used, given in Tab. 20.55.



Fig. 20.65 Vertical spin component of a few particles accelerated from 3.6 MeV to 3 GeV. This plot is obtained using zpop, which reads data from [b_]zgobui.fai: menu 7; 1/2 to open b_zgoubi.fai; 2/[20,23] to select S_Z versus turn; 7 to plot



Fig. 20.66 Average vertical spin component of a 200 particle bunch, accelerated from 3.6 MeV to 3 GeV

Table 20.55 A gnuplot script to plot the average vertical spin component of the 200 particle set, along the acceleration ramp (Fig. 20.66). The average is prior computed by an awk script, which reads the necessary data from zgoubi.fai.

data[yval] = 1; ; END{ average(curr, data); }

- (g) Crossing an isolated intrinsic depolarizing resonance.
- ⁶⁶⁹⁶ The simulation uses the file given in Tab. 20.54, with the following changes:
- Under OBJET:
- ⁶⁶⁹⁸ 1st line, change the reference rigidity to *BORO* = 4.08807740024 T m ⁶⁶⁹⁹ which corresponds to an initial $G\gamma$ = 2.65, 448.604769 MeV (in lieu of ⁶⁷⁰⁰ 0.274426548 T m, 3.6 MeV), which is conveniently upstream of $G\gamma_R$ = ⁶⁷⁰¹ 4 - $v_Z \approx 3.1$,
- 3rd line, request a single particle ("1 1 1", in lieu of "1 11 1" which distributes
 11 particles on the vertical invariant),
- $_{6704}$ 6th line, set the invariant ϵ_Z/π to the desired value,
- change the dipole field accordingly under DIPOLE, to 4.85560248505 T (in lieu of 0.3259493638 T), which maintains the expected curvature radius $\rho_0 = BORO/B = 8.4193 \text{ m}$ (Tab. 9.2),
- under REBELOTE, set the number of turns. A total of 15,000, about 7,500 turns upstream and as many downstream of the resonance, is convenient for the present peak voltage $\hat{V} = 20 \text{ kV}$ (acceleration rate 20 keV/turn).

6711 *Changing the particle invariant value*

Particle spin motion through the isolated resonance for five different invariant values, $\epsilon_Z/\pi = 2, 10, 20, 40, 200 \,\mu\text{m}$, is displayed in Fig.20.67.

Fig. 20.67 Turn by turn spin motion through the isolated resonance $G\gamma_R = 4 - \nu_Z$, observed at the beginning of the optical sequence (FAISTORE location, Tab. 20.54) for six different values of the particle invariant from $2 \,\mu$ m to $200 \,\mu$ m where full spin flip occurs. A graph obtained using zpop, which reads particle data from b_polarLand.fai (as specified under FAISTORE): menu 7; 1/8 to choose b_polarLand.fai; 2/[59,23] to select S_Z versus $G\gamma$; 7 to plot



⁶⁷¹⁴ The intrinsic resonance strength satisfies $\epsilon_R^2 = A \epsilon_Z$, with A a factor which characterizes the lattice. On the other hand, from the Froissart-Stora formula (Eq. 9.45) one gets

$$\epsilon_R^2 = \frac{2\alpha}{\pi} \ln\left(\frac{2}{1 + S_{Z,f}/S_{Z,i}}\right) \xrightarrow{S_{Z,i} \ll 1} \frac{\alpha}{\pi} (1 - \frac{S_{Z,f}}{S_{Z,i}})$$
(20.15)

with α , crossing speed, a constant. Thus one expects to find $\ln\left(\frac{2}{1+S_{Z,f}/S_{Z,i}}\right) / \epsilon_Z$ constant. This property appears to be satisfied by the tracking outcomes, Tab. 20.56. Note that numerical results slowly depart from that rule if $P_f/P_I \rightarrow 1$ or when approaching full flip, explaining this variation requires closer inspection of the theory and dedicated simulations.

⁶⁷²² Calculation of the resonance strength from the P_f/P_i tracking results, using ⁶⁷²³ Eq. 20.15, requires the crossing speed, namely,

$$\alpha = \frac{1}{2\pi} \frac{\Delta E}{M} = \frac{1}{2\pi} \frac{20 \times 10^3 \times \sin 30^o \,[\text{eV/turn}]}{938.27208 \times 10^6 \,[\text{eV}]} = 1.696 \times 10^{-6} \tag{20.16}$$

Table 20.56, rightmost column, displays the ratio $|\epsilon_R|/\sqrt{\epsilon_Z/\pi}$ so obtained. The resonance strength $|\epsilon_R|$ appears to be proportional to $\sqrt{\epsilon_Z/\pi}$, this is the expected result [6].

Table 20.56 Relationship between the invariant value ϵ_Z/π and the quantity $\ln\left(\frac{2}{1+S_{Z,f}/S_{Z,i}}\right) \propto |\epsilon_R|$. $\hat{V} = 20 \text{ kV}$, here. $S_{Z,f}/S_{Z,i}$ (col. 2) is from raytracing, ϵ_R^2 results, using Eq. 20.15 with crossing speed $\alpha = 1.696 \times 10^{-6}$ (Eq. 20.16). The resulting ratio $\epsilon_R^2 / \epsilon_Z/\pi$, rightmost column, appears to be about constant, this is the expected result (Eq. 20.15)

ϵ_Z/π	$\frac{S_{Z,f}}{S_{Z,i}} \equiv S_{Z,f}$	$\ln \frac{2}{1+S_{Z,f}}$	$\frac{\ln \frac{2}{1+S_{Z,f}}}{\epsilon_Z/\pi}$	$\frac{\epsilon_R^2}{\epsilon_R/\pi}$
$(\times 10^{-4} \mu m)$			c_Z/π	$(\times 10^{-8})$
1	+0.9	0.0513	0.0513	5.5
2	+0.822	0.0932	0.0466	5.0
10	+0.27	0.454	0.0454	4.9
20	-0.19	0.904	0.0452	4.9
40	-0.68	1.833	0.0458	4.9
50	-0.775	2.19	0.438	4.7
80	-0.963	3.297	0.0412	4.5

6726

6727 Changing the crossing speed

⁶⁷²⁸ A crossing case using $\hat{V} = 10 \,\text{kV}$ is displayed in Fig. 20.68. an acceleration ⁶⁷²⁹ rate/resonance crossing speed twice as slow compared to the previous ones.

⁶⁷³⁰ Comparison with the case $\hat{V} = 20 \text{ kV}$ is given in Tab. 20.57. From what ⁶⁷³¹ precedes, with the crossing speed $\alpha \propto \hat{V} \propto \Delta E/\text{turn}$, one expects to find ⁶⁷³² $\frac{\hat{V}}{\epsilon_Z/\pi} \times \ln\left(\frac{2}{1+S_{Z,f}/S_{Z,i}}\right)$ =constant. This property appears to be satisfied by track-⁶⁷³³ ing outcomes, Tab. 20.57.

Fig. 20.68 Turn by turn spin motion through the isolated resonance $G\gamma_R = 4 - \nu_Z$, observed at the beginning of the optical sequence (FAIS-TORE location, Tab. 20.54) for $\epsilon_Z/\pi = 1 \,\mu$ m and 20 μ m. $\hat{V} = 20 \,\text{kV}$. A graph obtained using zpop, which reads particle data from b_polarLand.fai (as specified under FAIS-TORE): menu 7; 1/8 to choose b_polarLand.fai; 2/[59,23] to select S_Z versus $G\gamma$; 7 to plot



Table 20.57 Relationship between the acceleration rate $\Delta E \propto \hat{V}$ and the quantity $\ln\left(\frac{2}{1+S_{Z,f}/S_{Z,i}}\right)$. Normalized to ϵ_Z/π , their product (rightmost column) appears about constant, this is the expected result. Explaining the observed $\approx 10\%$ variation requires closer inspection of the theory and dedicated simulations

ϵ_Z/π	Ŷ	$\frac{S_{Z,f}}{S_{Z,i}} \equiv S_{Z,f}$	$\ln \frac{2}{1+S_{Z,f}}$	$\frac{\hat{V}}{\epsilon_Z/\pi} \times \ln \frac{2}{1+S_{Z,f}}$
$(\times 10^{-4} \mu m)$	(kV)			
1	10	+0.79	0.111	1.1
20	10	-0.763	2.133	1.07
1	20	+.9	0.051	1.0
2	20	+0.82	0.094	0.94

(h) Spin motion across a weak depolarizing resonance.

The goal is to check numerical outcomes against the Fresnel integral model (Eq. 9.51). A weak resonance is obtained using small amplitude vertical motion and fast crossing. A single particle is raytraced, in the following conditions:

• resonance to be crossed: $G\gamma_R = 4 - v_y \approx 3.1115$,

- acceleration: peak voltage $\hat{V} = 100$ kV, synchronous phase $\phi_s = 30^\circ$,

- particle invariant $\epsilon_Z/\pi = 10^{-6} \,\mu\text{m}.$

The initial rigidity is taken a few hundred turns upstream of the resonance, namely, $B\rho_{ref} = 4.0880774 \text{ Tm}, 605226550 \text{ MeV}, G\gamma = 2.94931241$, a distance to $G\gamma_R$ of $4 - \nu_Z - 2.949312415 \approx 0.16223$. Tracking extends a few thousand turns beyond $G\gamma_R$ so that S_Z reaches its asymptotic value, from which the resonance strength $|\epsilon_R|$ can be calculated, using Eq. 20.15.

The simulation file is given in Tab. 20.58. Note the new setting of the SCALING 6747 factor SCL: DIPOLE field was set for a curvature radius $\rho_0 = 8.4193$ m, given a 6748 reference rigidity $B\rho_{ref} \equiv BORO = 0.274426548 \text{ Tm}$ (Tab. 20.46). However the 6749 reference rigidity is now changed to $B\rho_{ref} = 4.0880774 \,\mathrm{T}\,\mathrm{m}$, thus maintaining ρ_0 6750 requires scaling the field in DIPOLE by 4.0880774/0.274426548 = 14.8968 at turn 6751 1: this is the new factor, SCL = 14.8968, under SCALING (Tab. 20.58). Option 6752 NT=-1 under SCALING ensures that the scaling factor will automatically follow, 6753 turn-by-turn, the rigidity boost by CAVITE so preserving constant curvature radius 6754 $\rho_0 = 8.4193 \,\mathrm{m}.$ 6755

The resulting turn-by-turn spin motion is displayed in Fig. 20.69. The Fresnel integral model (Eq. 9.51) has been superimposed. Parameters in the latter are as follows:

- crossing speed
$$\alpha = \frac{1}{2\pi} \frac{\Delta E}{M} = \frac{1}{2\pi} \frac{10^3 \times \sin 30^6 \text{ [eV/turn]}}{938.27208 \times 10^6 \text{[eV]}} = 8.4812 \times 10^{-6},$$

- asymptotic $S_{Z,f} = 0.999780$, whereas initial $S_{Z,i} = 1$, thus (Eq. 20.15)

$$\epsilon_R^2 = 5.939 \times 10^{-10}$$

- orbital angle origin set at the location of $G\gamma_R$, which is turn 1699.





Table 20.58 Simulation input data file: track a particle launched on a vertical invariant $\epsilon_y/\pi = 10^{-6} \mu m$, with horizontal motion indifferent, taken zero here. The INCLUDE adds the Saturne I cell four times, the latter is defined in Tab. 20.48 and Fig. 9.23

```
      Saturne I ring. Crossing Ggamma=4-nu_Z, weak resonance case (small vertical invariant)

      'MARKER'
      SatIWeakXing_S

      '0BJET'
      ! Just for edition purposes.

      4.08507740924e3
      ! Reference Erho: 605226550 MeV proton.

      8
      ! Create a (set of) particle(s) on a given invariant.

      1
      1

      0.
      0.
      0.

      1.
      .

      1.
      .

      0.
      0.
      0.

      1.
      .
      ! create a single particle.

      0.
      1.
      .
      ! Horizontal invariant is null.

      0.
      1.4.426
      .
      ! Horizontal invariant value.

      0.
      1.
      .
      ! Nomentum spread.

      'PARTICUL'
      PROTON
      ! Necessary data in order to allow (i) spin trackingand. and (ii) acceleration

                                                    ! Necessary data in order to allow (i) spin trackingand, and (ii) acceleration.
! Switch on spin tracking,
! nitial spin vertical.
  PROTON
'SPNTRK'
  3
   'FAISCEAU'
 'FAISCEAU
'FAISTORE'
xing4-Qy.fai
                                                                                                                                                 ! Log particle data in xing.fai, turn-by-turn.
'SCALING'
1 1
DIPOLE
                                                                             ! Causes field increase in DIPOLE to follow rigidity increase by CAVITE.
! Relative rigidities at turn 1.
 -1
14.8968
  1
 ! 4 cells follow.
'INCLUDE'
 4* ./SatI_cell.inc[S_SatI_cell:E_SatI_cell]
  'CAVITE'
 0 0
100e3 0.523598775598
                                                                                                                                                         ! Acceleration rate is 200*0.5=100keV/turn.
'REBELOTE'
3999 0.3 99
                                                                                                                                                                                        ! A total of 3999+1=4000 turns.
  'FAISCEAU'
                           SatIWeakXing E
                                                                                                                                                                                               ! Just for edition purposes.
  'MARKER'
'SPNPRT'
  'END'
```

(i) Static spin motion near a resonance

The simulation input data file of Tab. 20.58 can be used for these fixed energy trials, however OBJET[KOBJ=8] has to be changed, as follows:

- OBJET[KOBJ=1] is used instead, as it allows to define a set of particles with sampled momentum offset,

- the reference rigidity in OBJET is set closest to the resonance, $G\gamma_R = 4 - v_y \approx 3.1115$ (considering that $v_Z \approx 0.8885$ (Fig. 20.64), thus $BORO \equiv B\rho_{ref} = 4.439362178$. Note that $G\gamma_R = 4 - v_Z$ is only known at the accuracy that v_Z is, from prior first order mapping computation, or Fourier analysis; finding the resonant $G\gamma_R$ from spin motion will bring its value

- the SCALING factor is set to SCL = 1 and concurrently the DIPOLE field is set to $B_0 = 5.27283999437$,

- since half an oscillation of $S_Z(turn)$ is enough to determine $\langle S_Z \rangle$, a number of turns *IPASS* \approx a few thousand, under REBELOTE, is enough considering the vertical motion amplitude considered. considering the initial Z=3 cm in this simulations (thus $\epsilon_Z/\pi = Z^2/\beta_Z = 79 \, 10^{-6}$ with $\beta_Z = 11.411$ (Tab. 20.49)), *IPASS* = 3000 is enough.

```
To conclude on this updating of the input data file of Tab. 20.58:
6778
             - substitute the following to OBJET:
6779
             'OBJET
6780
                                       ! BORO taken as close to resonant G.gamma as prior knowledge of nu_Z allows.
! Create a set of particles.
             4.4393621786553803e3
6781
6782
            1
1 1 1 1 1 41
0. 0. 0. 0. 0. 0. .00001
0. 0. 3. 0. 0. 1.
                                                                          ! 41 particles sampling a
! momentum offset, in -20*1e-5< D-1 < 20*1e-5<
! All particles have initial Z=3cm.</pre>
6783
6784
6785
6786

    substitute the following to SCALING:

6787
6788
6789
             'SCALING
             DIPOLE
6790
6791
                                                          ! Causes field increase in DIPOLE, in correlation to particle
6792
             1.
                                                                                            ! rigidity increase by CAVITE.
6793
6794
             - substitute the following for B_0 (3rd line) under DIPOLE:
6795
             22.5 5.27283999437 -0.6 0. 0.
                                                   ! Reference angle ACENT set to AT/2; Bo field at RM; radial index.
6796
6797
6798
```

The turn-by-turn values of the vertical component of the spins as they precess at fixed energy are displayed in Fig. 20.70. A quick, and accurate enough, approximation to the vertical component of the precession axis is $\langle S_Z \rangle|_{period} = \frac{1}{2} \{\min[S_Z(\theta)] + \max[S_Z(\theta)]\}$, it yields the $\langle S_Z \rangle (\Delta)$ graph of Fig. 20.71.

A match of the $\langle S_Z \rangle$ values by (Eq. 9.49)

$$S_y(\Delta) = \frac{\Delta}{\sqrt{\Delta^2 + |\epsilon_R|^2}}$$

given $G\gamma_R = 4 - nu_Z$, yield vertical tune and resonance strength values, respectively,

$$v_Z = 0.88845$$
 and $|\epsilon_R| = 2.77 \times 10^{-4}$

Satisfactorily, v_Z is consistent with earlier results, and $|\epsilon_R| = 2.77 \times 10^{-4}$ for $\epsilon_Z/\pi = 79 \times 10^{-6}$ here, is consistent in order of magnitude with $|\epsilon_R| = 2.44 \times 10^{-5}$ for

Fig. 20.70 Turn-by-turn value of the vertical component of spins precessing at fixed energy in Saturne I synchrotron, observed at the beginning of the sequence, where spins start vertical ($S_Z = 1$). The greater (respectively smaller) the distance to the resonance, the closer the precession axis to the vertical axis (resp., to the bend plane), and the greater (resp. the smaller) the oscillation frequency



Fig. 20.71 Vertical component of the spin precession axis as a function of $G\gamma$, in the vicinity of the resonance

 $\epsilon_{Z}/\pi = 10^{-6}$ in the previous question (h). The difference deserves further inspection, a possible additional question in this exercise.

6806 **9.2**

Construct the ZGS. Spin Resonances

A photo taken in the ZGS ring can be found in Fig. 9.4. A schematic layout is given in Fig. 9.24 and a sketch of the double dipole cell in Fig. 9.25. These figures introduce to the geometry and, in complement to Tab. 9.3, display some of the parameters of the synchrotron, and will referred to in building the ZGS ring in the following.

6813 (a) A model of ZGS synchrotron.

⁶⁸¹⁴ DIPOLE is used to simulate the cell dipoles. It is necessary to have Fig. 20.55 at ⁶⁸¹⁵ hand (in addition to the users' guide), when filling up the data list under DIPOLE. ⁶⁸¹⁶ Some comments regarding these data:

• DIPOLE is defined in a cylindrical coordinate system.

- The bending sector is 45 degrees, however the field region extent AT has to encompass the fringe fields, at both ends of the 45 deg sector. A large 5 deg extension is taken, for a total AT=55 deg which ensures absence of truncation of the fringe fields at the AT sector boundaries, over the all radial excursion of the beam.
- RM is given the curvature radius value, $RM = B\rho/B = 1.035270_{[T m]}/0.04986851_{[T]} = 20.76 m$, this makes magnet positioning and closed orbit checks easier (see below). Note that the field and reference rigidity are, as for the simulation input data, for 50 MeV, the injection energy, this is an arbitrary choice.
- ACENT=27.5 deg is the reference azimuth, for the positioning of the entrance and exit effective field boundaries (EFB). It is taken in the middle of the AT range, an arbitrary choice.
- The entrance radius in the AT sector is $RE = RM/\cos(AT \omega^+) = RM/\cos(5^\circ)$, with $\omega^+ = 22.5$ deg the positioning of the entrance EFB with respect to ACENT. And similarly for the positioning of the exit reference frame, $RS = RM/\cos(AT - (ACENT - \omega^-)) = RM/\cos(5^\circ)$ with $\omega^- = -22.5$ deg the positioning of the exit EFB. Note that $\omega^+ - \omega^- = 45^\circ$, the value of the bend angle.
- The entrance angle TE identifies with the extension to the 45 deg sector, namely, TE=5 deg. And similarly for the positioning of exit frame, 5 deg downstream of the exit EFB, thus TS=5 deg.

In order to build the cell, and in the first place the two cell dipole (the two dipoles are mirror symmetric, thus build one, the other follows straightforwardly), it is a good idea to proceed by steps:

(i) first build a 45 deg deviation sector in the hard edge model (Tab. 20.59). FAISCEAU located next to DIPOLE indicates that a trajectory entering DIPOLE at radius R=RM, normally to the EFB (thus, $Y_0 = 0$ and $T_0 = 0$ in OBJET) exits with Y=0 and T=0. Data validation at this stage can be performed by comparing DIPOLE's transport matrix computed with MATRIX, and the theoretical expectation (after Eq. ??, Sec. 19.5, numerical values truncated to 4th decimal)

	$(\cos \alpha)$	$\rho \sin \alpha$	0	0	0	$\rho(1 - \cos \alpha)$		(0.7071	14.6795	0	0	0	6.0804)
	$-\frac{1}{\alpha}\sin\alpha$	$\cos \alpha$	0	0	0	$\sin \alpha$	$\alpha = \pi/4$,	-	-0.03406	0.7071	0	0	0	0.7071
-	^P 0	0	1	ρα	0	0	$\rho = 20.76$		0	0	1	16.3048	0	0
1 =	0	0	0	1	0	0	\longrightarrow		0	0	0	1	0	0
	$\sin \alpha$	0	0	0	1	$\rho(\alpha - \sin \alpha)$			0.7071	0	0	0	1	1.6253
	0	0	0	0	0	(1)		l	0	0	0	0	0	1)

⁶⁸⁴⁷ MATRIX computation outcomes from raytracing can be found for comparison in Tab. 20.60.

Table 20.59 Simulation input data file: a 45 degree sector bend in the hard edge model. The reference trajectory has equal entrance and exit position, and opposite sign angles. It coincides with the arc R=RM. MATRIX computes the transport matrix of the dipole (bottom of this Table), for comparison with the fringe field model, and possible comparison with matrix codes outcomes

```
ZGS. Hard edge dipole model. Transport matrix.
'OBJET'

        1.63527036749193e3
        ! Reference Brho: 50 MeV proton.

        5
        ! Create an 11 particle set, proper for MATRIX computation.

        .001.01.001.01.0001
        ! Coordinates sampling.

        0.0.0.0.0.1.'o'
        ! Reference trajectory: all initial coordinates nul, relative rigidity D=1.

      ' IDIPOLE'
      ! Analytical moueling of a dipole import
20
      ! IL=2 here, to log trajectory coordinates in zgoubi.plt, at integration steps.

      25. 2076.
      ! Field region angle=45; reference radius set to curvature radius value.

      22.5 0.4986851481175 0. 0. 0. ! Reference angle ACENT set to AT/2; Bo field at RR; indices, all zero.
0 0.
      ! EFB 1 with fringe field extent.

  DIPOLE'
                                                                                                                  ! Analytical modeling of a dipole magnet.
4 .1455 2.2670 -.6395 1.1558 0. 0. 0.
22.5 0. 1.E6 -1.E6 1.E6 1.E6
                                                                                                                                                                     ae coefficients
                                                                                                       ! Angle to ACENT; face angle; face is straight
                                                                                                                                                                                    I FFR 2
     0.
.1455 2.2670 -.6395 1.1558 0. 0. 0.
2.5 0. 1.E6 -1.E6 1.E6 1.E6
4 .1455 2.2005 ...

-22.5 0. 1.E6 -1.E6 1.E6 1.E6

0.0.

0 0. 0. 0. 0. 0. 0. 0. 0. 0.

0 0. 0. 1.E6 -1.E6 1.E6 1.E6 0.
                                                                                                                                                                    ! EFB 3. Unused.
                                                                                    2 2076. 0. 2076. 0.
   reference frames.
                                                                                                                                            ! Local particle coordinates
 'FAISCEAU'
                                                                                        ! Compute transport matrix, from trajectory coordinates
 'MATRIX'
1 0
 'END'
```

6848

(ii) next, add fringe fields, including the 5 deg extensions that add to AT 6849 (Tab. 20.61). Note that negative drifts with length $RM \tan(5^{\circ}) = 181.62646548 \text{ cm}$ 6850 have been added at both ends, this recovers the actual length of the trajec-6851 tory across the 45 deg sector, for comparison with the hard-edge case, namely, 6852 s = 16.30486 \pm 10⁻⁵ m under FAISCEAU in both cases. A FIT procedure finds 6853 the field value necessary for recovering the exact deviation, as the latter is changed 6854 when fringe fields are introduced. Again, FAISCEAU allows checking the correct-6855 ness of DIPOLE data: exit coordinates come out to be Y=0 and T=0; however the 6856 path across the dipole is changed under the effect of the fringe fields, thus its length 6857 (s=1630.459 cm) is slightly different, compared to the hard edge case (an arc of 6858 radius radius RM=2076 cm and length 1630.487 cm) 6859

(iii) next, add the EFB angles: the sector is closing (wedge angles $\epsilon_1 > 0$ and $\epsilon_2 > 0$ by convention) thus the EFB tilt angle θ under DIPOLE if positive at entrance, negative at exit (Fig. 20.55). In order to reach proper wave number values (this is addressed below), the wedge angles are taken to be $\epsilon_1 = 13^{\circ}$ and $\epsilon_2 = 8^{\circ}$. These considerations result in the following:

Table 20.60 Outcomes of the simulation file of Tab. 20.59

An excerpt from zgoubi.res. Coordinates of the first particle (the reference trajectory) and its path length under FAISCEAU, at OBJET on the left hand side below, locally on the right hand side:

	3	Keyword,	label(s)	:	FAISCEAU									IPASS= 1
						TRACE	DU FAISO	CEAU						
						(follows	element	# 3	2)					
						11	TRAJECTO	DIRES						
					OBJET						FAISCE	AU		
		D	Y(cm)	Т	(mr) Z(cm	n) P(m	r)	S(cm)	D-1	Y(cm)	T(mr)	Z(cm)	P(mr)	S(cm)
0 1	1	.0000	0.000	0	.000 0.00	0.0	00 0	0000.0	0.0000	-0.000	-0.000	0.000	0.000	1.630487E+03

Transport matrix of a 45 degree sector, hard edge model, two difference cases of integration step size, namely, 4 cm and 2 m (an excerpt of MATRIX computation, from zgoubi.res). It can be checked against matrix transport expectations. The "first order symplectic conditions" are very small in the 4 cm step size case, which is an indication of accurate numerical integration of the trajectories across DIPOLE; the reference trajectory (first one) exits better aligned (reference coordinates, before change of frame for MATRIX computation, are closer to zero):

	~	C 4			•
-	Case	of 4	cm	step	size:

4 Keyword, labe	l(s) : MATRIX					1	PASS= 1
Reference, before ch	ange of frame (pa	article # 1 - D-	-1,Y,T,Z,s,time) :			
0.0000000E+00 -3.	25144356E-10 -4	.13789229E-10 0.	.00000000E+00	0.0000000E+00	1.63048659E+03	5.43871783E-02	
TRAN	SFER MATRIX ORI	DRE 1 (MKSA unit	ts)				
0.707107	14.6795	0.00000	0.00000	0.00000	6.08046		
-3.406102E-02	0.707107	0.00000	0.00000	0.00000	0.707107		
0.00000	0.00000	1.00000	16.3049	0.00000	0.00000		
0.00000	0.00000	7.285552E-16	1.00000	0.00000	0.00000		
0.707107	6.08046	0.00000	0.00000	1.00000	1.62533		
0.00000	0.00000	0.00000	0.00000	0.00000	1.00000		
DetY-1 =	0.000000025,	DetZ-1 =	0.000000002				
T12=0 at -	20.76 m,	T34=0 at -16	6.30 m				
First order symp	lectic conditions	s (expected values	s = 0) :				
2.5100E-09	2.3381E-10 0	0.00 0.00	90.00	0 0.000			
4 Keyword, la	bel(s) : MATRIX						IPASS= 1
Reference, before ch 0.00000000E+00 -2.	ange of frame (pa 01277929E-03 -2	article # 1 - D- .51514609E-03 0.	-1,Y,T,Z,s,time .00000000E+00) : 0.00000000E+00	1.63048722E+03	5.43871994E-02	
TRAN	SFER MATRIX ORI	DRE 1 (MKSA unit	ts)				
0.707105	14.6795	0.00000	0.00000	0.00000	6.08056		
-3.406102E-02	0.707108	0.00000	0.00000	0.00000	0.707120		
0.00000	0.00000	1.00000	16.3051	0.00000	0.00000		
0.00000	0.00000	1.457135E-17	1.00003	0.00000	0.00000		
0.707109	6.08048	0.00000	0.00000	1.00000	1.62531		
0.00000	0.00000	0.00000	0.00000	0.00000	1.00000		
DetY-1 =	-0.0000010903,	DetZ-1 =	0.0000286273				
R12=0 at -	20.76 m,	R34=0 at -16	6.30 m				
First order symp	lectic conditions	s (expected values	s = 0) :				
-1.0903E-06	2.8627E-05	0.000 0.00	0.00	0 0.000			

- the entrance (respectively exit) EFB of the upstream dipole of the cell (Fig. 9.25) is tilted with respect to the reference orbit by an angle $\theta = +13^{\circ}$ (resp. $\theta = -8^{\circ}$),

- the entrance (resp. exit) EFB of the downstream dipole is tilted with respect to the reference orbit by an angle $\theta = +8 \text{ deg}$ (resp. $\theta = -13^{\circ}$).

This final step requires again re-adjusting the radial positioning of the dipole (RE and RS, entrance and exit radius respectively), and field. In that aim the FIT procedure in Tab. 20.61 is added a variable: the RE and RS radii, coupled, and a constraint: the reference orbit has zero radial excursion at exit of the dipole. This FIT results in

Table 20.61 Simulation input data file: ZGS 45deg sector bend, with entrance and exit EFBs wedge angles and fringe fields. The reference trajectory has equal entrance and exit position, and opposite sign angles. It runs closely to the arc R=RM, not strictly coinciding with the latter due to the fringe fields. MATRIX computes the transport matrix of the dipole, for comparison with the hard edge model. Negative drifts with length $RM \tan(5^{\circ}) = 181.62646548$ cm are added to recover the hard edge path length

ZGS. Simplfied model. Find centered orbit in DIPOLE. 1.03527036749193e3 ! Reference Brho: 50 MeV proton. ! Create an 11 particle set, proper for MATRIX computation. 5 .001 .01 .001 .01 .001 .0001 ! Coordinate sampling. 0. 0. 0. 0. 0. 1. 'o' ! Reference trajectory: all initial coordinates nul, relative rigidity D=1. 'DRIFT' -181.62646548 .62646548 OLE' ! Analytical modeling of a dipole magnet. ! IL=2 here, to log trajectory coordinates in zgoubi.plt, at integration steps. 2076. ! Field region angle=45; reference radius set to curvature radius value. 0.49860858 0.0.0. ! Reference angle ACENT set to AT/2; Bo field at RR; indices, all zero. 0. ! EFB 1 with fringe field extent. 1555 2 2670 - 6305 1 1558 0.0.0 'DIPOLE' 0 55. 2076. 60. 0. ! EFB 2 with fringe field extent. 4 .1455 2.2670 -.6395 1.1558 0.0. 0. -22.5 -8. 1.E6 -1.E6 1.E6 1.E6 ! EFB angle to ACENT; -8 deg EFB tilt angle; EFB is straight. 2 3 5 0 .1 ! Vary DIPOLE field. 3 64 3.66 .1 2 1e-15 999 ! Request nul trajcory position at exit of DIPOLE. ! Request nul trajcory angle at exit of DIPOLE. 3 1 2 #End 0. 1. 0 3 1 3 #End 0. 1. 0 'FAISCEAU' ! Local particle coordinates. 'MATRIX' ! Compute transport matrix, from trajectory coordinates. 1 0 'END'

An excerpt from zgoubi.res. Coordinates of the first particle (the reference trajectory) and its path length, under FAISCEAU, at OBJET on the left hand side, locally on the right hand side:

	5 Keywor	rd, label(s) : FAISC	EAU								IPASS= 1	
				(f	TRACE DU F ollows elem 11 TRAJ	AISCEAU ent # ECTOIRES	4)						
				OBJET					FAISO	EAU			
0 1	D 1.0000	Y(cm) 0.000	T(mr) 0.000	Z(cm) 0.000	P(mr) 0.000	S(cm) 0.0000	D-1 0.0000	Y(cm) -0.000	T(mr) -0.000	Z(cm) 0.000	P(mr) 0.000	S(cm) 1.630459E+03	1

Transport matrix of ZGS 45 degree sector with EFB wedge angles and fringe fields (an excerpt of MATRIX computation, from zgoubi.res). It is not be checked against matrix transport expectations. The "first order symplectic conditions" are small, which is an indication of accurate numerical integration of the trajectories across DIPOLE:

/ Reyword, faber(3) . Infikin

IPASS= 1

Reference, before change of frame (particle # 1 - D-1,Y,T,Z,s,time) : 0.00000000E+00 -2.19331903E-08 -2.24434360E-08 0.00000000E+00 0.00000000E+00 1.63080750E+03 6.65146963E-02

	TRANSFER	MATRIX O	RDRE 1	(MKSA :	units)					
0.8703	65	14.6806	0.	00000	0.0	00000	0.0000	00	6.08068	
-2.03022	24E-02 (0.806503	0.	00000	0.0	00000	0.0000	00	0.748209	
0.000	00	0.00000	0.8	27040	16	.3143	0.0000	00	0.00000	
0.000	90	0.00000	-1.5	80329E-	02 0.89	97394	0.0000	00	0.00000	
0.7746	66	6.08004	0.	00000	0.0	00000	1.0000	00	1.63006	
0.000	00	0.00000	0.	00000	0.0	00000	0.000	00	1.00000	
DetY-1	= -0	.0000003451	, Det	Z-1 =	0.000	00000379				
T12=0 a	at -18.20	9 m,	T34:	=0 at	-18.18	m				
irst order	r symplect:	ic conditio	ns (expe	cted va	lues = 0)	:				
-3 4507F	-07 3.7	861E-08	0.000		0.000.0	0.000		0,000		

-3.4507E-07 3.7861E-08 0.000 0.000 0.000

re-adjusted magnetic field and RE, RS positioning, with the respective values

 $B_0 = 0.49860858 \, kG$ and $RE = RS = 2084.5090 \, cm$

⁶⁸⁶⁰ This is the values used in the ZGS cell simulation in Tab. 20.62,

(iv) and, finally, assemble this dipole and its mirror symmetric, in a cell (Fig. 9.25
and Tab. 20.62). The mirror symmetric is obtained by just permuting the entrance
and exit wedge angles. The cell includes a half long-drift at each end, and a short
drift between the dipoles. The three have been taken equal for simplification, 3.37 m
long.

Table 20.62 Simulation input data file: ZGS cell simplified model, obtained by assembling DIPOLE taken from Tab. 20.61 and its mirror symmetric (which means, permuting entrance and exit EFB tilt angles θ), and adding drift spaces. This input data file defines the ZGS cell sequence segment S_ZGS_cell to E_ZGS_cell, for possible use in INCLUDE statements in subsequent exercises. It also defines, for the same purpose, the dipoles sequence segment S_ZGS-DIP_UP to E_ZGS-DIP_UP to E_ZGS-DIP_UP (first dipole of the cell) and S_ZGS-DIP_DW to E_ZGS-DIP_DW (second dipole of the cell). In these possible INCLUDE statements, this file is used under the name ZGS_cell.inc

File ZGS_cell.INC.dat. ! ZGS. Simplfied model, 8-periodic. 'MARKER' ZGSCellMATRIX_S ! Just for edition purposes. 'OP 157

 1.03527036749193e3
 ! Reference Brho: 50 MeV proton.

 5
 ! Create an 11 particle set, proper for MATRIX computation.

 .001 .01 .001 .001 .0001
 ! Coordinate sampling.

 0. 0. 0. 0. 1.
 ! Reference trajectory: all initial coordinates nul, relative rigidity D=1.

 1

 'MARKER' S_ZGS_cell 'DRIFT' half_longDrift ! ZGS cell begins here. 337. 'MARKER' S_ZGS-DIP_UP 1 1st dipole of cell begins here. 'DRIFT' -181 62646548 ! Analytical modeling of a dipole magnet.
 ! IL=2 here, to log trajectory coordinates in zgoubi.plt, at integration steps.
 ! Field region angle=45; reference radius set to curvature radius value.
 ! Reference angle ACENT set to AT/2; BO field at RK; indices, all zero.
 ! EFB 1 with fringe field extent.
 ! EFG 2 officients 'DIPOLE' DIP_UP 55 2076
 55. 2076.
 ! Field region :

 27.5 0.49860858 0. 0. 0.
 ! Reference and

 60.
 0.

 4.1455
 2.2670 -.6395

 22.5
 13.

 1.E6
 -1.E6

 1.E6
 ! Enge coefficients ! EFB angle to ACENT; EFB tilt angle; EFB is straight 60. 0. 4 .1455 2.2670 -.6395 1.1558 0. 0. 0. -22.5 -8. 1.E6 -1.E6 1.E6 1.E6 ! EFB 2 with fringe field extent. ! EFB angle to ACENT; EFB tilt angle; EFB is straight. 0. 0. EFB 3. Unu Pogree of interpolation polynomial; flying grid sizing is step, proper for accuracy. Integration step size. 087266462599717 2084.5090 0.087266462599717 Positioning of entrance and exit. 212.0 2 2084.5090 -0.087266462599717 2084.5090 0.087266462599717 'DRIFT' -181.62646548 'MARKER' E_ZGS-DIP_UP ! 1st dipole of cell ends here 'DRIFT' shortDrift 337. 'MARKER' S_ZGS-DIP_DW ! 2nd dipole of cell begins here. 'DRIFT -181.62646548
 'DIPOLE'
 IPIP_DW
 ! Analytical modeling of a dipole magnet.

 2
 ! IL=2 here, to log trajectory coordinates in zgoubi.plt, at integration steps.

 25. 2076.
 ! Field region angle=45; reference radius set to curvature radius value.

 27.5 0.49860858 0.0.0.
 ! Reference angle ACENT set to AT/2; Bo field at RM; indices, all zero.

 60.0.
 ! Reference angle ACENT set to AT/2; Bo field at RM; indices, all zero.

 61.155
 2.2670 - .6395

 7.5
 0.49860858

 7.5
 0.49860858

 8.1.26
 -1.66

 9.0
 ! EFB angle to ACENT; EFB tilt angle; EFB is straight.

 60.0.
 ! EFB 2 with fringe field extent.

 4
 1455
 2.2670 - .6395

 1.1558
 0.0.0.
 'DIPOLE' DIP DW ! Analytical modeling of a dipole magnet. 60. 0. 4 .1455 2.2670 -.6395 1.1558 0. 0. 0. -22.5 -13. 1.E6 -1.E6 1.E6 1.E6 ! EFB angle to ACENT; EFB tilt angle; EFB is straight. ! EFB 3. Unused 0. 0. 0 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 1.E6 -1.E6 1.E6 1.E6 0. .E6 1.E6 1.E6 0. ! Degree of interpolation polynomial; flying grid sizing is step, proper for accuracy. ! Integration step size. 087266462599717 2084.5090 0.087266462599717 ! Positioning of entrance and exit. 2 1 2.0 2 2084.5090 -0.087266462599717 2084.5090 0.087266462599717 'DRIFT -181.62646548 'MARKER' E_ZGS-DIP_DW ! 2nd dipole of cell ends here. 'DRIFT' half_longDrift 337. 'MARKER' E_ZGS_cell ! ZGS cell ends here. 'FAISCEAU' ! Local particle coordinates. 'TWISS' 1. 1. 'MARKER' ZGSCellMATRIX_E ! Just for edition purposes. 'END ZGS. Simplfied model, 8-periodic.

An excerpt from zgoubi.res. Coordinates of the first particle (the reference trajectory) and its path length, under FAISCEAU, at OBJET on the left hand side, locally on the right hand side:

	1	8 Keyword	l, label(s)) : FAIS	CEAU								IPASS= 4
						TRACE DU F	FAISCEAU						
					(follows elem	nent #	4)					
						11 TRAJ	ECTOIRES						
					OBJET					FAISO	EAU		
		D	Y(cm)	T(mr)	Z(cm)	P(mr)	S(cm)	D-1	Y(cm)	T(mr)	Z(cm)	P(mr)	S(cm)
0	1	1.0000	0.000	0.000	0.000	0.000	0.0000	0.0000	-0.000	-0.000	0.000	0.000	4.272614E+03

The TWISS command down the sequence (Tab. 20.62) produces the periodic beam matrix results shown in Tab. 20.63

 Table 20.63
 Results obtained running the simulation input data file of Tab. 20.62, ZGS cell - an excerpt from zgoubi.res

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The TWISS command also produces a zgoubi.TWISS.out file which details the optical functions along the sequence (at the downstream end of the optical elements. The header of that file details the optical parameters of the structure (Tab. 20.64).

⁶⁸⁶⁶ Lattice parameters

Table 20.64 An excerpt of zgoubi.TWISS.out file resulting from the execution of the ZGS cell simulation input data file of Tab. 20.62. Note that the ring (4-period) wave numbers are 4 times the cell values Q1, Q2 displayed here. Optical functions (betatron function and derivative, orbit, phase advance, etc.) along the optical sequence are listed as part of zgoubi.TWISS.out following the header. The top part and last line of that listing are given below

Q	LENGTH	%le	42.72614305			
Q	ALFA	%le	1.412693458			
Q	ORBIT5	%le	-0			
0	GAMMATR	%le	0.8413487096			
Q	Q1	%le	0.2123591260	[fractional]		
0	Q2	%le	0.1928670550	[fractional]		
Q	DQ1	%le	0.4709865847E-01			
0	DQ2	%le	0.4457456345E-01			
Q	DXMAX	%le	3.81566835E+01	@ DXMIN	%le	3.68534544E+01
0	DYMAX	%le	0.0000000E+00	@ DYMIN	%le	0.0000000E+00
Q	XCOMAX	%le	3.68530296E-01	@ XCOMIN	%le -	1.59240732E-07
0	YCOMAX	%le	0.0000000E+00	@ YCOMIN	%le	0.0000000E+00
Q	BETXMAX	%le	3.25272034E+01	@ BETXMIN	%le	2.86307346E+01
0	BETYMAX	%le	3.73198843E+01	@ BETYMIN	%le	3.50936471E+01
Q	XCORMS	%le	8.67153286E-04			
0	YCORMS	%le	0.			
Q	DXRMS	%le	6.22665688E-01			
0	DYRMS	%le	0.0000000E+00			

Top and bottom four lines (truncated) of zgoubi.TWISS.out optical functions listing, including the periodic β_x , β_y (β_Y , β_Z in zgoubi notations) and D_x (η_Y in zgoubi notations) values at cell ends:

# alfx	btx	alfy	bty	alfl	btl	Dx	Dxp	
-2.2668087e-6	2.8636996e+1	-1.0203802e-6	3.7013045e+1	0.0000000e+0	0.0000000e+0	3.6856253e+1	1.2733594e-4	etc.
-2.2589191e-6	2.8636995e+1	-9.9937511e-7	3.7013042e+1	0.0000000e+0	0.0000000e+0	3.6853463e+1	2.6012859e-6	etc.
-2.2589191e-6	2.8636995e+1	-9.9937511e-7	3.7013042e+1	0.0000000e+0	0.0000000e+0	3.6853463e+1	2.6012859e-6	etc.
-1.1768220e-1	2.9033592e+1	-9.1049986e-2	3.7319884e+1	0.0000000e+0	0.0000000e+0	3.6853472e+1	2.6012859e-6	etc.
1.1775697e-1	2.9027748e+1	9.1140989e-2	3.7313084e+1	0.0000000e+0	0.0000000e+0	3.6853454e+1	2.6012859e-6	etc.
1.1775697e-1	2.9027748e+1	9.1140989e-2	3.7313084e+1	0.0000000e+0	0.0000000e+0	3.6853454e+1	2.6012859e-6	etc.
5.1297527e-5	2.8630735e+1	7.3912348e-5	3.7005690e+1	0.0000000e+0	0.0000000e+0	3.6853463e+1	2.6012859e-6	etc.
5.1297527e-5	2.8630735e+1	7.3912348e-5	3.7005690e+1	0.0000000e+0	0.0000000e+0	3.6853463e+1	2.6012859e-6	etc.

(b) Betatron functions of the ZGS cell.

Among the various ways to produce the betatron functions along the sequence (and throughout the DIPOLEs), here are two possibilities, based on the storage of particle coordinates in zgoubi.plt during stepwise raytracing:

 a direct way consists in using OBJET[KOBJ=5] and transport the 11-particle set so obtained across the sequence. Then, betaFromPlt from zgoubi toolbox [2] can be used to compute the transport matrix, step by step across the sequence, from the coordinate values logged in zgoubi.plt during the stepwise integration;

2. an indirect way consists in launching a few particles on a common invariant (horizontal and/or vertical) and subsequently plot the s-dependent quantities $\hat{Y}^2(s)/\epsilon_Y$ and/or $\hat{Z}^2(s)/\epsilon_Z$. The maximum value of the latter, a function of the distance s, is the betatron function along the sequence, $\beta_{Y,Z}(s)$.

The second method is used here (this is an arbitrary choice. Exercises may be found in the various Chapters, that use the first method and may be referred to, if desired).

The input data file to derive the betatron function following method (2) above is given in Tab. 20.65. The initial ellipse parameters (under OBJET) are the periodic values, namely, $\alpha_Y = \alpha_Z = 0$, $\beta_Y = 28.63$ m, $\beta_Z = 37.01$ m, they are a sub-product of the TWISS procedure performed in (a), to be found in zgoubi.TWISS.out (Tab. 20.64). The resulting envelopes and their squared value are shown in Fig. 20.72. Note that this raytracing also provides the coordinates of the 60 particles on their common upright invariant

$$x^2/\beta_x + \beta_x x'^2 = \epsilon_x/\pi$$

at start and at the end of the cell (with x standing for either Y or Z, and $\epsilon_{Y,Z}/\pi$ =

 10^{-4} , here). This allows checking that the initial ellipse parameters (under OBJET,

Tab. 20.65) are effectively periodic values, and that the raytracing went correctly,

namely by observing that the initial and final ellipses do superimpose (Fig. 20.73).

 Table 20.65
 Simulation input data file: raytrace 60 particles across ZGS cell to generate beam envelopes. Store particle data in zgoubi.plt, along DRIFTs and DIPOLEs. The INCLUDE file and segments are defined in Tab. 20.62



Fig. 20.72 Left: horizontal and vertical envelopes as generated by plotting the coordinates Y(s) (thick lines, red, along the drifts only) or Z(s) (thin lines, blue) across the ZGS cell, of 60 particles evenly distributed on a common $10^{-4} \pi \mu m$ invariant, either horizontal or vertical (while the other invariant is zero). Right: a plot of $Y^2(s)/\epsilon_Y$ and $Y^2(s)/\epsilon_Y$: the extrema identify with $\beta_Y(s)$ and $\beta_Z(s)$, respectively. The extrema extremorum values are $\hat{\beta}_Y = 32.5 \text{ m}$ and $\hat{\beta}_Z = 37.1 \text{ m}$, respectively. These plots are obtained using zpop, which reads stepwise particle data from zgoubi.plt



Fig. 20.73 Sixty particles evenly distributed on a common periodic invariant (of value either $\epsilon_Y = 10^{-4}\pi\mu$ m and $\epsilon_Z = 0$, left plot, or the reverse, right plot) have been tracked from start to end of the cell. These periodic invariants are defined assuming the periodic ellipse parameters determined from prior TWISS, given in Tab. 20.64; values resulting from an *rms* match of the coordinates are given in the figure, and do agree with those TWISS data. The figure shows the good superposition of the start and end invariants (the start and end *rms* match ellipse parameters show negligible difference), which confirms the correct value of the periodic ellipse parameters, namely, left graph: horizontal phase space at start (crosses) and end (dots) of the cell; right graph: vertical phase space at start (crosses) and end (dots) of the cell

6891 Dispersion function

Raytracing off-momentum particles on their chromatic closed orbit provides the periodic dispersion function. In order to do so, the input data file of Tab. 20.65 can be used, it just requires changing OBJET to the following:

 6895
 '0BJET'

 6896
 1.0352703749193e3
 ! Reference Brho: 50 MeV proton.

 6897
 2
 ! Create particles individually'

 6898
 3 1
 ! three particles.

 6899
 +36.85e-10.0.0.0.0.1.001 'p' ! Chromatic orbit coordinates Y0 and T0 for D=1.001 relative rigidity.

 6900
 0.0.0.0.0.1. 'o'
 ! On-momentum orbit.

 6901
 -36.85e-10.0.0.0.0.999 'm' ! Chromatic orbit coordinates Y0 and T0 for D=0.999 relative rigidity.

 6902
 1 1

The position and angle of the chromatic particles, which are offset by $\Delta p/p = \pm 10^{-3}$, are drawn from the value of the periodic dispersion $\eta_Y = 36.85$ m and its derivative $\eta'_Y \approx 0$ (Tab. 20.64), namely, $Y_0 = \eta_Y \Delta p/p = \pm 3.685$ cm and $T_0 = \eta_Y \Delta p/p = 0$.

Running Tab. 20.65 simulation file with this new OBJET produces the following coordinates at FAISCEAU, located at the end of the sequence (an excerpt from zgoubi.res):

910		1	8 Keyword,	label(s)	: FAISO	EAU								IPASS= 1
911							TRACE DU F	AISCEAU						
912						(f	ollows elem	ent#	L7)					
913							3 TRAJ	ECTOIRES						
914						OBJET					FAISO	EAU		
915			D	Y(cm)	T(mr)	Z(cm)	P(mr)	S(cm)	D-1	Y(cm)	T(mr)	Z(cm)	P(mr)	S(cm)
916	р	1	1.0010	3.685	0.000	0.000	0.000	0.0000	0.0010	3.685	0.000	0.000	0.000	4.278650E+03
917	0	1	1.0000	0.000	0.000	0.000	0.000	0.0000	0.0000	0.000	0.000	0.000	0.000	4.272614E+03
918	m	1	0.9990	-3.685	0.000	0.000	0.000	0.0000	-0.0010	-3.685	-0.000	0.000	0.000	4.266579E+03

The local coordinates Y, T (under FAISCEAU, right hand side) are equal to the 6919 initial coordinates Y_0 , T_0 (under OBJET, left hand side), to better than 5 μ m, 0.5 μ rad 6920 accuracy respectively (zgoubi.fai can be consulted for greater precision on these 6921 values), so confirming the periodicity of these chromatic trajectories. Figure 20.74 6922 shows the particle trajectories through the two DIPOLEs. A difference between the 6923 on- and off-momentum trajectories yields as expected a quasi-constant $\eta_Y \approx 36.8$ m 6924 whereas $\eta'_Y \approx 0$. η_Y departs from exactly zero due to the fringe fields and to the 6925 wedge focusing. 6926

Fig. 20.74 A plot of the radial

excursion, within DIPOLE range (namely, AT=55° extent, Tab. 20.62), of an onmomentum particle (its radial position in the dipole body is $R_0 \approx 20.7628$ m, corresponding to Y=0 in this graph) and two particles at respectively $dp/p = \pm 10^{-3}$. The diverging parts at DIPOLE ends are in the 5 deg fringe field regions. A graph obtained using zpop, which reads stepwise particle data from zgoubi.plt: menu 7; 1/1 to open zgoubi.plt; 2/[6,2] to select Y versus distance; 7 to plot



6927 Orbit difference

The method can be used to compute the dispersion function, just like in machine operation. This requires tracking a particle with +dp/p momentum offset, save its zgoubi.plt data (say, in zgoubi.plt+dpp), and repeat with -dp/p (zgoubi.plt-dpp). A gnuplot script can compute and plot the orbit difference, and normalize to dp/p; the result is the periodic dispersion, displayed in Fig. 20.75.





6932

⁶⁹³³ (c) Some verifications regarding the model.

The field along large excursion orbits can be logged in zgoubi.plt, using option IL=2 (or 20, or 200, etc. for printout every 10, or 100, etc. integration step) under DIPOLE.

The simulation file of Tab. 20.65 is used to raytrace five particles, with OBJET changed to the following:

939	OBJEI	
940	1.03527036749193e3	! Reference Brho: 50 MeV proton.
941	2	! Create particles individually,
942	5 1	! five particles.
943	+36.85e-1 0. 0. 0. 0. 1.01 'p'	! Chromatic orbit coordinates for D=1.01 relative rigidity.
944	0. 0. 0. 0. 0. 1. '0'	! On-momentum closed orbit.
945	-36.85e-1 0. 0. 0. 0. 0.99 'm'	! Chromatic orbit coordinates for D=0.99 relative rigidity.
946	0. 0. 5. 0. 0. 1. 'm'	! Initial vertial excursion is Z0= 5 cm off-mid-plane.
947	0. 0. 20. 0. 0. 1. 'm'	! Initial vertial excursion is Z0=20 cm off-mid-plane.
948	1 1 1 1 1	

Apart from the on-momentum particle (2nd in the list) this OBJET defines two particles on $\Delta p/p = \pm 1\%$ chromatic orbit (1st and 3rd in the list), this is an excursion of a few tens of centimeters, large as requested, as $\Delta x \approx 38 \times dp/p$. OBJET also defines 2 particles launched into the cell at respectively $Z_0 = 5$ cm and $Z_0 = 20$ cm.

The magnetic field as a function of the azimuthal angle in DIPOLE frame, along these trajectories across the upstream DIPOLE of the cell, is shown in Fig. 20.76. The field curves for the first four trajectories essentially superimpose except for the fringe field regions (Fig. 20.76), due to the wedge angles. This behaves as expected. Detail inspection is possible, from the detailed particle coordinate and field data in zgoubi.plt - this is out of the scope of the present question.

The field along the 5th particle trajectory features overshoots (Fig. 20.76), this is due to the very large vertical excursion ($Z \approx 20$ cm in the entrance fringe field region). It looks reasonable, however it may be an artifact in the case that the high order derivatives of the field in that region are large, resulting from the truncated Taylor series method used for off mid-plane field extrapolation [1, Sec. 1.3.3].



Fig. 20.76 Magnetic field along 5 different trajectories across the upstream DIPOLE, including four large horizontal and vertical excursion cases, and a zoom in on the entrance fringe field region

(d) Sinusoidal approximation of the betatron motion

The approximation

$$y(\theta) = A \cos(v_Z \theta + \phi)$$

is checked here considering the vertical motion (considering the horizontal motion
 leads to similar conclusions). The value of the various parameters in that expression
 are determined as follows:

- the particle raytraced for comparison is launched with an initial excursion $Z_0(\theta = 0) = 5 \text{ cm}$ (4th particle in OBJET, above). At the launch point (middle of the long drift) the beam ellipse is upright (Fig. 20.73), whereas phase space motion is clockwise, thus take

$$A = 5 \,\mathrm{cm}$$
 and $\phi = \pi/2$

- the vertical betatron tune of the 4-cell ring is (Tab. 20.64)

$$v_{\rm Z} = 4 \times 0.192869 = 0.77147$$

- $\theta = s/R$ and $R = \oint ds/2\pi$ with (Tab. 20.64)

$$2\pi R$$
 = circumference = 4 × 42.72614331 = 170.90457 m

The comparison with a trajectory obtained from raytracing is given in Fig. 20.77 and confirms the validity of the sinusoidal approximation.





(e) An acceleration cycle. Symplecticity checks. 6970

Eleven particles are launched for 65,000 turn tracking at a rate of

$$\Delta W = q\hat{V}\cos\phi_s = 400 \times \sin 150^0 = 200 \,\text{keV/turn}$$

 $(E: 0.05 \rightarrow 13.05 \text{ GeV})$, all evenly distributed on the same initial vertical invariant 6971

$$Z^2/\beta_Z + \beta_Z Z'^2 = \epsilon_Z/\pi \tag{20.17}$$

6972

with $\epsilon_Z/\pi = 10^{-4}$ m, or, normalized, $\beta \gamma \epsilon_Z/\pi = 0.33078 \times 10^{-4}$ m. The simulation file is given in Tab. 20.66. CAVITE[IOPT=3] is used, it provides an RF phase independent boost

$$\Delta W = q\hat{V}\sin\phi_s$$

as including synchrotron motion is not necessary here, even better, this ensures 6973 constant depolarizing resonance crossing speed, so precluding any possibility of 6974 multiple crossing (it can be referred to [3] regarding that effect). 6975

Table 20.66 Simulation input data file: track 11 particles launched on the same vertical invariant, with quasi-zero horizontal invariant. The INCLUDE adds the ZGS cell four times, the latter is defined in Tab. 20.62 and Fig. 9.25. An MCOBJET is commented, it is used in a subsequent spin tracking exercise

ZGS ring. Polarization landscape. 'MARKER' ZGSPolarLand_S 'OBJET' ! Just for edition purposes. 'UBJET' ! Reference Brho: 50 MeV proton. 1.03527036749193e3 ! Reference Brho: 50 MeV proton. 8 ! Create an 11 particle set, proper for MATRIX computation. 1 11 1 ! Define 9 particles, all with ~0 horiz. inveariant, evenly spread on same vertical invariant. 0. 0. 0. 0. 1. 'o' ! Reference trajectory: all initial coordinates nul, relative rigidity D=1. 0. 28.63 0. ! Horiz. invariant taken zero. Nominal would be 0.14mm_m norm. i.e. 4.6e=8 non-normalized. 0. 37.01 150e-6 ! epsilon_Z/pi = beta.gamma * epsilon_norm, latter =0.05e-6 m, beta.gamma=0.3308. 0. 1. 0. 0. ! All paricls are on-momentum. 1.03527036749193e3 !'MCOBJET' !1.03527036749193e3 ! Commented. ! Reference Brho: 50 MeV proton. ! Create an 11 particle set, proper for MATRIX computation.

 11.032/030/4919363

 13

 1200

 12.2 2 2 2 2

 10.0.0.0.0.1.

 10.28.63 25e-6 3

 10.37.01 10e-6 3

 10.1.e-8 3

 11.23456 234567 345678

 ! Periodic alpha_Y, beta_Y, and invariant value; ! Periodic alpha_Z, beta_Z, and invariant value. 'PARTICUL' PROTON 'SPNTRK' ! Necessary data in order to allow (i) spin trackingand, and (ii) acceleration. ! Switch on spin tracking, ! all initial spins vertical. 3 'FAISCEAU' 'FAISTORE' b_polarLand.fai 7 ! Log particle data in b_polarLand.fai, turn-by-turn; "b_" imposes ! binary write, which results in faster i/o. 'SCALING' 1 1 DIPOLE -1 ! Causes field increase in DIPOLE, in correlation to particle ! rigidity increase by CAVITE. 1. ! 4 cells follow. 'INCLUDE' 1 4* ./ZGS_cell.inc[S_ZGS_cell:E_ZGS_cell] 'CAVITE' 0 0 400e3 0.523598775598 ! Acceleration rate is 400*0.5=200keV/turn. 'REBELOTE' 87000 0.3 99 'FAISCEAU' 'MARKER' 'SPNPRT' ZGSPolarLand_E ! Just for edition purposes. 'END'

6976 Betatron damping

Figure 20.78 shows the damped vertical motion of the individual particles, over the acceleration range, together with the initial and final distributions of the 11 particles on elliptical invariants. Departure from the matching ellipse at the end of the acceleration cycle, 13 GeV (Eq. 20.17 with $\epsilon_Z/\pi = 2.2244 \times 10^{-7}$ m), is marginal.



Fig. 20.78 Left: damped vertical motion, from 50 MeV to 13.05 GeV, 65,001 turns. Right: the initial coordinates of the 11 particles (squares) are taken on a common invariant $\epsilon_Z(0) = 10^{-5} \pi m$ (at 50 MeV, $\beta \gamma = 0.33078$, thus $\beta \gamma \epsilon_Z(0) = 0.33078 \times 10^{-5} \pi m$); the final coordinates after 65,000 turns (crosses) appear to still be (with negligible departure) on a common invariant of value $\epsilon_Z(\text{final}) = 2.2244 \times 10^{-7} \pi m$ (at 13 GeV, $\beta \gamma = 14.869842$) thus $\beta \gamma \epsilon_Z(\text{final}) = 0.33076 \times 10^{-5} \pi m$, equal to the initial value

6981 Degree of non-symplecticity of the numerical integration

The degree of non-symplecticity as a function of integration step size is illustrated in Fig. 20.79. The initial motion is taken paraxial, vertical motion is considered as it resorts to off-mid plane Taylor expansion of fields [1, DIPOLE Sec.], a stringent test as the latter is expected to deteriorate further the non-simplecticity inherent to the Lorentz equation integration method (a truncated Taylor series method [1, Eq. 1.2.4]).



Fig. 20.79 Turn-by-turn evolution of the normalized invariant, $\beta\gamma\epsilon_Z(\text{turn})/\beta\gamma\epsilon_Z(0)$ (initial $\epsilon_Z(0)$ taken paraxial), for four different integration step size values: 1, 2 and 4 cm

6988 Evolution of the wave numbers.

The Fortran tool tunesFromFai_iterate can be used to computes tunes as a function 6989 of turn number or energy, it reads turn by turn particle data from zgoubi.fai and 6990 computes a discrete Fourier transform over so many turns (a few tens, for instance), 6991 every so many turns [4]. Typical results are displayed in Fig. 20.80, tunes have the 6992 expected values: $v_Y = 0.849$, $v_Z = 0.771$. An acceleration rate of 200 keV/turn has 6993 been taken (namely, $\hat{V} = 400 \,\text{kV}$ and still $\phi_s = 150^0$), to save on computing time. 6994 Note that turn-by-turn raytracing allows determining the tune value at all γ along the 6995 acceleration cycle (and thus for instance the γ values at which the resonance occurs, 6996 see (f)). In these simulations anyway the horizontal and vertical tunes are essentially 6997 constant over the all cycle: it is determined by the wedge angle, which won't charge 6998 as long as the reference orbit isn't changed. The latter holds here, as SCALING with 6999 option NTIM=-1 causes the magnet field to strictly follow the momentum boost by 7000 7001 CAVITE.

Fig. 20.80 Horizontal ring tune (left vertical axis), $v_Y \approx$ 0.8494, and vertical ring tune (right vertical axis), $v_Z \approx 0.77147$, as a function of turn number, over 65,000 turns ($E : 0.05 \rightarrow 13$ GeV at a rate of 200 keV/turn). The graph displays results for 3 different integration step sizes, namely, 1, 2 and 4 cm, essentially converged



⁷⁰⁰² (f) Spin tracking. Bunch polarization.

Spin depolarizing resonances in the ZGS are located at

 $G\gamma_R = kP \pm v_Z = 4 - v_Z, 4 + v_Z, 8 - v_Z, 8 + v_Z, 12 - v_Z,$ etc.

with P=4 the superperiodicity of the ring, and $v_Z = 0.77147$ taken from Tab. 20.64, or from Fig. 20.80. $G\gamma_R$ is bounded, in the present simulation, by $G\gamma(17.4 \, GeV) =$ $35.0 < 9P - v_Z$. Resonances are expected to be stronger at $G\gamma_R = 2 \times 4k \pm v_Z =$ $8 - v_Z$, $8 + v_Z$, $16 - v_Z$, etc., with the additional factor 2 the number of cells per superperiod [6, Sec. 3.II].

The simulation data file to track through these resonances is the same as in question (e), Tab. 20.66, except for the substitution of MCOBJET (to be uncommented) to OBJET (to be commented). MCOBJET creates a 200 particle bunch with Gaussian transverse and longitudinal densities, with the following *rms* values at 50 MeV:

$$\epsilon_Y/\pi = 25 \,\mu m, \quad \epsilon_Z/\pi = 10 \,\mu m, \quad \frac{dp}{p} = 10^{-4}$$

which are presumably close to ZGS polarized proton runs [7]. CAVITE accelerates that bunch from 50 MeV to 17.4 GeV about, at a rate of $q\hat{V}\sin(\phi_s) = 200$ keV/turn $(\hat{V} = 400 \, kV, \phi_s = 30^\circ)$, in 87,000 turns about.

Figure 20.81 shows sample S_Z spin components of a few particles taken among the 200 tracked. Figure 20.82 displays $\langle S_Z \rangle$, the vertical polarization component of the 200 particle set. A gnuplot script is used, given in Tab. 20.67.



Fig. 20.81 Individual vertical spin component of 20 particles accelerated in ZGS from 50 MeV to 17.4 GeV, at a rate of 200 keV/turn. This plot is obtained using zpop, which reads data from [b_]zgobui.fai: menu 7; 1/2 to open b_zgoubi.fai; 2/[20,23] to select S_Z versus turn; 7 to plot



Table 20.67 A gnuplot script to plot the average vertical spin component of the 200 particle set, along the acceleration ramp (Fig. 20.82). The average is prior computed by an awk script, which reads the necessary data from zgoubi.fai.

```
# gnuplot_avrgFromFai.gnu
set x2label "E [GeV]"; set xlabel "G(/Symbol g)"; set ylabel "<S_Z>|_{200 prtcls}"
set x2ics: set x2ics; set ylics; set format y '%0.2f'; set grid
M=938.27208; Ei = 50.; G = 1.79284735; Qy = 0.7115; dE = 0.2 # MeV/turn
fName = 'zgoubi.fai'; plotCond(col_num)=sprinff('< gawk -f average.awk -v col_num=%d %s', col_num,fName)
do for [intgr=1:9] { set arrow nohead from 4*intgr+Qy, 0.7 to 4*intgr+Qy, 1.01 lw .6 dt 3
set arrow nohead from 4*intgr-Qy, 0.7 to 4*intgr-Qy, 1.01 lw .6 dt 3 }
do for [intgr=8:32:8] { set label " ".intgr."-Qy" at intgr-Qy, 0.71 rotate by 90
set label " ".intgr."+Qy" at intgr-Qy, 0.71 rotate by 90 }
set x2r [0:19.]; set xr [0:19000./M*G]; set yr [:1.01]
plot plotCmd(22) u (G/M*(Ei+($1-1.)*dE +M)):2 w l lw 2 lc rgb 'dark-red' t "<col.22> vs col.38"
average(awk script to compute (S<sub>Z</sub>) [5]:
function average(x, data){
```

```
function average(x, data){
    n = 0;wean = 0;
    val_min = 0;val_max = 0;
    for(val in data){
        n += 1;
        delta = val - mean;
        mean += delta/n;
        val_min = (n == 1)?val:((val < val_min)?val:val_min);
        val_max = (n == 1)?val:((val > val_max)?val:val_max);
    }
    if(n > 0){
        print x, mean, val_min, val_max;
    }}
    {
        curr = $38;
        yval = $(col_num);
        if((NR==1 || prev != curr){
            average(prev, data);
            delte data;
            prev = curr; }
        data[yval] = 1; }
END{
        average(curr, data); }
    }
```

⁷⁰¹⁴ (g) Crossing an intrinsic depolarizing resonance.

The simulation data files of question (f) can be used here, Tab. 20.66, *mutatis mutandis*, and the methodology in (f) can be followed. In particular, the following changes are needed:

- ⁷⁰¹⁸ Under OBJET:
- 1 st line, change the reference rigidity *BORO* to the proper value, some distance
 upstream of the resonance to be crossed,
- 3rd line, request a single particle ("1 1 1", in lieu of "1 11 1" which distributes
 11 particles on the vertical invariant),
 - 6th line, set the invariant ϵ_Z/π to the desired value,
- change the dipole field accordingly under DIPOLE, to maintains the expected curvature radius $\rho_0 = BORO/B = 20.76$ m (Tab. 9.3,
- under CAVITE, provide the desired peak voltage \hat{V} ,
- under REBELOTE, set the number of turns: a few thousands of turns upstream and downstream of the resonance.
- ⁷⁰²⁹ On the other hand, similar simulations are performed in questions (f)-(i) of ⁷⁰³⁰ exercise 9.1. Please refer to the solutions of these Saturne I simulations.

⁷⁰³¹ (h) Study of an imperfection depolarizing resonance.

The simulation data files of question (g) can be used here, *mutatis mutandis*, and the methodology in (g) can be followed.

On the other hand, similar simulations are performed in questions (f)-(i) of exercise 9.1, as well as in the "Strong Focusing Synchrotron" Chapter, Sec. 20.5. Please refer to the solutions of these simulations.

7037 **References**

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