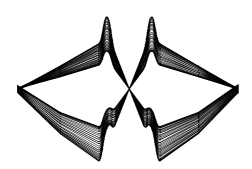
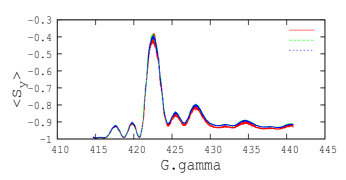
ZGOUBI USERS' GUIDE

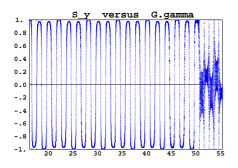
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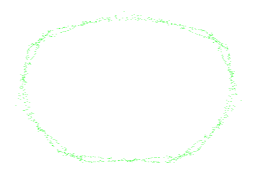
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Collider-Accelerator Department
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July 3, 2012









${}^{\scriptscriptstyle{0}}\textbf{Cover figures}:$

upper left: collision optics at ATLAS and CMS,

upper right: polarization upon crossing of 393+ Q_y resonance in RHIC,

lower left: spin-flipping with partial snakes along AGS cycle,

lower right: dynamic aperture in the Neutrino Factory muon decay ring.

Table of contents

PA	ART A	A Description of software contents	5
	GLOS	SSARY OF KEYWORDS	7
	OPTI	ICAL ELEMENTS VERSUS KEYWORDS	9
	PREF	FACE TO THE BNL EDITION (2012)	11
	INTR	CODUCTION TO THE 4th EDITION (1997), EXCERPT	13
	INTR	CODUCTION	15
1	NUM	ERICAL CALCULATION OF MOTION AND FIELDS	17
		Integration of the Lorentz Equation	
		1.2.1 Integration in magnetic fields	19
		1.2.2 Integration in electric fields	19
		1.2.3 Integration in combined electric and magnetic fields	22
		1.2.4 Calculation of the time of flight	23
	1.3	Calculation of $ec{E}$ and $ec{B}$ fields and their Derivatives $\dots \dots \dots$	23
		1.3.1 Extrapolation from 1-D axial field map	24
		1.3.2 Extrapolation From Analytical Models of Median Plane Fields	24
		1.3.3 Extrapolation from arbitrary 2-D Field Maps	25
		1.3.4 Interpolation in 3-D Field Maps	25
		1.3.5 2-D Analytical Field Models and Extrapolation	25
		1.3.6 3-D Analytical Models of Fields	25
		Calculation of \vec{E} and \vec{B} from Field Maps	27
		1.4.1 1-D Axial Map, with Cylindrical Symmetry	27
		1.4.2 2-D Median Plane Map, with Median Plane Antisymmetry	28
		1.4.3 Arbitrary 2-D Map, no Symmetry	31
		1.4.4 3-D Field Map	32
2	SPIN	TRACKING	35
3		CHROTRON RADIATION	37
	3.1	Energy loss and related dynamical effects	37
		Spectral-angular radiated densities	39
		3.2.1 Calculation of the radiated electric field	39
		3.2.2 Calculation of the Fourier transform of the electric field	42
4		CRIPTION OF THE AVAILABLE PROCEDURES	43
		Introduction	43
		Definition of an Object	43
		Declaration of options	54
		1	76
		Output Procedures	
		Complementary Features	
		4.6.1 Backward Ray-tracing	
		4.6.2 Checking Fields and Trajectories inside Optical Elements	
		4.6.3 Labeling keywords	
		4.6.4 Multiturn tracking in circular machines	
		4.6.5 Positioning, (mis-)alignement, of optical elements and field maps	
	4	4.6.6 Coded integration step	130

	4.6.7 4.6.8 4.6.9	Ray-tracing of an arbitrarily large number of particles	. 156
	4.0.9	regative figitity	. 130
P	ART B K	eywords and input data formatting	157
	GLOSSAR	Y OF KEYWORDS	159
	OPTICAL	ELEMENTS VERSUS KEYWORDS	161
	INTRODU	CTION	163
P	ART C E	xamples of input data files and output result files	257
	INTRODU	CTION	259
1	MONTE C	ARLO IMAGES IN SPES 2	261
2	TRANSFE	R MATRICES ALONG A TWO-STAGE SEPARATION KAON BEAM LINE	264
3	IN-FLIGH	T DECAY IN SPES 3	267
4	USE OF TH	HE FITTING PROCEDURE	270
5	MULTITU	RN SPIN TRACKING IN SATURNE 3 GeV SYNCHROTRON	272
6	MICRO-BI	EAM FOCUSING WITH ELECTROMAGNETIC QUADRUPOLES	274
P	ART D R	unning zgoubi and its post-processor/graphic interface zpop	279
	INTRODU	CTION	281
1	1.1 Makin 1.1.1 1.1.2 1.2 Runnin	TO RUN zgoubi AND zpop g the executable files zgoubi and zpop . The transportable package zgoubi The post-processor and graphic interface package zpop	. 281 . 281 . 281
2	STORAGE	FILES	281
	REFEREN	CES	284
	INDEX		285

PART A

Description of software contents

Glossary of Keywords

AGSMM	AGS main magnet	77
AGSQUAD	AGS quadrupole	78
AIMANT	Generation of dipole mid-plane 2-D map, polar frame	79
AUTOREF	Automatic transformation to a new reference frame	84
BEND	Bending magnet, Cartesian frame	85
BINARY	BINARY/FORMATTED data converter	55
BREVOL	1-D uniform mesh magnetic field map	86
CARTEMES	2-D Cartesian uniform mesh magnetic field map	87
CAVITE	Accelerating cavity	89
CHAMBR	Long transverse aperture limitation	
CHANGREF	Transformation to a new reference frame	
CIBLE	Generate a secondary beam from target interaction	
COLLIMA	Collimator	
DECAPOLE	Decapole magnet	
DIPOLE	Dipole magnet, polar frame	
DIPOLE-M	Generation of dipole mid-plane 2-D map, polar frame	
DIPOLES	Dipole magnet N -tuple, polar frame	
DODECAPO	Dodecapole magnet	
DRIFT	Field free drift space	
EBMULT	Electro-magnetic multipole	
EL2TUB	Two-tube electrostatic lens	
ELMIR	Electrostatic N-electrode mirror/lens, straight slits	
ELMIRC	Electrostatic N-electrode mirror/lens, circular slits	
ELMULT	Electric multipole	
ELREVOL	1-D uniform mesh electric field map	
EMMA	2-D Cartesian or cylindrical mesh field map for EMMA FFAG	
END	End of input data list; see FIN	
ESL	Field free drift space	
FAISCEAU	Print particle coordinates	
FAISCNL	Store particle coordinates in file FNAME	
FAISTORE	Store coordinates every IP other pass at labeled elements	
FFAG	FFAG magnet, N-tuple	
FFAG-SPI	Spiral FFAG magnet, N-tuple	
FIN	End of input data list	
FIT	Fitting procedure	
FIT2	Fitting procedure	
FOCALE	Particle coordinates and horizontal beam dimension at distance XL .	
FOCALEZ	Particle coordinates and vertical beam dimension at distance $XL\ldots$	
GASCAT	Gas scattering	
HISTO	1-D histogram	144

DAACE	T 1' ' 1' ' 1' ' 1' ' 1' ' ' 1' ' ' ' 1' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' '	1.40
IMAGE	Localization and size of horizontal waist	
IMAGES DAAGESZ	Localization and size of horizontal waists	
IMAGESZ	Localization and size of vertical waists	
IMAGEZ	Localization and size of vertical waist	
MAP2D	2-D Cartesian uniform mesh field map - arbitrary magnetic field	
MAP2D-E	2-D Cartesian uniform mesh field map - arbitrary electric field	
MARKER	Marker	
TRANSMAT	Matrix transfer	
MATRIX	Calculation of transfer coefficients, periodic parameters	
MCDESINT	Monte-Carlo simulation of in-flight decay	
MCOBJET	Monte-Carlo generation of a 6-D object	
MULTIPOL	Magnetic multipole	
OBJET	Generation of an object	
OBJETA	Object from Monte-Carlo simulation of decay reaction	
OCTUPOLE	Octupole magnet	
OPTICS	Write out optical functions	
ORDRE	Taylor expansions order	
PARTICUL	Particle characteristics	
PICKUPS	Beam centroid path; closed orbit	
PLOTDATA	Intermediate output for the PLOTDATA graphic software	
POISSON	Read magnetic field data from POISSON output	124
POLARMES	2-D polar mesh magnetic field map	125
PS170	Simulation of a round shape dipole magnet	
QUADISEX	Sharp edge magnetic multipoles	127
QUADRUPO	Quadrupole magnet	128
REBELOTE	'Do it again'	69
RESET	Reset counters and flags	70
SCALING	Time scaling of power supplies and R.F	71
SEPARA	Wien Filter - analytical simulation	130
SEXQUAD	Sharp edge magnetic multipole	127
SEXTUPOL	Sextupole magnet	131
SOLENOID	Solenoid	132
SPNPRNL	Store spin coordinates into file FNAME	149
SPNPRT	Print spin coordinates	149
SPNSTORE	Store spin coordinates every IP other pass at labeled elements	149
SPNTRK	Spin tracking	73
SRLOSS	Synchrotron radiation loss	75
SRPRNT	Print SR loss statistics	150
SYNRAD	Synchrotron radiation spectral-angular densities	76
TARGET	Generate a secondary beam from target interaction; see CIBLE	94
TOSCA	2-D and 3-D Cartesian or cylindrical mesh field map	
TRAROT	Translation-Rotation of the reference frame	
TWISS	Calculation of optical parameters; periodic parameters	151
UNDULATOR	Undulator magnet	
UNIPOT	Unipotential cylindrical electrostatic lens	
VENUS	Simulation of a rectangular shape dipole magnet	
WIENFILT	Wien filter	
YMY	Reverse signs of Y and Z reference axes	

Optical elements versus keywords

This glossary gives a list of keywords suitable for the simulation of common optical elements. These are classified in three categories: magnetic, electric and combined electro-magnetic elements.

Field map procedures are also listed; they provide a means for ray-tracing through measured or simulated electric and/or magnetic fields.

MAGNETIC ELEMENTS

AGS main magnet AGSMM

Decapole DECAPOLE, MULTIPOL

Dipole[s] AIMANT, BEND, DIPOLE[S], DIPOLE-M, MULTIPOL, QUADISEX

Dodecapole DODECAPO, MULTIPOL

FFAG magnets DIPOLES, FFAG, FFAG-SPI, MULTIPOL, EMMA

Helical dipole HELIX

Multipole MULTIPOL, QUADISEX, SEXQUAD

Octupole OCTUPOLE, MULTIPOL, QUADISEX, SEXQUAD

Quadrupole QUADRUPO, MULTIPOL, SEXQUAD

Sextupole SEXTUPOL, MULTIPOL, QUADISEX, SEXQUAD

Skew multipoles MULTIPOL Solenoid SOLENOID Undulator UNDULATOR

Using field maps

1-D, cylindrical symmetry BREVOL

2-D, mid-plane symmetry CARTEMES, POISSON, TOSCA

2-D, no symmetry
2-D, polar mesh, mid-plane symmetry
3-D, no symmetry
TOSCA

ELECTRIC ELEMENTS

2-tube (bipotential) lens EL2TUB 3-tube (unipotential) lens **UNIPOT** Decapole **ELMULT** Dipole **ELMULT** Dodecapole **ELMULT** Multipole **ELMULT** N-electrode mirror/lens, straight slits **ELMIR** N-electrode mirror/lens, circular slits **ELMIRC** Octupole **ELMULT** Quadrupole **ELMULT** R.F. (kick) cavity **CAVITE** Sextupole **ELMULT** Skew multipoles ELMULT

Using field maps

1D, cylindrical symmetry ELREVOL 2-D, no symmetry MAP2D-E

ELECTRO-MAGNETIC ELEMENTS

Decapole **EBMULT** Dipole **EBMULT** Dodecapole **EBMULT** Multipole **EBMULT** Octupole **EBMULT** Quadrupole **EBMULT** Sextupole **EBMULT** Skew multipoles **EBMULT**

Wien filter SEPARA, WIENFILT

PREFACE TO THE BNL EDITION (2012)

The previous release of the Zgoubi Users' Guide as a Lab. report dates from 1997, making the present one the last in a series of five [1]-[4].

The Introductory section of the Guide deserved updating, since so much developments have been accomplished these last years in the frame of a number of project design and beam dynamics studies, as the neutrino factory, lepton and hadron colliders, spin studies at AGS and RHIC, etc.

So did it go with the list of optical elements and the compendium of numerical methods, so-called "Glossary of keywords" list, pp. 7 and 159, that has stretched with new simulation and computing procedures, ranging from fitting to overlapping magnetic fields capabilities via spin and other radiation damping tools.

The code has been installed and made fully, and freely, available on SourceForge with the collaboration of J. S. Berg, on Sep 17, 2007 [5]. The SourceForge package evolves with the various projects dealt with and is continuously maintained. It includes the sources, the postprocessor progam **zpop**, as well as many examples with template input data files ("zgoubi.dat") and reference output result files ("zgoubi.res").

A series of computing tools have been developped in addition, planned to be made available on **zgoubi** SourceForge site in near future, that render the designer's life easier, as, search for closed orbits in periodic machines, computation of optical functions and parameters, tune scans, dynamic aperture scans, gnuplot graphic scripts, etc., including dedicated ones regarding, *e.g.*, FFAG R/D, AGS R/D, spin dynamics. In the same spirit, "python" interfaces are being developped, some made browsable on web by their authors.

Worth stressing, this manual is intended to describe the contents of the most recent version of **zgoubi**. Both the code and its Guide are far from being "finished products", though.

INTRODUCTION TO THE 4th EDITION (1997), EXCERPT

The initial version of **zgoubi**, dedicated to ray-tracing in magnetic fields, was developed by D. Garreta and J.C. Faivre at CEN-Saclay in the early 1970's. It was perfected for the purpose of studying the four spectrometers SPES I, II, III, IV at the Laboratoire National Saturne (CEA-Saclay, France), and SPEG at Ganil (Caen, France). It is being used since long in several national and foreign laboratories.

The first manual was in French [1]. Accounting for many developments and improvements, and in order to facilitate access to the program an English version of the manual was written at TRIUMF with the assistance of J. Doornbos. P. Stewart prepared the manuscript for publication [2]

An updating was necessary for accompanying the third version of the code which included developments regarding spin tracking and ray-tracing in combined electric and magnetic fields; this was done with the help of D. Bunel (SATURNE Laboratory, Saclay) for the preparation of the document and lead to the third release [3].

In the mid-1990s, the computation of synchrotron radiation electromagnetic impulse and spectra was introduced. In the mean time, several new optical elements were added, such as electro-magnetic and other electrostatic lenses. Used since several years for special studies in periodic machines (*e.g.*, SATURNE at Saclay, COSY at Julich, LEP and LHC at CERN), **zgoubi** has also undergone extensive developments regarding storage ring related features.

These developments of **zgoubi** have strongly benefited of the environment of the Groupe Théorie, Laboratoire National SATURNE, CEA/DSM-Saclay.

The graphic interface to **zgoubi** (addressed in Part D) has also undergone concomitent extensive developments, which make it a performing tool for the post-processing of **zgoubi** outputs.

This manual is intended only to describe the details of the most recent version of **zgoubi**, which is far from being a "finished product".

INTRODUCTION

The computer code **zgoubi** calculates trajectories of charged particles in magnetic and electric fields. At the origin specially adapted to the definition and adjustment of beam lines and magnetic spectrometers, it has so evolved that it allows the study of systems including complex sequences of optical elements such as dipoles, quadrupoles, arbitrary multipoles, FFAG magnets and other magnetic or electric devices, and is able as well to handle periodic structures. Compared to other codes, it presents several peculiarities, as follows - a non-exhaustive list:

- a numerical method for integrating the Lorentz equation, based on Taylor series, which optimizes computing time and provides high accuracy and strong symplecticity,
- spin tracking, using the same numerical method as for the Lorentz equation,
- full account of stochastic photon emission, and its effects on particle dynamics,
- calculation of the synchrotron radiation electric field and spectra in arbitrary magnetic fields, from the ray-tracing outcomes,
- the possibility of using a mesh, which allows ray-tracing from simulated or measured (1-D, 2-D or 3-D) electric and magnetic field maps,
- numerous Monte Carlo procedures : unlimited number of trajectories, in-flight decay, stochastic radiation, etc.
- built-in fitting procedures allowing arbitrary variables and a large variety of constraints, easily expandable,
- multiturn tracking in circular accelerators including features proper to machine parameter calculation and survey,
- simulation of time-varying power supplies,
- simulation of arbitrary radio-frequency programs.

The initial version of **zgoubi** was dedicated to ray-tracing in magnetic elements, beam lines, spectrometers. It was perfected for the purpose of studying, and operating, the four spectrometers SPES I, II, III, IV at the Laboratoire National Saturne (CEA-Saclay, France), and, later, SPEG at Ganil (Caen, France).

Developments regarding spin tracking and ray-tracing in combined electric and magnetic fields were implemented, in the late 1980s and early 1990s respectively.

In the mid-1990s, the computation of synchrotron radiation electromagnetic impulse and spectra was introduced, for the purpose of synchrotron radiation diagnostic R&D at LEP, and further applied to the design of the SR diagnostics installations at LHC in the early 2000s. In the mean time, several new optical elements were added, such as electro-magnetic and other electrostatic lenses. Used since several years for special studies in periodic machines (*e.g.*, SATURNE at Saclay, COSY at Julich, LEP and LHC at CERN), **zgoubi** has also undergone extensive developments regarding storage ring related features.

Many developments have been accomplished since the early 2000s in the frame of a number of project design and beam dynamics studies, as the neutrino factory, lepton and hadron colliders, spin studies at AGS and RHIC, etc. As a consequence the list of optical elements and the compendium of numerical methods, so-called "Glossary of Keywords" list, pp. 7 and 159, has stretched with new simulation and computing procedures, ranging from fitting to overlapping magnetic fields capabilities via spin and other radiation damping tools.

The graphic interface to **zgoubi** (**zpop**, Part D) has also been subject to extensive developments, making it a convenient companion tool to the use of **zgoubi**.

1 NUMERICAL CALCULATION OF MOTION AND FIELDS

1.1 zgoubi Frame

The reference frame of **zgoubi** is presented in Fig. 1. Its origin is in the median plane on a reference curve which coincides with the optical axis of optical elements.

1.2 Integration of the Lorentz Equation

The Lorentz equation, which governs the motion of a particle of charge q, relativistic mass m and velocity \vec{v} in electric and magnetic fields \vec{e} and \vec{b} , is written

$$\frac{d(m\vec{v})}{dt} = q\left(\vec{e} + \vec{v} \times \vec{b}\right) \tag{1.2.1}$$

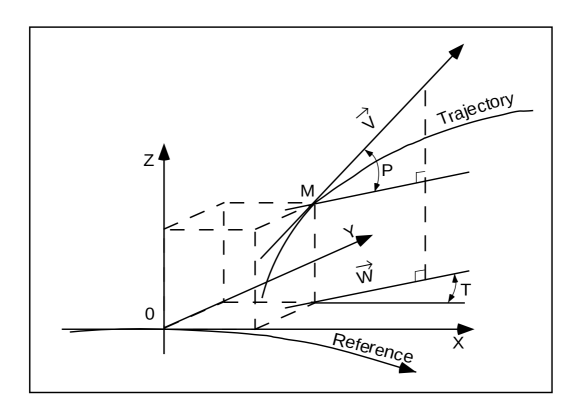


Figure 1: Reference frame and coordinates (Y, T, Z, P) in **zgoubi**.

OX: in the plane of the reference curve in the direction of motion,

OY: in the plane of the reference curve, normal to OX,

OZ: orthogonal to the (X, Y) plane,

 \vec{W} : projection of the velocity, \vec{v} , in the (X,Y) plane,

T =angle between \vec{W} and the X-axis,

 $P = \text{angle between } \vec{W} \text{ and } \vec{v}.$

Taking

$$\vec{u} = \frac{\vec{v}}{v}, \quad ds = v \, dt, \quad \vec{u}' = \frac{d\vec{u}}{ds}, \quad m\vec{v} = mv\vec{u} = q \, B\rho \, \vec{u}$$
 (1.2.2)

where $B\rho$ is the rigidity of the particle, this equation can be rewritten

$$(B\rho)'\vec{u} + B\rho\,\vec{u}' = \frac{\vec{e}}{v} + \vec{u} \times \vec{b} \tag{1.2.3}$$

From position $\vec{R}(M_0)$ and unit velocity $\vec{u}(M_0)$ at point M_0 , position $\vec{R}(M_1)$ and unit velocity $\vec{u}(M_1)$ at point M_1 following a displacement Δs , are obtained from truncated Taylor expansions (Fig. 2)

$$\vec{R}(M_1) \approx \vec{R}(M_0) + \vec{u}(M_0) \, \Delta s + \vec{u}'(M_0) \, \frac{\Delta s^2}{2!} + \dots + \vec{u}'''''(M_0) \, \frac{\Delta s^6}{6!}$$

$$\vec{u}(M_1) \approx \vec{u}(M_0) + \vec{u}'(M_0) \, \Delta s + \vec{u}''(M_0) \, \frac{\Delta s^2}{2!} + \dots + \vec{u}'''''(M_0) \, \frac{\Delta s^5}{5!}$$
(1.2.4)

The rigidity at M_1 is obtained in the same way from

$$(B\rho)(M_1) \approx (B\rho)(M_0) + (B\rho)'(M_0)\Delta s + \dots + (B\rho)''''(M_0)\frac{\Delta s^4}{4!}$$
 (1.2.5)

The equation of time of flight is written in a similar manner

$$T(M_1) \approx T(M_0) + \frac{dT}{ds}(M_0) \Delta s + \frac{d^2T}{ds^2}(M_0) \frac{\Delta s^2}{2} + \frac{d^3T}{ds^3}(M_0) \frac{\Delta s^3}{3!} + \frac{d^4T}{ds^4}(M_0) \frac{\Delta s^4}{4!}$$
 (1.2.6)

The derivatives $\vec{u}^{(n)} = \frac{d^n \vec{u}}{ds^n}$ and $(B\rho)^{(n)} = \frac{d^n (B\rho)}{ds^n}$ involved in these expressions are calculated as described in the next sections. For the sake of computing speed, three distinct software procedures are involved, depending on whether \vec{e} or \vec{b} is zero, or \vec{e} and \vec{b} are both non-zero.

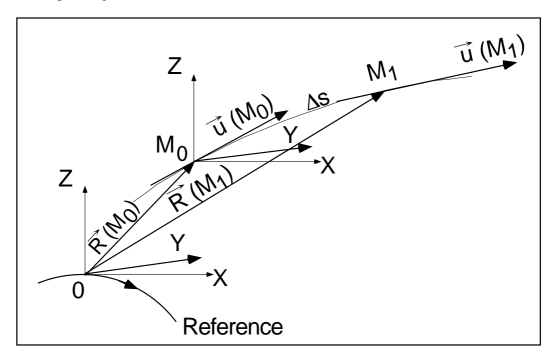


Figure 2: Position and velocity of a particle in the reference frame.

1.2.1 Integration in magnetic fields

Admitting that $\vec{e}=0$, and noting $\vec{B}=\frac{\vec{b}}{B\rho}$, eq. (1.2.3) reduces to

$$\vec{u}' = \vec{u} \times \vec{B}$$

The successive derivatives $\vec{u}^{(n)}=\frac{d^n\vec{u}}{ds^n}$ of \vec{u} needed in the Taylor expansions (eqs. 1.2.4) are calculated by differentiating $\vec{u}'=\vec{u}\times\vec{B}$

$$\vec{u}'' = \vec{u}' \times \vec{B} + \vec{u} \times \vec{B}'$$

$$\vec{u}''' = \vec{u}'' \times \vec{B} + 2\vec{u}' \times \vec{B}' + \vec{u} \times \vec{B}''$$

$$\vec{u}'''' = \vec{u}''' \times \vec{B} + 3\vec{u}'' \times \vec{B}' + 3\vec{u}' \times \vec{B}'' + \vec{u} \times \vec{B}'''$$

$$\vec{u}'''''' = \vec{u}'''' \times \vec{B} + 4\vec{u}''' \times \vec{B}' + 6\vec{u}'' \times \vec{B}'' + 4\vec{u}' \times \vec{B}''' + \vec{u} \times \vec{B}''''$$
(1.2.7)

where $\vec{B}^{(n)} = \frac{d^n \vec{B}}{ds^n}$.

From $d\vec{B} = \frac{\partial \vec{B}}{\partial X} dX + \frac{\partial \vec{B}}{\partial Y} dY + \frac{\partial \vec{B}}{\partial Z} dZ = \sum_{i=1,3} \frac{\partial \vec{B}}{\partial X_i} dX_i$, and by successive differentiation, we get

$$\vec{B}' = \sum_{i} \frac{\partial \vec{B}}{\partial X_{i}} u_{i}$$

$$\vec{B}'' = \sum_{ij} \frac{\partial^{2} \vec{B}}{\partial X_{i} \partial X_{j}} u_{i} u_{j} + \sum_{i} \frac{\partial \vec{B}}{\partial X_{i}} u'_{i}$$

$$\vec{B}''' = \sum_{ijk} \frac{\partial^{3} \vec{B}}{\partial X_{i} \partial X_{j} \partial X_{k}} u_{i} u_{j} u_{k} + 3 \sum_{ij} \frac{\partial^{2} \vec{B}}{\partial X_{i} \partial X_{j}} u'_{i} u_{j} + \sum_{i} \frac{\partial \vec{B}}{\partial X_{i}} u''_{i}$$

$$\vec{B}'''' = \sum_{ijkl} \frac{\partial^{4} \vec{B}}{\partial X_{i} \partial X_{j} \partial X_{k} \partial X_{l}} u_{i} u_{j} u_{k} u_{l} + 6 \sum_{ijk} \frac{\partial^{3} \vec{B}}{\partial X_{i} \partial X_{j} \partial X_{k}} u'_{i} u_{j} u_{k}$$

$$+ 4 \sum_{ij} \frac{\partial^{2} \vec{B}}{\partial X_{i} \partial X_{j}} u''_{i} u_{j} + 3 \sum_{ij} \frac{\partial^{2} \vec{B}}{\partial X_{i} \partial X_{j}} u'_{i} u'_{j} + \sum_{i} \frac{\partial \vec{B}}{\partial X_{i}} u''_{i}$$

$$(1.2.8)$$

From the knowledge of $\vec{u}(M_0)$ and $\vec{B}(M_0)$ at point M_0 of the trajectory, we calculate alternately the derivatives of $\vec{u}(M_0)$ and $\vec{B}(M_0)$, by means of eqs. (1.2.7) and (1.2.8), and inject it in eq. (1.2.4) to get $\vec{R}(M_1)$ and $\vec{u}(M_1)$.

1.2.2 Integration in electric fields [6]

Admitting that $\vec{b} = 0$, eq. (1.2.3) reduces to

$$(B\rho)'\vec{u} + B\rho\vec{u}' = \frac{\vec{e}}{v} \tag{1.2.9}$$

which, by successive differentiations, gives the recursive relations

$$(B\rho)'\vec{u} + B\rho\vec{u}' = \frac{\vec{e}}{v}$$

$$(B\rho)''\vec{u} + 2(B\rho)'\vec{u}' + B\rho\vec{u}'' = \left(\frac{1}{v}\right)'\vec{e} + \frac{\vec{e}'}{v}$$

$$(B\rho)'''\vec{u} + 3(B\rho)''\vec{u}' + 3(B\rho)'\vec{u}'' + B\rho\vec{u}''' = \left(\frac{1}{v}\right)''\vec{e} + 2\left(\frac{1}{v}\right)''\vec{e}' + \left(\frac{1}{v}\right)''\vec{e}'' + \left(\frac{1}{v}\right)''\vec{e}'' + \left(\frac{1}{v}\right)''\vec{e}'' + 4(B\rho)''\vec{u}''' + 4(B\rho)''\vec{u}''' + B\rho\vec{u}'''' = \left(\frac{1}{v}\right)'''\vec{e} + 3\left(\frac{1}{v}\right)'''\vec{e}'' + 3\left(\frac{1}{v}\right)''\vec{e}'' + \frac{1}{v}\vec{e}''''$$

$$(1.2.10)$$

that provide the derivatives $\frac{d^n \vec{u}}{ds^n}$ needed in the Taylor expansions (eq. 1.2.4)

where $\vec{E} = \frac{\vec{e}}{B\rho}$, and ()⁽ⁿ⁾ $|_{B\rho}$ denotes differentiation at constant $B\rho$: $\vec{E}^{(n)}$ $|_{B\rho} = \frac{1}{B\rho} \frac{d^n \vec{e}}{ds^n}$. These derivatives of the electric field are obtained from the total derivative

$$d\vec{E} = \frac{\partial \vec{E}}{\partial X} dX + \frac{\partial \vec{E}}{\partial Y} dY + \frac{\partial \vec{E}}{\partial Z} dZ$$
 (1.2.12)

by successive differentiations

$$\vec{E}' = \sum_{i} \frac{\partial \vec{E}}{\partial X_{i}} u_{i}$$

$$\vec{E}'' = \sum_{ij} \frac{\partial^{2} \vec{E}}{\partial X_{i} \partial X_{j}} u_{i} u_{j} + \sum_{i} \frac{\partial \vec{E}}{\partial X_{i}} u'_{i}$$

$$\vec{E}''' = \sum_{ijk} \frac{\partial^{3} \vec{E}}{\partial X_{i} \partial X_{j} \partial X_{k}} u_{i} u_{j} u_{k} + 3 \sum_{ij} \frac{\partial^{2} \vec{E}}{\partial X_{i} \partial X_{j}} u'_{i} u_{j} + \sum_{i} \frac{\partial \vec{E}}{\partial X_{i}} u''_{i}$$

$$(1.2.13)$$

etc. as in eq. 1.2.8. These eqs. (1.2.11), as well as the calculation of the rigidity, following eq. (1.2.5), involve derivatives $(B\rho)^{(n)} = \frac{d^n(B\rho)}{ds^n}$, which are obtained in the following way. Considering that

$$\frac{dp^2}{dt} = \frac{d\vec{p}^2}{dt} \quad i.e., \quad \frac{dp}{dt}p = \frac{d\vec{p}}{dt}\vec{p}$$
 (1.2.14)

with $\frac{d\vec{p}}{dt} = q \left(\vec{e} + \vec{v} \times \vec{b} \right)$ (eq. 1.2.1), we obtain

$$\frac{dp}{dt}p = q(\vec{e} + v \times \vec{b}) \cdot \vec{p} = q\vec{e} \cdot \vec{p}$$
(1.2.15)

since $(\vec{v} \times \vec{b}) \cdot \vec{p} = 0$. Normalizing as previously with $\vec{p} = p\vec{u} = qB\rho\vec{u}$ and ds = vdt, and by successive differentiations, eq. (1.2.15) leads to the $(B\rho)^{(n)}$

$$(B\rho)' = \frac{1}{v} (\vec{e} \cdot \vec{u})$$

$$(B\rho)'' = \left(\frac{1}{v}\right)' (\vec{e} \cdot \vec{u}) + \frac{1}{v} (\vec{e} \cdot \vec{u})'$$

$$(B\rho)''' = \left(\frac{1}{v}\right)'' (\vec{e} \cdot \vec{u}) + 2\left(\frac{1}{v}\right)' (\vec{e} \cdot \vec{u})' + \frac{1}{v} (\vec{e} \cdot \vec{u})''$$

$$(B\rho)'''' = \left(\frac{1}{v}\right)''' (\vec{e} \cdot \vec{u}) + 3\left(\frac{1}{v}\right)'' (\vec{e} \cdot \vec{u})' + 3\left(\frac{1}{v}\right)' (\vec{e} \cdot \vec{u})'' + \frac{1}{v} (\vec{e} \cdot \vec{u})'''$$

$$(1.2.16)$$

Note that the derivatives $(\vec{e} \cdot \vec{u})^{(n)} = \frac{d^n(\vec{e} \cdot \vec{u})}{ds^n}$ can be related to the derivatives of the kinetic energy W by $dW = \frac{d\vec{p}}{dt} \cdot \vec{v} \, dt = q\vec{e} \cdot \vec{v} \, dt$ which leads to

$$\frac{d^{n+1}W}{ds^{n+1}} = q \frac{d^n(\vec{e} \cdot \vec{u})}{ds^n}$$
 (1.2.17)

Finally, the derivatives $\left(\frac{1}{v}\right)^{(n)} = \frac{d^n\left(\frac{1}{v}\right)}{ds^n}$ involved in eqs. (1.2.11,1.2.16) are obtained from $p = \frac{v}{c} \frac{W + m_0 c^2}{c}$, (m_0 is the rest mass) by successive differentiations, that give the recursive relations

$$\begin{pmatrix} \frac{1}{v} \end{pmatrix} = \frac{1}{c^2} \frac{W + m_0 c^2}{q B \rho}
\begin{pmatrix} \frac{1}{v} \end{pmatrix}' = \frac{1}{c^2} \frac{(\vec{e} \cdot \vec{u})}{B \rho} - \frac{1}{v} \frac{(B \rho)'}{B \rho}
\begin{pmatrix} \frac{1}{v} \end{pmatrix}'' = \frac{1}{c^2} \frac{(\vec{e} \cdot \vec{u})'}{B \rho} - 2 \left(\frac{1}{v} \right)' \frac{(B \rho)'}{B \rho} - \frac{1}{v} \frac{(B \rho)''}{B \rho}
\begin{pmatrix} \frac{1}{v} \end{pmatrix}''' = \frac{1}{c^2} \frac{(\vec{e} \cdot \vec{u})''}{B \rho} - 3 \left(\frac{1}{v} \right)'' \frac{(B \rho)'}{B \rho} - 3 \left(\frac{1}{v} \right)' \frac{(B \rho)''}{B \rho} - \frac{1}{v} \frac{(B \rho)'''}{B \rho}$$
(1.2.18)

1.2.3 Integration in combined electric and magnetic fields

When both \vec{e} and \vec{b} are non-zero, the complete eq. (1.2.3) must be considered. Recursive differentiations give the following relations

$$(B\rho)'\vec{u} + B\rho\vec{u}' = \frac{\vec{e}}{v} + \vec{u} \times \vec{b}$$

$$(B\rho)''\vec{u} + 2(B\rho)'\vec{u}' + B\rho\vec{u}'' = \left(\frac{1}{v}\right)'\vec{e} + \left(\frac{1}{v}\right)\vec{e}' + (\vec{u} \times \vec{b})'$$

$$(B\rho)'''\vec{u} + 3(B\rho)''\vec{u}' + 3(B\rho)'\vec{u}'' + B\rho\vec{u}''' = \left(\frac{1}{v}\right)''\vec{e} + 2\left(\frac{1}{v}\right)''\vec{e}' + \left(\frac{1}{v}\right)\vec{e}'' + (\vec{u} \times \vec{b})''$$

$$(B\rho)''''\vec{u} + 4(B\rho)'''\vec{u}' + 6(B\rho)''\vec{u}'' + 4(B\rho)'\vec{u}''' + B\rho\vec{u}'''' = \left(\frac{1}{v}\right)'''\vec{e}' + 3\left(\frac{1}{v}\right)''\vec{e}'' + 3\left(\frac{1}{v}\right)''\vec{e}'' + \frac{1}{v}\vec{e}''' + (\vec{u} \times \vec{b})'''$$

$$(1.2.19)$$

that provide the derivatives $\frac{d^n \vec{u}}{ds^n}$ needed in the Taylor expansions (1.2.4)

$$\vec{u}' = \left(\frac{1}{v}\right) \vec{E} + (\vec{u} \times \vec{B}) - \frac{(B\rho)'}{B\rho} \vec{u}$$

$$\vec{u}'' = \left(\frac{1}{v}\right)' \vec{E} + \left(\frac{1}{v}\right) \vec{E}' \mid_{B\rho} + (\vec{u} \times \vec{B}')' \mid_{B\rho} - 2\frac{(B\rho)'}{B\rho} \vec{u}' - \frac{(B\rho)''}{B\rho} \vec{u}$$

$$\vec{u}''' = \left(\frac{1}{v}\right)'' \vec{E} + 2\left(\frac{1}{v}\right)' \vec{E}' \mid_{B\rho} + \frac{1}{v} \vec{E}'' \mid_{B\rho} + (\vec{u} \times \vec{B})'' \mid_{B\rho} - 3\frac{(B\rho)'}{B\rho} \vec{u}'' - 3\frac{(B\rho)''}{B\rho} \vec{u}'' - \frac{(B\rho)'''}{B\rho} \vec{u}$$

$$\vec{u}'''' = \left(\frac{1}{v}\right)''' \vec{E} + 3\left(\frac{1}{v}\right)'' \vec{E}' \mid_{B\rho} + 3\left(\frac{1}{v}\right)' \vec{E}'' \mid_{B\rho} + \left(\frac{1}{v}\right) \vec{E}''' \mid_{B\rho}$$

$$+ (\vec{u} \times \vec{B})''' \mid_{B\rho} - 4\frac{(B\rho)'}{B\rho} \vec{u}''' - 6\frac{(B\rho)''}{B\rho} \vec{u}'' - 4\frac{(B\rho)'''}{B\rho} \vec{u}' - \frac{(B\rho)''''}{B\rho} \vec{u}'$$

$$(1.2.20)$$

where $\vec{E} = \frac{\vec{e}}{B\rho}$, $\vec{B} = \frac{\vec{b}}{B\rho}$, and $^{(n)}$ $|_{B\rho}$ denotes differentiation at constant $B\rho$

$$\vec{E}^{(n)} \mid_{B\rho} = \frac{1}{B\rho} \frac{d^n \vec{e}}{ds^n} \text{ and } (\vec{u} \times \vec{B})^{(n)} \mid_{B\rho} = \frac{1}{B\rho} (\vec{u} \times \vec{b})^{(n)}.$$
 (1.2.21)

These derivatives $\vec{E}^{(n)}$ and $\vec{B}^{(n)}$ of the electric and magnetic fields are calculated from the vector fields $\vec{E}(X,Y,Z)$, $\vec{B}(X,Y,Z)$ and their derivatives $\frac{\partial^{i+j+k}\vec{E}}{\partial X^i\partial Y^j\partial Z^k}$ and $\frac{\partial^{i+j+k}\vec{B}}{\partial X^i\partial Y^j\partial Z^k}$, following eqs. (1.2.8) and (1.2.13).

1.2.4 Calculation of the time of flight

The time of flight eq. (1.2.6) involves the derivatives dT/ds = 1/v, $d^2T/ds^2 = d(1/v)/ds$, etc. that are obtained from eq. (1.2.18). In the absence of electric field eq. (1.2.7) however reduces to the simple form

$$T(M_1) = T(M_0) + \Delta s/v \tag{1.2.22}$$

1.3 Calculation of \vec{E} and \vec{B} fields and their Derivatives

In this section, unless otherwise stated, $\vec{B} = (B_X(X,Y,Z), B_Y(X,Y,Z), B_Z(X,Y,Z))$ stands indifferently for electric field \vec{E} or magnetic field \vec{B} .

 $\vec{B}(X,Y,Z)$ and derivatives are calculated in various ways, depending whether field maps or analytic representations of optical elements are used. The basic means are the following.

1.3.1 Extrapolation from 1-D axial field map [7]

A cylindrically symmetric field (e.g., using BREVOL, ELREVOL) can be described by an axial 1-D field map of its longitudinal component $B_X(X,r=0)$ ($r=(Y^2+Z^2)^{1/2}$), while the radial component on axis $B_r(X,r=0)$ is assumed to be zero. $B_X(X,r=0)$ is obtained at any point along the X-axis by a polynomial interpolation from the map mesh (see section 1.4.1). Then the field components $B_X(X,r)$, $B_r(X,r)$ at the position of the particle, (X,r) are obtained from Taylor expansions truncated at the fifth order in r (hence, up to the fifth order derivative $\frac{\partial^5 B_X}{\partial X^5}(X,0)$), assuming cylindrical symmetry

$$B_X(X,r) = B_X(X,0) - \frac{r^2}{4} \frac{\partial^2 B_X}{\partial X^2} (X,0) + \frac{r^4}{64} \frac{\partial^4 B_X}{\partial X^4} (X,0)$$

$$B_r(X,r) = -\frac{r}{2} \frac{\partial B_X}{\partial X} (X,0) + \frac{r^3}{16} \frac{\partial^3 B_X}{\partial X^3} (X,0) - \frac{r^5}{384} \frac{\partial^5 B_X}{\partial X^5} (X,0)$$
(1.3.1)

Then, by differentiation with respect to X and r, up to the second order, these expressions provide the derivatives of $\vec{B}(X,r)$. Finally a conversion from the (X,r) coordinates to the (X,Y,Z) Cartesian coordinates of **zgoubi** is performed, thus providing the expressions $\frac{\partial^{i+j+k}\vec{B}}{\partial X^i\partial Y^j\partial Z^k}$ needed in the eq. (1.2.8).

1.3.2 Extrapolation From Analytical Models of Median Plane Fields

In the median plane, $B_Z(X, Y, 0)$ and its derivatives with respect to X or Y may be deriveded from analytical models (e.g., in Venus magnet - VENUS, and sharp edge multipoles SEXQUAD and QUADI-SEX) or numerically by polynomial interpolation from 2-D field maps (e.g., CARTEMES, TOSCA).

Median plane antisymmetry is assumed, which results in

$$B_X(X, Y, 0) = 0$$

$$B_Y(X, Y, 0) = 0$$

$$B_X(X, Y, Z) = -B_X(X, Y, -Z)$$

$$B_Y(X, Y, Z) = -B_Y(X, Y, -Z)$$

$$B_Z(X, Y, Z) = B_Z(X, Y, -Z)$$
(1.3.2)

Accommodated with Maxwell's equations, this results in Taylor expansions below, for the three components of \vec{B} (here, B stands for $B_Z(X,Y,0)$)

$$B_{X}(X,Y,Z) = Z \frac{\partial B}{\partial X} - \frac{Z^{3}}{6} \left(\frac{\partial^{3}B}{\partial X^{3}} + \frac{\partial^{3}B}{\partial X \partial Y^{2}} \right)$$

$$B_{Y}(X,Y,Z) = Z \frac{\partial B}{\partial Y} - \frac{Z^{3}}{6} \left(\frac{\partial^{3}B}{\partial X^{2} \partial Y} + \frac{\partial^{3}B}{\partial Y^{3}} \right)$$

$$B_{Z}(X,Y,Z) = B - \frac{Z^{2}}{2} \left(\frac{\partial^{2}B}{\partial X^{2}} + \frac{\partial^{2}B}{\partial Y^{2}} \right) + \frac{Z^{4}}{24} \left(\frac{\partial^{4}B}{\partial X^{4}} + 2 \frac{\partial^{4}B}{\partial X^{2} \partial Y^{2}} + \frac{\partial^{4}B}{\partial Y^{4}} \right)$$
(1.3.3)

which are then differentiated one by one with respect to X, Y, or Z, up to second or fourth order (depending on optical element or IORDRE option, see section 1.4.2) so as to get the expressions involved in eq. (1.2.8).

1.3.3 Extrapolation from arbitrary 2-D Field Maps

2-D field maps that give the three components $B_X(X,Y,Z_0)$, $B_Y(X,Y,Z_0)$ and $B_Z(X,Y,Z_0)$ at each node (X,Y) of a Z_0 Z-elevation map may be used. \vec{B} and its derivatives at any point (X,Y,Z) are calculated by polynomial interpolation followed by Taylor expansions in Z, without any hypothesis of symmetries (see section 1.4.3 and keywords MAP2D, MAP2D-E).

1.3.4 Interpolation in 3-D Field Maps [8]

In 3-D field maps \vec{B} and its derivatives up to the second order with respect to X, Y, or Z are calculated by means of a second order polynomial interpolation, from 3-D $3 \times 3 \times 3$ -point grid (see section 1.4.4).

1.3.5 2-D Analytical Field Models and Extrapolation

Several optical elements such as *BEND*, *WIENFILT* (that uses the *BEND* procedures), *QUADISEX*, *VENUS*, etc., are defined from the expression of the field and derivatives in the median plane. 3-D extrapolation of these off the median is drawn from Taylor expansions.

1.3.6 3-D Analytical Models of Fields

In many optical elements such as QUADRUPO, SEXTUPOL, MULTIPOL, EBMULT, etc., the three components of \vec{B} and their derivatives with respect to X, Y or Z are obtained at any step along trajectories from analytical expression drawn from the scalar potential V(X,Y,Z) following

$$B_X = \frac{\partial V}{\partial X}, \quad B_Y = \frac{\partial V}{\partial Y}, \quad B_Z = \frac{\partial V}{\partial Z}, \quad \frac{\partial B_X}{\partial X} = \frac{\partial^2 V}{\partial X^2}, \quad \frac{\partial B_X}{\partial Y} = \frac{\partial^2 V}{\partial X \partial Y}, \quad \text{etc.}$$
 (1.3.4)

Multipoles

The scalar potential used for the calculation of $\frac{\partial^{i+j+k}\vec{B}_n(X,Y,Z)}{\partial X^i\partial Y^i\partial Z^k}$ (i+j+k=0 to 4) in the case of magnetic and electro-magnetic multipoles with 2n poles (namely, QUADRUPO (n=2) to DODE-CAPO (n=6), MULTIPOL (n=1 to 10), EBMULT (n=1 to 10)) is [9]

$$V_n(X,Y,Z) = (n!)^2 \left(\sum_{q=0}^{\infty} (-1)^q \frac{G^{(2q)}(X)(Y^2 + Z^2)^q}{4^q q! (n+q)!} \right) \left(\sum_{m=0}^n \frac{\sin\left(m\frac{\pi}{2}\right) Y^{n-m} Z^m}{m! (n-m)!} \right)$$
(1.3.5)

where G(X) is the longitudinal gradient, defined at the entrance or exit of the optical element by

$$G(s) = \frac{G_0}{1 + \exp(P(s))}, \quad G_0 = \frac{B_0}{R_0^n}$$
 (1.3.6)

wherein

$$P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^2 + C_3 \left(\frac{s}{\lambda}\right)^3 + C_4 \left(\frac{s}{\lambda}\right)^4 + C_5 \left(\frac{s}{\lambda}\right)^5$$

and s is the distance to the EFB.

Skew multipoles

A multipole component with arbitrary order n can be tilted independently of the others by an arbitrary angle A_n around the X-axis. If so, the calculation of the field and derivatives in the rotated axis (X, Y_R, Z_R) is done in two steps. First, they are calculated at the rotated position (X, Y_R, Z_R) , in the (X, Y, Z) frame, as derived from expression (1.3.5) above. Second, \vec{B} and its derivatives at (X, Y_R, Z_R) in the (X, Y, Z) frame are transformed to the rotated (X, Y_R, Z_R) frame by a rotation of the same angle A_n .

In particular a skew 2n-pole component is created by taking $A_n=\pi/2n$.

A Note on Electrostatic Multipoles

A right electric multipole has the same field equations as a as the like-order skew magnetic multipole. Therefore, calculation of right or skew electric or electro-magnetic multipoles (ELMULT, EBMULT, ELMULT) uses the same eq. (1.3.5) together with the rotation process as described in section 1.3.6. The same method is used, for arbitrary rotation of arbitrary multipole component around the X-axis.

1.4 Calculation of \vec{E} and \vec{B} from Field Maps

In this section, unless otherwise stated, $\vec{B} = (B_X(X,Y,Z), B_Y(X,Y,Z), B_Z(X,Y,Z))$ stands indifferently for electric field \vec{E} or magnetic field \vec{B} .

1.4.1 1-D Axial Map, with Cylindrical Symmetry

Let B_i be the value of the longitudinal component $B_X(X, r=0)$ of the field \vec{B} , at node i of a uniform mesh that defines a 1-D field map along the symmetry X-axis, while $B_r(X, r=0)$ is assumed to be zero $(r=(Y^2+Z^2)^{1/2})$. The field component $B_X(X, r=0)$ is calculated by a polynomial interpolation of the fifth degree in X, using a 5 points grid centered at the node of the 1-D map which is closest to the actual coordinate X of the particle.

The interpolation polynomial is

$$B(X,0) = A_0 + A_1X + A_2X^2 + A_3X^3 + A_4X^4 + A_5X^5$$
(1.4.1)

and the coefficients A_i are calculated by expressions that minimize the quadratic sum

$$S = \sum_{i} (B(X,0) - B_i)^2$$
 (1.4.2)

Namely, the source code contains the explicit analytical expressions of the coefficients A_i solutions of the normal equations $\partial S/\partial A_i=0$.

The derivatives $\frac{\partial^n B}{\partial X^n}(X,0)$ at the actual position X, as involved in eqs. (1.3.1), are then obtained by differentiation of the polynomial (1.4.1), giving

$$\frac{\partial B}{\partial X}(X,0) = A_1 + 2A_2X + 3A_3X^2 + 4A_4X^3 + 5A_5X^4$$

$$\frac{\partial^2 B}{\partial X^2}(X,0) = 2A_2 + 6A_3X + 12A_4X^2 + 20A_5X^3$$
...
$$\frac{\partial^5 B}{\partial X^5}(X,0) = 120A_5$$
(1.4.3)

1.4.2 2-D Median Plane Map, with Median Plane Antisymmetry

Let B_{ij} be the value of $B_Z(X,Y,0)$ at the nodes of a mesh which defines a 2-D field map in the (X,Y) plane while $B_X(X,Y,0)$ and $B_Y(X,Y,0)$ are assumed to be zero. Such a map may have been built or measured in either Cartesian or polar coordinates. Whenever polar coordinates are used, a change to Cartesian coordinates (described below) provides the expression of \vec{B} and its derivatives as involved in eq. (1.2.8).

zgoubi provides three types of polynomial interpolation from the mesh (option *IORDRE*); namely, a second order interpolation, with either a 9- or a 25-point grid, or a fourth order interpolation with a 25-point grid (Fig. 3).

If the 2-D field map is built up from simulation, the grid simply aims at interpolating the field at a given point from its 9 or 25 neighbors. If the map results from measurements, the grid also smoothes field measurement fluctuations.

The mesh may be defined in Cartesian coordinates, (Figs. 3A and 3B) or in polar coordinates (Fig. 3C). The interpolation grid is centered on the node which is closest to the projection in the (X, Y) plane of the actual point of the trajectory.

The interpolation polynomial is

$$B(X,Y,0) = A_{00} + A_{10}X + A_{01}Y + A_{20}X^2 + A_{11}XY + A_{02}Y^2$$
(1.4.4)

in second order, or

$$B(X,Y,0) = A_{00} + A_{10}X + A_{01}Y + A_{20}X^{2} + A_{11}XY + A_{02}Y^{2}$$

$$+ A_{30}X^{3} + A_{21}X^{2}Y + A_{12}XY^{2} + A_{03}Y^{3}$$

$$+ A_{40}X^{4} + A_{31}X^{3}Y + A_{22}X^{2}Y^{2} + A_{13}XY^{3} + A_{04}Y^{4}$$

$$(1.4.5)$$

in fourth order. The coefficients A_{ij} are calculated by expressions that minimize, with respect to A_{ij} , the quadratic sum

$$S = \sum_{ij} (B(X, Y, 0) - B_{ij})^2$$
(1.4.6)

The source code contains the explicit analytical expressions of the coefficients A_{ij} solutions of the normal equations $\partial S/\partial A_{ij}=0$.

The A_{ij} may then be identified with the derivatives of B(X,Y,0) at the central node of the grid

$$A_{ij} = \frac{1}{i!j!} \frac{\partial^{i+j} B}{\partial X^i \partial Y^j} (0, 0, 0)$$

$$(1.4.7)$$

The derivatives of B(X, Y, 0) with respect to X and Y, at the actual point (X, Y, 0) are obtained by differentiation of the interpolation polynomial, which gives (e.g., from (1.4.4) in the case of second order interpolation)

$$\frac{\partial B}{\partial X}(X, Y, 0) = A_{10} + 2A_{20}X + A_{11}Y
\frac{\partial B}{\partial Y}(X, Y, 0) = A_{01} + A_{11}X + 2A_{02}Y
\text{etc.}$$
(1.4.8)

This allows stepping to the calculation of $\vec{B}(X,Y,Z)$ and its derivatives as described in subsection 1.3.2 (eq. 1.3.3).

The special case of polar maps

It is necessary to change from polar map frame (R, α, Z) to the Cartesian moving frame (X, Y, Z). This is done as follows.

In second order calculations the correspondence is (we note $B \equiv B_Z(Z=0)$)

$$\frac{\partial B}{\partial X} = \frac{1}{R} \frac{\partial B}{\partial \alpha}
\frac{\partial B}{\partial Y} = \frac{\partial B}{\partial R}
\frac{\partial^2 B}{\partial X^2} = \frac{1}{R^2} \frac{\partial^2 B}{\partial \alpha^2} + \frac{1}{R} \frac{\partial B}{\partial R}
\frac{\partial^2 B}{\partial X \partial Y} = \frac{1}{R} \frac{\partial^2 B}{\partial \alpha \partial R} - \frac{1}{R^2} \frac{\partial B}{\partial \alpha}
\frac{\partial^2 B}{\partial Y^2} = \frac{\partial^2 B}{\partial R^2}
\frac{\partial^3 B}{\partial X^3} = \frac{3}{R^2} \frac{\partial^2 B}{\partial \alpha \partial R} - \frac{2}{R^3} \frac{\partial B}{\partial \alpha}
\frac{\partial^3 B}{\partial X^2 \partial Y} = \frac{-2}{R^3} \frac{\partial^2 B}{\partial \alpha^2} - \frac{1}{R^2} \frac{\partial B}{\partial R} + \frac{1}{R} \frac{\partial^2 B}{\partial R^2}
\frac{\partial^3 B}{\partial X \partial Y^2} = \frac{2}{R^3} \frac{\partial B}{\partial \alpha} - \frac{2}{R^2} \frac{\partial^2 B}{\partial \alpha \partial R}
\frac{\partial^3 B}{\partial Y^3} = 0$$
(1.4.9)

In fourth order calculations the relations are the same up to second order, and then

$$\frac{\partial^{3}B}{\partial X^{3}} = \frac{1}{R^{3}} \frac{\partial^{3}B}{\partial \alpha^{3}} + \frac{3}{R^{2}} \frac{\partial^{2}B}{\partial \alpha \partial R} - \frac{2}{R^{3}} \frac{\partial B}{\partial \alpha}$$

$$\frac{\partial^{3}B}{\partial X^{2}\partial Y} = \frac{1}{R^{2}} \frac{\partial^{3}B}{\partial \alpha^{2}\partial R} - \frac{2}{R^{3}} \frac{\partial^{2}B}{\partial \alpha^{2}} - \frac{1}{R^{2}} \frac{\partial B}{\partial R} + \frac{1}{R} \frac{\partial^{2}B}{\partial R^{2}}$$

$$\frac{\partial^{3}B}{\partial X\partial Y^{2}} = \frac{1}{R} \frac{\partial^{3}B}{\partial \alpha \partial R^{2}} + \frac{2}{R^{3}} \frac{\partial B}{\partial \alpha} - \frac{2}{R^{2}} \frac{\partial^{2}B}{\partial \alpha \partial R}$$

$$\frac{\partial^{3}B}{\partial Y^{3}} = \frac{\partial^{3}B}{\partial R^{3}}$$

$$\frac{\partial^{4}B}{\partial X^{4}} = \frac{1}{R^{4}} \frac{\partial^{4}B}{\partial \alpha^{4}} - \frac{8}{R^{4}} \frac{\partial^{2}B}{\partial \alpha^{2}} + \frac{6}{R^{3}} \frac{\partial^{3}B}{\partial \alpha^{2}\partial R} + \frac{3}{R^{2}} \frac{\partial^{2}B}{\partial R^{2}} - \frac{3}{R^{3}} \frac{\partial B}{\partial R}$$

$$\frac{\partial^{4}B}{\partial X^{3}\partial Y} = \frac{1}{R^{3}} \frac{\partial^{4}B}{\partial \alpha^{3}\partial R} - \frac{3}{R^{4}} \frac{\partial^{3}B}{\partial \alpha^{3}\partial R} - \frac{2}{R^{2}} \frac{\partial^{3}B}{\partial \alpha \partial R^{2}} - \frac{8}{R^{3}} \frac{\partial^{2}B}{\partial \alpha \partial R} + \frac{6}{R^{4}} \frac{\partial B}{\partial \alpha}$$

$$\frac{\partial^{4}B}{\partial X^{2}2Y^{2}} = \frac{1}{R^{4}} \frac{\partial^{2}B}{\partial \alpha^{2}} - \frac{4}{R^{3}} \frac{\partial^{3}B}{\partial \alpha^{2}\partial R} - \frac{2}{R^{2}} \frac{\partial^{2}B}{\partial R^{2}} + \frac{2}{R^{3}} \frac{\partial B}{\partial R} + \frac{1}{R^{2}} \frac{\partial^{4}B}{\partial \alpha^{2}\partial R^{2}} + \frac{1}{R} \frac{\partial^{3}B}{\partial R^{3}}$$

$$\frac{\partial^{4}B}{\partial X^{2}Y^{3}} = \frac{1}{R} \frac{\partial^{4}B}{\partial \alpha \partial R^{3}} - \frac{3}{R^{2}} \frac{\partial^{3}B}{\partial \alpha \partial R^{2}} + \frac{6}{R^{3}} \frac{\partial^{2}B}{\partial \alpha \partial R} - \frac{6}{R^{4}} \frac{\partial^{4}B}{\partial \alpha^{4}}$$

$$\frac{\partial^{4}B}{\partial Y^{4}} = \frac{\partial^{4}B}{\partial R^{4}}$$

$$\frac{\partial^{4}B}{\partial R^{4}} = \frac{\partial^{4}B}{\partial R^{4}}$$

NOTE: If a particle goes beyond the limits of the field map, the field and its derivatives will be extrapolated by means of the same calculations, from the border grid which is the closest to the actual position of the particle. Its flag EX is given the value -1 (see section 4.6.8).

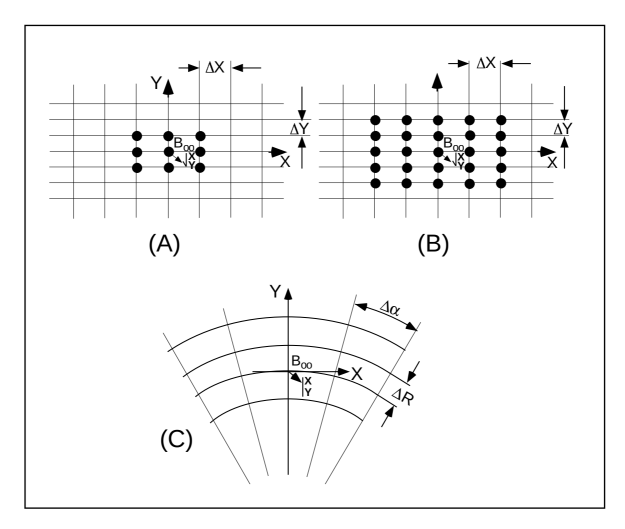


Figure 3: Mesh in the (X,Y) plane in Cartesian coordinates. The grid is centered

on the node which is closest to the actual position of the particle.

A: 9-point interpolation grid.

B: 25-point interpolation grid.

C: Mesh in the (X, Y) plane in polar coordinates.

1.4.3 Arbitrary 2-D Map, no Symmetry

The map is assumed to describe the field $\vec{B}(B_X, B_Y, B_Z)$ in the (X, Y) plane at elevation Z_0 . It provides the components $B_{X,ij}$, $B_{Y,ij}$, $B_{Z,ij}$ at each node (i,j) of a 2-D mesh.

The value of \vec{B} and its derivatives at the projection (X,Y,Z_0) of the actual position (X,Y,Z) of a particle is obtained by means of (parameter IORDRE in keyword data list - see for instance MAP2D, MAP2D-E) either a second degree polynomial interpolation from a 3×3 points grid (IOR-DRE=2), or a fourth degree polynomial interpolation from a 5×5 points grid (IORDRE=4), centered at the node (i,j) closest to the position (X,Y).

To second order for instance

$$B_{\ell}(X, Y, Z_0) = A_{00} + A_{10}X + A_{01}Y + A_{20}X^2 + A_{11}XY + A_{02}Y^2$$
(1.4.11)

where B_{ℓ} stands for any of the three components B_X , B_Y or B_Z . Differentiating then gives the derivatives

$$\frac{\partial B_{\ell}}{\partial X}(X, Y, Z_0) = A_{10} + 2A_{20}X + A_{11}Y$$

$$\frac{\partial^2 B_{\ell}}{\partial X \partial Y}(X, Y, Z_0) = A_{11}$$
etc.
$$(1.4.12)$$

Then follows the procedure of extrapolation from (X, Y, Z_0) to the actual position (X, Y, Z). No special symmetry is assumed, which allows the treatment of arbitrary field distribution (e.g., solenoid, helical snake).

1.4.4 3-D Field Map

The vector field $\vec{B}(X,Y,Z)$ and its derivatives necessary for the calculation of position and velocity of the particle are now defined using a 3-D field map, through second degree polynomial interpolation

$$B_{\ell}(X,Y,Z) = A_{000} + A_{100}X + A_{010}Y + A_{001}Z + A_{200}X^{2} + A_{020}Y^{2} + A_{002}Z^{2} + A_{110}XY + A_{101}XZ + A_{011}YZ$$

$$(1.4.13)$$

 B_{ℓ} stands for any of the three components, B_X , B_Y or B_Z . By differentiation of B_{ℓ} one gets

$$\frac{\partial B_{\ell}}{\partial X} = A_{100} + 2A_{200}X + A_{110}Y + A_{101}Z$$

$$\frac{\partial^2 B_{\ell}}{\partial X^2} = 2A_{200}$$
(1.4.14)

and so on for first and second order derivatives with respect to X, Y or Z.

The interpolation involves a $3 \times 3 \times 3$ -point parallelipipedic grid (Fig. 4), the origin of which is positioned at the node of the 3-D field map which is closest to the actual position of the particle.

Let B_{ijk}^{ℓ} be the value of the — measured or computed — magnetic field at each one of the 27 nodes of the 3-D grid (B^{ℓ} stands for B_X , B_Y or B_Z), and $B_{\ell}(X,Y,Z)$ be the value at a position (X,Y,Z) with respect to the central node of the 3-D grid. Thus, any coefficient A_i of the polynomial expansion of B_{ℓ} is obtained by means of expressions that minimize, with respect to A_i , the sum

$$S = \sum_{ijk} (B_{\ell}(X, Y, Z) - B_{ijk}^{\ell})^{2}$$
(1.4.15)

where the indices i, j and k take the values -1, 0 or +1 so as to sweep the 3-D grid. The source code contains the explicit analytical expressions of the coefficients A_{ijk} solutions of the normal equations $\partial S/\partial A_{ijk}=0$.

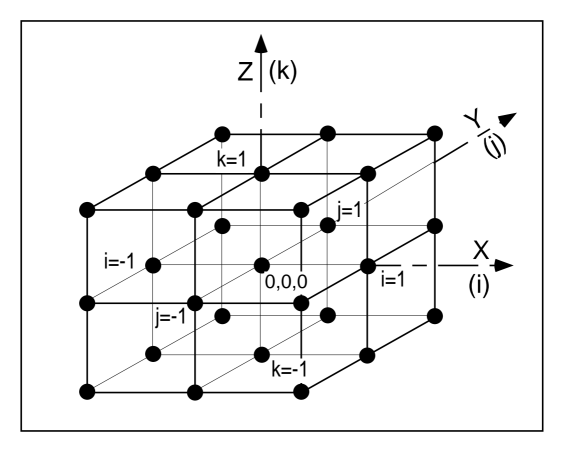


Figure 4: A 3-D 27-point grid is used for interpolation of \vec{B} and its derivatives up to second order. The central node of the grid (i=j=k=0) is the closest to the actual position of the particle.

2 SPIN TRACKING [10]

The depolarization of a particle beam travelling in a magnetic field \vec{b} takes its origin in the spin precession undergone by each particle. This motion of the spin \vec{S} is governed by the Thomas-BMT first order differential equation [11]

$$\frac{d\vec{S}}{dt} = \frac{q}{m}\vec{S} \times \vec{\Omega} \tag{2.16}$$

where

$$\vec{\Omega} = (1 + \gamma G)\vec{b} + G(1 - \gamma)\vec{b}_{//} \tag{2.17}$$

 q,m,γ and G are respectively the charge, mass, Lorentz relativistic factor, and anomalous magnetic moment of the particle. $\vec{b}_{/\!\!/}$ is the component of \vec{b} which is parallel to the velocity \vec{v} of the particle. These equations are normalized by introducing the same notation as previously. Let $b=\parallel\vec{b}\parallel$ and $v=\parallel\vec{v}\parallel$; ds=vdt is the differential path, $\frac{\gamma mv}{q}=B\rho$ is the rigidity of the particle; $\vec{S}'=\frac{d\vec{S}}{ds}=\frac{1}{v}\frac{d\vec{S}}{dt}$ is the derivative of the spin with respect to the path.

Introducing also $\vec{B} = \frac{\vec{b}}{B\rho}$, $\vec{B}_{\parallel} = \frac{\vec{b}_{\parallel}}{B\rho}$ and $\vec{\omega} = \frac{\vec{\Omega}}{B\rho} = (1 + \gamma G)\vec{B} + G(1 - \gamma)\vec{B}_{\parallel}$ (2.18)

eq. (2.16) can be re-written in a normalized way

$$\vec{S}' = \vec{S} \times \vec{\omega} \tag{2.19}$$

This equation is then solved in the same way as the reduced Lorentz equation (1.2.3). From the values of the magnetic factor $\vec{\omega}(M_0)$ and the spin $\vec{S}(M_0)$ of the particle at position M_0 of its trajectory, the spin $\vec{S}(M_1)$ at position M_1 , following a displacement Δs (fig. 2), is obtained from truncated Taylor expansion

$$\vec{S}(M_1) \approx \vec{S}(M_0) + \frac{d\vec{S}}{ds}(M_0)\Delta s + \frac{d^2\vec{S}}{ds^2}(M_0)\frac{\Delta s^2}{2} + \frac{d^3\vec{S}}{ds^3}(M_0)\frac{\Delta s^3}{3!} + \frac{d^4\vec{S}}{ds^4}(M_0)\frac{\Delta s^4}{4!}$$
(2.20)

The derivatives $\vec{S}^{(n)} = \frac{d^n \vec{S}}{ds^n}$ of \vec{S} at M_0 are obtained by differentiating eq. (2.19)

$$\vec{S}' = \vec{S} \times \vec{\omega}$$

$$\vec{S}'' = \vec{S}' \times \vec{\omega} + \vec{S} \times \vec{\omega}'$$

$$\vec{S}''' = \vec{S}'' \times \vec{\omega} + 2\vec{S}' \times \vec{\omega}' + \vec{S} \times \vec{\omega}''$$

$$\vec{S}'''' = \vec{S}''' \times \vec{\omega} + 3\vec{S}'' \times \vec{\omega}' + 3\vec{S}' \times \vec{\omega}'' + \vec{S} \times \vec{\omega}'''$$
(2.21)

where the derivatives $\vec{\omega}^{(n)}$ are obtained from eq. (2.18).

36 2 SPIN TRACKING

The last point consists in getting $\vec{B}_{/\!\!/}$ and its derivatives. This can be done in the following way. Let $\vec{u} = \frac{\vec{v}}{v}$ be the normalized velocity of the particle, then,

$$\vec{B}_{/\!/} = (\vec{B} \cdot \vec{u}) \vec{u}$$

$$\vec{B}_{/\!/}' = (\vec{B}' \cdot \vec{u} + \vec{B} \cdot \vec{u}') \vec{u} + (\vec{B} \cdot \vec{u}) \vec{u}'$$

$$\vec{B}_{/\!/}'' = (\vec{B}'' \cdot \vec{u} + 2\vec{B}' \cdot \vec{u}' + \vec{B} \cdot \vec{u}'') \vec{u} + 2(\vec{B}' \cdot \vec{u} + \vec{B} \cdot \vec{u}') \vec{u}' + (\vec{B} \cdot \vec{u}) \vec{u}''$$
etc. (2.22)

The quantities \vec{u} , \vec{B} and their n-th derivatives as involved in these equations are picked up from eqs. (1.2.7, 1.2.8).

3 SYNCHROTRON RADIATION

zgoubi allows the simulation of two types of synchrotron radiation (SR) related effects namely, on the one hand energy loss by stochastic emission of photon and the ensuing perturbation on particle dynamics and, on the other hand calculation of the radiated spectral-angular energy densities as observed in the lab.

3.1 Energy loss and related dynamical effects

Most of the content in the present section is drawn from Refs. [12].

Given a particle wandering in the magnetic field of an arbitrary optical element or field map, **zgoubi** computes the energy loss undergone, and its effect on the particle motion. The energy loss is calculated in a classical manner, by calling upon two random processes that accompany the emission of a photon namely,

- the probability of emission,
- the energy of the photon.

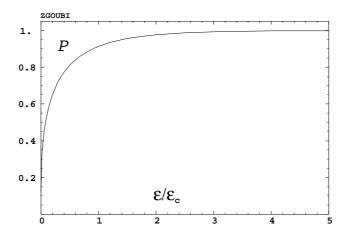


Figure 5: Cumulative distribution $\mathcal{P}(\epsilon/\epsilon_c)$.

The effects on the dynamic of the emitting particle is either limited to the alteration of the energy, or extended to angular kick effect, following user requested working options; particle position is supposed not to change upon emission of a photon. These calculations and ensuing dynamics corrections are performed after each integration step. In a practical manner, this means every centimer or tens of centimers in smoothly varying magnetic fields.

Main aspects of the method are developped in the following.

Probability of emission of a photon

Given that the number of photons emitted within a step Δs can be very low (units or fractions of unit)¹ a Poisson probability law

$$p(k) = \frac{\lambda^{-k}}{k!} \exp(-k) \tag{3.1.1}$$

is considered. k is the number of photons emitted over a $\Delta\theta$ (circular) arc of trajectory such that, the mean number of photons per radian expresses as²

$$\lambda = \frac{20er_0}{8\bar{h}\sqrt{3}}\beta^2 B\rho \Delta s \tag{3.1.2}$$

where $r_0 = e^2/4\pi\epsilon_0 m_0 c^2$ is the classical radius of the particle of rest-mass m_0 , e is the elementary charge, $\bar{h} = h/2\pi$, h is the Planck constant, $\beta = v/c$, $B\rho$ is the particle stiffness. λ is evaluated at each integration step from the current values β , $B\rho$ and Δs , then a value of k is drawn by a rejection method [41, routine POIDEV].

Energy of the photons

These k photons are assigned energies $\epsilon = h\nu$ at random, in the following way. The cumulative distribution of the energy probability law $p(\epsilon/\epsilon_c)d\epsilon/\epsilon_c$ writes

$$\mathcal{P}(\epsilon/\epsilon_c) = \frac{3}{5\pi} \int_0^{\epsilon/\epsilon_c} \int_{\epsilon/\epsilon_c}^{\infty} K_{5/3}(x) dx$$
 (3.1.3)

where $K_5/3$ is a modified Bessel function and, $\epsilon_c = \bar{h}\omega_c$ with $\omega_c = 2\pi 3\gamma^3 c/2\rho$ being the critical frequency of the radiation in constant field with bending radius ρ ; ω_c is evaluated at each integration step from the current values γ and ρ , in other words, this energy loss calculation assumes constant magnetic field 3 over the trajectory arc Δs . In the low frequency region $(\epsilon/\epsilon_c \ll 1)$ it can be approximated by

$$\mathcal{P}(\epsilon/\epsilon_c) = \frac{12\sqrt{3}}{52^{1/3}\Gamma(\frac{1}{2})} (\frac{\epsilon}{\epsilon_c})^{1/3}$$
(3.1.4)

About 40 values of $\mathcal{P}(\epsilon/\epsilon_c)$ computed from eq. 3.1.3 [42], honnestly spread over a range $\epsilon/\epsilon_c \leq 10$ are tabulated in **zgoubi** source file (see figure). In order to get ϵ/ϵ_c , first a random value $0 < \mathcal{P} < 1$ is generated uniformly, then ϵ/ϵ_c is drawn either by simple inverse linear interpolation of the tabulated values if $\mathcal{P} > 0.26$ (corresponding to $\epsilon/\epsilon_c > 10^{-2}$), or, if $\mathcal{P} < 0.26$ from eq. 3.1.4 that directly gives $\epsilon/\epsilon_c = \left(\frac{5 \, 2^{1/3} \Gamma(\frac{1}{3})}{12\sqrt{3}\mathcal{P}}\right)^3$ with precision no less than 1% at $\mathcal{P} \to 0.26$.

 $^{^1}$ For instance, a 1 GeV electron will emit about 20.6 photons per radian ; an integration step size $\Delta s = 0.1$ m upon $\rho = 10$ m bending radius results in 0.2 photons per step.

²This leads for instance, in the case of electrons, to the classical formula $\lambda/\Delta\theta \approx 129.5 \text{E(GeV)}/2\pi \approx \gamma/94.9$.

³From a practical viewpoint, note that the value of the magnetic field first computed for a one-step push of the particle (eqs. 1.2.4,1.2.7) is next used to obtain ρ and perform SR loss corrections afterwards.

Upon request of SR loss tracking, several optical elements that contain dipole magnetic field component (*e.g.*, *MULTIPOL*) provide a printouot of various quantities related to SR emission, as drawn from classical theoretical expressions, such as for instance,

- energy loss per particle $\Delta E(eV) = \frac{2}{3}r_0c\gamma^3B(T)\Delta\theta$, (B is the dipole field, exclusive of any other multipole component or non-linearity in the magnet; $\Delta\theta$ is the total deviation as calculated from B, the magnet length, and the reference rigidity BORO(as defined with, e.g., OBJET)
- energy $\epsilon_c(eV)=\frac{3\gamma^3c}{2\rho}\frac{\bar{h}}{e}$, with $\rho=BORO/B$
- energy of radiated photons $<\epsilon>=\frac{8}{15\sqrt{3}}\epsilon_c,$
- r.m.s. energy of radiated photons $\epsilon_{rms} = 0.5591\epsilon_c$,
- number of radiated photons per particle $N=\Delta E/<\epsilon>$.

This is done in order to facilitate verifications, since on the other hand statistics regarding those values are drawn from the tracking and printed upon use of the dedicated keyword SYNPRNL.

Finally, upon user's request as well, SR loss can be limited to particular classes of optical elements, for instance dipole fields alone, or dipole + quadrupole magnets, etc. These tricks are made available in order to permit deeper insight, or easier comparison with other codes, for instance.

3.2 Spectral-angular radiated densities

Most of the content in the present section is drawn from Refs. [13, 14].

The ray-tracing procedures provide the ingredients necessary for the determination of the electric field radiated by the particle subject to acceleration, as shown in Fig. 6 (section 3.2.1). This allows calculation⁴ of spectral-angular densities radiated by particles in magnetic fields (section 3.2.2).

3.2.1 Calculation of the radiated electric field

The expression for the radiated electric field $\vec{\mathcal{E}}(\vec{n},\tau)$ as seen by the observer in the long distance approximation is [15]

$$\vec{\mathcal{E}}(\vec{n},\tau) = \frac{q}{4\pi\varepsilon_0 c} \frac{\vec{n}(t) \times \left[\left(\vec{n}(t) - \vec{\beta}(t) \right) \times d\vec{\beta}/dt \right]}{r(t) \left(1 - \vec{n}(t) \cdot \vec{\beta}(t) \right)^3}$$
(3.2.1)

where t is the time in which the particle motion is described and τ is the observer time. Namely, when at position $\vec{r}(t)$ with respect to the observer [or as well at position $\vec{R}(t) = \vec{X} - \vec{r}(t)$ in the (O, x, y, z) frame] the particle emits a signal which reaches the observer at time τ , such that $\tau = t + r(t)/c$ where r(t)/c is the delay necessary for the signal to travel from the emission point to the observer, which also leads by differentiation to the well-known relation

⁴These procedures are for the moment implemented in the post-processor **zpop**

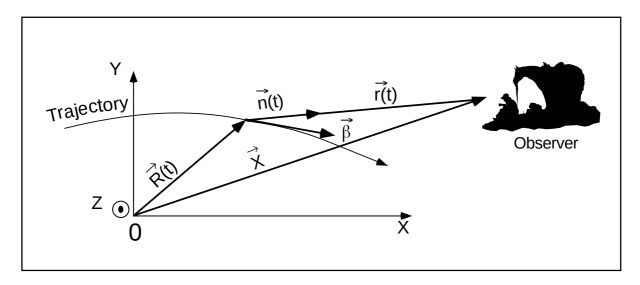


Figure 6: A scheme of the reference frame in **zgoubi** together with the vectors entering in the definition of the electric field radiated by the accelerated particle:

(x, y): horizontal plane; z: vertical axis.

 $\vec{R}(t)$ = particle position in the fixed frame (O, x, y, z);

 \vec{X} (time-independent) = position of the observer in the (O, x, y, z) frame;

 $\vec{r}(t) = \vec{X} - \vec{R}(t)$ = position of the particle with respect to the observer;

 $\vec{n}(t)$ = (normalized) direction of observation = $\vec{r}(t)/|\vec{r}(t)|$;

 $\vec{\beta}$ = normalized velocity vector of the particle $\vec{v}/c = (1/c)d\vec{R}/dt$.

$$d\tau/dt = 1 - \vec{n}(t) \cdot \vec{\beta}(t) \tag{3.2.2}$$

The vectors $\vec{R}(t)$ and $\vec{\beta}(t) = \frac{v}{c}\vec{u}$ (eq. 1.2.2) that describe the motion are obtained from the ray-tracing (eqs. 1.2.4). The acceleration is calculated from (eq. 1.2.1)

$$d\vec{\beta}/dt = (q/m) \vec{\beta}(t) \times \vec{b}(t) \tag{3.2.3}$$

Then, given the observer position \vec{X} in the fixed frame, it is possible to calculate

$$\vec{r}(t) = \vec{X} - R(t) \text{ and } \vec{n}(t) = \vec{r}(t)/|\vec{r}(t)|$$
 (3.2.4)

The calculation of $\vec{n} - \vec{\beta}$ and $1 - \vec{n} \cdot \vec{\beta}$

Owing to computer precision the crude computation of $\vec{n}-\vec{\beta}$ and $1-\vec{n}\cdot\vec{\beta}$ may lead to

$$\vec{n} - \vec{\beta} = 0$$
 and $1 - \vec{n} \cdot \vec{\beta} = 0$

since the preferred direction of observation is generally almost parallel to $\vec{\beta}$ (exactly parallel in the sense of computer precision), while $\beta \approx 1$ as soon as particle energies of a few hundred times the rest mass are concerned.

It is therefore necessary to express $\vec{n} - \vec{\beta}$ and $1 - \vec{n} \cdot \vec{\beta}$ in an adequate form for achieving accurate software computation.

The expression for \vec{n} is

$$\vec{n} = (n_x, n_y, n_z) = (\cos \psi \cos \phi, \cos \psi \sin \phi, \sin \psi)$$

$$= \left[1 - 2(\sin^2 \phi/2 + \sin^2 \psi/2) + 4\sin^2 \phi/2\sin^2 \psi/2, \sin \phi(1 - 2\sin^2 \psi/2), \sin \psi\right] \quad (3.2.5)$$

where ϕ and ψ are the observation angles, given by

$$\phi = \operatorname{Atg}\left(\frac{r_y}{r_x}\right) \text{ and } \psi = \operatorname{Atg}\left(\frac{r_z}{\sqrt{r_x^2 + r_y^2}}\right)$$
 (3.2.6)

with $\vec{r} = (r_x, r_y, r_z)$, while $\vec{\beta}$ can be written under the form

$$\vec{\beta} = (\beta_x, \beta_y, \beta_z) = \left[\sqrt{(\beta^2 - \beta_y^2 - \beta_z^2)}, \beta_y, \beta_z \right]$$

$$= \left[\sqrt{(1 - 1/\gamma^2 - \beta_y^2 - \beta_z^2)}, \beta_y, \beta_z \right] = (1 - a/2 + a^2/8 - a^3/16 + \dots, \beta_y, \beta_z)$$
 (3.2.7)

where $a=1/\gamma^2+\beta_y^2+\beta_z^2.$ This leads to

$$n_x = 1 - \varepsilon_x$$
 and $\beta_x = 1 - \xi_x$

with

$$\varepsilon_x = 2(\sin^2 \phi/2 + \sin^2 \psi/2) - 4\sin^2 \phi/2\sin^2 \psi/2$$

and

$$\xi_x = a/2 - a^2/8 + a^3/16 + \dots$$

All this provides, on the one hand,

$$\vec{n} - \vec{\beta} = (-\varepsilon_x + \xi_x, n_y - \beta_y, n_z - \beta_z), \qquad (3.2.8)$$

whose components are combinations of terms of the same order of magnitude (ε_x and $\xi_x \sim 1/\gamma^2$ while n_y, β_y, n_z and $\beta_z \sim 1/\gamma$) and, on the other hand,

$$1 - \vec{n} \cdot \vec{\beta} = \varepsilon_x + \xi_x - n_y \beta_y - n_z \beta_z - \varepsilon_x \xi_x , \qquad (3.2.9)$$

that combines terms of the same order of magnitude $(\varepsilon_x, \xi_x, n_y \beta_y)$ and $n_z \beta_z \sim 1/\gamma^2$, plus $\varepsilon_x \beta_x \sim 1/\gamma^4$.

The precision of these expressions is directly related to the order at which the series

$$\xi_x = a/2 - a^2/8 + a^3/16 + \dots \qquad (a = 1/\gamma^2 + \beta_y^2 + \beta_z^2)$$

is pushed, however the convergence is fast since $a \sim 1/\gamma^2 \ll 1$.

3.2.2 Calculation of the Fourier transform of the electric field

The Fourier transforms

$$FT_{\omega}[\vec{\mathcal{E}}(\tau)] = \int \vec{\mathcal{E}}(\tau)e^{-i\omega\tau}d\tau$$

of the σ and π electric field components provide the spectral angular energy density

$$\partial^3 W/\partial \phi \, \partial \psi \, \partial \omega = 2r^2 \left| FT_\omega \left(\vec{\mathcal{E}}(\tau) \right) \right|^2 / \mu_0 c$$
 (3.2.10)

They are calculated in a regular way, without use of FFT technics, namely from

$$FT_{\omega}\left[\vec{\mathcal{E}}(\tau)\right] \approx \sum \vec{\mathcal{E}}(\tau_k) e^{-i\omega\tau_k} \Delta\tau_k$$
 (3.2.11)

for two reasons. On the one hand, the number of integration steps Δs that define the trajectory (eqs. 1.2.4), is arbitrary and therefore in general not of order 2^n . On the other hand, the integration step defines a constant time differential element $\Delta t_k = \Delta s/\beta c$ which results in the observer differential time element $\Delta \tau_k$, which is also the differential element of the Fourier transform, being non-constant, since both are related by eq. 3.2.2 in which $\vec{\beta}$ and \vec{n} vary as a function of the integration step number k.

Another major point is that $\Delta \tau_k$ may reach drastically small values in the region of the central peak of the electric impulse emitted in a dipole $(1 - \vec{n}(t) \cdot \vec{\beta}(t) \to 1/2\gamma^2)$, whereas the total integrated time $\sum_{k=1}^{N} \Delta \tau_k$ may be several orders of magnitude larger. In terms of the physical phenomenon, the total duration of the electric field impulse as seen by the observer corresponds to the time delay $\sum_{k=1}^{N} \Delta \tau_k$ that separates photons emitted at the entrance of the magnet from photons emitted at the exit, but the significant part of it (in terms of energy density) which can be represented by the width

$$2\tau_c = \frac{2(1+\gamma^2\psi^2)^{3/2}}{3\gamma^3}\frac{2\rho}{c}$$
 of the radiation peak [16], is a very small fraction of $\sum_{k=1}^N \Delta \tau_k$.

The consequence is that, once again in relation with computer precision, the differential element $\Delta \tau_k$ involved in the computation of eq. 3.2.11 cannot be derived from such relation as $\Delta \tau_k = \sum_{k=1}^n \Delta \tau_k - \sum_{k=1}^{n-1} \Delta \tau_k$ but instead must be stored as such beforehand in the couorse of the ray-tracing process.

4 DESCRIPTION OF THE AVAILABLE PROCEDURES

4.1 Introduction

This chapter gives a detailed description of how the **zgoubi** procedures work, and their associated keywords. It has been split into several sections. Sections 4.2 to 4.5 explain the underlying content and functioning of all available keywords. Section 4.6 is dedicated to the description of some general procedures that may be accessed by means of special data or flags (such as negative integration steps), or through the available keywords (such as multiturn tracking with *REBELOTE*).

4.2 Definition of an Object

The description of the object, *i.e.*, initial coordinates of the beam, must be the first element of the input data to **zgoubi**.

Several types of automatically generated objects are available, as described in the following pages.

MCOBJET: Monte-Carlo generation of a 6-D object

MCOBJET generates a set of up to 10^4 random 6-D initial conditions. It can be used in conjunction with the keyword REBELOTE, which moreover allows generating an arbitrarily high number of initial conditions.

The first datum is the reference rigidity (negative value allowed)

$$BORO = \frac{p_0}{q} \text{ (kG.cm)}$$

Depending on the value of the next datum, KOBJ, the $IMAX (\le 10^4)$ particles have their initial random conditions Y, T, Z, P, X and D (relative momentum) generated on 3 different types of supports, as described below.

Next come the data

that specify the type of probability density for the 6 coordinates. KY, KT, KZ, KP, KX can take the following values :

- 1. uniform density, $p(x) = 1/2\delta x$ if $-\delta x \le x \le \delta x$, p(x) = 0 elsewhere,
- 2. Gaussian density, $p(x) = \frac{1}{\delta x \sqrt{2\pi}} e^{-\frac{x^2}{2\delta x^2}}$,
- 3. parabolic density, $p(x) = \frac{3}{4\delta x}(1 \frac{x^2}{\delta x^2})$ if $-\delta x \le x \le \delta x$, p(x) = 0 elsewhere.

KD can take the following values :

- 1. uniform density, $p(D) = 1/2\delta D$ if $-\delta D \le D \le \delta D$, p(D) = 0 elsewhere,
- 2. exponential density, $p(D) = N_0 \exp(C_0 + C_1 l + C_2 l^2 + C_3 l^3)$ with $0 \le l \le 1$ and $-\delta D \le D \le \delta D$,
- 3. p(D) is determined by a kinematic relation, namely, with T = horizontal angle, $D = \delta D * T$.

Next come the central value for the random sorting,

$$Y_0, T_0, Z_0, P_0, X_0, D_0$$

namely, the probability density laws p(x) (x = Y, T, Z, P or X) and p(D) described above apply to the variables $x - x_0$ ($\equiv Y - Y_0, T - T_0, ...$) and $D - D_0$ respectively. Negative value for D_0 is allowed (see section 4.6.9).

KOBJ = 1: Random generation of *MAX* particles in a hyper-window with widths (namely the half-extent for uniform or parabolic distributions (KY, KT, ... = 1 or 3), and the r.m.s. width for Gaussian distributions (KY, KT, ... = 2))

$$\delta Y$$
, δT , δZ , δP , δX , δD

Then follow the cut-off values, in units of the r.m.s. widths δY , δT , ... (used only for Gaussian distributions, KY, KT, ... = 2)

$$N_{\delta Y}$$
, $N_{\delta T}$, $N_{\delta Z}$, $N_{\delta P}$, $N_{\delta X}$, $N_{\delta D}$

The last data are the parameters

$$N_0, C_0, C_1, C_2, C_3$$

needed for generation of the D coordinate upon option KD=2 (unused if $KD=1,\ 3$) and a set of three integer seeds for initialization of random sequences,

$$IR1$$
, $IR2$, $IR3$ (all $\simeq 10^6$)

All particles generated by MCOBJET are tagged with a (non-S) character, for further statistic purposes (e.g., with HISTO and MCDESINT).

KOBJ = 2: Random generation of IY * IT * IZ * IP * IX * ID particles (maximum 10^4) in a hyper-grid. The input data are the number of bars in each coordinate

the spacing of the bars

the width of each bar

$$\delta Y$$
, δT , δZ , δP , δX , δD

the cut-offs, used with Gaussian densities (in units of the r.m.s. widths)

$$N_{\delta Y}$$
, $N_{\delta T}$, $N_{\delta Z}$, $N_{\delta P}$, $N_{\delta X}$, $N_{\delta D}$

This is illustrated in Fig. 7.

The last two sets of data in this option are the parameters

$$N_0, C_0, C_1, C_2, C_3$$

needed for generation of the D coordinate upon option KD= 2 (unused if KD= 1, 3) and a set of three integer seeds for initialization of random sequences, IR1, IR2, and IR3 (all $\simeq 10^6$).

All particles generated by MCOBJET are tagged with a (non-S) character, for further statistic purposes (see HISTO and MCDESINT).

KOBJ = 3: Distribution of *IMAX* particles inside a 6-D ellipsoid defined by the three sets of data (one set per 2-D phase-space)

$$\begin{array}{lll} \alpha_{Y}, & \beta_{Y}, & \frac{\varepsilon_{Y}}{\pi}, & N_{\varepsilon_{Y}} & [, & N_{\varepsilon_{Y}}', \text{ if } N_{\varepsilon_{Y}} < 0] \\ \alpha_{Z}, & \beta_{Z}, & \frac{\varepsilon_{Z}}{\pi}, & N_{\varepsilon_{Z}} & [, & N_{\varepsilon_{Z}}', \text{ if } N_{\varepsilon_{Z}} < 0] \\ \alpha_{X}, & \beta_{X}, & \frac{\varepsilon_{X}'}{\pi}, & N_{\varepsilon_{X}} & [, & N_{\varepsilon_{X}}', \text{ if } N_{\varepsilon_{X}} < 0] \end{array}$$

where α , β are the ellipse parameters and ε/π the emittance, corresponding to an elliptical frontier $\frac{1+\alpha_Y^2}{\beta_Y}Y^2+2\alpha_YYT+\beta_YT^2=\varepsilon_Y/\pi$ (idem for the (Z,P) or (X,D) planes). N_{ε_Y} , N_{ε_Z} and N_{ε_X} are the sorting cut-offs (used only for Gaussian distributions, KY,KT,...=2).

The sorting is uniform in surface (for KY=1, or KZ=1 or KX=1) or Gaussian (KY=2 or KZ=2), and so on, as described above. A uniform sorting has the ellipse above for support. A Gaussian sorting has the ellipse above for r.m.s. frontier, leading to $\sigma_Y=\sqrt{\beta_Y\varepsilon_Y/\pi}$,

$$\sigma_T = \sqrt{\frac{(1+\alpha_Y^2)}{\beta_Y}} \varepsilon_Y/\pi$$
, and similar relations for σ_Z , σ_X .

If N_{ε} is negative, thus the sorting fills the elliptical ring that extends from $|N_{\varepsilon}|$ to N'_{ε} (rather than the inner region determined by the N_{ε} cut-off, as addressed above).

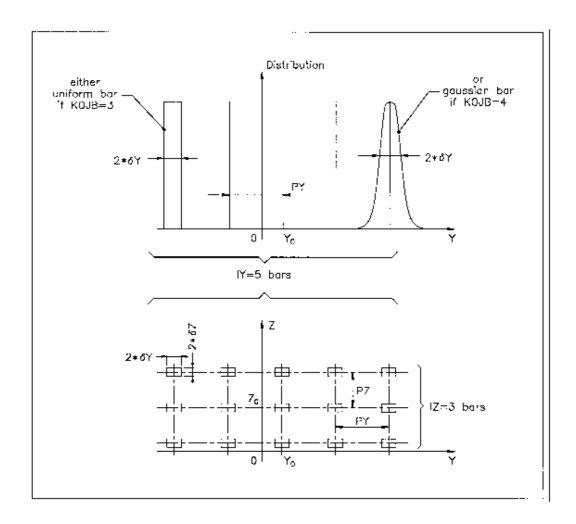


Figure 7: Scheme of the input parameters to MCOBJET when KOBJ=

3, 4

A : A distribution of the Y coordinate

B : A 2-D grid in (Y, Z) space.

OBJET: Generation of an object

OBJET is dedicated to the determination of the initial coordinates, in several ways.

The first datum is the reference rigidity (a negative value is allowed)

$$BORO = \frac{p_0}{q}$$

At the object, the beam is defined by a set of particles (maximum 10^4) with the initial conditions (Y, T, Z, P, X, D) where D is the relative momentum.

Depending on the value of the next datum *KOBJ*, these initial conditions may be generated in six different ways:

KOBJ = 1: Defines a grid in the Y, T, Z, P, X, D space. One gives the number of points desired,

(maximum 41 in each coordinate : $IY \le 41 \dots ID \le 41$ and such that $IY * IT * \dots * ID \le 10^4$) and the sampling size

zgoubi then generates $IY * IT * IZ * IP * IX * ID (\leq 10^4)$ initial conditions with the following coordinates

$$0, \pm PY, \pm 2 * PY, \dots, \pm IY/2 * PY, \\ 0, \pm PT, \pm 2 * PT, \dots, \pm IT/2 * PT, \\ 0, \pm PZ, \pm 2 * PZ, \dots, \pm IZ/2 * PZ, \\ 0, \pm PP, \pm 2 * PP, \dots, \pm IP/2 * PP, \\ 0, \pm PX, \pm 2 * PX, \dots, \pm IX/2 * PX, \\ 0, \pm PD, \pm 2 * PD, \dots, \pm ID/2 * PD,$$

In this option relative momenta will be classified automatically for the purpose of the use of *IMAGES* for momentum analysis.

The particles are tagged with an index IREP possibly indicating a symmetry with respect to the (X,Y) plane, as explained in option KOBJ=3. If two trajectories have mid-plane symmetry, only one will be ray-traced, while the other will be deduced using the mid-plane symmetries. This is done for the purpose of saving computing time. It may be incompatible with the use of some procedures $(e.g.\ MCDESINT)$, which involves random processes).

The last datum is the reference of the problem (YR, TR, ZR, PR, XR, DR). For instance the reference rigidity is DR*BORO, resulting in the rigidity of a particle of initial condition I*PD to be (DR+I*PD)*BORO.

KOBJ = **1.01**: Same as KOBJ = 1 except for the Z symmetry. The initial Z and P conditions are the following

0,
$$\pm PZ$$
, $\pm 2 * PZ$, ..., $\pm (IZ - 1) * PZ$, 0, $\pm PP$, $\pm 2 * PP$, ..., $\pm (IP - 1) * PP$,

This object results in shorter outputs/CPU-time when studying problems with Z symmetry.

KOBJ = 2: Next data: IMAX, IDMAX. Initial coordinates are entered explicitly for each trajectory. IMAX is the total number of particles ($IMAX \le 10^4$). These may be classified in groups of equal number for each value of momentum, in order to fulfill the requirements of image calculations by IMAGES. IDMAX is the number of groups of momenta. The following initial conditions defining a particle are specified for each one of the IMAX particles

$$Y$$
, T , Z , P , X , D , $'A'$

where D * BORO is the rigidity (negative value allowed) and 'A' is a (arbitrary) tagging character.

The last record **I**EX (I=1, **I**MAX) contains **I**MAX times either the string "1" (which indicates that the particle will be tracked) or the string "-9" (indicates that the particle should not be tracked).

This option KOBJ = 2 may be be useful for the definition of objects including kinematic effects.

KOBJ = 3: This option allows the reading of initial conditions from an external input file *FNAME*. The next three data lines are :

IT1, IT2, ITStep
IP1, IP2, IPStep
YF,TF,ZF,PF,SF,DPF,TiF,TAG
YR,TR,ZR,PR,SR,DPR,TiR
InitC

followed by the storage file name FNAME.

IT1, IT2, ITStep tell the code to read coordinates of particles number IT1 through IT2 by step ITStep.

IP1, IP2, IPStep tell the code to read coordinates belonging in the sole pass IP1 through IP2 by step IPStep. Indeed, IP2 > IP1 assumes prior filling of *FNAME* in the course of a run (*e.g.*, multiturn tracking) involving the keyword *REBELOTE*.

YF, TF, ZF, PF, SF, DPF, TiF are scaling factors whereas YR, TR, ZR, PR, SR, DPR, TiR are references added to the values of respectively Y, T, Z, P, S, DP as read from FNAME, so that any coordinate C = Y, T, Z... is changed into CF*C + CR. In addition a flag character TAG allows retaining only particles with identical tagging letter LET, unless TAG='** in which case it has no selection effect for instance TAG='S' can be used to retain only secondary particles following in-flight decay simulations.

If InitC= 1 ray-tracing starts from the current coordinates F(J, I), if InitC= 0 ray-tracing starts from the initial coordinates FO(J, I) as read from FNAME.

The file *FNAME* must be formatted in the appropriate manner. The following *FORTRAN* sequence is an instance, details and possible updates are to be found in the source file 'obj3.f':

```
OPEN (UNIT = NL, FILE = FNAME, STATUS = 'OLD')
    DO 1 I = 1, IMAX
                    {\tt READ} \  \, ({\tt NL}\,,100) \  \, {\tt LET} \  \, ({\tt I}) \, , \  \, {\tt IEX(I)} \, , \  \, ({\tt FO(J,I)}\,,{\tt J=1,6}) \, , \  \, ({\tt F(J,I)}\,,{\tt J=1,6}) \, , \  \, {\tt I} \, , \  \, {\tt IREP(I)} \, , \\  \, ({\tt REP(I)}\,,{\tt IREP(I)}\,,{\tt IREP(I)}\,,{\tt IREP(I)}\,, \  \, {\tt IREP(I)}\,, \  \, {\tt IREP(I)}\,, \  \, {\tt IREP(I)}\,, \  \, {\tt IREP(I)}\,, \\ \, ({\tt IREP(I)}\,,{\tt IREP(I)}\,,{\tt IREP(I)}\,,{\tt IREP(I)}\,, \  \, {\tt IREP(I)}\,
                                                 LET(I), IEX(I), -1.D0+FO(1,I), (FO(J,I), J=2, MXJ),
                                                      -1.D0+F(1,I),F(2,I),F(3,I),
                                                       (F(J,I),J=4,MXJ),ENEKI,
                                                   ID,I,IREP(I), SORT(I),D,D,D,D,RET(I),DPR(I),
                          D, D, D, BORO, IPASS, KLEY, LBL1, LBL2, NOEL
FORMAT(1X,
     100
                    LET(IT), KEX, 1.D0-FO(1,IT), (FO(J,IT), J=2,MXJ),
C1
                       1
                                                                          A1,1X,I2,1P,7E16.8,
                       1.D0-F(1,IT),(FO(J,IT),J=2,MXJ),
                                                                  /,3E24.16,
                                                                                                        TAR,
                  Z,P*1.D3,SAR,
                                                                   /,4E24.16,E16.8,
C4
                   KART, IT, IREP(IT), SORT(IT), X, BX, BY, BZ, RET(IT), DPR(IT),
                                                                    /,I1,2I6,7E16.8,
                                           EX,EY,EZ, BORO, IPASS, KLEY, (LABEL(NOEL,I),I=1,2),NOEL,4E16.8, I6,1X, A8,1X, 2A10, I5)
C5
                      5 /,4E16.8,
                                      CONTINUE
```

where the meaning of the parameters (apart from D=dummy real, ID=dummy integer) is the following

LET(I) : one-character string (for tagging)

IEX(I): flag, see KOBJ = 2

FO(1-6,I): coordinates D, Y, T, Z, P and path length of the particle

number I, at the origin. D * BORO = rigidity

F(1-6,I): idem, at the current position.

IREP is an index which indicates a symmetry with respect to median plane. For instance, if Z(I+1) = -Z(I), then normally IREP(I+1) = IREP(I). Consequently the coordinates of particle I+1 will not be obtained from ray-tracing but instead deduced without ray-tracing from those of particle I by simple symmetry. This results in gain of computing time.

KOBJ = 3 can be used directly for reading files filled by FAISCNL, FAISTORE. If more than 10^4 particles are to be read from a file, use $IMAX \le 10^4$ in conjunction with REBELOTE.

In this case (but not KOBJ = 3.01 or KOBJ = 3.02), particles will not have the reference charge and PARTICUL will not assign a mass or charge.

KOBJ = 3.01: Same as KOBJ = 3, except for the formatting of trajectory coordinate data in *FNAME* which is much simpler, namely, according to the following *FORTRAN* sequence

```
OPEN (UNIT = NL, FILE = FNAME, STATUS = 'OLD')

CONTINUE

READ (NL,*,END=10,ERR=99) Y, T, Z, P, S, D

GOTO 1

CALL ENDFIL

CALL ERREAD
```

KOBJ = 3.02: As for KOBJ = 3.01, except the format is

```
READ(NL,*) X,Y,Z,PX,PY,PZ
```

where PX, PY, and PZ, are the momenta in MeV/c. Note that DPR will be ignored in this case.

KOBJ = 3.03: As for KOBJ=3.01, except for the format :

where MASS is the mass in MeV/c and CHARGE is the charge divided by e. In this case, particles will not have the reference charge and PARTICUL will not assign a mass or charge.

KOBJ = 5: Mostly dedicated to the calculation of first order transfer matrix and various other optical parameters, using for instance *MATRIX* or *TWISS*. The input data are the stepsizes

The code generates 11 particles

$$0, \pm PY, \pm PT, \pm PZ, \pm PP, \pm PX, \pm PD$$

These values should be small enough, so that the paraxial ray approximation be valid.

The last data are the initial coordinates of the reference trajectory [normally (YR, TR, ZR, PR, XR, DR) = (0, 0, 0, 0, 0, 1)]. The reference rigidity is DR * BORO (negative value allowed).

KOBJ = **5.01**: Same as **KOBJ** = **5**, except for an additional data line giving initial beam ellipse parameters α_Y , β_Y , α_Z , β_Z , α_X , β_X , for further transport of these using *MATRIX*, or for possible use by the *FIT* procedure.

KOBJ = **5.NN**: Like **KOBJ** = **5**, except that instead of just one set of initial coordinates, the input file contains **NN** sets of initial coordinates. For example, to have 6 sets of initial coordinates, **KOBJ** should be **5.06**.

KOBJ = **6:** Mostly dedicated to the calculation of first, second and other higher order transfer coefficients and various other optical parameters, using for instance *MATRIX* or *TWISS*. The input data are the step sizes

to allow the building up of an object containing 61 particles. The last data are the initial coordinates of the reference trajectory [normally (YR,TR,ZR,PR,XR,DR)=(0,0,0,0,0,1)]. The reference rigidity of the beam is DR*BORO.

KOBJ = 7: Object with kinematics

The data and functioning are the same as for KOBJ = 1, except for the following

- *ID* is not used,
- PD is the kinematic coefficient, such that for particle number I, the initial relative momentum D_I is calculated from the initial angle T_I following

$$D_I = DR + PD * T_I$$

while T_I is in the range

$$0, \pm PT, \pm 2 * PT, \dots, \pm IT/2 * PT$$

as stated under KOBJ = 1

KOBJ = **8**: Generation of phase-space coordinates on ellipses.

The ellipses are defined by the three sets of data (one set per ellipse)

$$\begin{array}{lll} \alpha_Y, & \beta_Y, & \varepsilon_Y/\pi \\ \alpha_Z, & \beta_Z, & \varepsilon_Z/\pi \\ \alpha_X, & \beta_X, & \varepsilon_X/\pi \end{array}$$

where α , β are the ellipse parameters and ε/π is the emittance encompassed, corresponding to an ellipse with equation $\frac{1+\alpha_Y^2}{\beta_Y}Y^2+2\alpha_YYT+\beta_YT^2=\varepsilon_Y/\pi$ (idem for the (Z,P) or (X,D) planes).

The ellipses are centered respectively on (Y_0, T_0) , (Z_0, P_0) , (X_0, D_0) .

The number of samples per plane is respectively IX, IY, IZ. If that value is zero, the central value above is assigned.

OBJETA: Object from Monte-Carlo simulation of decay reaction [17]

This generator simulates the reactions

$$M_1 + M_2 \longrightarrow M_3 + M_4$$

and then

$$M_4 \longrightarrow M_5 + M_6$$

where M_1 is the mass of the incoming body; M_2 is the mass of the target; M_3 is an outgoing body; M_4 is the rest mass of the decaying body; M_5 and M_6 are decay products. Example:

$$p + d \longrightarrow^3 \text{He} + \eta$$

 $\eta \longrightarrow \mu^+ + \mu^-$

The first input data are the reference rigidity

$$BORO = \frac{p_0}{q}$$

an index IBODY which specifies the particle to be ray-traced, namely M3 (IBODY = 1), M5 (IBODY = 2) or M6 (IBODY = 3). In this last case, initial conditions for M6 must be generated by a first run of OBJETA with IBODY = 2; they are then stored in a buffer array, and restored as initial conditions at the next occurrence of OBJETA with IBODY = 3. Note that **zgoubi** by default assumes positively charged particles.

Another index, KOBJ specifies the type of distribution for the initial transverse coordinates Y, Z; namely either uniform (KOBJ= 1) or Gaussian (KOBJ= 2). The other three coordinates T, P and D are deduced from the kinematic of the reactions.

The next data are the number of particles to be generated, **MAX**, and the masses involved in the two previous reactions.

$$M_1, M_2, M_3, M_4, M_5, M_6$$

and the kinetic energy T_1 of the incoming body (M_1) .

Then one gives the central value of the distribution for each coordinate

$$Y_0, T_0, Z_0, P_0, D_0$$

and the width of the distribution around the central value

$$\delta Y$$
, δT , δZ , δP , δD

so that only those particles in the range

$$Y_0 - \delta Y \le Y \le Y_0 + \delta Y$$
 ... $D_0 - \delta D \le D \le D_0 + \delta D$

will be retained. The longitudinal initial coordinate is uniformly sorted in the range

$$-XL \le X_0 \le XL$$

The random sequences involved may be initialized with different values of the two integer seeds IR_1 and IR_2 ($\simeq 10^6$).

PARTICUL will not change the masses that are assigned by OBJETA.

4.3 Declaration of options

These options allow the control of procedures that affect certain functions of the code. Some options are normally declared right after the object definition (e.g. SPNTRK - spin tracking, MCDESINT - in-flight decay), others are normally declared at the end of the data pile (e.g. END - end of a problem, REBELOTE - for tracking more than 10^4 particles or for multi-turn tracking, FIT - fitting procedure).

BINARY: BINARY/FORMATTED data converter

This procedure translates field map data files from "BINARY" to "FORMATTED" – in the FORTRAN sense, or the other way.

The keyword is followed, next line, by NF.NCOL ($NF \le 9$, $NCOL \le 9$), the number of files to be translated and of data columns in the file. NCOL should be consistant with the following FORTRAN READ statement:

```
READ (unit=ln,*) (X7(I),I=1,NCOL)
```

The first data line in a field map file is a header, and contains NCOL reals as reference X coordinate, mesh step δX , reference Y coordinate, mesh step δY , and three others (see the FMAPW FORTRAN procedure for more datails).

Then follow, line per line, the NF names of the files to be translated.

Iff a file name begins with the prefix "B_" or "b_", it is presumed "binary", and hence converted to "formatted", and given the same name after suppression of the prefix "B_" or "b_". Conversely, *iff* the file name does not begin with "B_" or "b_", the file is presumed "formatted" and hence translated to "binary", and is given the same name after addition of the prefix "B_".

In its present state, the procedure *BINARY* only supports a limited number of output formatting, *e.g.* from *TOSCA* magnet code (see keyword *TOSCA*).

END or FIN: End of input data list; see FIN

The end of a problem, or of a set of several problems stacked in the data file, should be stated by means of the keywords *FIN* or *END*.

Any information following these keywords will be ignored.

FIT, FIT2: Fitting procedure

The keywords FIT, FIT2 allow the automatic adjustment of up to 20 variables, for fitting up to 20 constraints.

They are compatible with the use of (i.e., can be encompassed in) REBELOTE for successive FIT trials using various sets of parameters (option K = 22 in REBELOTE).

FIT has been implemented recently [18] and may have some advantages over the original method. The earlier FIT2 was drawn from the matrix transport code BETA [19]. One or the other may converge faster depending on the problem.

Any physical parameter of any element (i.e., keyword) may be varied. Available constraints are, amongst others: any of the 6×6 coefficients of the first order transfer matrix $[R_{ij}]$ as defined in the keyword MATRIX, and its horizontal $(R_{11}R_{22}-R_{12}R_{21})$ and vertical $(R_{33}R_{44}-R_{34}R_{43})$ determinants; horizontal and vertical tunes (if periodical structure); any of the $6\times 6\times 6$ coefficients of the second order array $[T_{ijk}]$ as defined in MATRIX; any of the 2×4 coefficients of the σ -matrix as defined by

$$[\sigma_{ij}] = \begin{pmatrix} \sigma_{11} & \sigma_{12} & & \\ \sigma_{21} & \sigma_{22} & & \\ & & \sigma_{33} & \sigma_{34} \\ & & \sigma_{43} & \sigma_{44} \end{pmatrix}$$

and any trajectory coordinates F(J, I) as defined in *OBJET* (I = particle number, J = coordinate number = 1 to 6 for respectively D, Y, T, Z, P or S =path length).

Tunes $\nu_{Y,Z}$ and periodic betatron functions $\beta_{Y,Z}$, $\alpha_{Y,Z}$, $\gamma_{Y,Z}$ are adjustable as well; they are defined by identification of the transfer matrix of the full optical structure, $[R_{ij}]$, with the form $Icos(2\pi\nu_{Y,Z}) + \frac{1}{2\pi} \left(\frac{1}{2\pi} \frac{1}{2\pi} \frac{1}{$

$$Jsin(2\pi\nu_{Y,Z})$$
, wherein $J=\begin{pmatrix} \alpha & \beta \\ -\gamma & -\alpha \end{pmatrix}$.

VARIABLES

The first input data in *FIT* are the number of variables *NV*, and for each one of them, the following parameters

IR = number of the varied element in the structure

IP = number of the physical parameter to be varied in this element

XC= coupling parameter. Normally XC=0. If $XC\neq 0$, coupling will occur (see below). followed by, either

DV = allowed relative range of variation of the physical parameter IP

or

[Vmin, Vmax] = allowed interval of variation of the physical parameter IP

Numbering of the elements (IR):

The elements (DIPOLE, QUADRUPO, etc.) are numbered following their sequence in the **zgoubi** input data file, for the purpose of the FIT procedure. The number of any element just identifies with its position in the data sequence. However, a simple way to get IR is to make a preliminary run: **zgoubi** will then print the whole structure into the file zgoubi.res with all elements numbered.

Numbering of the physical parameters (IP):

In the elements *DIPOLE*, *AIMANT* and *EBMULT*, *ELMULT*, *MULTIPOL*, the numbering of the physical parameters just follows their sequence, as it is shown here after for *DIPOLE-M*: the left column below represents the input data, the right one the corresponding numbering to be used for the *FIT* procedure.

Input data	Numbering for FIT
DIPOLE-M	
NFACE, IC, IL	1, 2, 3
IAMAX, IRMAX	4, 5
B_0, N, B, G	6, 7, 8, 9
AT, ACENT, RM, RMIN, RMAX	10, 11, 12, 13, 14
λ , ξ	15,16
$NC, C_0, C_1, C_2, C_3, C_4, C_5$ shift	17, 18, 19, 20, 21, 22, 23, 24
ω , θ , R_1 , U_1 , U_2 , R_2	25, 26, 27, 28, 29, 30
etc.	etc.

Parameters in SCALING also have a specific numbering, as follows.

Input data	Numbering for FIT
SCALING	
IOPT, NFAM	
NAMEF	
NT_1	
$SCL(I), I = 1, NT_1$	$10 [,, 10 + NT_1]$
$TIM(I), I = 1, NT_1$	$10[,, 10 + 2 * NT_1]$
NAMEF	
NT_2	
$SCL(I), I = 1, NT_2$	$20 [,, 20 + NT_2]$
$TIM(I), I = 1, NT_2$	$20[,, 20 + 2 * NT_2]$
•••	
etc. up to <i>NFAM</i>	etc.

For all other keywords, the parameters are numbered in the following way

Numbering for FIT
1, 2, 3,
10, 11, 12, 13,
a line of comments is skipped
20, 21, 22,
30, 31, 32, 33,

The examples of QUADRUPO (quadrupole) and TOSCA (Cartesian or cylindrical mesh field map) are given below.

Numbering for FIT
1
10, 11, 12
20, 21
30, 31, 32, 33, 34, 35, 36
40, 41
50, 51, 52, 53, 54, 55, 56
60
70, 71, 72, 73
1, 2
10, 11 [, 12, 13]
This is text
20, 21, 22, 23
This is text
30, 31, 32, 33 [34, 35, 36 [, 37, 38, 39] if $ID \ge 2$]
40
50
60,61,62,63

Coupled variables (XC)

Coupling a variable parameter to any other parameter in the structure is possible. This is done by giving XC a value of the form $r \cdot ppp$ where the integer part r is the number of the coupled element in the structure (equivalent to IR, see above), and the decimal part ppp is the number of its parameter of concern (equivalent to IP, see above) (if the parameter number is in the range 1, 2, ..., 9 (resp. 10, 11, ... 19 or 100, ...), then ppp must take the form 00p (resp. 0pp, ppp)). For example, $XC = 20 \cdot 010$ is a request for coupling with the parameter number 10 of element number 20 of the structure, while $XC = 20 \cdot 100$ is a request for coupling with the parameter number 100 of element 20.

An element of the structure which is coupled (by means of $XC \neq 0$) to a variable declared in the data list of the *FIT* keyword, needs not appear as one of the *NV* variables in that data list (this would be redundant information).

XC can be either positive or negative. If XC>0, then the coupled parameter will be given the same value as the variable parameter (for example, symmetric quadrupoles in a lens triplet will be given the same field). If XC<0, then the coupled parameter will be given a variation opposite to that of the variable, so that the sum of the two parameters stays constant (for example, an optical element can be shifted while preserving the length of the structure, by coupling together its upstream and downstream drift spaces).

Variation range (DV)

For a parameter IP of initial value p, the FIT procedure is allowed to explore the range $p(1 \pm DV)$.

IC = type of constraint (see table below).

 $I, J = \text{constraint } (i.e., R_{ij}, \text{determinant, tune }; T_{ijk}; \sigma_{ij}; \text{trajectory}$

#I and coordinate #J)

IR = number of the element in the **zgoubi** input data file, right

after which the constraint applies

V = desired value of the constraint

W =weight of the constraint (smaller W for higher weight)

CONSTRAINTS

The next input data in *FIT* are the number of constraints, *NC*, and for each one of them the following parameters.

IC=0: The coefficients σ_{11} (σ_{33}) = horizontal (vertical) beta values and σ_{22} (σ_{44}) = horizontal (vertical) derivatives ($\alpha = -\beta'/2$) are obtained by transport of their initial values at line start as introduced using for instance *OBJET*, *KOBJ=5.1*.

IC=0.1: Periodic optical functions: $\sigma_{11}=\beta_Y, \sigma_{12}=\sigma_{21}=-\alpha_Y, \sigma_{22}=\gamma_Y, \sigma_{33}=\beta_Z, \sigma_{34}=\sigma_{43}=-\alpha_Z, \sigma_{44}=\gamma_Z$; periodic dispersion: $\sigma_{16}=D_Y, \sigma_{26}=D_Y', \sigma_{36}=D_Z, \sigma_{46}=D_Z'$, all quantities derived by assuming periodic structure and identifying the first order transfer matrix with the form $Icos\mu+Jsin\mu$.

Type of constraint				Parameters defining the constraints					Object definition
Type of constraint	IC	I	J	Constraint	#	Parameter(s) values)	(recommended)
σ -matrix	0	1 - 6	1 - 6	σ_{IJ} $(\sigma_{11}=\beta_Y,\sigma_{12}=\sigma_{21}=\alpha_Y,\text{etc.})$					OBJET/KOBJ=5,6
Beam matrix (N=1-9 for MATRIX block 1-9))	0.N	1 - 6 7 8 9 10	1 - 6 any any any any	$\sigma_{IJ} (\sigma_{11} = \cos \mu_Y + \alpha_Y \sin \mu_Y, \text{ etc.})$ $Y \text{-tune} = \mu_Y / 2\pi$ $Z \text{-tune} = \mu_Z / 2\pi$ $\cos(\mu_Y)$ $\cos(\mu_Z)$					OBJET/KOBJ=5,6
First order parameters	1	$ \begin{array}{c c} 1 - 6 \\ 7 \\ 8 \end{array} $	$ \begin{array}{c c} 1-6 \\ i \\ j \end{array} $	Transport coeff. R_{IJ} $i \neq 8$: YY-determinant; i=8: YZ-det. $j \neq 7$: ZZ-determinant; j=7: ZY-det.					OBJET/KOBJ=5
Second order parameters	2	1-6	11 – 66	Transport coeff. $T_{I,j,k}$ $(j = [J/10], k = J - 10[J/10])$					OBJET/KOBJ=6
Trajectory coordinates	3.1 3.2 3.3 3.4	1 - MAX -1 -2 -3 1 - MAX 1 - MAX 1 - MAX 1 - MAX	$ \begin{array}{r} 1 - 7 \\ 1 - 7 \\ 1 - 7 \\ 1 - 7 \\ 1 - 7 \\ 1 - 7 \\ 1 - 7 \\ 1 - 7 \\ 1 - 7 \\ \end{array} $	$F(J,I) < F(J,i) >_{i=1,\text{MAX}} \\ Sup(F(J,i))_{i=1,\text{MAX}} \\ Dist F(J,I) _{i=1,I2,dI} \\ F(J,I) - FO(J,I) \\ F(J,I) + FO(J,I) \\ \min. \ (1) \ \text{or max.} \ (2) \ \text{value of} \ F(J,I) \\ F(J,I) - F(J,K) \ \ (K=1-\text{MAX})$	3 1 1	11 1-2 K	12	dI	[MC]OBJET
Matched ellipse parameters	4	1 - 6	1 – 6	σ_{IJ} ($\sigma_{11}=\beta_Y, \sigma_{12}=\sigma_{21}=\alpha_Y,$ etc.)					OBJET/KOBJ=8; MCOBJET/KOBJ=3
Number of particles	5	$ \begin{array}{r} -1 \\ 1 - 3 \\ 4 - 6 \end{array} $	any any any	$N_{survived}/ extbf{MAX} \ N_{in~\epsilon_{Y,Z,X}}/N_{survived} \ N_{in~best~\epsilon_{Y,Z,X},rms}/N_{survived}$	1	ϵ/π ϵ/π			OBJET MCOBJET MCOBJET
Spin	10 10.1	$1 - \mathbf{I}\mathbf{M}\mathbf{A}\mathbf{X}$ $1 - \mathbf{I}\mathbf{M}\mathbf{A}\mathbf{X}$	$ \begin{array}{c c} 1-4 \\ 1-3 \end{array} $	$ S_{X,Y,Z}(I), \vec{S}(I) \\ S_{X,Y,Z}(I) - SO_{X,Y,Z}(I) $					[MC]OBJET +SPNTRK

IC=1, 2: The coefficients R_{ij} and T_{ijk} are calculated following the procedures described in MATRIX, option IFOC=0. The fitting of the $[R_{ij}]$ matrix coefficients or determinants supposes the tracking of particles having initial coordinates sampled as described in MATRIX (these particles are normally defined with OBJET, KOBJ=5 or 6). The same is true for the T_{ijk} second order coefficients (Initial coordinates normally defined with OBJET, KOBJ=6).

IC=3: If 1 < I < IMAX then the value of coordinate type J (J=1,6 for respectively D, Y, T, Z, P, S) of particle number I (1 < I < IMAX) is constrained.

Case I = -1: the constraint is the mean value of coordinate of type J.

Case I = -1: the constraint is the max. value of coordinate of type J.

Case I = -1: the constraint is the J-distance for two different particles.

IC=3.1: Difference between final and initial J-coordinate of particle I (convenient e.g. for closed orbit search).

IC=4: The coefficients σ_{11} (σ_{33}) = horizontal (vertical) beta values and σ_{22} (σ_{44}) = horizontal (vertical) derivatives ($\alpha = -\beta'/2$) are derived from an ellipse match of the current particle population (as generated for instance using MCOB-JET, KOBJ=3).

The fitting of the $[\sigma_{ij}]$ coefficients supposes the tracking of a relevant population of particles within an adequate emittance.

IC=5: If I=-1 then the constraint value is the ratio of particles still on the run. If $I \ge 1$ then the constraint value is the ratio of particles encompassed within a given I-type (I=1-3 for respectively Y, Z, D) phase-space surface.

IC=10: If 1 < I < IMAX then the value of coordinate type J (J=1,3 for respectively S_X, S_Y, S_Z) of particle number I (1 < I < IMAX) is constrained.

IC=10.1: Difference between final and initial J-spin coordinate of particle I (convenient e.g. for closed orbit search).

OBJECT DEFINITION

Depending on the type of constraint (see Table), constraint calculations are performed either from transport coefficient calculation and in such case need OBJET with either KOBJ = 5 or KOBJ = 6, or from particle distributions and in this case need object definition using for instance OBJET with KOBJ = 8, MCOBJET with either KOBJ = 3.

THE FITTING METHODS [18, 19]

The FIT procedure has been implemented recently [18] and has various advantages over the original method, as converging speed.

The older FIT2 was drawn from the matrix transport code BETA [19]. The numerical procedure is a direct sequential minimization of the quadratic sum of all errors (i.e., differences between desired and actual values of the NC constraints), each normalized by its specified weight W (the smaller W, the stronger the constraint).

The step sizes for the variation of the physical parameters depend on their initial values, and cannot be accessed by the user. At each iteration, the optimum value of the step size, as well as the optimum direction of variation, is determined for each one of the *NV* variables. Then follows an iterative global variation of all *NV* variables, until the minimization fails which results in a next iteration on the optimization of the step sizes.

GASCAT: Gas scattering

Modification of particle momentum and velocity vector, performed at each integration step, under the effect of scattering by residual gas.

To be documented

MCDESINT: Monte-Carlo simulation of in-flight decay[20]

As soon as MCDESINT appears in a structure (normally, after OBJET or after CIBLE), in-flight decay simulation starts. It must be preceded by PARTICUL for the definition of mass M_1 and COM lifetime τ_1 . The two-body decay simulated is

$$1 \longrightarrow 2 + 3$$

The decay is isotropic in the center of mass. 1 is the incoming particle, with mass M_1 , momentum $p_1 = \gamma_1 M_1 \beta_1 c$ (relative momentum $D_1 = \frac{p_1}{q} \frac{1}{BORO}$ with BORO= reference rigidity, see OBJET), and position Y_1, Z_1 in the **zgoubi** frame. 2 and 3 are decay products with respective masses and momenta M_2, M_3 and $p_2 = \gamma_2 M_2 \beta_2 c$, $p_3 = \gamma_3 M_3 \beta_3 c$. The decay length s_1 of particle 1 is related to its center of mass lifetime τ_1 by

$$s_1 = c\tau_1 \sqrt{\gamma_1^2 - 1}$$

The path length s up to the decay point is then calculated from a random number $0 < R_1 \le 1$ by using the exponential decay formula

$$s = -s_1 \ell n R_1$$

After decay, particle 2 will be ray-traced with assumed positive charge, while particle 3 is discarded. Its scattering angles in the center of mass θ^* and ϕ are generated from two other random numbers R_2 and R_3 .

 ϕ is a relativistic invariant, and θ in the laboratory frame (Fig. 8) is given by

$$\tan \theta = \frac{1}{\gamma_1} \frac{\sin \theta^*}{\frac{\beta_1}{\beta_2^*} + \cos \theta^*}$$

 β_2^* and momentum p_2 are given by

$$\gamma_2^* = \frac{M_1^2 + M_2^2 - M_3^2}{2M_1M_2}$$

$$\beta_2^* = \left(1 - \frac{1}{\gamma^2}\right)^{1/2}$$

$$\gamma_2 = \gamma_1\gamma_2^* \left(1 + \beta_1\beta_2^* \cos \theta^*\right)$$

$$p_2 = M_2\sqrt{\gamma_2^2 - 1}$$

Finally, θ and ϕ are transformed into the angles T_2 and P_2 in the **zgoubi** frame, and the relative momentum takes the value $D_2 = \frac{p_2}{q} \frac{1}{BORO}$ (where BORO is the reference rigidity, see OBJET), while the starting position of M_2 is $Y_2 = Y_1$ and $Z_2 = Z_1$.

The decay simulation by **zgoubi** obeys the following procedures. In optical elements and field maps, after each integration step *XPAS*, the actual path length of the particle, F(6,I), is compared to its limit path length s. If s is passed, then the particle is considered as having decayed at $F(6,I) - \frac{XPAS}{2}$, at a position obtained by a linear translation from the position at F(6,I). [Presumably, the smaller *XPAS*, the smaller the error on position and angles at the decay point].

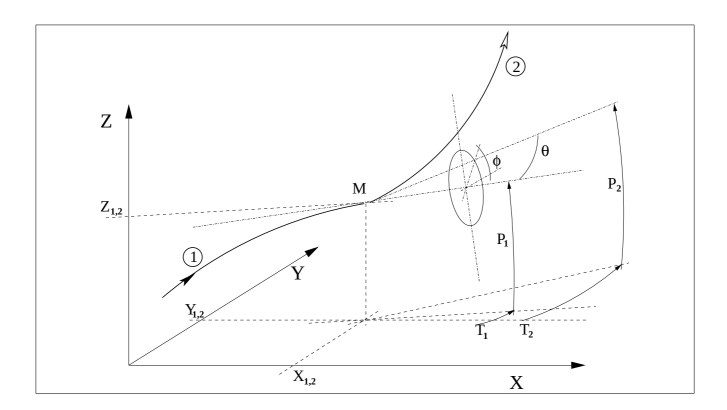


Figure 8: At position $M(X_1,Y_1,Z_1)$, particle 1 decays into 2 and 3; **zgoubi** then calculates the trajectory of 2, while 3 is discarded. θ and ϕ are the scattering angles of particle 2 relative to the direction of the incoming particle 1; they transform to T_2 and P_2 in **zgoubi** frame.

In ESL and CHANGREF, F(6, I) is compared to s at the end of the element. If the decay occurs inside the element, the particle is considered as having decayed at its actual limit path length s, and its coordinates at s are recalculated by translation.

The limit path length of all particles $(I=1, M\!A\!X)$ is stored in the array FDES(6,I), for further statistical purposes. For the same purpose (e.g., use of HISTO), any particle of type 2 (resulting from decay of 1) will be tagged with an S standing for "secondary". When a particle decays, its coordinates D, Y, T, Z, P at the decay point are stored in FDES(J,I), J=1, 5.

NOTE on negative drifts:

The use of negative drifts with MCDESINT is allowed and correct. For instance, negative drifts may occur in a structure for some of the particles when using CHANGREF (due to the Z-axis rotation or negative XCE), or when using DRIFT with XL < 0. Provision has been made to take it into account during the MCDESINT procedure, as follows.

If, due to a negative drift, a secondary particle reaches back the decay spot of the primary particle from which it originated, then that primary particle is regenerated with its original coordinates at that spot. Then the secondary particle is discarded while ray-tracing resumes in a regular way for the primary particle which is again susceptible of decay at the same time-of-flight. This procedure is made possible by prior storage of the coordinates of the primary particles (in array FDES(J, I)) each time a decay occurs.

Negative steps (XPAS < 0) in optical elements are not compatible with MCDESINT.

OPTICS: Write out optical functions

OPTICS normally appears next to object definition, it normally works in conjunction with element label(s). *OPTICS* causes the transport and write out, in zgoubi.res, of the 6×6 beam matrix, following options *KOPT* and 'label', below.

IF KOPT=0: Off

IF KOPT=1: Will transport the optical functions with initial values as specified in OBJET, option KOBJ=5.01.

Note: The initial values in *OBJET[KOBJ=5.01]* may be the periodic ones, as obtained, for instance, from a first run using *MATRIX[IFOC=11]*.

A second argument, 'label', allows

- if *label* = *all* : printing out, into zgoubi.res, after all keywords of the zgoubi.dat structure,
- otherwise, printing out at all keyword featuring $LABEL \equiv label$ as a first label (see section 4.6.3, page 153, regarding the labelling of keywords).

A third argument, IMP=1, will cause saving of the transported beta functions into file zgoubi.OPTICS.out.

ORDRE: Taylor expansions order

The position \vec{R} and velocity \vec{u} of a particle are obtained from Taylor expansions as described in eq. (1.2.4). By default, these expansions are up to the fourth order derivative of \vec{u} ,

$$\vec{R}_1 \approx \vec{R}_0 + \vec{u}_0 \Delta s + \dots + \vec{u}_0^{(4)} \frac{\Delta s^5}{5!}$$

 $\vec{u}_1 \approx \vec{u}_0 + \vec{u}_0' \Delta s + \dots + \vec{u}_0^{(4)} \frac{\Delta s^4}{4!}$

which corresponds to third order derivatives of \vec{B} , since (eq. (1.2.7))

$$\vec{u}^{(4)} = \vec{u}^{""} \times \vec{B} + 3\vec{u}^{"} \times \vec{B}^{"} + 3\vec{u}^{"} \times \vec{B}^{"} + \vec{u} \times \vec{B}^{""}$$

and to the third order derivatives of \vec{E} (eq. (1.2.11)) as well.

However \vec{B}''' , or \vec{E}''' , and higher order derivatives may be zero in second order type optical elements, for instance in a sharp edge quadrupole. Also, in several elements, no more than first and second order field derivatives are implemented in the code. One may also wish to fasten calculations by limiting the time-consuming calculation of lengthy (while possibly ineffective in terms of accuracy) Taylor expansions.

In that spirit, the purpose of *ORDRE*, option IO = 2-5, is to allow for expansions to the $\vec{u}_0^{(IO)}$ term in eq. 1.2.4. Default functionning is IO = 4.

Note the following:

As concerns the optical elements

QUADRUPO, SEXTUPOL, OCTUPOLE, DECAPOLE, DODECAPO, MULTIPOL, ELMULT, EBMULT

magnetic field derivatives (see eq. 1.2.8) have been installed in the code according to $\vec{u}_0^{(5)}$ developement order; it may not be as complete for some other optical elements, as well as for the possible electric field component whose field derivatives may not be provided to more than second order.

In electric optical elements field derivatives (eq. 1.2.13) are usually provided to no more than second order, which justifies saving computing time by not pushing Taylor expansions as high as $\vec{u}_0^{(5)}$.

NOTE: see also the option *IORDRE* in field map declarations (*DIPOLE-M*, *TOSCA*, etc.).

PARTICUL: Particle characteristics

PARTICUL allows the definition of several characteristics of the particles (mass, charge, gyromagnetic factor and lifetime in the center of mass), that are needed in various procedures as,

MCDESINT: mass, COM life-timeSPNTRK: mass, gyromagnetic factor

SRLOSS: mass, chargeSYNRAD: mass, chargeElectric and Electro-Magnetic elements: mass, charge

The declaration of *PARTICUL* must **precede** these keywords.

The charge is needed for computing energy gain in electric fields. The mass is needed for computing time of flight. If PARTICUL is not used, the charge defaults to +e and the mass defaults to zero.

PARTICUL sets a reference charge annd mass, but individual paritles have charges and masses as well. The reference charge relates $B\rho$ to the particle momentum $(B\rho=p/q)$; the particle charge relates the fields $(\vec{b}$ and \vec{e}) to the scaled fields $(\vec{B}=\vec{b}/(B\rho)$ and $\vec{E}=\vec{e}/(B\rho)$).

Most object definitions assign each particle the reference charge and mass (cases where they do not are documented). In cases where the reference charge and mass have been assigned to a particle, *PARTICUL* will change the particle charge and mass to the new reference charge and mass. For the few objects where each particle was assigned its own mass and charge, *PARTICUL* will not affect those values. In most cases (where all particles have the reference charge), the charge assigned by *PARTICUL* will not affect the results for systems having only magnetic fields.

REBELOTE: 'Do it again'

When REBELOTE is encountered in the input data file, the code execution jumps,

- either back to the beginning of the data file the default behavior,
- or (option K=99.1 or K=99.2) back to a particular *LABEL*.

Then NPASS-1 passes (from LABEL to REBELOTE) follow.

As to the last pass, number *NPASS+1*, there are two possibilities :

- either it also encompasses the whole *LABEL* to *REBELOTE* range,
- or, upon request (option K=99.2), execution may exit that pass at a particular second dedicated LABEL placed at arbitrary location between the first above mentionned LABEL, and REBELOTE. In both cases, following the end of this "multiple-pass" procedure, the execution continues from the keyword which follows REBELOTE, until 'END' is encountered.

REBELOTE can be used for Monte Carlo simulations when more than IMAX particles are to be tracked. In this case, when the following random procedures are used: MCOBJET, OBJETA, MCDESINT, SPNTRK (KSO = 5), their random seeds are not reset and independent statistics will add up.

REBELOTE can be used for multi-turn tracking in circular machines (e.g. Synchrotron accelerators, FFAGs, etc.). For instance, using option described K=99.2 above, a full "injection line + ring + extraction line" installation can be simulated - kicker firing and other magnet ramping can be simulated using SCALING.

Monte Carlo simulations: normally K=0. *NPASS* runs through the same structure will follow, resulting in the calculation of (1+NPASS)*IMAX trajectories.

Circular machines: normally K=99. NPASS turns in the same structure will follow, resulting in the tracking of MAX particles over 1+NPASS turns (Note: for the simulation of pulsed power supplies, synchrotron motion, and other Q-jump manipulation, see SCALING).

Using the double- LABEL method discussed above with option K=99.2, it is possible to encompass the ring between an injection line section (namely, with the element sequence of the latter extending from OBJET to the first LABEL), and an extraction line (its description will then follow REBELOTE), whereas the ring description extends from to the first LABEL to REBELOTE, with possible extraction, at the last pass, at the location of the second LABEL.

Output prints over NPASS+1 runs might result in a prohibitively big zgoubi.res file. They may be switched on/off by means of the option KWRIT=i.j, with i=1/0 respectively. The j flag commands printing pass number and some other information onto the video output, every 10^{j-1} turns if j>0; output is switched off if j=0.

REBELOTE also provides informations: statistical calculations and related data on particle decay (MCDESINT), spin tracking (SPNTRK), stopped particles (CHAMBR, COLLIMA), etc.

RESET: Reset counters and flags

Piling up problems in **zgoubi** input data file is allowed, with normally no particular precaution, except that each new problem must begin with a new object definition (with *MCOBJET*, *OBJET*, etc.). Nevertheless, when calling upon certain keywords, flags, counters or integrating procedures are involved. It may therefore be necessary to reset them. This is the purpose of *RESET* which normally appears right after the object definition and causes each problem to be treated as a new and independent one.

The keywords or procedures of concern and the effect of RESET are the following

CHAMBR : NOUT = number of stopped particles = 0; CHAMBR option switched off

COLLIMA: NOUT = number of stopped particles = 0

HISTO: Histograms are emptied

INTEG : *NRJ* = number of particles out of range = 0 (*INTEG* is the numerical integration subroutine;

NRJ is incremented when a particle goes out of a field map)

MCDESINT: Decay in flight option switched off

SCALING : Scaling options disabled

SPNTRK : Spin tracking option switched off

SCALING: Time scaling of power supplies and R.F.

SCALING acts as a function generator dedicated to varying fields in optical elements, or potentials in electrostatic devices, or frequency in *CAVITE*. It is normally intended to be declared right after the object definition, and used in conjunction with *REBELOTE*, for the simulation of multiturn tracking - possibly including acceleration cycles.

SCALING acts on families of elements, a family being designated by its name that coincides with the keyword of the corresponding element. For instance, declaring MULTIPOL as to be varied will result in the same timing law being applied to all MULTIPOL's in the **zgoubi** optical structure data file. Subsets can be selected by labeling keywords in the data file (section 4.6.3, page 153) and adding the corresponding LABEL('s) in the SCALING declarations (two LABEL's maximum). The family name of concern, as well as the field versus timing scaling law of that family (or frequency versus timing in the case of CAVITE) are given as input data to the keyword SCALING. Up to NF = 9 families can be declared as subject to a scaling law; a scaling law can be made of up to NT = 10 successive timings; between two successive timings, the variation law is linear.

An example of data formatting is given in the following.

SCALING		- Scaling
1 4		Active. $NF = 4$ families of elements are concerned, as listed below
QUADRUPO QFA QFB		- Quadrupoles labeled 'QFA' and Quadrupoles labeled 'QFB'
2		NT = 2 timings
18131.E-3	24176.E-3	The field increases (linearly) from 18131E-3* B_0 to 24176E-3* B_0
1	6379	from turn 1 to turn 6379
MULTIPOL QDA QDB		- Multipoles labeled 'QDA' and Multipoles labeled 'QDB'
2		
18131.E-3	24176.E-3	Fields increase from 18131E-3* B_i to 24176E-3* B_i ($\forall i = 1, 10$ poles)
1	6379	from turn 1 to turn 6379
BEND		- All BEND's (regardless of any LABEL)
2		
18131.E-3	24176.E-3	Same scaling
1	6379	
CAVITE		- Accelerating cavity
2		
1 1.22	1.33352	The synchronous rigidity $(B\rho)_s$ increases,
1 1200	6379	from $(B\rho)_{s_o}$ to $1.22*(B\rho)_{s_o}$ from turn 1 to 1200, and
		from $1.22 * (B\rho)_{s_o}$ to $1.33352 (B\rho)_{s_o}$ from turn 1200 to 6379

The timing is in unit of turns. In this example, TIMING = 1 to 6379 (turns). Therefore, at turn number N, B and B_i are updated in the following way. Let SCALE(TIMING = N) be the updating scale factor

$$SCALE(N) = 18.131 \frac{24.176 - 18.131}{1 + 6379 - 1} (N - 1)$$

and then

$$B(N) = SCALE(N)B_0$$

 $B_i(N) = SCALE(N)B_{i0}$

The R.F. frequency is computed using

$$f_{RF} = \frac{hc}{\mathcal{L}} \frac{q(B\rho)_s}{(q^2(B\rho)_s^2 + (Mc^2)^2)^{1/2}}$$

where the rigidity is updated in the following way. Let $(B\rho)_{s_o}$ be the initial rigidity (namely, $(B\rho)_{s_o}=BORO$ as defined in the keyword OBJET for instance). Then, at turn number N,

$$\begin{split} &\text{if } 1 \leq N \leq 1200 \text{ then, } \textit{SCALE}(N) = 1 + \frac{1.22 - 1}{1 + 1200 - 1} \left(N - 1\right) \\ &\text{if } 1200 \leq N \leq 6379 \text{ then, } \textit{SCALE}(N) = 1.22 + \frac{1.33352 - 1.22}{1 + 6379 - 1200} \left(N - 1200\right) \end{split}$$

and then,

$$(B\rho)_s(N) = SCALE(N) \cdot (B\rho)_{s_0}$$

from which value the calculations of $f_{RF}(N)$ follow.

NT can take negative values, then acting as an option switch (rather than giving number of timings), as follows:

- NT=-1: this is convenient for synchrotron acceleration using trivial RF law $f_{RF}=h/T_{rev}$. In this case the next two lines both contain a single data (as for NT=1), respectively the starting scaling factor value, and 1. The current field scaling factor can then be updated (computed) from the energy kick by the cavity if for instance CAVITE/IOPT=2 is used.
- \bullet NT=-2: this is convenient for reading an RF law for *CAVITE* from an external data file, including usage for acceleration in fixed field accelerators. To be documented.

Note: It may happen that some optical elements won't scale, for source code developement reasons. This should be paid attention to.

SPNTRK: Spin tracking

The keyword *SPNTRK* permits switching on the spin tracking option. It also permits the attribution of an initial spin component to each one of the *IMAX* particles of the beam, following a distribution that depends on the option index *KSO*. It must be preceded by *PARTICUL* for the definition of mass and gyromagnetic factor.

KSO = 1 (respectively 2, 3): the IMAX particles of the beam are given a longitudinal (1,0,0) spin component (respectively transverse horizontal (0,1,0), vertical (0,0,1)).

KSO = 4: initial spin components are entered explicitly for each one of the MAX particles of the beam.

KSO = **4.1**: three initial spin components S_X , $S_Y n S_Z$ are entered explicitly just once, they are then assigned to each one of the *IMAX* particles of the beam.

KSO = 5: random generation of *MAX* initial spin conditions as described in Fig. 9. Given a mean polarization axis (S) defined by its angles T_0 and P_0 , and a cone of angle A with respect to this axis, the *MAX* spins are sorted randomly in a Gaussian distribution

$$p(a) = \exp\left[-\frac{(A-a)^2}{2\delta A^2}\right]/\delta A\sqrt{2\pi}$$

and within a cylindrical uniform distribution around the (S) axis. Examples of simple distributions available by this mean are given in Fig. 10.

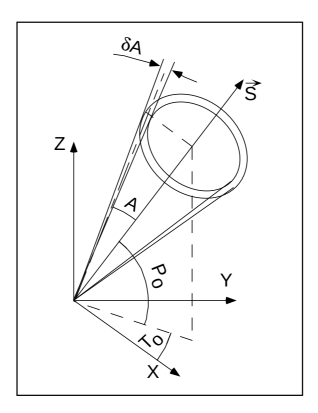


Figure 9: Spin distribution as obtained with option KSO = 5. The spins are distributed within an annular strip δA (standard deviation) at an angle A with respect to the axis of mean polarization (S) defined by T_0 and P_0 .

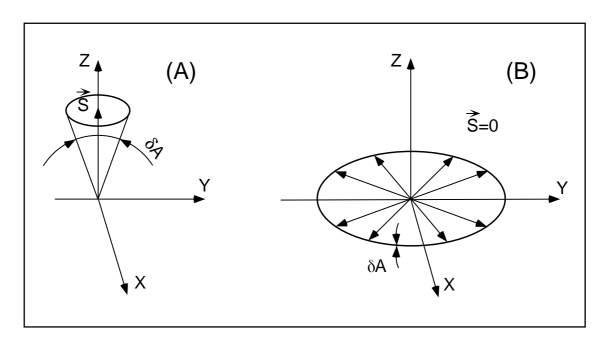


Figure 10: Examples of the use of KSO = 5.

A : Gaussian distribution around a mean vertical polarization axis, obtained with T_0 = arbitrary, $P_0 = \pi/2$, A = 0 and $\delta A \neq 0$.

B : Isotropic distribution in the median plane, obtained with $P_0=\pm\pi/2,\,A=\pi/2,$ and $\delta A=0.$

SRLOSS: Synchrotron radiation loss [12]

The keyword SRLOSS allows activating or stopping (option KSR=1,0 respectively) stepwise tracking of energy loss by emission of photons in magnetic fields and the ensuing particle energy perturbation, following the method described in section 3.1.

SRLOSS must be preceded by PARTICUL for defining mass and charge values as they enter in the definition of SR parameters.

Statistics on SR parameters are perform while tracking, results of which can be obtained by means of keyword SRPRNT.

SYNRAD: Synchrotron radiation spectral-angular densities

The keyword *SYNRAD* enables (or disables) the calculation of synchrotron radiation (SR) electric field and spectral angular energy density. It must be preceded by *PARTICUL* for defining mass and charge values, as they enter in the definition of SR parameters.

SYNRAD is supposed to appear a first time at the location where SR calculations should start, with the first data KSR set to 1. It results in on-line storage of the electric field vector and other relevant quantities in zgoubi.sre, as step by step integration proceeds. The observer position (XO, YO, ZO) is specified next to KSR.

Data stored in zgoubi.sre:

```
(ELx, ELy, ELz): \text{ electric field vector } \vec{\mathcal{E}} \text{ (eq. 3.2.1)} (btx, bty, btz) = \vec{\beta} = \frac{1}{c} \times \text{ particle velocity} (gx, gy, gz) = \frac{d\vec{\beta}}{dt} = \text{particle acceleration (eq. 3.2.3)} \Delta \tau = \text{observer time increment (eq. 3.2.2)} t' = \tau - r(t')/c = \text{retarded (particle) time} (rtx, rty, rtz) : \vec{R}(t), \text{ particle to observer vector (eq. 3.2.4)} (x, y, z) = \text{particle coordinates} \Delta s = \text{step size in the magnet (fig. 2)} NS = \text{step number} I = \text{particle number} LET(I) = \text{tagging letter}  \textit{EX}(I) = \text{stop flag (see section 4.6.8)}
```

SYNRAD is supposed to appear a second time at the location where SR calculations should stop, with KSR set to 2. It results in the output of the angular energy density $\int_{\nu_1}^{\nu_2} \partial^3 W/\partial\phi \,\partial\psi \,\partial\nu$ (eq. 3.2.11) as calculated from the Fourier transform of the electric field (eq. 3.2.11). The spectral range of interest and frequency sampling (ν_1, ν_2, N) are specified next to KSR.

Note that KSR = 0 followed by a dummy line of data allows temporary inhibition of SR procedures.

4.4 Optical Elements and related numerical procedures

AGSMM: AGS main magnet

The AGS main magnet is a combined function dipole with straight axis (curves of constant field are straight lines). The simulation of *AGSMM* works like *MULTIPOL*, with the following three particularities:

- the dipole field B_0 in AGSMM is drawn for the reference rigidity, $B\rho_{ref}$ so to preserve $\rho = B\rho_{ref}/B_0$ and the orbit deviation L/ρ . In particular,
 - in the absence of acceleration, $B\rho_{ref} \equiv BORO$, with BORO the quantity appearing in the object definition using OBJET, MCOBJET,
 - in pessence of acceleration using CAVITE, $B\rho_{ref} \equiv BORO(1+D)$, with D the relative momentum increase, a quantity that **zgoubi** updates at cavity traversal.
- the field indices, quadrupole K1 and sextupole K2, are derived from the reference rigidity, $B\rho_{ref}$, via so-called "transfer functions" momentum-dependent polynomials.
 - A consequence of the previous two items is that no field value is required in defining the AGS main magnets in zgoubi data list "zgoubi.dat".
- the AGS main dipole has backleg windings, used for instance for injection and extraction orbit bumps. Windings turn number and Ampere-turns are part of the data in the input data list. The intensity in the windings is accounted for in the calculation of the transfer function from coil current to magnetic field in AGSMM.

AGSQUAD: AGS quadrupole

The AGS quadrupoles are regular quadrupoles. The simulation of *AGSQUAD* works like *MULTIPOL*. However some of the AGS quadrupoles have two superimposed coil circuits, with separate power supplies. It has been dealt with this particularity by allowing for an additional set of multipole data in *AGSQUAD*, compared to *MULTIPOL*.

The field in AGSQUAD is computed using transfer functions from the intensity in the coils to the magnetic field, accounting for non-linearities.

AIMANT: Generation of dipole mid-plane 2-D map, polar frame

The keyword *AIMANT* provides an automatic generation of a dipole median plane field map in polar coordinates. A more recent and improved version will be found in *DIPOLE-M*. The extent of the map is defined by the following parameters, as shown in Figs. 11A and 11B,

AT : total angular aperture

RM: mean radius used for the positioning of field boundaries RMIN, RMAX: minimum and maximum radial boundaries of the map

The 2 or 3 effective field boundaries (EFB) inside the map are defined from geometric boundaries, the shape and position of which are determined by the following parameters,

ACENT: arbitrary angle, used for the positioning of the EFB's.

 ω : azimuth of an EFB with respect to ACENT

 θ : angle of a boundary with respect to its azimuth (wedge angle)

 R_1, R_2 : radius of curvature of an EFB U_1, U_2 : extent of the linear part of the EFB.

At any node of the map mesh, the value of the Z component of the field is calculated as

$$B_Z = \mathcal{F} * B_0 * \left(1 + N * \left(\frac{R - RM}{RM}\right) + B * \left(\frac{R - RM}{RM}\right)^2 + G * \left(\frac{R - RM}{RM}\right)^3\right)$$
(4.4.1)

where N, B and G are respectively the first, second and third order field indices and \mathcal{F} is the fringe field coefficient.

Calculation of the Fringe Field Coefficient

With each EFB a realistic extent of the fringe field, λ , is associated (Figs. 11A and 11B), and a fringe field coefficient F is calculated. In the following λ stands for either λ_E (Entrance), λ_S (Exit) or λ_L (Lateral EFB).

If a node of the map mesh is at a distance of the EFB larger than λ , then F=0 outside the field map and $\mathcal{F}=1$ inside. If a node is inside the fringe field zone, then F is calculated as follows.

Two options are available, for the calculation of F, depending on the value of ξ .

If $\xi \geq 0$, F is a second order type fringe field (Fig. 12) given by

$$F = \frac{1}{2} \frac{(\lambda - s)^2}{\lambda^2 - \xi^2} \quad \text{if } \xi \le s \le \lambda$$
 (4.4.2)

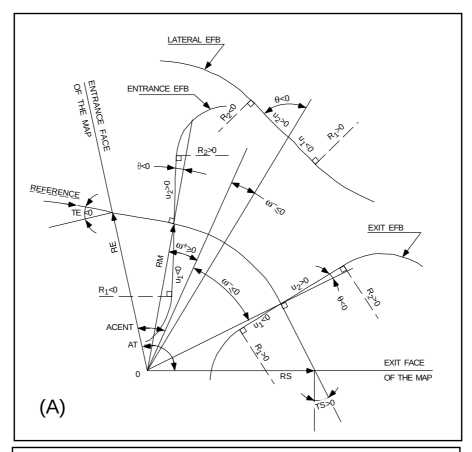
$$F = 1 - \frac{1}{2} \frac{(\lambda - s)^2}{\lambda^2 - \xi^2} \quad \text{if} \quad -\lambda \le s \le -\xi$$
 (4.4.3)

where s is the distance to the EFB, and

$$F = \frac{1}{2} + \frac{s}{\lambda + \xi}$$
 if $0 \le s \le \xi$ (4.4.4)

$$F = \frac{1}{2} - \frac{s}{\lambda + \xi}$$
 if $-\xi \le s \le 0$ (4.4.5)

This simple model allows a rapid calculation of the fringe field, but may lead to erratic behavior of the field when extrapolating out of the median plane, due to the discontinuity of d^2B/ds^2 at $s=\pm\xi$ and $s=\pm\lambda$. For more accuracy it is better to use the next option.



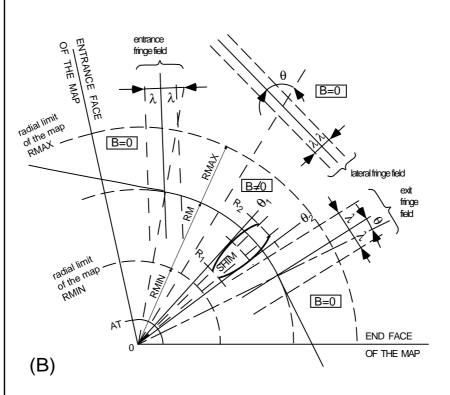


Figure 11: A : Parameters used to define the field map and geometric boundaries. B : Parameters used to define the field map and fringe fields.

If $\xi = -1$, F is an exponential type fringe field (Fig. 12) given by [21]

$$F = \frac{1}{1 + \exp P(s)} \tag{4.4.6}$$

where s is the distance to the EFB, and

$$P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^2 + C_3 \left(\frac{s}{\lambda}\right)^3 + C_4 \left(\frac{s}{\lambda}\right)^4 + C_5 \left(\frac{s}{\lambda}\right)^5 \tag{4.4.7}$$

The values of the coefficients C_0 to C_5 should be such that the derivatives of B_Z with respect to s be negligible at $s=\pm\lambda$, so as not to perturb the extrapolation of \vec{B} out of the median plane (this restriction no longer holds in the improved version DIPOLE-M).

It is also possible to simulate a shift of the EFB, by giving a non zero value to the parameter *SHIFT*. s is then changed to s- SHIFT in the previous equation. This allows small variations of the total magnetic length.

Let F_E (respectively F_S , F_L) be the fringe field coefficient attached to the entrance (respectively exit, lateral) EFB following eqs. above. At any node of the map mesh, the resulting value of the fringe field coefficient (eq. 4.4.1) is (Fig. 13)

$$\mathcal{F} = F_E * F_S * F_L$$

 $(F_L = 1 \text{ if no lateral EFB is requested}).$

The Mesh of the Field Map

The magnetic field is calculated at the nodes of a mesh with polar coordinates, in the median plane. The radial step is given by

$$\delta R = \frac{RMAX - RMIN}{IRMAX - 1}$$

and the angular step by

$$\delta\theta = \frac{AT}{IAMAX - 1}$$

where, RMIN and RMAX are the lower and upper radial limits of the field map, and AT is its total angular aperture (Fig. 11B). IRMAX and IAMAX are the total number of nodes in the radial and angular directions.

Simulating Field Defects and Shims

Once the initial map is calculated, it is possible to modify it by means of the parameter *NBS*, so as to simulate field defects or shims.

If NBS = -2, the map is globally modified by a perturbation proportional to $R - R_0$, where R_0 is an arbitrary radius, with an amplitude $\Delta B_Z/B_0$, so that B_Z at the nodes of the mesh is replaced by

$$B_Z * \left(1 + \frac{\Delta B_Z}{B_0} \frac{R - R_0}{RMAX - RMIN}\right)$$

If NBS = -1, the perturbation is proportional to $\theta - \theta_0$, and B_Z is replaced by

$$B_Z * \left(1 + \frac{\Delta B_Z}{B_0} \frac{\theta - \theta_0}{AT}\right)$$

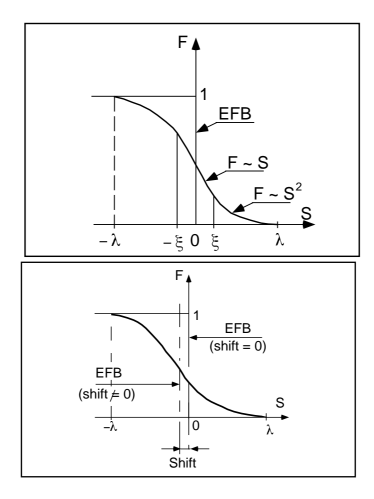


Figure 12: Second order type fringe field (upper plot) and exponential type fringe field (lower plot).

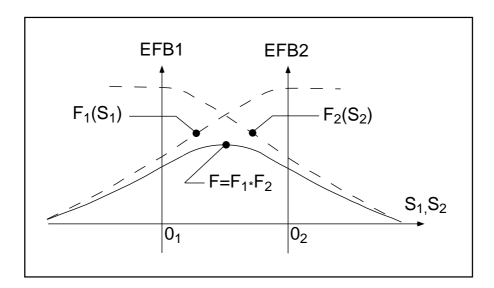


Figure 13: Effective value of \mathcal{F} for overlapping fringe fields F_1 and F_2 centered at O_1 and O_2 .

If NBS \geq 1, then NBS shims are introduced at positions $\frac{R_1 + R_2}{2}$, $\frac{\theta_1 + \theta_2}{2}$ (Fig. 14) [22] The initial field map is modified by shims with second order profiles given by

$$\theta = \left(\gamma + \frac{\alpha}{\mu}\right) \beta \, \frac{X^2}{\rho^2}$$

where X is shown in Fig. 14, $\rho=\frac{R_1+R_2}{2}$ is the central radius, α and γ are the angular limits of the shim, β and μ are parameters.

At each shim, the value of B_Z at any node of the initial map is replaced by

$$B_Z * \left(1 + F\theta * FR * \frac{\Delta B_Z}{B_0}\right)$$

where $F\theta=0$ or FR=0 outside the shim, and $F\theta=1$ and FR=1 inside.

Extrapolation Off Median Plane

The vector field \vec{B} and its derivatives in the median plane are calculated by means of a second or fourth order polynomial interpolation, depending on the value of the parameter IORDRE (IORDRE=2, 25 or 4, see section 1.4.2). The transformation from polar to Cartesian coordinates is performed following eqs. (1.4.9 or 1.4.10). Extrapolation off median phase is then performed by means of Taylor expansions following the procedure described in section 1.3.2.

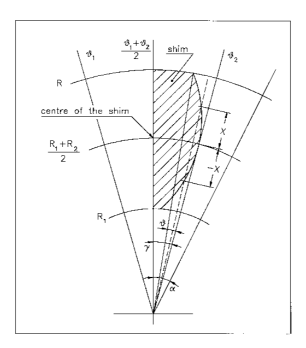


Figure 14: A second order profile shim. The shim is centered at $\frac{(R_1 + R_2)}{2}$ and $\frac{(\theta_1 + \theta_2)}{2}$.

AUTOREF: Automatic transformation to a new reference frame

AUTOREF positions the new reference frame following 3 options :

If I = 1, AUTOREF is equivalent to

$$CHANGREF[XCE = 0, YCE = Y(1), ALE = T(1)]$$

so that the new reference frame is at the exit of the last element, with particle 1 at the origin with its horizontal angle set to T=0.

If I = 2, it is equivalent to

so that the new reference frame is at the position (XW, YW) of the waist (calculated automatically in the same way as for IMAGE) of the three rays number 1, 4 and 5 (compatible for instance with OBJET, KOBJ = 5, 6 together with the use of MATRIX) while T(1) is set to zero.

If I = 3, it is equivalent to

CHANGREF[XW, YW, T(I1)]

so that the new reference frame is at the position (XW, YW) of the waist (calculated automatically in the same way as for IMAGE) of the three rays number I1, I2 and I3 specified as data, while T(1) is set to zero.

BEND: Bending magnet, Cartesian frame

BEND is one of the several keywords available for the simulation of dipole magnets. It presents the interest of easy handling, and is well adapted for the simulation of synchrotron dipoles and such other regular dipoles as sector magnets with wedge angles.

The field in *BEND* is defined in a Cartesian coordinate frame (unlike for instance *DIPOLE[S]* that uses a polar frame). As a consequence, having particle coordinates at entrance or exit of the magnet referring to the curved main direction of motion may require using *KPOS*, in particular *KPOS=3* (in a circular machine cell for instance).

The dipole simulation is performed from the magnet geometrical length XL, from the skew angle (rotation wrt. the X axis, useful for obtaining vertical deviation magnet), and from the field B1 such that in absence of fringe field the deviation θ satisfies $XL=2\frac{BORO}{B1}\sin(\frac{\theta}{2})$.

Then follows the description of the entrance and exit EFB's and fringe fields. The wedge angles W_E (entrance) and W_S (exit) are defined with respect to the sector angle, with the signs as described in Fig. 15. Within a distance $\pm X_E(\pm X_S)$ on both sides of the entrance (exit) EFB, the fringe field model is used; elsewhere, the field is supposed to be uniform.

If λ_E (resp. λ_S) is zero sharp edge field model is assumed at entrance (resp. exit) of the magnet and X_E (resp. X_S) is set to zero. In this case, the wedge angle vertical first order focusing effect (if $\vec{B}1$ is non zero) is simulated at magnet entrance and exit by a kick $P_2 = P_1 - Z_1 \tan(\epsilon/\rho)$ applied to each particle (P_1 , P_2 are the vertical angles upstream and downstream the EFB, Z_1 the vertical particle position at the EFB, ρ the local horizontal bending radius and ϵ the wedge angle experienced by the particle; ϵ depends on the horizontal angle T).

Magnet (mis-)alignement is assured by KPOS. KPOS also allows some degrees of automatic alignement useful for periodic structures (section 4.6.5).

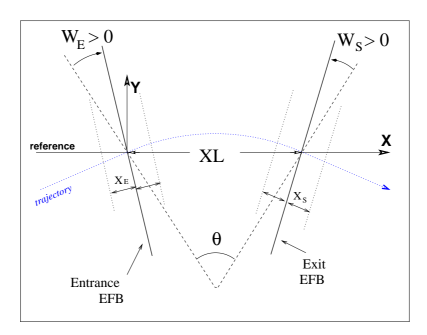


Figure 15: Geometry and parameters of *BEND* in its Cartesian frame : XL = length, $\theta = \text{deviation}$, W_E , W_S are the entrance and exit wedge angles.

BREVOL: 1-D uniform mesh magnetic field map

BREVOL reads a 1-D axial field map from a storage data file, whose content must fit the following FORTRAN reading sequence

where IX is the number of nodes along the (symmetry) X-axis, X(I) their coordinates, and BX(I) the values of the X component of the field. BX is normalized with BNORM factor prior to ray-tracing, as well X is normalized with a XNORM coefficient (usefull to convert to centimeters, the working units in zgoubi). For binary files, FNAME must begin with 'B-' or 'b-', a flag 'BINARY' will thus be set to '.TRUE.'.

X-cylindrical symmetry is assumed, resulting in BY and BZ taken to be zero on axis. $\vec{B}(X,Y,Z)$ and its derivatives along a particle trajectory are calculated by means of a 5-point polynomial fit followed by second order off-axis Taylor series extrapolation (see sections 1.3.1, 1.4.1).

Entrance and/or exit integration boundaries may be defined in the same way as in *CARTEMES* by means of the flag ID and coefficients A, B, C, etc.

CARTEMES: 2-D Cartesian uniform mesh magnetic field map

CARTEMES was originally dedicated to the reading and processing of the measured median plane field maps of the QDD spectrometer SPES2 at Saclay. However, it can be used for the reading of any other 2-D median plane maps, provided that the format of the field data storage file fits the following FORTRAN sequence

where, IX and JY are the number of longitudinal and transverse horizontal nodes of the uniform mesh, and X(I), Y(J) their coordinates. FNAME is the file containing the field data. For binary files, FNAME must begin with 'B_' or 'b_', a flag 'BINARY' will thus be set to '.TRUE.'.

The measured field BMES is normalized with BNORM,

$$B(I, J) = BMES(I, J) \times BNORM$$

As well the longitudinal coordinate X is normalized with a XNORM coefficient (usefull to convert to centimeters, the working units in **zgoubi**.

The vector field, \vec{B} , and its derivatives out of the median plane are calculated by means of a second or fourth order polynomial interpolation, depending on the value of the parameter IORDRE (IORDRE = 2, 25 or 4, see section 1.4.2).

In case a particle exits the mesh, its EX flag is set to -1 (see section 4.6.8 on page 156), however it is still tracked with the field being *extrapolated* from the closest mesh nodes of the map. Note that such extrapolation process may induce eratic behavior if the distance from the mesh gets too large.

Entrance and/or exit integration boundaries can be defined with the flag ID, as follows (Fig. 16).

If ID = 1: the integration in the field is terminated on a boundary with equation A'X + B'Y + C' = 0, and then the trajectories are extrapolated linearly onto the exit end of the map.

If ID = -1: an entrance boundary is defined, with equation A'X + B'Y + C' = 0, up to which trajectories are first extrapolated linearly from the map entrance end, prior to being integrated in the field.

If $ID \ge 2$: one entrance boundary, and ID-1 exit boundaries are defined, as above. The integration in the field terminates on the last (ID-1) exit boundary. No extrapolation onto the map exit end is performed in this case.

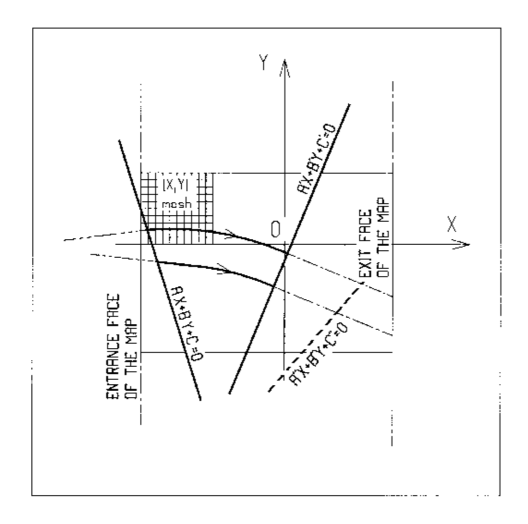


Figure 16: OXY is the coordinate system of the mesh. Integration boundaries may be defined, using $ID \neq 0$: particle coordinates are extrapolated linearly from the entrance face of the map, onto the boundary A'X + B'Y + C' = 0; after ray-tracing inside the map and terminating on the boundary AX + BY + C = 0, coordinates are extrapolated linearly onto the exit face of the map if ID = 2, or terminated on the last (ID - 1) boundary if ID > 2.

CAVITE: Accelerating cavity

CAVITE provides an simulation of a (zero length) accelerating cavity; it can be used in conjunction with keywords REBELOTE and SCALING for the simulation of multiturn tracking with synchrotron acceleration (see section 4.6.7). It must be preceded by PARTICUL for the definition of mass M and charge q.

If IOPT = 0: CAVITE is switched off.

If IOPT = 1: CAVITE simulates the R.F. cavity of a synchrotron accelerator. Normally the keyword CAVITE appears at the end of the optical structure (the periodic motion over IT = 1, NPASS + 1 turns is simulated by means of the keyword REBELOTE, option K = 99 while R.F. and optical elements timings are simulated by means of SCALING — see section 4.6.7). The synchrotron motion of any of the MAX particles of a beam is obtained by solving the following mapping

$$\begin{cases} \phi_2 - \phi_1 = 2\pi f_{RF} \left(\frac{\ell}{\beta c} - \frac{\mathcal{L}}{\beta_s c} \right) \\ W_2 - W_1 = q\hat{V} \sin \phi_1 \end{cases}$$

where

 $\phi = \text{R.F. phase}$; $\phi_2 - \phi_1 = \text{variation of } \phi \text{ between two traversals}$

W = kinetic energy; $W_2 - W_1 = \text{energy gain at a traversal of } CAVITE$

 \mathcal{L} = length of the synchronous closed orbit (to be calculated by prior ray-tracing,

see the bottom NOTE)

 ℓ = orbit length of the particle between two traversals

 $\beta_s c$ = velocity of the (virtual) synchronous particle

 βc = velocity of the particle

 \hat{V} = peak R.F. voltage

q = particle electric charge.

The R.F. frequency f_{RF} is a multiple of the synchronous revolution frequency, and is obtained from the input data, following

$$f_{RF} = \frac{hc}{\mathcal{L}} \frac{q(B\rho)_s}{\sqrt{q^2(B\rho)_s^2 + (Mc)^2}}$$

where

h = harmonic number of the R.F

M = mass of the particle

c = velocity of light.

The current rigidity $(B\rho)_s$ of the synchronous particle is obtained from the timing law specified by means of *SCAL-ING* following $(B\rho)_s = BORO \cdot SCALE(TIMING)$ (see *SCALING* for the meaning and calculation of the scale factor *SCALE(TIMING)*). If *SCALING* is not used, $(B\rho)_s$ is assumed to keep the constant value *BORO* given in the object description (see *OBJET* for instance).

The velocity βc of a particle is calculated from its current rigidity

$$\beta = \frac{q(B\rho)}{\sqrt{q^2(B\rho)^2 + (Mc)^2}}$$

The velocity $\beta_s c$ of the synchronous particle is obtained in the same way from

$$\beta_s = \frac{q(B\rho)_s}{\sqrt{q^2(B\rho)_s^2 + (Mc)^2}}$$

The kinetic energies and rigidities involved in these formulae are related by

$$q(B\rho) = \sqrt{W(W + 2Mc^2)}$$

Finally, the initial conditions for the mapping, at the first turn, are the following

- For the (virtual) synchronous particle

$$\phi_1 = \phi_s = \text{synchronous phase}$$
 $(B\rho)_{1s} = BORO$

- For any of the I=1, IMAX particles of the beam

$$\phi_{1I} = \phi_s = \text{synchronous phase}$$

 $(B\rho)_{1I} = BORO*D_I$

where the quantities BORO and D_I are given in the object description.

Calculation of the coordinates

Let $p_I = \left[p_{XI}^2 + p_{YI}^2 + p_{ZI}^2\right]^{1/2}$ be the momentum of particle I at the exit of the cavity, while $p_{I_0} = \left[p_{XI_0}^2 + p_{YI_0}^2 + p_{ZI_0}^2\right]^{1/2}$ is its momentum at the entrance. The kick in momentum is assumed to be fully longitudinal, resulting in the following relations between the coordinates at the entrance (denoted by the index zero) and at the exit

$$p_{XI} = \left[p_I^2 - (p_{I_0}^2 - p_{XI_0}^2)\right]^{1/2}$$
 $p_{YI} = p_{YI_0}$, and $p_{ZI} = p_{ZI_0}$ (longitudinal kick)
 $X_I = X_{I_0}$, $Y_I = Y_{I_0}$ and $Z_I = Z_{I_0}$ (zero length cavity)

and for the angles (see Fig. 1)

$$T_I = \operatorname{Atg}\left(\frac{p_{YI}}{p_{XI}}\right)$$

$$P_I = \operatorname{Atg}\left(\frac{P_{ZI}}{(p_{XI}^2 + p_{YI}^2)^{1/2}}\right)$$
 (damping of the transverse motion)

If IOPT = 2: the same simulation of a synchrotron R.F. cavity, as for IOPT = 1, is performed, except that the keyword *SCALING* (family *CAVITE*) is not taken into account in this option: the increase in kinetic energy at each traversal, for the synchronous particle, is

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

where the synchronous phase ϕ_s is given in the input data. From this, the calculation of the law $(B\rho)_s$ and the R.F. frequency f_{RF} follows, according to the formulae given in IOPT = 1.

If IOPT = 3: acceleration without synchrotron motion. Any particle will be given a kick

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

where \hat{V} and ϕ_s are input data.

NOTE: Calculation of the closed orbit.

Due to the fringe fields, the horizontal closed orbit may not coincide with the ideal axis of the optical elements. One way to calculate it at the beginning of the structure (i.e., where the initial particle coordinates have to be defined) is to ray-trace a single particle over a sufficiently large number of turns, starting with the initial condition $(Y_0 = T_0 = Z_0 = P_0 = 0)$, and so as to obtain a statistically well-defined phase-space ellipse. The initial conditions of the closed orbit then correspond to the coordinates Y_c and T_c of the center of this ellipse. Next, ray-tracing over one turn a particle starting with the initial condition $(Y_c, T_c, Z_0 = P_0 = 0)$ will provide the length \mathcal{L} (namely, the F(6,1) coordinate) of the closed orbit.

CHAMBR: Long transverse aperture limitation

CHAMBR causes the identification, counting and stopping of particles that reach the transverse limits of the vacuum chamber. The chamber can be either rectangular (IFORM = 1) or elliptic (IFORM = 2). The chamber is centered at YC, ZC and has transverse dimensions $\pm YL$ and $\pm ZL$ such that any particle will be stopped if its coordinates Y, Z satisfy

$$(Y-YC)^2 \ge YL^2 \text{ or } (Z-ZC)^2 \ge ZL^2 \quad \text{if} \quad \textit{IFORM} = 1$$

$$\frac{(Y-YC)^2}{YL^2} + \frac{(Z-ZC)^2}{ZL^2} \ge 1 \quad \text{if} \quad \textit{IFORM} = 2$$

The conditions introduced with CHAMBR are valid along the optical structure until the next occurrence of the keyword CHAMBR. Then, if IL=1 the aperture is possibly modified by introducing new values of YC, ZC, YL and ZL, or, if IL=2 the chamber ends and information is printed concerning those particles that have been stopped.

The testing is done in optical elements at each integration step, between the *EFB*'s. For instance, in *QUADRUPO* there will be no testing from $-X_E$ to 0 and from XL to $XL + X_S$, but only from 0 to XL; in *DIPOLE*, there is no testing as long as the *ENTRANCE EFB* is not reached, and testing is stopped as soon as the *EXIT* or *LATERAL EFB*'s are passed.

In polar coordinate optical elements Y stands for the radial coordinate (e.g. with DIPOLE, see Figs. 3C and 11). Therefore, centering CHAMBR at YC=RM simulates a chamber curved with radius RM, and having a radial acceptance $RM \pm YL$. The testing is done in ESL (DRIFT) at the beginning and the end, and only for positive drifts. There is no testing in CHANGREF.

When a particle is stopped, its index **E**X (see *OBJET* and section 4.6.8) is set to the value -4, and its actual path length is stored in the array *SORT* for possible further statistical purposes.

CHANGREF: Transformation to a new reference frame

The "old style" CHANGREF transports the particles to a new (O, Y, Z) reference plane. It can be used anywhere in a structure. The new particle coordinates Y_2 , T_2 , Z_2 and P_2 path length S_2 are deduced from the old ones Y_1 , T_1 , Z_1 , P_1 and S_1 by

$$T_{2} = T_{1} - ALE$$

$$Y_{2} = \frac{(Y_{1} - YCE)\cos T_{1} + XCE\sin T_{1}}{\cos T_{2}}$$

$$DL^{2} = (XCE - Y_{2}\sin ALE)^{2} + (YCE - Y_{1} + Y_{2}\cos ALE)^{2}$$

$$Z_{2} = Z_{1} + DL\operatorname{tg}P_{1}$$

$$S_{2} = S_{1} + \frac{DL}{\cos P_{1}}$$

$$P_{2} = P_{1}$$

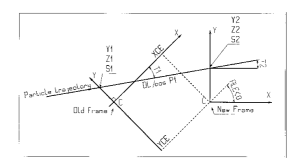


Figure 17: Scheme of the CHANGREF procedure.

where, XCE and YCE are shifts in the horizontal plane along,respectively, X- and Y-axis, and ALE is a rotation around the Z-axis. DL is given the sign of $XCE - Y_2 \sin(ALE)$.

This keyword may for instance be used for positioning optical elements, or for setting a reference frame at the entrance or exit of field maps, or to simulate misalignements (see also *KPOS*).

Effects of *CHANGREF* on spin tracking, particle decay and gas-scattering are taken into account (but not on synchrotron radiation).

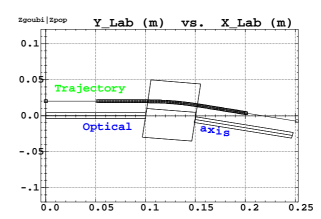
The example below shows the use of *CHANGREF* for the symmetric positioning of a combined function dipole+quadrupole magnet in a drift-bend-drift geometry with 12.691 degrees deviation (obtained upon combined effect of a dipole component and of quadrupole axis shifted 1 cm off optical axis).

Zgoubi data file:

```
Using CHANGREF
'OBJET'
51.71103865921708
                                                One particle, with
2. 0. 0.0 0.0 0.0 1. 'R'
                                Y 0=2 cm, other coordinates zero.
'MARKER' BE(
'DRIFT'

10.
'CHANGREF'

0. 0. -6.34165
'CHANGREF'
           BEG
                                        -> list into zgoubi.plt
                                             First half Z-rotate.
                                                    Next Y-shift.
'MULTIPOL'
               Combined function multipole, dipole + quadrupole
   step size
 'CHANGREF'
0. -1. -6.34165
'DRIFT'
                          First Y-shift back, next half Z-rotate.
 'MARKER
           END
                    .plt
                                        -> list into zgoubi.plt.
 'FAISCEAU'
```



Note: The square markers scheme the stepwise integration in case of ± 5 cm additional fringe field extent upstream and downstream of the 5 cm long multipole.

The "new style" *CHANGREF* transports particles as well to a new (O, Y, Z) reference plane, in a similar manner, however it allows all 6 degrees of freedom, namely, X-, Y-, Z-shift, X-, Y-, Z-rotation.

CHANGREF "new style" allows up to 9 successive such elementary transformations, in arbitrary order. The previous example is transposed into "new style", below.

Zgoubi data file:

```
Using CHANGREF 'New Style
'OBJET'
51.71103865921708 Electron, Ekin=15MeV.
2
11 One particle, with
2. 0. 0.0 0.0 0.0 1. 'R' Y_0=2 cm, other coordinates zero.
1 1 1 1 1 1 1
'MARKERF' BEG .plt -> list into zgoubi.plt.
'DBIFF'
10.
'CHANGREF'
ZR -6.34165 YS 1. First half Z-rotate, Next Y-shift.
'CHANGREF'
0. 1. 0.
'MULTIPOL' Combined function multipole, dipole + quadrupole.
2 -> list into zgoubi.plt.
5 10. 2.064995867082342 2. 0. 0. 0. 0. 0. 0. 0. 0. 0.
0 0 5. 1.1 1.00 1.00 1.00 1.00 1.00 1. 1. 1. 1.
4 .1455 2.2670 -.6395 1.1558 0. 0. 0.
0 0 5. 1.1 1.00 1.00 1.00 1.00 1.00 1. 1. 1. 1.
4 .1455 2.2670 -.6395 1.1558 0. 0. 0.
0 0 0 0 0 0 0 0 0 0 0 0
.1 step size
1 0. 0. 0.
'CHANGREF'
YS -1. ZR -6.341 First Y-shift back, next half Z-rotate.
'PBIFF'
10.
'FAISCEAU'
'END'
```

CIBLE or TARGET: Generate a secondary beam from target interaction

The reaction is $1+2 \longrightarrow 3+4$ with the following parameters

The geometry of the interaction is shown in Fig. 18.

The angular sampling at the exit of the target consists of the NT coordinates $0, \pm TS, \pm 2*TS... \pm (NT-1)*TS/2$ in the median plane, and the NP coordinates $0, \pm PS, \pm 2*PS... \pm (NP-1)*PS/2$ in the vertical plane.

The position of B downstream is deduced from that of A upstream by a transformation equivalent to two transformations using CHANGREF, namely

$$CHANGREF(XCE = YCE = 0, \quad ALE = \beta)$$

followed by

CHANGREF(
$$XCE = YCE = 0$$
, $ALE = \theta - \beta$).

Particle 4 is discarded, while particle 3 continues. The energy loss Q is related to the variable mass M_4 by

$$Q = M_1 + M_2 - (M_3 + M_4)$$
 and $dQ = -dM_4$

The momentum sampling of particle 3 is derived from conservation of energy and momentum, according to

$$M_1c^2 + W_2 = W_3 + W_4$$

 $p_4^2 = p_2^2 + p_3^2 - 2p_2p_3\cos(\theta - T)$

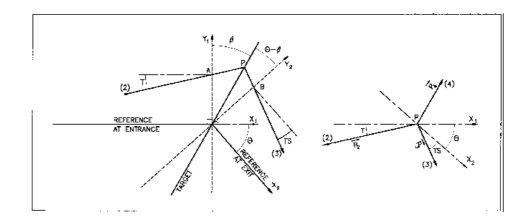


Figure 18: Scheme of the principles of CIBLE (TARGET)

A,T= position, angle of incoming particle 2 in the entrance reference frame P= position of the interaction

B, T =position, angle of the secondary particle in the exit reference frame

 θ = angle between entrance and exit frames

 β = tilt angle of the target

COLLIMA: Collimator

COLLIMA acts as a mathematical aperture of zero length. It causes the identification, counting and stopping of particles that reach the aperture limits.

Physical aperture

A physical aperture can be either rectangular (IFORM = 1) or elliptic (IFORM = 2). The collimator is centered at YC, ZC and has transverse dimensions $\pm YL$ and $\pm ZL$ such that any particle will be stopped if its coordinates Y, Z satisfy

$$\begin{split} (Y-YC)^2 \geq YL^2 \text{ or } (Z-ZC)^2 \geq ZL^2 & \text{ if } & \textit{IFORM} = 1 \\ \frac{(Y-YC)^2}{YL^2} + \frac{(Z-ZC)^2}{ZL^2} \geq 1 & \text{ if } & \textit{IFORM} = 2 \end{split}$$

Longitudinal collimation

COLLIMA can act as a longitudinal phase-space aperture, coordinates acted on are selected with IFORM.J. Any particle will be stopped if its horizontal (h) and vertical (v) coordinates satisfy

$$(h \le h_{min} \text{ or } h \ge h_{max}) \text{ or } (v \le v_{min} \text{ or } v \ge v_{max})$$

wherein, h is either path length S if IFORM=6 or time if IFORM=7, and v is either 1+DP/P if J=1 or kinetic energy if J=2 (provided mass and charge have been defined using the keyword PARTICUL).

If IFORM=11 (respectively 12) then ϵ_Y/π (respectively ϵ_Z/π) is to be specified by the user as well as $\alpha_{Y,Z}$, $\beta_{Y,Z}$. If IFORM=14 (respectively 15) then α_Y and β_Y (respectively α_Z , β_Z) are computed by **zgoubi** by prior matching of the particle population, only $\epsilon_{Y,Z}/\pi$ need be specified by the user.

Phase-space collimation

COLLIMA can act as a phase-space aperture. Any particle will be stopped if its coordinates satisfy

$$\gamma_Y Y^2 + 2\alpha_Y YT + \beta_Y T^2 \ge \epsilon_Y/\pi$$
 if $\mathit{IFORM} = 11$ or 14 $\gamma_Z Z^2 + 2\alpha_Z ZP + \beta_Z P^2 \ge \epsilon_Z/\pi$ if $\mathit{IFORM} = 12$ or 15

If IFORM=11 (respectively 12) then ϵ_Y/π (respectively ϵ_Z/π) is to be specified by the user as well as $\alpha_{Y,Z}$, $\beta_{Y,Z}$. If IFORM=14 (respectively 15) then α_Y and β_Y (respectively α_Z , β_Z) are computed by **zgoubi** by prior matching of the particle population, only $\epsilon_{Y,Z}/\pi$ need be specified by the user.

When a particle is stopped, its index **E**X (see *OBJET* and section 4.6.8) is set to the value -4, and its actual path length is stored in the array **SORT** for possible further statistical purposes (*e.g.* with *HISTO*).

DECAPOLE: Decapole magnet (Fig. 19)

The meaning of parameters for DECAPOLE is the same as for QUADRUPO.

In fringe field regions the magnetic field $\vec{B}(X,Y,Z)$ and its derivatives up to fourth order are derived from the scalar potential approximated to the 5th order in Y and Z

$$V(X,Y,Z) = G\left(Y^4Z - 2Y^2Z^3 + \frac{Z^5}{5}\right)$$
 with $G_0 = \frac{B_0}{R_0^4}$

Outside fringe field regions, or everywhere in sharp edge decapole ($\lambda_E=\lambda_S=0$) , $\vec{B}(X,Y,Z)$ in the magnet is given by

$$B_X = 0$$

 $B_Y = 4G_0(Y^2 - Z^2)YZ$
 $B_Z = G_0(Y^4 - 6Y^2Z^2 + Z^4)$

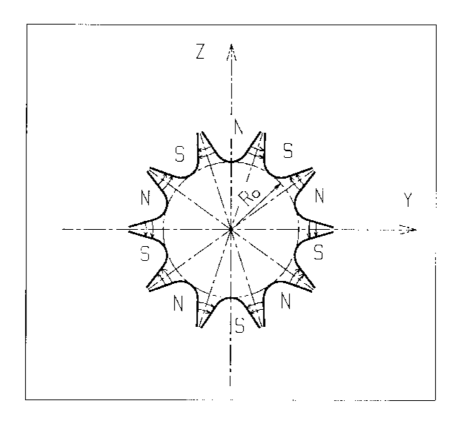


Figure 19: Decapole magnet

DIPOLE: Dipole magnet, polar frame

DIPOLE provides a model of a dipole field, possibly with transverse indices. The field along a particle trajectory is computed as the particle motion proceeds, straightforwardly from the dipole geometrical parametres. To make it more precise, field simulation model in DIPOLE is the same as used in DIPOLE-M and AIMANT for computing a field map; the main difference in DIPOLE is in its skipping that intermediate step of field map generation found in DIPOLE-M and AIMANT.

These are handled as follows. The dimensionning of the magnet is defined by

AT: total angular aperture

RM: mean radius used for the positioning of field boundaries

The 2 or 3 effective field boundaries (EFB), from which the dipole field is drawn, are defined from geometric boundaries, the shape and position of which are determined by the following parameters.

ACENT: arbitrary inner angle, used for EFB's positioning ω : azimuth of an EFB with respect to ACENT

 θ : angle of an EFB with respect to its azimuth (wedge angle)

 R_1, R_2 : radius of curvature of an EFB U_1, U_2 : extent of the linear part of an EFB.

The magnetic field is calculated in polar coordinates. At any position (R, θ) along the particle trajectory the value of the vertical component of the mid-plane field is calculated by

$$B = \mathcal{F}(\mathcal{R}, \theta) * B_0 * \left(1 + N * \left(\frac{R - RM}{RM}\right) + B * \left(\frac{R - RM}{RM}\right)^2 + G * \left(\frac{R - RM}{RM}\right)^3\right)$$
(4.4.8)

where N, B and G are respectively the first, second and third order field indices and $\mathcal{F}(R,\theta)$ is the fringe field coefficient.

Calculation of the Fringe Field Coefficient

With each EFB a realistic extent of the fringe field, λ (normally equal to the gap size), is associated and a fringe field coefficient F is calculated. In the following λ stands for either λ_E (Entrance), λ_S (Exit) or λ_L (Lateral EFB).

F is an exponential type fringe field (Fig. 12) given by [21]

$$F = \frac{1}{1 + \exp P(s)}$$

wherein s is the distance to the EFB and depends on (R, θ) , and

$$P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^2 + C_3 \left(\frac{s}{\lambda}\right)^3 + C_4 \left(\frac{s}{\lambda}\right)^4 + C_5 \left(\frac{s}{\lambda}\right)^5$$

It is also possible to simulate a shift of the *EFB*, by giving a non zero value to the parameter *SHIFT*. s is then changed to s-SHIFT in the previous equation. This allows small variations of the magnetic length.

Let F_E (respectively F_S , F_L) be the fringe field coefficient attached to the entrance (respectively exit, lateral) EFB. At any position on a trajectory the resulting value of the fringe field coefficient (eq. 4.4.9) is

$$\mathcal{F}(R,\theta) = F_E * F_S * F_L$$

 $(F_L = 1 \text{ if no lateral EFB is requested}).$

Simulating Field Defects and Shims

Not provisionned in the present version.

Calculation of the mid-plane field derivatives

This is performed using the numerical interpolation method based on a flying grid as described in the *DIPOLES* procedure (page 103).

Extrapolation Off Median Plane

From the vector field \vec{B} and derivatives in the median plane, first a transformation from polar to Cartesian coordinates is performed, following eqs (1.4.9 or 1.4.10), then, extrapolation off median plane is performed by means of Taylor expansions, following the procedure described in section 1.3.2.

DIPOLE-M: Generation of dipole mid-plane 2-D map, polar frame

DIPOLE-M is a more recent, simpler and improved version of AIMANT.

The keyword *DIPOLE-M* provides an automatic generation of a dipole field map in polar coordinates. The extent of the map is defined by the following parameters, as shown in Figs. 11A and 11B.

AT : total angular aperture

RM : mean radius used for the positioning of field boundaries

RMIN, RMAX: minimum and maximum radii

The 2 or 3 effective field boundaries (EFB) inside the map are defined from geometric boundaries, the shape and position of which are determined by the following parameters.

ACENT: arbitrary inner angle, used for EFB's positioning ω : azimuth of an EFB with respect to ACENT

 θ : angle of an EFB with respect to its azimuth (wedge angle)

 R_1, R_2 : radius of curvature of an EFB U_1, U_2 : extent of the linear part of an EFB.

At any node of the map mesh, the value of the field is calculated as

$$B = \mathcal{F} * B_0 * \left(1 + N * \left(\frac{R - RM}{RM}\right) + B * \left(\frac{R - RM}{RM}\right)^2 + G * \left(\frac{R - RM}{RM}\right)^3\right)$$
(4.4.9)

where N, B and G are respectively the first, second and third order field indices and \mathcal{F} is the fringe field coefficient.

Calculation of the Fringe Field Coefficient

With each EFB a realistic extent of the fringe field, λ (normally equal to the gap size), is associated and a fringe field coefficient F is calculated. In the following λ stands for either λ_E (Entrance), λ_S (Exit) or λ_L (Lateral EFB).

F is an exponential type fringe field (Fig. 12) given by [21]

$$F = \frac{1}{1 + \exp P(s)}$$

where s is the distance to the EFB, and

$$P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^2 + C_3 \left(\frac{s}{\lambda}\right)^3 + C_4 \left(\frac{s}{\lambda}\right)^4 + C_5 \left(\frac{s}{\lambda}\right)^5$$

It is also possible to simulate a shift of the *EFB*, by giving a non zero value to the parameter *SHIFT*. s is then changed to s-SHIFT in the previous equation. This allows small variations of the total magnetic length.

Let F_E (respectively F_S , F_L) be the fringe field coefficient attached to the entrance (respectively exit, lateral) EFB. At any node of the map mesh, the resulting value of the fringe field coefficient (eq. 4.4.9) is

$$\mathcal{F} = F_E * F_S * F_L$$

 $(F_L = 1 \text{ if no lateral EFB is requested}).$

The Mesh of the Field Map

The magnetic field is calculated at the nodes of a mesh with polar coordinates, in the median plane. The radial step is given by

$$\delta R = \frac{RMAX - RMIN}{IRMAX - 1}$$

and the angular step by

$$\delta\theta = \frac{AT}{IAMAX - 1}$$

where, RMIN and RMAX are the lower and upper radial limits of the field map, and AT is its total angular aperture (Fig. 11B). IRMAX and IAMAX are the total number of nodes in the radial and angular directions.

Simulating Field Defects and Shims

Once the initial map is calculated, it is possible to modify it by means of the parameter *NBS*, so as to simulate field defects or shims.

If NBS = -2, the map is globally modified by a perturbation proportional to $R - R_0$, where R_0 is an arbitrary radius, with an amplitude $\Delta B_Z/B_0$, so that B_Z at the nodes of the mesh is replaced by

$$B_Z * \left(1 + \frac{\Delta B_Z}{B_0} \frac{R - R_0}{RMAX - RMIN}\right)$$

If NBS = -1, the perturbation is proportional to $\theta - \theta_0$, and B_Z is replaced by

$$B_Z * \left(1 + \frac{\Delta B_Z}{B_0} \frac{\theta - \theta_0}{AT}\right)$$

If NBS \geq 1, then NBS shims are introduced at positions $\frac{R_1 + R_2}{2}$, $\frac{\theta_1 + \theta_2}{2}$ (Fig. 14) [22] The initial field map is modified by shims with second order profiles given by

$$\theta = \left(\gamma + \frac{\alpha}{\mu}\right) \beta \, \frac{X^2}{\rho^2}$$

where X is shown in Fig. 12, $\rho = \frac{R_1 + R_2}{2}$ is the central radius, α and γ are the angular limits of the shim, β and μ are parameters.

At each shim, the value of B_Z at any node of the initial map is replaced by

$$B_Z * \left(1 + F\theta * FR * \frac{\Delta B_Z}{B_0}\right)$$

where $F\theta = 0$ or FR = 0 outside the shim, and $F\theta = 1$ and FR = 1 inside.

Extrapolation Off Median Plane

The vector field \vec{B} and its derivatives in the median plane are calculated by means of a second or fourth order polynomial interpolation, depending on the value of the parameter IORDRE (IORDRE=2, 25 or 4, see section 1.4.2). The transformation from polar to Cartesian coordinates is performed following eqs (1.4.9 or 1.4.10). Extrapolation off median plane is then performed by means of Taylor expansions, following the procedure described in section 1.3.2.

DIPOLES: Dipole magnet N-tuple, polar frame [27, 28]

DIPOLES works much like DIPOLE as to the field modelling, yet with the particularity that it allows positioning up to 5 such dipoles within the angular sector with full aperture AT thus allowing accounting for overlapping fringe fields This is done in the following way⁵.

The dimensionning of the magnet is defined by

AT: total angular aperture

RM: mean radius used for the positioning of field boundaries

For each one of the N=1 to 5 dipoles of the N-tuple, the 2 effective field boundaries (entrance and exit EFBs) from which the dipole field is drawn (eq. 4.4.11) are defined from geometric boundaries, the shape and position of which are determined by the following parameters (in the same manner as in DIPOLE, DIPOLE-M) (see Fig. 11-A page 80, and Fig. 20)

 ACN_i : arbitrary inner angle, used for EFB's positioning

 ω : azimuth of an EFB with respect to ACN

 θ : angle of an EFB with respect to its azimuth (wedge angle)

 $R_1,\,R_2$: radius of curvature of an EFB $U_1,\,U_2$: extent of the linear part of an EFB

Calculation of the field from a single dipole

The magnetic field is calculated in polar coordinates. At all (R, θ) in the median plane (z = 0), the magnetic field due a single one (index i) of the dipoles of a N-tuple magnet is written

$$B_{zi}(R,\theta) = B_{z0,i} \mathcal{F}_i(R,\theta) \left(1 + b_{1i}(R - RM_i) / RM_i + b_{2i}(R - RM_i)^2 / RM_i^2 + \dots \right)$$
(4.4.10)

wherein $B_{z0,i}$ is a reference field, at reference radius RM_i , and $\mathcal{F}(R,\theta)$ is the fringe field coefficient, see below. This field model is proper to simulate for instance chicane dipoles, isochronous or superconducting FFAG magnets, etc.

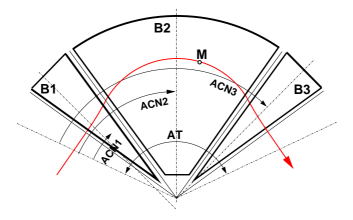


Figure 20: Definition of a dipole triplet using the DIPOLES or FFAG procedures.

Calculation of the fringe field coefficient

In a dipole, with each EFB a realistic extent of the fringe field, g, is associated and a fringe field coefficient F is calculated.

 $^{^5}$ FFAG can be referred to as another instance of a procedure based on such method.

F is an exponential type fringe field (Fig. 12, page 82) given by [21]

$$F = \frac{1}{1 + \exp P(d)}$$

wherein d is the distance to the EFB and depends on (R, θ) , and

$$P(d) = C_0 + C_1 \left(\frac{d}{g}\right) + C_2 \left(\frac{d}{g}\right)^2 + C_3 \left(\frac{d}{g}\right)^3 + C_4 \left(\frac{d}{g}\right)^4 + C_5 \left(\frac{d}{g}\right)^5$$

In addition, q is made dependent of R (a way to simulate the effect of variable gap size on field fall-off), under the form

$$g(R) = g_0 (RM/R)^{\kappa}$$

This dependence is accounted for rigorously if the interpolation method is used, to zero order (derivatives of g(R) are not considered) if the analytic method is used.

Let F_E (respectively F_S) be the fringe field coefficient attached to the entrance (respectively exit) EFB; at any position on a trajectory the resulting value of the fringe field coefficient is taken to be

$$\mathcal{F}_i(R,\theta) = F_E * F_S \tag{4.4.11}$$

Calculation of the full field from all N dipoles

Now, accounting for N neighboring dipoles in an N-tuple, the mid-plane field and field derivatives are obtained by addition of the contributions of the N dipoles taken separately, namely

$$B_{z}(R,\theta) = \sum_{i=1,N} B_{zi}(R,\theta) = \sum_{i=1,N} B_{z0,i} \mathcal{F}_{i}(R,\theta) \mathcal{R}_{i}(r)$$

$$\frac{\partial^{k+l} \vec{B_{z}}(R,\theta)}{\partial \theta^{k} \partial r^{l}} = \sum_{i=1,N} \frac{\partial^{k+l} \vec{B_{zi}}(R,\theta)}{\partial \theta^{k} \partial r^{l}}$$
(4.4.12)

with $\mathcal{R}_i(R)$ as defined in Eq. 4.4.10. Note that, in doing so it is not meant that field superposition does apply in reality, it is just meant to provide the possibility of obtaining a realistic field shape, that would for instance closely match (using appropriate $C_0 - C_5$ sets of coefficients) 3-D field simulations obtained from magnet codes.

Calculation of the mid-plane field derivatives

Two methods have been implemented to calculate the field derivatives in the median plane (Eq. 4.4.12), based on either analytical expressions derived from the magnet geometrical description, or classical numerical interpolation.

The first method has the merit of insuring best symplecticity in principle and fastest tracking. The interest of the second method is in its facilitating possible changes in the mid-plane magnetic field model $B_z(R, \theta)$, for instance if simulations of shims, defects, or special R, θ field dependence need to be introduced.

Analytical method:

The starting ingredients are, on the one hand distances to the EFBs,

$$d(R,\theta) = \sqrt{(x(R,\theta) - x_0(R,\theta))^2 + (y(R,\theta) - y_0(R,\theta))^2}$$

to be computed for the two cases d_{Entrance} , d_{Exit} , and on the other hand the expressions of the coordinates of particle position M and its projection P on the EFB in terms of the magnet geometrical parameters, namely

$$\begin{array}{rcl} x(R,\theta) & = & \cos(ACN - \theta) - RM \\ y(R,\theta) & = & R\sin(ACN - \theta) \\ x_P(R,\theta) & = & \sin(u) \left(y(R,\theta) - y_b \right) / 2 + x_b \sin^2(u) + x(R,\theta) \cos^2(u) \\ y_P(R,\theta) & = & \sin(u) \left(x(R,\theta) - x_b \right) / 2 + y_b \cos^2(u) + y(R,\theta) \sin^2(u) \end{array}$$

with $x_b,\ y_b,\ u$ parameters drawn from the magnet geometry (sector angle, wedge angle, face curvatures, etc.). These ingredients allow calculating the derivatives $\frac{\partial^{u+v}x(R,\theta)}{\partial \theta^u\partial r^v}$, $\frac{\partial^{u+v}y(R,\theta)}{\partial \theta^u\partial r^v}$, $\frac{\partial^{u+v}x_0(R,\theta)}{\partial \theta^u\partial r^v}$, $\frac{\partial^{u+v}y_0(R,\theta)}{\partial \theta^u\partial r^v}$, which, in turn, intervene in the derivatives of the compound functions $\frac{\partial^{u+v}F(R,\theta)}{\partial \theta^u\partial r^v}$, $\frac{\partial^{u+v}p(R,\theta)}{\partial \theta^u\partial r^v}$, $\frac{\partial^{u+v}d(R,\theta)}{\partial \theta^u\partial r^v}$.

Interpolation method:

The expression $B_z(R, \theta)$ in Eq. 4.4.12 is, in this case, computed at the $n \times n$ nodes (n = 3 or 5 in practice) of a "flying" interpolation grid in the median plane centered on the projection m_0 of the actual particle position M_0 as schemed in Fig. 21. A polynomial interpolation is involved, of the form

$$B_z(R,\theta) = A_{00} + A_{10}\theta + A_{01}r + A_{20}\theta^2 + A_{11}\theta r + A_{02}r^2$$

that yields the requested derivatives, using

$$A_{kl} = \frac{1}{k!l!} \frac{\partial^{k+l} B}{\partial \theta^k \partial r^l}$$

Note that, the source code contains the explicit analytical expressions of the coefficients A_{kl} solutions of the normal equations, so that the operation is not CPU time consuming.

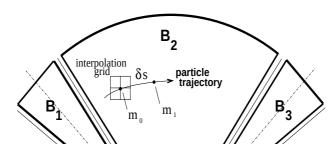


Figure 21: Interpolation method. m_0 and m_1 are the projections in the median plane of particle positions M_0 and M_1 separated by one integration step δs .

Extrapolation Off Median Plane

From the vector field \vec{B} and derivatives in the median plane, first a transformation from polar to Cartesian coordinates is performed, following eqs (1.4.9 or 1.4.10), then, extrapolation off median plane is performed by means of Taylor expansions, following the procedure described in section 1.3.2.

Sharp edge

Sharp edge field fall-off at a field boundary can only be simulated if the following conditions are fulfilled:

- entrance (resp. exit) field boundary coincides with entrance (resp. exit) dipole limit (it means in particular, see Fig. 11, $\omega^+ = ACENT$ (resp. $\omega^- = -(AT ACENT)$), together with $\theta = 0$ at entrance (resp. exit) EFBs),
 - analytical method for calculation of the mid-plane field derivatives is used.

DODECAPO: Dodecapole magnet (Fig. 22)

The meaning of parameters for DODECAPO is the same as for QUADRUPO.

In fringe field regions the magnetic field $\vec{B}(X,Y,Z)$ and its derivatives up to fourth order are derived from the scalar potential approximated to the 6th order in Y and Z

$$V(X,Y,Z)=G\left(Y^4-\frac{10}{3}Y^2Z^2+Z^4\right)YZ$$
 with $G_0=\frac{B_0}{R_0^5}$

Outside fringe field regions, or everywhere in sharp edge dodecapole ($\lambda_E = \lambda_S = 0$) , $\vec{B}(X,Y,Z)$ in the magnet is given by

$$B_X = 0$$

$$B_Y = G_0(5Y^4 - 10Y^2Z^2 + Z^4)Z$$

$$B_Z = G_0(Y^4 - 10Y^2Z^2 + 5Z^4)Y$$

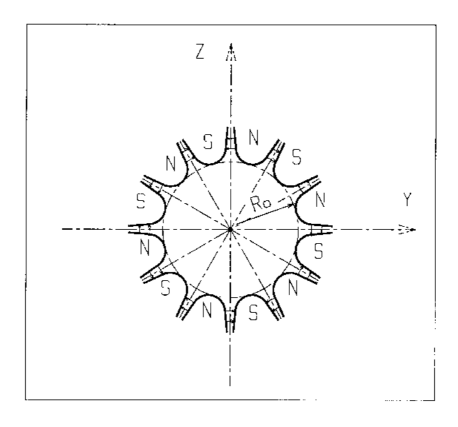


Figure 22: Dodecapole magnet

DRIFT or **ESL**: Field free drift space

DRIFT or ESL allow introduction of a drift space with length XL with positive or negative sign, anywhere in a structure. The associated equations of motion are (Fig. 23)

$$\begin{aligned} Y_2 &= Y_1 + XL * \mathsf{tg}T \\ Z_2 &= Z_1 + \frac{XL}{\cos T} \, \mathsf{tg}P \\ SAR_2 &= SAR_1 + \frac{XL}{\cos T * \cos P} \end{aligned}$$

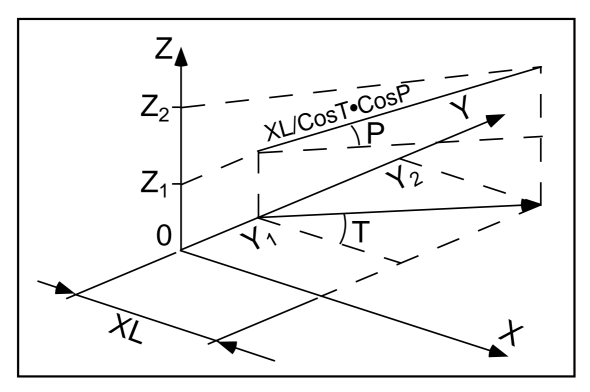


Figure 23: Transfer of particles in a drift space.

EBMULT: Electro-magnetic multipole

EBMULT simulates an electro-magnetic multipole, by addition of electric (\vec{E}) and magnetic (\vec{B}) multipole components (dipole to 20-pole). \vec{E} and its derivatives $\frac{\partial^{i+j+k}\vec{E}}{\partial X^i\partial Y^j\partial Z^k}$ $(i+j+k\leq 4)$ are derived from the general expression of the multipole scalar potential (eq. 1.3.5), followed by a $\frac{\pi}{2n}$ rotation $(n=pole\ order)$, as described in section ?? (see also ELMULT). \vec{B} and its derivatives are derived from the same general potential, as described in section 1.3.6 (see also MULTIPOL).

The entrance and exit fringe fields of the \vec{E} and \vec{B} components are treated separately, in the same way as described under ELMULT and MULTIPOL, for each one of these two fields. Wedge angle correction is applied in sharp edge field model if $\vec{B}1$ is non zero, as in MULTIPOL. Any of the \vec{E} or \vec{B} multipole field component can be rotated independently of the others.

Use PARTICUL prior to EBMULT, for the definition of particle mass and charge.

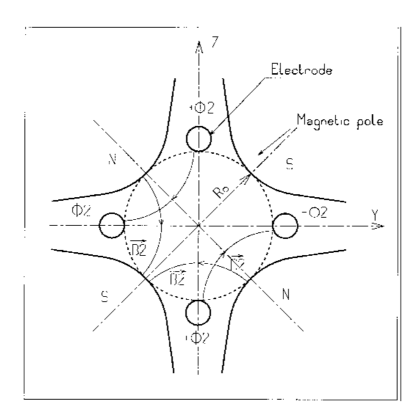


Figure 24: An example of \vec{E} , \vec{B} multipole : the achromatic quadrupole (known for its allowing null second order chromatic aberrations [23]).

EL2TUB: Two-tube electrostatic lens

The lens is cylindrically symmetric about the X-axis.

The length and potential of the first (resp. second) electrode are X1 and V1 (X2 and V2). The distance between the two electrodes is D, and their inner radius is R_0 (Fig. 25). X-axis cylindrical symmetry is assumed. The model for the electrostatic potential along the axis is [25]

$$V(X) = \frac{V_2 - V_1}{2} \operatorname{th} \frac{\omega x}{R_0} \left[+ \frac{V_1 + V_2}{2} \right]$$
 if $D = 0$

$$V(X) = \frac{V_2 - V_1}{2} \frac{1}{2\omega D/R_0} \ln \frac{\operatorname{ch} \omega}{\operatorname{ch} \omega} \frac{x + D}{R_0} \left[+ \frac{V_1 + V_2}{2} \right] \quad \text{if } D \neq 0$$

(x = distance from half-way between the electrodes; $\omega = 1.318$; th = hyperbolic tangent; ch = hyperbolic cosine) from which the field $\vec{E}(X,Y,Z)$ and its derivatives are derived following the procedure described in section ?? (note that they don't depend on the constant term $\left\lceil \frac{V_1 + V_2}{2} \right\rceil$ which disappears when differentiating).

Use *PARTICUL* prior to *EL2TUB*, for the definition of particle mass and charge.

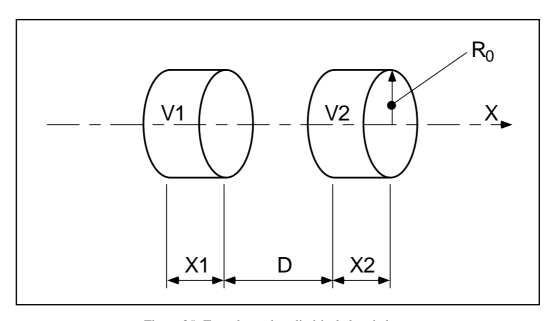


Figure 25: Two-electrode cylindrical electric lens.

ELMIR: Electrostatic N-electrode mirror/lens, straight slits

The device works as mirror or lens, horizontal or vertical. It is made of N 2-plate electrodes and has mid-plane symmetry.

Electrode lengths are L1, L2, ..., LN. D is the mirror/lens gap. The model for the Y-independent electrostatic potential is (after Ref. [26, p.412])

$$V(X,Z) = \sum_{i=2}^{N} \frac{Vi - Vi - 1}{\pi} \arctan \frac{\sinh(\pi(X - Xi - 1)/D)}{\cos(\pi Z/D)}$$

where Vi are the potential at the N electrodes (and normally V1=0 refers to the incident beam energy), Xi are the locations of the slits, X is the distance from the origin taken at the first slit (located at $X1\equiv 0$ between the first and second electrodes). From V(X,Z) the field $\vec{E}(X,Y,Z)$ and derivatives are deduced following the procedure described in section $\ref{eq:potential}$? (page $\ref{eq:potential}$).

The total X-extent of the mirror/lens is $L = \sum_{i=1}^{N} Li$.

In the mirror mode (i.e., option flag MT=11 for vertical mid-plane or 12 for horizontal mid-plane) stepwise integration starts at X=-L1 (entrance of the first electrode) and terminates either when back to X=-L1 or when reaching X=L-L1 (end of the N-th electrode). In the latter case particles are stopped with their index IEX set to -8 (see section 4.6.8 on page 156). Normally X1 should exceed 3D (possibly sensibly, so that V(X < X1) have negligible effect in terms of trajectory behavior).

In the lens mode (i.e., option flag MT=21 for vertical mid-plane or 22 for horizontal mid-plane) stepwise integration starts at X=-L1 (entrance of the first electrode) and terminates either when reaching X=L-L1 (end of the N-th electrode) or when the particle deflection exceeds $\pi/2$. In the latter case the particle is stopped with their index IEX set to -3.

Use PARTICUL prior to ELMIR, for the definition of particle mass and charge.

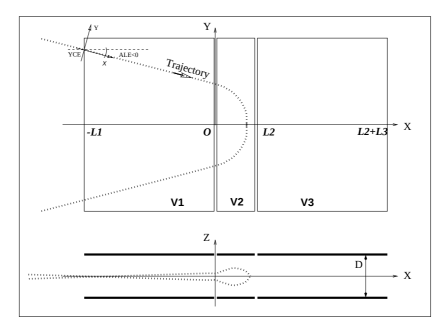


Figure 26: Electrostatic N-electrode mirror/lens, straight slits, in the case N=3, in horizontal mirror mode (MT=11).

Possible non-zero entrance quantities YCE, ALE should be specified using CHANGREF, or using KPOS=3 with YCE=pitch, ALE=half-deviation.

ELMIRC: Electrostatic N-electrode mirror/lens, circular slits [26]

The device works as mirror or lens, horizontal or vertical. It is made of N 2-plate electrodes and has mid-plane symmetry⁶.

Electrode slits are circular, concentric with radii R1, R2, ..., RN-1, D is the mirror/lens gap. The model for the mid-plane (Z=0) radial electrostatic potential is (after Ref. [26, p.443])

$$V(r) = \sum_{i=2}^{N} \frac{Vi - Vi - 1}{\pi} \arctan\left(\sinh\frac{\pi(r - Ri - 1)}{D}\right)$$

where Vi are the potential at the N electrodes (and normally V1 = 0 refers to the incident beam energy). r is the current radius

The mid-plane field $\vec{E}(r)$ and its r-derivatives are first derived by differentiation, then $\vec{E}(r,Z)$ and derivatives are obtained from Taylor expansions and Maxwell relations. Eventually a transformation to the rotating frame provides $\vec{E}(X,Y,Z)$ and derivatives as involved in eq. 1.2.13.

Stepwise integration starts at entrance (defined by RE, TE) of the first electrode and terminates when rotation of the reference rotating frame (RM, X, Y) has reached the value AT. Normally, R1 - RE and R1 - RS should both exceed 3D (possibly sensibly, so that V(r < RE) and V(r < RS) have negligible effect in terms of trajectory tails).

Positioning of the element is performed by means of KPOS (see section 4.6.5).

Use PARTICUL prior to ELMIRC, for the definition of particle mass and charge.

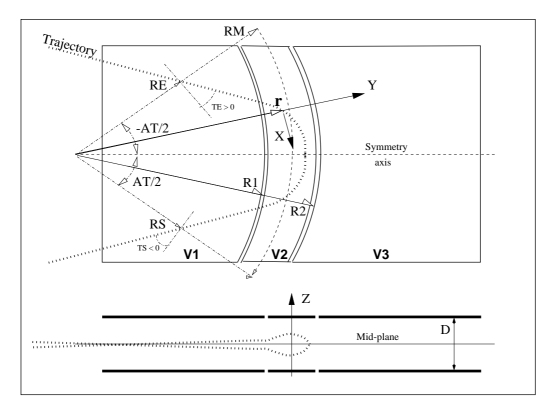


Figure 27: Electrostatic N-electrode mirror/lens, circular slits, in the case N=3, in horizontal mirror mode.

 $^{^6}$ NOTE: in the present version of the code, the sole horizontal mirror mode is operational, and N is limited to 3.

ELMULT: Electric multipole

The simulation of multipolar electric field \vec{M}_E proceeds by addition of the dipolar $(\vec{E}1)$, quadrupolar $(\vec{E}2)$, sextupolar $(\vec{E}3)$, etc., up to 20-polar $(\vec{E}10)$ components, and of their derivatives up to fourth order, following

$$\begin{split} \vec{M}_E &= \vec{E}1 + \vec{E}2 + \vec{E}3 + \ \dots \ + \vec{E}10 \\ \frac{\partial \vec{M}_E}{\partial X} &= \frac{\partial \vec{E}1}{\partial X} + \frac{\partial \vec{E}2}{\partial X} + \frac{\partial \vec{E}3}{\partial X} + \ \dots \ + \frac{\partial \vec{E}10}{\partial X} \\ \frac{\partial^2 M_E}{\partial X \partial Z} &= \frac{\partial^2 \vec{E}1}{\partial X \partial Z} + \frac{\partial^2 \vec{E}2}{\partial X \partial Z} + \frac{\partial^2 \vec{E}3}{\partial X \partial Z} + \ \dots \ + \frac{\partial^2 \vec{E}10}{\partial X \partial Z} \\ \text{etc.} \end{split}$$

The independent components $\vec{E}1$ to $\vec{E}10$ and their derivatives up to the second order are calculated by differentiating the general multipole potential given in eq. 1.3.5 (page 26), followed by a $\frac{\pi}{2n}$ rotation about the X-axis, so that the so defined right electric multipole of order n, and of strength [23, 24]

$$K_n = \frac{1}{2} \frac{\gamma}{\gamma^2 - 1} \frac{V_n}{R_0^n}$$

 $(V_n = \text{potential at the electrode}, R_0 = \text{radius at pole tip}, \gamma = \text{relativistic Lorentz factor of the particle})$ has the same focusing effect than the right magnetic multipole of order n and strength $K_n = \frac{B_n}{R_0^{n-1}B\rho}$ (B_n = field at pole tip, $B\rho$ = particle rigidity, see MULTIPOL).

Such $\frac{\pi}{2n}$ rotation of the multipole components is obtained following the procedure described in section ??.

The entrance and exit fringe fields are treated separately. They are characterized by the integration zone X_E at entrance and X_S at exit, as for QUADRUPO, and by the extent λ_E at entrance, λ_S at exit. The fringe field extents for the dipole component are λ_E and λ_S . The fringe field for the quadrupolar (sextupolar, ..., 20-polar) component is given by a coefficient E_2 (E_3 , ..., E_{10}) at entrance, and E_S (E_S) at exit, such that the fringe field extent is E_S (E_S) at exit, ..., E_S) at entrance and E_S (E_S) at exit.

If $\lambda_E = 0$ ($\lambda_S = 0$) the multipole lens is considered to have a sharp edge field at entrance (exit), and then, X_E (X_S) is forced to zero (for the mere purpose of saving computing time).

If $E_i = 0$ ($S_i = 0$) (i = 2, 10), the entrance (exit) fringe field for multipole component i is considered as a sharp edge field.

Overlapping of fringe fields inside the element is treated separately for each component, in the way described in *QUADRUPO*. Moreover, any multipole component $\vec{E}i$ can be rotated independently by an angle RXi around the longitudinal X-axis, for the simulation of positioning defects, as well as skew lenses.

Use PARTICUL prior to ELMULT, for the definition of particle mass and charge.

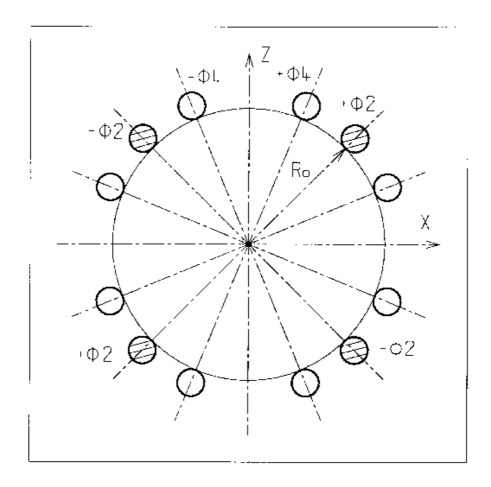


Figure 28: An electric multipole combining skew-quadrupole $(\vec{E}2 \neq \vec{0}, \ \vec{R}2 = \pi/4)$ and skew-octupole $(\vec{E}4 \neq \vec{0}, \ \vec{R}4 = \pi/8)$ components $(\vec{E}1 = \vec{E}3 = \vec{E}5 = \dots = \vec{E}10 = \vec{0})$ [24].

ELREVOL: 1-D uniform mesh electric field map

ELREVOL reads a 1-D axial field map from a storage data file, whose content must fit the following FORTRAN reading sequence

```
OPEN (UNIT = NL, FILE = FNAME, STATUS = 'OLD' [,FORM='UNFORMATTED'])
DO 1 I=1, IX
   IF (BINARY) THEN
        READ(NL) X(I), EX(I)
   ELSE
        READ(NL,*) X(I), EX(I)
   ENDIF
1 CONTINUE
```

where IX is the number of nodes along the (symmetry) X-axis, X(I) their coordinates, and EX(I) the values of the X component of the field. EX is normalized with ENORM prior to ray-tracing. As well the longitudinal coordinate X is normalized with a XNORM coefficient (usefull to convert to centimeters, the working units in zgoubi.

X-cylindrical symmetry is assumed, resulting in EY and EZ taken to be zero on axis. $\vec{E}(X,Y,Z)$ and its derivatives along a particle trajectory are calculated by means of a 5-points polynomial fit followed by second order off-axis Taylor series extrapolation (see sections $\ref{eq:taylor}$).

Entrance and/or exit integration boundaries may be defined in the same way as in *CARTEMES* by means of the flag ID and coefficients A, B, C, A', B', C'.

Use PARTICUL prior to ELREVOL, for the definition of particle mass and charge.

EMMA: 2-D Cartesian or cylindrical mesh field map for EMMA FFAG

EMMA is dedicated to the reading and treatment of 2-D or 3-D Cartesian or cylindrical mesh field maps as delivered by the *TOSCA* magnet computer code standard output.

That part of the manual to be provisionned...

where, IX (JY, KZ) is the number of longitudinal (transverse horizontal, vertical) nodes of the 3-D uniform mesh. For binary files, FNAME must begin with 'B_' or 'b_', a flag 'BINARY' will thus be set to '.TRUE.'.

A flag MOD determines wether Cartesian or Z-axis cylindrical mesh is used. MOD can take various values depending also on the map data file formatting. (To be documented - see FORTRAN subroutine FMAPW and its entries FMAPR, FMAPR2.)

The field $\vec{B} = (B_X, B_Y, B_Z)$ is normalized by means of *BNORM* in a similar way as in *CARTEMES*. As well the coordinates X (and Y, Z with 3-D field maps) is normalized with a *X-[Y-,Z-]NORM* coefficient (usefull to convert to centimeters, the working units in **zgoubi**.

At each step of the trajectory of a particle inside the map, the field and its derivatives are calculated

- in the case of 2-D map, by means of a second or fourth order polynomial interpolation, depending on IORDRE (IORDRE = 2, 25 or 4), as for CARTEMES,
- in the case of 3-D map, by means of a second order polynomial interpolation with a $3\times3\times3$ -point parallelipipedic grid, as described in section 1.4.4.

Entrance and/or exit integration boundaries between which the trajectories are integrated in the field may be defined, in the same way as in *CARTEMES*.

FFAG: FFAG magnet, N-tuple [27, 28]

FFAG works much like DIPOLES as to the field modelling, apart from the radial dependence of the field (so-called "scaling", $B = B_0(r/r_0)^k$. Note that DIPOLES could do the same job by using a multipole expansion of $B_0(r/r_0)^k$).

The FFAG procedure allows overlapping of fringe fields of neighboring dipoles, thus simulating in some sort the field in a dipole N-tuple - as for instance in an FFAG doublet or triplet. This is done in the way described below.

The dimensionning of the magnet is defined by

AT: total angular aperture

RM: mean radius used for the positioning of field boundaries

For each one of the N=1 to (maximum) 5 dipoles of the N-tuple, the two effective field boundaries (entrance and exit EFBs) from which the dipole field is drawn are defined from geometric boundaries, the shape and position of which are determined by the following parameters (in the same manner as in *DIPOLE*, *DIPOLE-M*) (see Fig. 11-A page 80, and Fig. 29)

 ACN_i : arbitrary inner angle, used for EFB's positioning

 ω : azimuth of an EFB with respect to ACN

 θ : angle of an EFB with respect to its azimuth (wedge angle)

 $R_1,\,R_2$: radius of curvature of an EFB $U_1,\,U_2$: extent of the linear part of an EFB

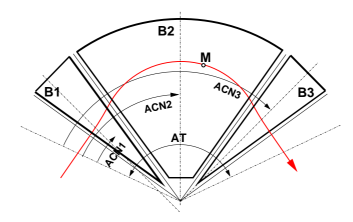


Figure 29: Definition of a dipole N-tuple (N=3, a triplet here) using the DIPOLES or FFAG procedures.

Calculation of the field from a single dipole

The magnetic field is calculated in polar coordinates. At all (R, θ) in the median plane (z = 0), the magnetic field due a single one (index i) of the dipoles of a N-tuple FFAG magnet is written

$$B_{zi}(R,\theta) = B_{z0,i} \mathcal{F}_i(R,\theta) \left(R/R_M \right)^{K_i}$$

wherein $B_{z_0,i}$ is a reference field, at reference radius RM_i , whereas $\mathcal{F}(R,\theta)$ is calculated as described below.

Calculation of $\mathcal{F}_i(R,\theta)$

The fringe field coefficient $\mathcal{F}_i(R,\theta)$ associated with a dipole is computed as in the procedure *DIPOLES* (eq. 4.4.11), including (rigorously if the interpolation method is used, to zero order if the analytic method is used) radial dependence of the gap size

$$g(R) = g_0 \left(RM/R \right)^{\kappa} \tag{4.4.13}$$

so to simulate the effect of gap shaping of $B_{zi}(R,\theta)$ on field fall-off, over the all radial extent of a scaling FFAG dipole (with normally - but not in practice - $\kappa = K_i$).

Calculation of the full field from all N dipoles

For the rest, namely, calculation of the full field at particle position from the N dipoles, analytical calculation or numerical interpolation of the mid-plane field derivatives, extrapolation off median plane, etc., things are performed exactly as in the case of the DIPOLES procedure (see page 102).

Sharp edge

Sharp edge field fall-off at a field boundary can only be simulated if the following conditions are fulfilled:

- entrance (resp. exit) field boundary coincides with entrance (resp. exit) dipole limit (it means in particular, see Fig. 11, $\omega^+ = ACENT$ (resp. $\omega^- = -(AT ACENT)$), together with $\theta = 0$ at entrance (resp. exit) EFBs),
 - analytical method for calculation of the mid-plane field derivatives is used.

FFAG-SPI : Spiral FFAG magnet, N-tuple [28, 29]

FFAG-SPI works much like FFAG as to the field modelling, apart from the axial dependence of the field.

The *FFAG* procedure allows overlapping of fringe fields of neighboring dipoles, thus simulating in some sort the field in a dipole *N*-tuple - as for instance in an FFAG doublet or triplet (Fig. 30). This is done in the way described below.

The dimensionning of the magnet is defined by

AT: total angular aperture

RM: mean radius used for the positioning of field boundaries

For each one of the N=1 to (maximum) 5 dipoles of the N-tuple, the two effective field boundaries (entrance and exit EFBs) from which the dipole field is drawn are defined from geometric boundaries, the shape and position of which are determined by the following parameters

 ACN_i : arbitrary inner angle, used for EFB's positioning

 ω : azimuth of an EFB with respect to ACN

 ξ : spiral angle

with ACN_i and ω as defined in Fig. 30 (similar to what can be found in Figs. 29 and 11-A).

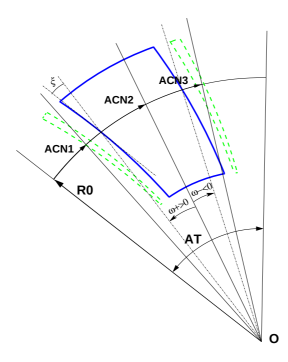


Figure 30: A N-tuple spiral sector FFAG magnet (N=3 here, simulating active field clamps at entrance and exit side of a central dipole).

Calculation of the field from a single dipole

The magnetic field is calculated in polar coordinates. At all (R, θ) in the median plane (z = 0), the magnetic field due a single one (index i) of the dipoles of a N-tuple FFAG magnet is written

$$B_{zi}(R,\theta) = B_{z0,i} \mathcal{F}_i(R,\theta) \left(R/R_M \right)^{K_i}$$

wherein $B_{z_{0,i}}$ is a reference field, at reference radius RM_i , whereas $\mathcal{F}(R,\theta)$ is calculated as described below.

Calculation of $\mathcal{F}_i(R,\theta)$

The fringe field coefficient $\mathcal{F}_i(R,\theta)$ associated with a dipole is computed as in the procedure *DIPOLES* (eq. 4.4.11), including radial dependence of the gap size

$$g(R) = g_0 \left(RM/R \right)^{\kappa} \tag{4.4.14}$$

so to simulate the effect of gap shaping of $B_{zi}(R,\theta)$ on field fall-off, over the all radial extent of a scaling FFAG dipole (with normally - but not in practice - $\kappa = K_i$).

Calculation of the full field from all N dipoles

For the rest, namely, calculation of the full field at particle position from the N dipoles, analytical calculation or numerical interpolation of the mid-plane field derivatives, extrapolation off median plane, etc., things are performed exactly as in the case of the DIPOLES procedure (see page 102).

MAP2D: 2-D Cartesian uniform mesh field map - arbitrary magnetic field [30]

MAP2D reads a 2-D field map that provides the three components B_X , B_Y , B_Z of the magnetic field at all nodes of a 2-D Cartesian uniform mesh in an (X,Y) plane. No particular symmetry is assumed, which allows the treatment of any type of field (e.g., dipole field with arbitrary Z elevation - the map needs not be a mid-plane map, solenoidal field, etc.). The field map data file has to be be filled with a format that fits the FORTRAN reading sequence (presumably compatible with TOSCA code outputs). The following is an instance, details and possible updates are to be found in the source file 'fmapw.f':

At each step of the trajectory of a particle, the field and its derivatives are calculated by a polynomial interpolation followed by a Z extrapolation (see sections 1.3.3, 1.4.3). Entrance and/or exit integration boundaries may be defined, in the same way as for *CARTEMES*.

MAP2D-E: 2-D Cartesian uniform mesh field map - arbitrary electric field

MAP2D-E reads a 2-D field map that provides the three components E_X , E_Y , E_Z of the electric field at all nodes of a 2-D Cartesian uniform mesh in an (X,Y) plane. No particular symmetry is assumed, which allows the treatment of any type of field (e.g., field) of a parallel-plate mirror with arbitrary Z elevation - the map needs not be a mid-plane map). The field map data file has to be be filled with a format that fits the FORTRAN reading sequence. The following is an instance, details and possible updates are to be found in the source file 'fmapw.f':

where IX (JY) is the number of longitudinal (transverse horizontal) nodes of the 2-D uniform mesh, Z(1) is the considered Z-elevation of the map. For binary files, FNAME must begin with 'E_' or 'b_', a flag 'BINARY' will thus be set to '.TRUE.'. The field $\vec{E} = (E_X, E_Y, E_Z)$ is next normalized with ENORM, prior to ray-tracing. As well the coordinates X, Y re normalized with X-, Y-NORM coefficients (usefull to convert to centimeters, the working units in **zgoubi**.

At each step of the trajectory of a particle, the field and its derivatives are calculated by a second or fourth degree polynomial interpolation followed by a Z extrapolation (see sections 1.3.3 page 25, 1.4.3 page 31). Interpolation grid is 3*3 for 2nd order (option IORDRE = 2) or 5*5 for 4th order (option IORDRE = 4).

Entrance and/or exit integration boundaries may be defined, in the same way as for CARTEMES.

MARKER: Marker

MARKER does nothing. Just a marker. No data.

As any other keyword, MARKER is allowed two LABELs. Using '.plt' as a second LABEL will cause storage of current coordinates into zgoubi.plt

TRANSMAT: Matrix transfer

TRANSMAT performs a matrix transfer of the particle coordinates in the following way

$$X_{i} = \sum_{j} R_{ij} X_{j}^{0} + \sum_{j,k} T_{ijk} X_{j}^{0} X_{k}^{0}$$

where, X_i stands for any of the current coordinates Y, T, Z, P, path length and dispersion, and X_i^0 stands for any of the initial coordinates. $[R_{ij}]$ ($[T_{ijk}]$) is the first order (second order) transfer matrix as usually involved in second order beam optics [19]. Second order transfer is optional. The length of the element represented by the matrix may be introduced for the purpose of path length updating. Note: MATRIX delivers $[R_{ij}]$ and $[T_{ijk}]$ matrices in a format suitable for straightforward use with TRANSMAT.

MULTIPOL: Magnetic multipole

The simulation of multipolar magnetic field \vec{M} by MULTIPOL proceeds by addition of the dipolar $(\vec{B1})$, quadrupolar $(\vec{B2})$, sextupolar $(\vec{B3})$, etc., up to 20-polar $(\vec{B10})$ components, and of their derivatives up to fourth order, following

$$\begin{split} \vec{M} &= \vec{B}1 + \vec{B}2 + \vec{B}3 + \ \dots \ + \vec{B}10 \\ \frac{\partial \vec{M}}{\partial X} &= \frac{\partial \vec{B}1}{\partial X} + \frac{\partial \vec{B}2}{\partial X} + \frac{\partial \vec{B}3}{\partial X} + \dots \ + \frac{\partial \vec{B}10}{\partial X} \\ \frac{\partial^2 \vec{M}}{\partial X \partial Z} &= \frac{\partial^2 \vec{B}1}{\partial X \partial Z} + \frac{\partial^2 \vec{B}2}{\partial X \partial Z} + \frac{\partial^2 \vec{B}3}{\partial X \partial Z} + \dots \ + \frac{\partial^2 \vec{B}10}{\partial X \partial Z} \\ \text{etc.} \end{split}$$

The independent components $\vec{B}1$, $\vec{B}2$, $\vec{B}3$, ..., $\vec{B}10$ and their derivatives up to the fourth order are calculated as described in section 1.3.6.

The entrance and exit fringe fields are treated separately. They are characterized by the integration zone X_E at entrance and X_S at exit, as for *QUADRUPO*, and by the extent λ_E at entrance, λ_S at exit. The fringe field extents for the dipole component are λ_E and λ_S . The fringe field for the quadrupolar (sextupolar, ..., 20-polar) component is given by a coefficient E_2 (E_3 , ..., E_{10}) at entrance, and S_2 (S_3 , ..., S_{10}) at exit, such that the extent is $\lambda_E * E_2$ ($\lambda_E * E_3$, ..., $\lambda_E * E_{10}$) at entrance and $\lambda_S * S_2$ ($\lambda_S * S_3$, ..., $\lambda_S * S_{10}$) at exit.

If $\lambda_E=0$ ($\lambda_S=0$) the multipole lens is considered to have a sharp edge field at entrance (exit), and then, X_E (X_S) is forced to zero (for the mere purpose of saving computing time). If $E_i=0$ ($S_i=0$) (i=2,10), the entrance (exit) fringe field for the multipole component i is considered as a sharp edge field. In sharp edge field model, the wedge angle vertical first order focusing effect (if $\vec{B}1$ is non zero) is simulated at magnet entrance and exit by a kick $P_2=P_1-Z_1\tan(\epsilon/\rho)$ applied to each particle (P_1,P_2 are the vertical angles upstream and downstream the EFB, Z_1 the vertical particle position at the EFB, ρ the local horizontal bending radius and ϵ the wedge angle experienced by the particle; ϵ depends on the horizontal angle T).

Overlapping of fringe fields inside the optical element is treated separately for each component, in the way described in *OUADRUPO*.

Any multipole component \vec{Bi} can be rotated independently by an angle RXi around the longitudinal X-axis, for the simulation of positioning defects, as well as skew lenses.

Magnet (mis-)alignement is assured by KPOS. KPOS also allows some degrees of automatic alignement useful for periodic structures (section 4.6.5).

OCTUPOLE: Octupole magnet (Fig. 31)

The meaning of parameters for OCTUPOLE is the same as for QUADRUPO. In fringe field regions the magnetic field $\vec{B}(X,Y,Z)$ and its derivatives up to fourth order are derived from the scalar potential approximated to the 8-th order in Y and Z

$$V(X,Y,Z) = \left(G - \frac{G''}{20} (Y^2 + Z^2) + \frac{G''''}{960} (Y^2 + Z^2)^2\right) (Y^3 Z - Y Z^3)$$
 with $G_0 = \frac{B_0}{R_0^3}$

Outside fringe field regions, or everywhere in sharp edge dodecapole ($\lambda_E = \lambda_S = 0$) , $\vec{B}(X,Y,Z)$ in the magnet is given by

$$B_X = 0$$

 $B_Y = G_0(3Y^2Z - Z^3)$
 $B_Z = G_0(Y^3 - 3YZ^2)$

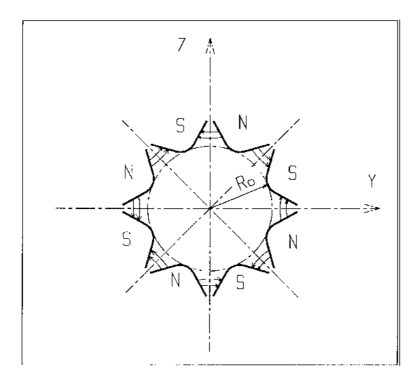


Figure 31: Octupole magnet

POISSON : Read magnetic field data from POISSON output

This keyword allows reading a field profile B(X) from *POISSON* output. Let *FNAME* be the name of this output file (normally, *FNAME* = outpoi.lis); the data are read following the *FORTRAN* statements hereunder

where X(I) is the longitudinal coordinate, and B(I) is the Z component of the field at a node (I) of the mesh. K's and R's are dummy variables appearing in the *POISSON* output file outpoi.lis but not used here.

From this field profile, a 2-D median plane map is built up, with a rectangular and uniform mesh; mid-plane symmetry is assumed. The field at each node (X_i, Y_j) of the map is $B(X_i)$, independent of Y_j (i.e., the distribution is uniform in the Y direction).

For the rest, *POISSON* works in a way similar to *CARTEMES*.

POLARMES : 2-D polar mesh magnetic field map

Similar to *CARTEMES*, apart from the polar mesh frame: IX is the number of angular nodes, JY the number of radial nodes; X(I) and Y(J) are respectively the angle and radius of a node (these parameters are similar to those entering in the definition of the map in DIPOLE-M).

PS170: Simulation of a round shape dipole magnet

PS170 is dedicated to a 'rough' simulation of CERN's PS170 dipole.

The field B_0 is constant inside the magnet, and zero outside. The pole is a circle of radius R_0 , centered on X axis. The output coordinates are generated at the distance XL from the entrance (Fig. 25).

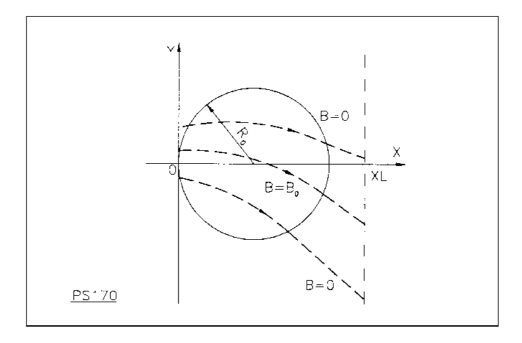


Figure 32: Scheme of the PS170 magnet simulation.

QUADISEX, SEXQUAD: Sharp edge magnetic multipoles

SEXQUAD defines in a simple way a sharp edge field with quadrupolar, sextupolar and octupolar components. QUADI-SEX adds a dipole component. The length of the element is XL. The vertical component $B \equiv B_Z(X,Y,Z=0)$ of the field and its derivatives in median plane are calculated at each step from the following expressions

$$\begin{split} B &= B_0 \left(U + \frac{N}{R_0} Y + \frac{B}{R_0^2} Y^2 + \frac{G}{R_0^3} Y^3 \right) \\ \frac{\partial B}{\partial Y} &= B_0 \left(\frac{N}{R_0} + 2 \frac{B}{R_0^2} Y + 3 \frac{G}{R_0^3} Y^2 \right) \\ \frac{\partial^2 B}{\partial Y^2} &= B_0 \left(2 \frac{B}{R_0^2} + 6 \frac{G}{R_0^3} Y \right) \\ \frac{\partial^3 B}{\partial Y^3} &= 6 B_0 \frac{G}{R_0^3} \end{split}$$

and then extrapolated out of the median plane by Taylor expansion in Z (see section 1.3.2).

With option SEXQUAD, U = 0, while with QUADISEX, U = 1.

QUADRUPO: Quadrupole magnet (Fig. 33)

The length of the magnet XL is the distance between the effective field boundaries (EFB). The field at the pole tip R_0 is B_0 .

The extent of the entrance (exit) fringe field is characterized by $\lambda_E(\lambda_S)$. The distance of ray-tracing on both sides of the EFB's, in the field fall off regions, will be $\pm X_E$ at the entrance, and $\pm X_S$ at the exit (Fig. 34), by prior and further automatic changes of frame.

In the fringe field regions $[-X_E, X_E]$ and $[-X_S, X_S]$ on both sides of the EFB's, $\vec{B}(X, Y, Z)$ and its derivatives up to fourth order are calculated at each step of the trajectory from the analytical expressions of the three components B_X , B_Y , B_Z obtained by differentiation of the scalar potential (see section 1.3.6) approximated to the 8th order in Y and Z.

$$\begin{split} V(X,Y,Z) &= \left(G - \frac{G''}{12} \left(Y^2 + Z^2\right) + \frac{G''''}{384} \left(Y^2 + Z^2\right)^2 - \frac{G''''''}{23040} \left(Y^2 + Z^2\right)^3\right) YZ \\ &\qquad \left(G'' = d^2G/dX^2, \ldots\right) \end{split}$$

where G is the gradient on axis [21]:

$$G(s) = \frac{G_0}{1 + \exp P(s)} \quad \text{with} \quad G_0 = \frac{B_0}{R_0}$$

and,

$$P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^2 + C_3 \left(\frac{s}{\lambda}\right)^3 + C_4 \left(\frac{s}{\lambda}\right)^4 + C_5 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^5 P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right)^5 P(s) =$$

where, s is the distance to the field boundary and λ stands for λ_E or λ_S (normally, $\lambda \simeq 2 * R_0$). When fringe fields overlap inside the magnet $(XL \le X_E + X_S)$, the gradient G is expressed as

$$G = G_E + G_S - 1$$

where, G_E is the entrance gradient and G_S is the exit gradient.

If $\lambda_E = 0$ ($\lambda_S = 0$), the field at entrance (exit) is considered as sharp edged, and then $X_E(X_S)$ is forced to zero (for the mere purpose of saving computing time).

Outside of the fringe field regions (or everywhere when $\lambda_E = \lambda_S = 0$) $\vec{B}(X,Y,Z)$ in the magnet is given by

$$B_X = 0$$

$$B_Y = G_0 Z$$

$$B_Z = G_0 Y$$

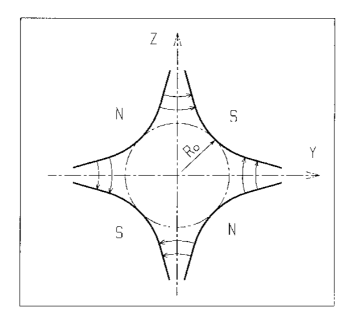


Figure 33: Quadrupole magnet

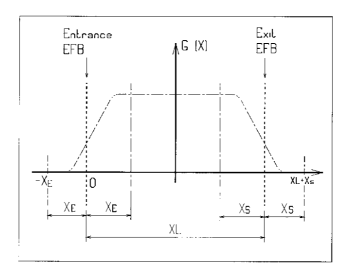


Figure 34: Scheme of the longitudinal field gradient G(X). (OX) is the longitudinal axis of the reference frame (0,X,Y,Z) of **zgoubi**. The length of the element is XL, but trajectories are ray-traced from $-X_E$ to $XL+X_S$, by means of prior and further automatic changes of frame.

SEPARA: Wien Filter - analytical simulation

SEPARA provides an analytic simulation of an electrostatic separator. Input data are the length L of the element, the electric field E and the magnetic field B. The mass m and charge q of the particles are entered by means of the keyword PARTICUL.

The subroutines involved in this simulation solve the following system of three equations with three unknown variables S, Y, Z (while $X \equiv L$), that describe the cycloidal motion of a particle in \vec{E}, \vec{B} static fields (Fig. 35).

$$X = -R\cos\left(\frac{\omega S}{\beta c} + \epsilon\right) - \frac{\alpha S}{\omega \beta c} + \frac{C_1}{\omega}$$
$$Y = R\sin\left(\frac{\omega S}{\beta c} + \epsilon\right) - \frac{\alpha}{\omega^2} - \frac{C_2}{\omega} + Y_0$$
$$Z = S\sin(P_0) + Z_0$$

where, S is the path length in the separator, $\alpha = -\frac{Ec^2}{\gamma}$, $\omega = -\frac{Bc^2}{m\gamma}$, $C_1 = \beta \sin(T_0)\cos(P_0)$ and $C_2 = \beta c\cos(T_0)\cos(P_0)$ are initial conditions. c = velocity of light, βc = velocity of the particle, $\gamma = (1-\beta^2)^{-\frac{1}{2}}$ and $\tan \epsilon = (C_2 + \frac{\alpha}{\omega})/C_1$. Y_0, T_0, Z_0, P_0 are the initial coordinates of the particle in the **zgoubi** reference frame. Here βc and γ are assumed constant, which is true as long as the change of momentum due to the electric field remains negligible all along the separator.

The index IA in the input data allows switching to inactive element (thus equivalent to ESL), horizontal or vertical separator. Normally, E, B and the value of β_W for wanted particles are related by

$$B(T) = -\frac{E\left(\frac{V}{m}\right)}{\beta_W \cdot c\left(\frac{m}{s}\right)}$$

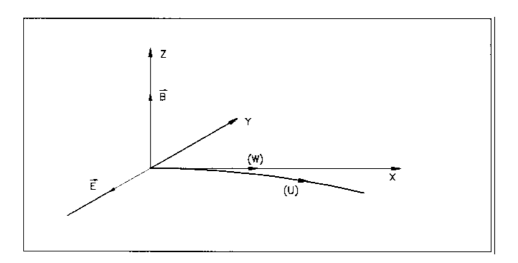


Figure 35: Horizontal separation between a wanted particle, (W), and an unwanted particle, (U). (W) undergoes a linear motion while (U) undergoes a cycloidal motion.

SEXTUPOL: Sextupole magnet (Fig. 36)

The meaning of parameters for SEXTUPOL is the same as for QUADRUPO.

In fringe field regions the magnetic field $\vec{B}(X,Y,Z)$ and its derivatives up to fourth order are derived from the scalar potential approximated to 7th order in Y and Z

$$\begin{split} V(X,Y,Z) &= \left(G - \frac{G''}{16} \left(Y^2 + Z^2\right) + \frac{G''''}{640} \left(Y^2 + Z^2\right)^2\right) \left(Y^2 Z - \frac{Z^3}{3}\right) \\ \text{with } G_0 &= \frac{B_0}{R_0^2} \end{split}$$

Outside fringe field regions, or everywhere in sharp edge sextupole ($\lambda_E = \lambda_S = 0$), $\vec{B}(X,Y,Z)$ in the magnet is given by

$$B_X = 0$$

$$B_Y = 2G_0YZ$$

$$B_Z = G_0(Y^2 - Z^2)$$

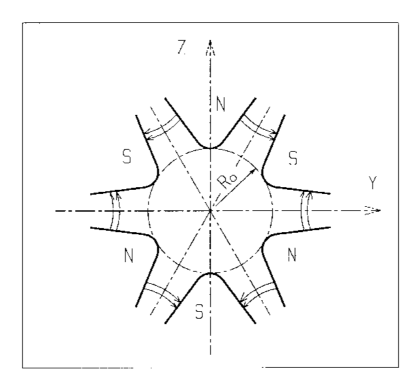


Figure 36: Sextupole magnet

SOLENOID: Solenoid (Fig. 37)

The solenoidal magnet has an effective length XL, a mean radius R_0 and an asymptotic field $B_0 = \mu_0 NI/XL$ (i.e., $\int_{-\infty}^{\infty} B_X(X,r) dX = \mu_0 NI$, $\forall r < R_0$), wherein B_X =longitudinal field component, NI = number of Ampere-Turns, $\mu_0 = 4\pi 10^{-7}$.

The distance of ray-tracing beyond the effective length XL, is X_E at the entrance, and X_S at the exit (Fig. 37).

The field $\vec{B}(X,r)$, $r=(Y^2+Z^2)^{1/2}$, and its derivatives up to the second order with respect to X,Y or Z are obtained after the method proposed in ref. [31], that involves the three complete elliptic integrals K,E and Π . These are calculated with the algorithm proposed in the same reference. Their derivatives are calculated by means of recursive relations [32].

This analytical model for the solenoidal field allows simulating an extended range of coil geometries (legnth and radius) provided that the coil thickness is small enough compared to the mean radius R_0 .

In particular the field on-axis writes (taking x = r = 0 as solenoid center)

$$B_X(x, r = 0) = \frac{\mu_0 NI}{2XL} \left[\frac{XL/2 - x}{\sqrt{(XL/2 - x)^2 + R_0^2}} + \frac{XL/2 + x}{\sqrt{(XL/2 + x)^2 + R_0^2}} \right]$$

and yields the magnetic length

$$L_{mag} \equiv \frac{\int_{-\infty}^{\infty} B_X(x, r < R_0) dx}{B_X(x = r = 0)} = XL \sqrt{1 + \frac{4R_0^2}{XL^2}} > XL$$

with in addition

$$B_X(\text{center}) \equiv B_X(x=r=0) = \frac{\mu_0 NI}{XL\sqrt{1+\frac{4R_0^2}{XL^2}}}.$$

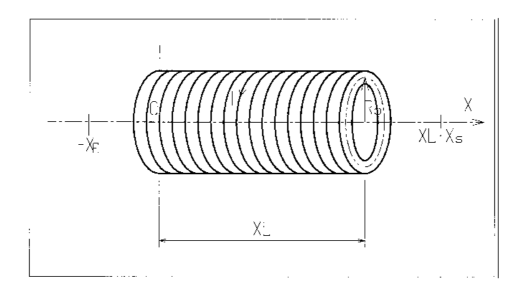


Figure 37: Solenoidal magnet.

TOSCA: 2-D and 3-D Cartesian or cylindrical mesh field map

TOSCA is dedicated to the reading and treatment of 2-D or 3-D Cartesian or cylindrical mesh field maps as delivered by the TOSCA magnet computer code standard output.

The total number of field data files to be read is given by the parameter IZ that appears in the data list following the keyword. Each file contains the field components B_X , B_Y , B_Z on an (X,Y) mesh at a given Z coordinate. IZ=1 for 2-D maps, and in this case B_X and B_Y are assumed zero all over the map⁷. For 3-D maps with mid-plane symmetry, $IZ \geq 2$, and thus, the first data file whose name follows in the data list is supposed to contain the median plane field (assuming Z=0 and $B_X=B_Y=0$), while the next file(s) contain the next maps in increasing Z order. For arbitrary 3-D maps (and in particular, contrary to what precedes without mid-plane symmetry assumption), following MOD value, see below, the total number of maps (whose names follow in the data list) is IZ, and map number [IZ/2]+1 is the Z=0 one.

The field map data file has to be be filled with a format that fits the *FORTRAN* reading sequence. The following is an instance, details and possible updates are to be found in the source file 'fmapw.f':

```
DO 1 K = 1, KZ

OPEN (UNIT = NL, FILE = FNAME, STATUS = 'OLD' [,FORM='UNFORMATTED'])

DO 1 J = 1, JY

DO 1 I = 1, IX

IF (BINARY) THEN

READ(NL) Y(J), Z(K), X(I), BY(J,K,I), BZ(J,K,I), BX(J,K,I)

ELSE

READ(NL,100) Y(J), Z(K), X(I), BY(J,K,I), BZ(J,K,I), BX(J,K,I)

100

FORMAT(1X,6E11.2)

ENDIF

1 CONTINUE
```

where, IX (JY, KZ) is the number of longitudinal (transverse horizontal, vertical) nodes of the 3-D uniform mesh. For binary files, FNAME must begin with 'B_' or 'b_', a flag 'BINARY' will thus be set to '.TRUE.'.

A flag MOD determines wether Cartesian or Z-axis cylindrical mesh is used. MOD can take various values depending also on the map data file formatting. (To be documented - see the FORTRAN subroutine FMAPW and its entries FMAPR, FMAPR2.)

The field $\vec{B} = (B_X, B_Y, B_Z)$ is normalized by means of *BNORM* in a similar way as in *CARTEMES*. As well the coordinates X (and Y, Z with 3-D field maps) is normalized with a X-[Y-,Z-]NORM coefficient (usefull to convert to centimeters, the working units in **zgoubi**.

At each step of the trajectory of a particle inside the map, the field and its derivatives are calculated

- in the case of 2-D map, by means of a second or fourth order polynomial interpolation, depending on IORDRE (IORDRE = 2, 25 or 4), as for CARTEMES,
- in the case of 3-D map, by means of a second order polynomial interpolation with a $3\times3\times3$ -point parallelipipedic grid, as described in section 1.4.4.

Entrance and/or exit integration boundaries between which the trajectories are integrated in the field may be defined, in the same way as in *CARTEMES*.

⁷Use MAP2D in case non-zero B_X , B_Y are to be taken into account in a 2-D map.

TRAROT: Translation-Rotation of the reference frame

UNDER DEVELOPEMENT. Check before use.

This procedure transports particles into a new frame by translation and rotation. Effect on spin tracking, particle decay and gas-scattering are taken into account (but not on synchrotron radiation).

UNDULATOR: Undulator magnet

UNDULATOR

To be documented

Figure 38: Undulator magnet.

UNIPOT: Unipotential cylindrical electrostatic lens

The lens is cylindrically symmetric about the X-axis.

The length of the first (resp. second, third) electrode is X1 (resp. X2, X3). The distance between the electrodes is D. The potentials are V1 and V2. The inner radius is R_0 (Fig. 39). The model for the electrostatic potential along the axis is [33]

$$V(x) = \frac{V2 - V1}{2\omega D} \left[\ln \frac{\cosh \frac{\omega \left(x + \frac{X2}{2} + D\right)}{R_0}}{\cosh \frac{\omega \left(x + \frac{X2}{2}\right)}{R_0}} + \ln \frac{\cosh \frac{\omega \left(x - \frac{X2}{2} - D\right)}{R_0}}{\cosh \frac{\omega \left(x - \frac{X2}{2}\right)}{R_0}} \right]$$

(x= distance from the center of the central electrode; $\omega=1,318$; cosh = hyperbolic cosine), from which the field $\vec{E}(X,Y,Z)$ and its derivatives are deduced following the procedure described in section ??. Use *PARTICUL* prior to *UNIPOT*, for the definition of particle mass and charge.

The total length of the lens is X1 + X2 + X3 + 2D; stepwise integration starts at entrance of the first electrode and terminates at exit of the third one.

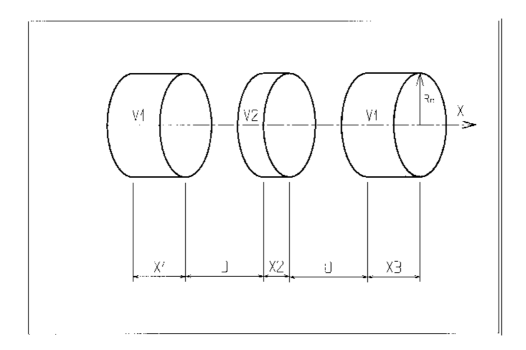


Figure 39: Three-electrode cylindrical unipotential lens.

VENUS: Simulation of a rectangular shape dipole magnet

VENUS is dedicated to a 'rough' simulation of Saturne Laboratory's VENUS dipole. The field B_0 is constant inside the magnet, with longitudinal extent XL and transverse extent $\pm YL$; outside these limits, $B_0=0$ (Fig. 40).

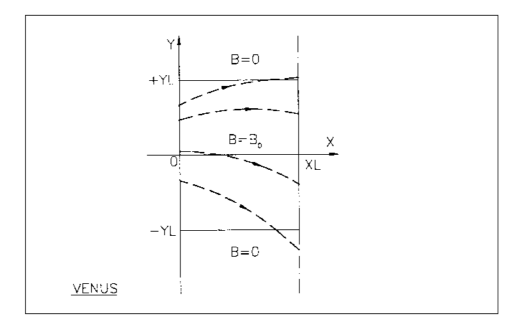


Figure 40: Scheme of VENUS rectangular dipole.

WIENFILT: Wien filter

WIENFILT simulates a Wien Filter, with transverse and orthogonal electric and magnetic fields \vec{E}_Y , \vec{B}_Z or \vec{E}_Z , \vec{B}_Y (Fig. 35). It must be preceded by PARTICUL for the definition of particle mass and charge.

The length XL of the element is the distance between its entrance and exit EFB's. The electric and magnetic field intensities E_0 and B_0 in the central, uniform field region, normally satisfy the relation

$$B_0 = -\frac{E_0}{\beta_W c}$$

for the selection of "wanted" particles of velocity $\beta_W c$. Ray-tracing in field fall-off regions extends over a distance X_E (X_S) beyond the entrance (exit) EFB by means of prior and further automatic changes of frame. Four sets of coefficients λ , C_0-C_5 allow the description of the entrance and exit fringe fields outside the uniform field region, following the model [21]

$$F = \frac{1}{1 + \exp(P(s))}$$

where P(s) is of the term

$$P(s) = C_0 + C_1 \left(\frac{s}{\lambda}\right) + C_2 \left(\frac{s}{\lambda}\right)^2 + C_3 \left(\frac{s}{\lambda}\right)^3 + C_4 \left(\frac{s}{\lambda}\right)^4 + C_5 \left(\frac{s}{\lambda}\right)^5$$

and s is the distance to the EFB. When fringe fields overlap inside the element (i.e., $XL \le X_E + X_S$), the field fall-off is expressed as

$$F = F_E + F_S - 1$$

where $F_E(F_S)$ is the value of the coefficient respective to the entrance (exit) EFB.

If $\lambda_E=0$ ($\lambda_S=0$) for either the electric or magnetic component, then both are considered as sharp edge fields and $X_E(X_S)$ is forced to zero (for the purpose of saving computing time). In this case, the magnetic wedge angle vertical first order focusing effect is simulated at entrance and exit by a kick $P_2=P_1-Z_1\tan(\epsilon/\rho)$ applied to each particle (P_1,P_2) are the vertical angles upstream and downstream the EFB, Z_1 the vertical particle position at the EFB, ρ the local horizontal bending radius and ϵ the wedge angle experienced by the particle; ϵ depends on the horizontal angle T). This is not done for the electric field, however it is advised not to use a sharp edge electric dipole model since this entails non symplectic mapping, and in particular precludes focusing effects of the non zero longitudinal electric field component.

$\mathbf{Y}\mathbf{M}\mathbf{Y}$: Reverse signs of Y and Z reference axes

YMY performs a 180° rotation of particle coordinates with respect to the X-axis, as shown in Fig. 41. This is done by means of a change of sign of Y and Z axes, and therefore coordinates, as follows

$$Y2 = -Y1$$
, $T2 = -T1$, $Z2 = -Z1$ and $P2 = -P1$

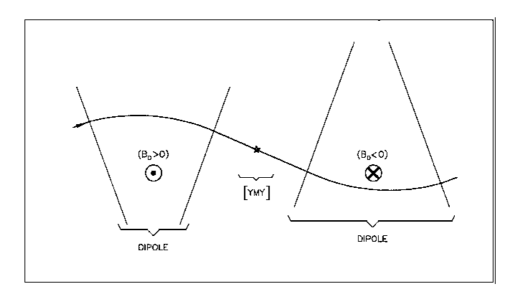


Figure 41: The use of YMY in a sequence of two identical dipoles of opposite signs.

4.5 Output Procedures

These procedures are dedicated to the printing of particle coordinates, histograms, spin coordinates, etc. They may be called for at any spot in the data pile.

FAISCEAU, FAISCNL, FAISTORE: Print/Store particle coordinates

FAISCEAU can be introduced anywhere in a structure. It produces a print (into zgoubi.res) of initial and actual coordinates of the IMAX particles at the location where it stands, together with their tagging indices and letters, following the same format as for FAISCNL (except for SORT(I) which is not printed).

FAISCNL has a similar effect, except that the information is stored in a dedicated file FNAME (advised name is FNAME = 'zgoubi.fai' (formatted write) or 'b_zgoubi.fai' (binary write) if post-processing with **zpop** should follow). This file may further on be read by means of OBJET, option KOBJ= 3, or used for other purposes such as graphics (see Part D of the Guide).

The data written to that file are formatted and ordered according to the *FORTRAN* sequence in the subroutine *impfai.f*, where details and possible updates are to found. The following is an instance :

```
OPEN (UNIT = NL, FILE = FNAME, STATUS = 'NEW')
        DO 1 I=1,IMAX
            P = BORO*CL9 *F(1,I) * AMQ(2,I)
            ENERG = SQRT(P*P + AMQ(1,I)*AMQ(1,I))
            ENEKI = ENERG - AMQ(1,I)
            WRITE(NFAI,110)
     1
            LET(I), IEX(I), -1.D0+FO(1,I), (FO(J,I),J=2,MXJ),
     2
            -1.D0+F(1,I),F(2,I),F(3,I),
     3
            (F(J,I),J=4,MXJ),ENEKI,
            I, IREP(I), SORT(I), (AMQ(J,I), J=1,5), RET(I), DPR(I), PS,
     5
            BORO, IPASS, KLEY, LBL1, LBL2, NOEL
        ENDDO
 110
        FORMAT(1X,1P,
C1
        LET(IT), KEX,
                          XXXO, (FO(J,IT), J=2,MXJ),
     1 A1, 1X, I2,
                           7E16.8,
C2
         XXX, Y, T*1.D3,
     2 /,3E24.16,
C3
        Z,P*1.D3,SAR,
                          TAR,
                                 ENEKI.
              /,4E24.16,E16.8,
        IT,IREP(IT), SORT(IT), (AMQ(J,I),J=1,5), RET(IT), DPR(IT), PS,
C4
                     9E16.8,
     4
       /,216,
C5
        BORO,
                 IPASS, KLEY, (LABEL(NOEL,I),I=1,2),NOEL
     5 /,E16.8, I6,1X, A8,1X, 2A8,
```

The meaning of the main data is the following (see the keyword *OBJET*)

```
: one-character string, for tagging particle number I
LET(I)
\mathbf{E}X, I, IREP(I)
                    : flag, particle number, index
FO(1-6,I)
                    : coordinates D, Y, T, Z, P and path length at the origin of the structure
F(1-6,I)
                    : idem, at the current position
SORT(I)
                    : path length at which the particle has possibly been stopped
                      (see CHAMBR or COLLIMA)
RET(I), DPR(I): synchrotron phase space coordinates; RET =phase (radian),
                      DPR = \text{momentum dispersion (MeV/c) (see } CAVITE)
IPASS
                    : turn number (see REBELOTE)
etc.
```

FAISTORE has an effect similar to FAISCNL, with two more features.

- On the first data line, *FNAME* may be followed by a series of up to 10 *LABEL*'s proper to the elements of the data file at the exit of which the print should occur; if there is no label, the print occurs by default at the location of *FAISTORE*; if there are labels the print occurs right downstream of all optical elements wearing those labels (and no longer at the *FAISTORE* location).
- The next data line gives a parameter, IP : printing will occur every IP other pass, if using REBELOTE with NPASS $\geq IP-1$.

For instance the following data input in zgoubi.dat:

```
FAISTORE
zgoubi.fai HPCKUP VPCKUP
12
```

will result in output prints into zgoubi.fai, every 12 other pass, each time elements of the zgoubi.dat data list labeled either *HPCKUP* or *VPCKUP* are encountered.

Note

Binary storage can be obtained from *FAISCNL* and *FAISTORE*. This for the sake of compactness and access speed, for instance in case voluminous amounts of data would have to be manipulated.

This is achieved by giving the storage file a name of the form *b_FNAME* or *B_FNAME* (*e.g.*, 'b_zgoubi.fai'). The *FORTRAN WRITE* list is the same as in the *FORMATTED* case above.

This is compatible with the READ statements in **zpop** that will recognize binary storage from that very radical 'b_' or 'B_'.

4.5 Output Procedures 143

FOCALE, IMAGE[S]: Particle coordinates and beam size; localization and size of horizontal waist

FOCALE calculates the dimensions of the beam and its mean transverse position, at a longitudinal distance XL from the position corresponding to the keyword FOCALE.

IMAGE computes the location and size of the closest horizontal waist.

IMAGES has the same effect as *IMAGE*, but, in addition, for a non-monochromatic beam it calculates as many waists as there are distinct momenta in the beam, provided that the object has been defined with a classification of momenta (see *OBJET*, *KOBJ*= 1, 2 for instance).

Optionally, for each of these three procedures, **zgoubi** can list a trace of the coordinates in the X, Y and in the Y, Z planes.

The following quantities are calculated for the N particles of the beam (IMAGE , FOCALE) or of each group of momenta (IMAGES)

• Longitudinal position:

$$\begin{split} \textit{FOCALE}: \quad X &= XL \\ \textit{IMAGE[S]}: \quad X &= -\frac{\sum_{i=1}^{N} Y_i * tgT_i - \left(\sum_{i=1}^{N} Y_i * \sum_{i=1}^{N} tgT_i\right) / N}{\sum_{i=1}^{N} tg^2T_i - \left(\sum_{i=1}^{N} tgT_i\right)^2 / N} \\ Y &= Y_1 + X * \mathsf{tg}T_1 \end{split}$$

where Y_1 and T_1 are the coordinates of the first particle of the beam (*IMAGE*, *FOCALE*) or the first particle of each group of momenta (*IMAGES*).

• Transverse position of the center of mass of the waist (IMAGE[S]) or of the beam (FOCALE), with respect to the reference trajectory

$$YM = \frac{1}{N} \sum_{i=1}^{N} (Y_i + X \operatorname{tg} T_i) - Y = \frac{1}{N} \sum_{i=1}^{N} Y M_i$$

ullet FWHM of the image (IMAGE[S]) or of the beam (FOCALE), and total width, respectively, W and WT

$$W = 2.35 \left(\frac{1}{N} \sum_{i=1}^{N} Y M_i^2 - Y M^2 \right)^{\frac{1}{2}}$$

$$WT = \max(YM_i) - \min(YM_i)$$

FOCALEZ, IMAGE[S]Z: Particle coordinates and beam size; localization and size of vertical waist

Similar to FOCALE and IMAGE[S], but the calculations are performed with respect to the vertical coordinates Z_i and P_i , in place of Y_i and T_i .

HISTO: 1-D histogram

Any of the coordinates used in **zgoubi** may be histogrammed, namely initial $Y_0, T_0, Z_0, P_0, S_0, D_0$ or actual Y, T, Z, P, S, D particle coordinates (S = path length; D may change in decay process simulation with MCDESINT, or when ray-tracing in \vec{E} fields), and also spin coordinates and modulus S_X, S_Y, S_Z and $\|\vec{S}\|$.

HISTO can be used in conjunction with MCDESINT, for statistics on the decay process, by means of TYP. TYP is a one-character variable. If it is set equal to 'S', only secondary particles will be histogrammed. If it is set equal to 'P', then only primary particles will be histogrammed. For no discrimination between S-econdary and P-rimary particles, TYP = 'Q' must be used.

The dimensions of the histogram (number of lines and columns) may be modified. It can be normalized with NORM = 1, to avoid saturation.

Histograms are indexed with the parameter NH. This allows making independent histograms of the same coordinate at several spots in a structure. This is also useful when piling up problems in an input data file (see also *RESET*). NH is in the range 1-5.

If REBELOTE is used, the statistics on the 1+NPASS runs in the structure will add up.

IMAGE[S][Z]: Localization and size of vertical waists

See FOCALE[Z].

4.5 Output Procedures 145

MATRIX: Calculation of transfer coefficients, periodic parameters

MATRIX causes the calculation of the transfer coefficients through the optical structure, from the OBJET up to the location where MATRIX is introduced in the structure, or, upon option, up to the horizontal focus closest to that location. In this last case the position of the focus is calculated automatically in the same way as the position of the waist in IMAGE. Depending on option IFOC, MATRIX also delivers the beam matrix and betatron phase advances or (case of periodic structure) periodic beam matrix and tunes, chromaticities and other global parameters.

Depending on the value of option IORD, different procedures follow

- If IORD = 0, MATRIX is inhibited (equivalent to FAISCEAU, whatever IFOC).
- If IORD = 1, the first order transfer matrix $[R_{ij}]$ is calculated, from a third order expansion of the coordinates. For instance

$$Y^{+} = \left(\frac{Y}{T_0}\right) T_0 + \left(\frac{Y}{T_0^2}\right) T_0^2 + \left(\frac{Y}{T_0^3}\right) T_0^3, \quad Y^{-} = -\left(\frac{Y}{T_0}\right) T_0 + \left(\frac{Y}{T_0^2}\right) T_0^2 - \left(\frac{Y}{T_0^3}\right) T_0^3$$

will yield, neglecting third order terms,

$$R_{11} = \left(\frac{Y}{T_0}\right) = \frac{Y^+ - Y^-}{2T_0}$$

In addition, if OBJET, KOBJ = 5.01 is used (hence introducing initial optical function values, $\alpha_{Y,Z}$, $\alpha_{Y,Z}$, $D_{Y,Z}$, $D'_{Y,Z}$), then, using the R_{ij} above, MATRIX will transport the optical functions and phase advances ϕ_Y , ϕ_Z , following

$$\begin{pmatrix} \beta \\ \alpha \\ \gamma \end{pmatrix}_{\text{ct MATRIX}} = \begin{pmatrix} R_{11}^2 & -2R_{11}R_{12} & R_{12}^2 \\ -R_{11}R_{21} & R_{12}R_{21} & R_{11}R_{12} \\ R_{21}^2 & -2R_{21}R_{22} & R_{22}^2 \end{pmatrix} \begin{pmatrix} \beta \\ \alpha \\ \gamma \end{pmatrix}_{\text{ct ORIET}}$$

$$\Delta\phi_{Y} = A \tan \frac{R_{12}}{(R_{11}\beta_{Y,objet} - R_{12}\alpha_{Y,objet})}, \quad \Delta\phi_{Z} = A \tan \frac{R_{34}}{(R_{33}\beta_{Z,objet} - R_{34}\alpha_{Z,objet})},$$

$$\phi_{Y,Z} \rightarrow \phi_{Y,Z} + 2\pi \quad \text{if } \phi_{Y,Z} < 0, \text{ given } [0,\pi] \text{Atan determination}$$

$$(4.5.1)$$

and print these out.

• If IORD = 2, fifth order Taylor expansions are used for the calculation of the first order transfer matrix $[R_{ij}]$ and of the second order matrix $[T_{ijk}]$. Other higher order coefficients are also calculated.

An automatic generation of an appropriate object for the use of MATRIX can be obtained by means

- if IORD = 1, of the procedure OBJET(KOBJ = 5[.I, I=1,9]) (pages 47, 219), that generates sets of up to 9*11 trajectories. In this case, up to nine matrices may be calculated, each one *wrt*. to the reference trajectory of concern as indicated using I in KOBJ = 5[.I, I=1,9];
- if IORD = 2, of the procedure OBJET(KOBJ = 6) that generates a set of 61 trajectories.

The next option, IFOC, acts as follows

- If IFOC = 0, the transfer coefficients are calculated at the location of MATRIX, and with respect to the reference trajectory. For instance, Y^+ and T^+ above are defined for particle number i as $Y^+ = Y^+(i) Y(Ref)$, and $T^+ = T^+(i) T(ref.)$.
- If IFOC = 1, the transfer coefficients are calculated at the horizontal focus closest to MATRIX (determined automatically), while the reference direction is that of the reference particle. For instance, Y^+ is defined for particle number i as $Y^+ = Y^+(i) Y_{\text{focus}}$, while T^+ is defined as $T^+ = T^+(i) T(\text{ref.})$).
- If IFOC = 2, no change of reference frame is performed : the coordinates refer to the current frame. Namely, $Y^+ = Y^+(i)$, $T^+ = T^+(i)$, etc.

Periodic structures

• If IFOC = 10 + NPeriod, MATRIX calculates periodic parameters characteristic of the structure such as optical functions and tune numbers, assuming that it is NPeriod-periodic; no change of reference is performed for these calculations. If IORD = 2 additional periodic parameters are computed such as chromaticities, beta-function momentum dependence, etc.

These quantities are derived from the first order perturbed and unperturbed transfer matrices as obtained in the way described above, and by identification $[R_{ij}] = I \cos \mu + J \sin \mu$.

Addition of *zgoubi.MATRIX.out* next to *IORD*, *IFOC* will cause stacking of *MATRIX* output data into zgoubi.MATRIX.out file (convenient for use with *e.g.* gnuplot type of data treatment software).

4.5 Output Procedures 147

PICKUPS: Beam centroid path; closed orbit

PICKUPS computes the beam centroid path, from average value of particle coordinates as observed at LABEL'ed keywords.

In conjunction with *REBELOTE*, this procedure computes by the same method the closed orbit in the periodic structure. The *LABEL* list of concern follows the keyword *PICKUPS*.

PLOTDATA: Intermediate output for the PLOTDATA graphic software [34]

To be documented

SPNPRNL, SPNSTORE, SPNPRT: Print/Store spin coordinates

SPNPRT can be introduced anywhere in a structure. It produces a listing (into zgoubi.res) of the initial and actual coordinates and modulus of the spin of the IMAX particles, at the location where it stands, together with their Lorentz factor γ , etc. The mean values of the spin components are also printed.

SPNPRNL has similar effect to SPNPRT, except that the information is stored in a dedicated file FNAME (should post-processing with **zpop** follow, advised name is FNAME = 'zgoubi.spn' (formatted write) or 'b_zgoubi.spn' (binary write)). The data are formatted and ordered according to the FORTRAN sequence found in the subroutine spnprn.f, with meaning of printed quantities as follows:

LET(I),IEX(I): tagging character and flag (see *OBJET*)

SI(1-4,I): spin components SX, SY, SZ and modulus, at the origin

SF(1-4,I) : idem, at the current position
GAMMA : Lorentz relativistic factor
I : particle number

IMAX : total number of particles ray-traced (see *OBJET*)

IPASS : turn number (see *REBELOTE*)

SPNSTORE has an effect similar to SPNPRNL, with two more features.

- On the first data line, *FNAME* may be followed by a series of up to 10 *LABEL*'s proper to the elements of the zgoubi.dat data file at the exit of which the print should occur; if no label is given, the print occurs by default at the very location of *SPNSTORE*; if there label(s) is (are) given print occurs right downstream of all optical elements wearing those labels (and no longer at the *SPNSTORE* location).

- The next data line gives a parameter, IP: printing will occur every IP other pass, when using REBELOTE with NPASS > IP - 1.

For instance the following data input in zgoubi.dat:

```
SPNSTORE zgoubi.spn HPCKUP VPCKUP 12
```

will result in output prints into zgoubi.spn, every 12 other pass, each time elements of the zgoubi.dat data list labeled either *HPCKUP* or *VPCKUP* are encountered.

Note

Binary storage can be obtained from *SPNPRNL* and *SPNSTORE*. This for the sake of compactness and I/O access speed by zgoubi or zpop, for instance in case voluminous amounts of data should be manipulated.

This is achieved by giving the storage file a name of the form *b_FNAME* or *B_FNAME* (*e.g.*, 'b_zgoubi.spn'). The *FORTRAN WRITE* ouput list is the same as in the *FORMATTED* case above.

SRPRNT: Print SR loss statistics

SRPRNT may be introduced anywhere in a structure. It produces a listing (into zgoubi.res) of current state of statistics on several parameters related to SR loss presumably activated beforehand with keyword *SRLOSS*.

4.5 Output Procedures 151

TWISS: Calculation of optical parameters; periodic parameters

TWISS causes the calculation of transport coefficients and various other parameters, in particular periodical quantities such as tunes, chromaticies, etc.

the object necessary for these calculations will be generated automatically if one uses *OBJET* with option *KOBJ*= 5.

The way *KTW* works is the following:

- It assumes that the reference particle (e.g., particle #1 of 11, when using OBJET, KOBJ=5) is located on the closed orbit. This condition has to be satisfied for TWISS to work consistently.
- A first pass through the structure allows computing the periodic beam matrix from the rays, including the periodic dispersions.
- The periodic dispersions are used to define chromatic closed orbits at $\pm \delta p/p$. A second and a third pass with chromatic objects centered respectively on $\pm \delta p/p$ chromatic orbits will compute the chromatic first order transport matrices. From these the chromaticities are deduced.

4.6 Complementary Features

4.6.1 Backward Ray-tracing

For the purpose of parameterization for instance, it may be interesting to ray-trace backward from the image toward the object. This can be performed by first reversing the position of optical elements in the structure, and then reversing the integration step sign in all the optical elements.

An illustration of this feature is given in the following Figure 42.

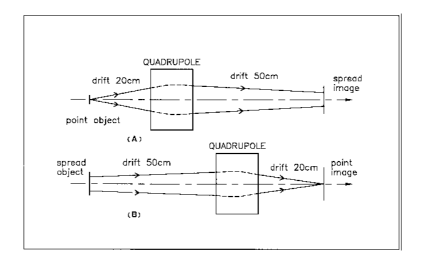


Figure 42: A. Regular forward ray-tracing, from object to image.

B. Same structure, with backward ray-tracing from image to object: negative integration step XPAS is used in the quadrupole.

4.6.2 Checking Fields and Trajectories inside Optical Elements

In all optical elements, an option index IL is available. It is normally set to 0 and in this case has no effect.

IL=1 causes a print in zgoubi.res of particle coordinates and field along trajectories in the optical element. In the meantime, a calculation and summation of the values of $\vec{\nabla} \cdot \vec{B}$, $\vec{\nabla} \times \vec{B}$ and $\nabla^2 \vec{B}$ (same for \vec{E}) at all integration steps is performed, which allows a check of the behavior of \vec{B} (or \vec{E}) in field maps (all these derivatives should normally be zero). IL=2 causes a print of particle coordinates and other informations in zgoubi.plt at each integration step; this information can further be processed with \mathbf{zpop}^8 . In order to limit the volume of that storage file (when dealing with small step size, large number of particles, etc.) it is possible to print out every other 10^n integration step by taking $IL=2\times10^n$ (for instance, IL=200 would cause output into zgoubi.plt every 100 other step).

When dealing with maps (e.g., CARTEMES, ELREVOL), another option index IC is available. It is normally set to 0 and in this case has no effect.

IC = 1 causes a print of the field map in zgoubi.res.

IC = 2 will cause a print of field maps in zgoubi.map which can further be processed with **zpop**.

⁸See Part D of the Guide.

4.6.3 Labeling keywords

Keywords in **zgoubi** data file zgoubi.dat can be *LABEL*'ed, for the purpose of the execution of such procedures as *PICKUPS*, *FAISCNL*, *FAISTORE*, *SCALING*, and also for the purpose of particle coordinate storage into zgoubi.plt (see Sections 4.6.2 and 2).

Each keyword accepts two *LABEL*'s, of which the first one is used for the above mentioned purposes. The keyword and related *LABEL*['s] should fit within a 80-character long string on a single line.

4.6.4 Multiturn tracking in circular machines

Multiturn tracking in circular machines can be performed by means of the keyword *REBELOTE*, put at the end of the optical structure with its argument NPASS+1 being the number of turns to be performed. In order that the IMAX particles of the beam start a new turn with the coordinates they have reached at the end of the previous one, the option K=99 has to be specified in REBELOTE.

Synchrotron acceleration can be simulated, following the procedure below

- CAVITE appears at the end of the structure (before REBELOTE), with option IOPT = 1
- the R.F. frequency of the cavity is given a timing law by means of SCALING, family CAVITE
- the magnets are given the same timing law $B\rho(T)$, (where T=1 to NPASS+1 is the turn number) by means of SCALING.

Eventually some families of magnets may be given a law which does not follow $B\rho(T)$, for the simulation of special processes (e.g., fast crossing of spin resonances with independent families of quadrupoles).

4.6.5 Positioning, (mis-)alignement, of optical elements and field maps

The last record in most optical elements and field maps is the positioning flag KPOS, followed by the parameterss XCE, YCE for translation and ALE for rotation. The positioning works in two different ways, depending whether they are defined in Cartesian (X, Y, Z) coordinates (e.g., QUADRUPO, TOSCA), or polar (R, θ, Z) coordinates (DIPOLE).

Cartesian Coordinates:

If KPOS = 1, the optical element is moved (shifted by XCE, YCE and Z-rotated by ALE) with respect to the incoming reference frame. Trajectory coordinates after traversal of the element refer the element frame.

If KPOS = 2, the shifts XCE and YCE, and the tilt angle ALE are taken into account, for mis-aligning the element with respect to the incoming reference, as shown in Fig. 43. The effect is equivalent to a CHANGREF(XCE,YCE,ALE) upstream of the optical element, followed by CHANGREF(XCS,YCS,ALS=-ALE) downstream of it, with computed XCS, YCS values as schemed in Fig. 43.

KPOS = 3 option is available for a number of magnets (e.g., BEND, MULTIPOL, AGSMM (AGS main magnet)); it is effective only if a non zero dipole component B1 is present, or if ALE is non-zero. It positions automatically the magnet in a symmetric manner with respect to the incoming and outgoing reference axis, convenient for periodic structures, as follows (Fig 44).

Both incoming and outgoing refernce frames are tilted w.r.t. the magnet,

- either, by an angle ALE if $ALE \neq 0$,
- or, if ALE=0 by half the Z-rotation $\theta_Z/2$ (such that $L=2\frac{BORO}{B1}\sin(\theta_Z/2)$ wherein L = geometrical length, BORO= reference rigidity as defined in OBJET).

Next, the optical element is Y-shifted by YCE (XCE is not used) in a direction orthogonal to the new magnet axis (i.e., at an angle $ALE + \pi/2$ wrt. the X axis of the incoming reference frame).

KPOS=4 applies to AGSMM (AGS main magnet). By default, it aligns the magnet in a way similar to KPOS=3, with reference frame Z-rotated by $\theta_Z/2$ as drawn from $L=2\frac{BORO}{B1}\sin(\theta_Z/2)$. However magnet mis-alignement (alignement errors) are handled in a specific way, as follows.

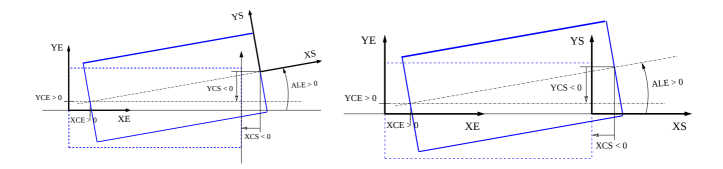


Figure 43: (X_E, Y_E) and (X_S, Y_S) are respectively the incoming and outgoing reference frames. Left: moving an optical element using KPOS = 1. Right: Mis-aligning an optical element using KPOS = 2.

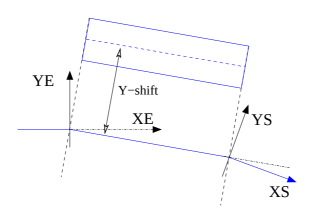


Figure 44: (X_E,Y_E) and (X_S,Y_S) are respectively the incoming and outgoing reference frames. Half-deviation alignement of a Cartesian coordinate bending element, using KPOS=3.

All 6 types of misalignements, namely, X-, Y-, Z-shift, X-, Y-, Z-rotation, can be accounted for, in an arbitrary order. They are specified using the "new style" *CHANGREF* method as described in page 92. Longitudinal rotation "XR" is taken *wrt*. the logitudinal axis, whereas radial and axial rotations, "YR" and "ZR", are taken around an axis going through the center of the magnet. Transformations are as follows, see Fig. 46:

ullet Y-rotation (pitch) by an angle φ : new coordinates (at M) as well as path lengthening, etc, derive from old ones (at M_0) following

 $X_{new} = 0$ by definition,

 $Y_{new} = Y_{old} + dS \cos P \sin T$, dS is the path lengthening, given below,

 $Z_{new} = Z_{old} \cos p / \cos(p - \varphi)$, ensuing from $\varphi = \frac{\pi}{2} - p$, and $\tan p = \tan P / \cos T$ (since $om \cos p = OM \cos P \cos T$ as well as $om \sin p = OM \sin P$),

 $dS = dL/\cos P/\cos T \text{ with } dL = sign(dL, -Z_{new}\varphi)\sqrt{Z_{new}^2\sin^2\varphi + (Z_{new}\cos\varphi - Z_{old})^2}.$

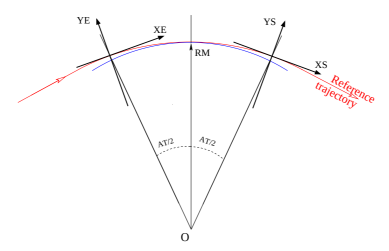


Figure 45: Positioning of a polar field map, using KPOS = 1.

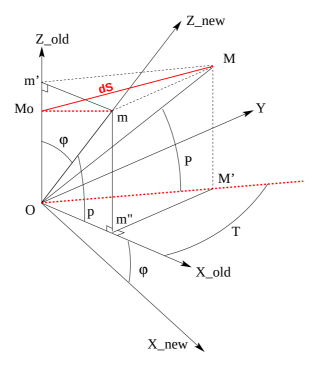


Figure 46: Pitch angle, φ , in the "YR" type of rotation, using KPOS=4. M is the new position, in the rotated plane (Y,Z_{new}) , of a particle with velocity $\vec{v}//\vec{M_o}M$ located at former position M_o in the old, (Y,Z_{old}) , plane. M', m, m' are projections of M, m'' is projection of M'. T and P are the horizontal and vertical angles as defined in Fig. 1. p is the projection of P.

Polar Coordinates

If KPOS = 1, the element is positioned automatically in such a way that a particle entering with zero initial coordinates and $1 + DP = B\rho/BORO$ relative momentum will reach position $(RM, \frac{AT}{2})$ in the element with T = 0 angle with respect to the moving frame in the polar coordinates system of the element (Fig. 45; see DIPOLE-M and POLARMES). If KPOS = 2, the map is positioned in such a way that the incoming reference frame is presented at radius RE with angle TE. The exit reference frame of **zgoubi** is positioned in a similar way with respect to the map, by means of the two parameters RS (radius) and TS (angle) (see Fig. 11A.).

4.6.6 Coded integration step

In several optical elements (e.g., all multipoles, BEND) the integration step (in general noted XPAS) can be coded under the form $XPAS = \#E \mid C \mid S$, where E is the number of steps taken in the entrance fringe field, C is the number of steps in the magnet body, and S is the number of steps in the exit fringe field.

4.6.7 Ray-tracing of an arbitrarily large number of particles

Monte Carlo multiparticle simulations involving an arbitrary number of particles can be performed by means of *REBE-LOTE*, put at the end of the optical structure, with its argument *NPASS* being the number of passes through *REBELOTE*, and (NPASS+1) * MAX the number of particles to be ray-traced. In order that new initial conditions (D, Y, T, Z, P, X) be generated at each pass, K=0 has to be specified in *REBELOTE*.

Statistics on coordinates, spins, and other histograms can be performed by means of such procedures as *HISTO*, *SPNTRK*, etc. that stack the information from pass to pass.

4.6.8 Stopped particles: the **EX** flag

As described in *OBJET*, each particle I=1, *MAX* is attached a value $I\!\!E\!X(I)$ of the $I\!\!E\!X$ flag. Normally, $I\!\!E\!X(I)=1$. Under certain circumstances, $I\!\!E\!X$ may take negative values, as follows

- -1: the trajectory happened to wander outside the limits of a field map
- -2: too many integration steps in an optical element
- -3: deviation happened to exceed $\frac{\pi}{2}$ in an optical element
- -4: stopped by walls (procedures CHAMBR, COLLIMA)
- -5: too many iterations in subroutine *DEPLA*
- -6: energy loss exceeds particle energy
- -7: field discontinuities larger than 50% wthin a field map
- -8: reached field limit in an optical element

Only in the case $I\!E\!X = -1$ will the integration not be stopped since in this case the field outside the map is extrapolated from the map data, and the particle may possibly get back into the map (see section 1.4.2 on page 28). In all other cases the particle of concern will be stopped.

4.6.9 Negative rigidity

zgoubi can handle negative rigidities $B\rho = p/q$. This is equivalent to considering either particles of negative charges (q < 0), or counter going particles (p < 0), or virtually reversed fields (w.r.t. the field sign that shows in the optical element data list).

Negative rigidities may be specified in terms of BORO < 0 or $D = B\rho/BORO < 0$ when defining the initial coordinates with OBJET and MCOBJET.

PART B

Keywords and input data formatting

Glossary of Keywords

AGSMM	AGS main magnet	165
AGSQUAD	AGS quadrupole	166
AIMANT	Generation of dipole mid-plane 2-D map, polar frame	167
AUTOREF	Automatic transformation to a new reference frame	171
BEND	Bending magnet, Cartesian frame	172
BINARY	BINARY/FORMATTED data converter	173
BREVOL	1-D uniform mesh magnetic field map	174
CARTEMES	2-D Cartesian uniform mesh magnetic field map	175
CAVITE	Accelerating cavity	177
CHAMBR	Long transverse aperture limitation	178
CHANGREF	Transformation to a new reference frame	179
CIBLE	Generate a secondary beam from target interaction	180
COLLIMA	Collimator	181
DECAPOLE	Decapole magnet	182
DIPOLE	Dipole magnet, polar frame	183
DIPOLE-M	Generation of dipole mid-plane 2-D map, polar frame	185
DIPOLES	Dipole magnet N-tuple, polar frame	187
DODECAPO	Dodecapole magnet	189
DRIFT	Field free drift space	190
EBMULT	Electro-magnetic multipole	191
EL2TUB	Two-tube electrostatic lens	193
ELMIR	Electrostatic N-electrode mirror/lens, straight slits	194
ELMIRC	Electrostatic N-electrode mirror/lens, circular slits	195
ELMULT	Electric multipole	196
ELREVOL	1-D uniform mesh electric field map	197
EMMA	2-D Cartesian or cylindrical mesh field map for EMMA FFAG	
END	End of input data list; see FIN	203
ESL	Field free drift space	190
FAISCEAU	Print particle coordinates	199
FAISCNL	Store particle coordinates in file FNAME	199
FAISTORE	Store coordinates every IP other pass at labeled elements	199
FFAG	FFAG magnet, N-tuple	200
FFAG-SPI	Spiral FFAG magnet, N-tuple	202
FIN	End of input data list	203
FIT	Fitting procedure	
FIT2	Fitting procedure	
FOCALE	Particle coordinates and horizontal beam dimension at distance XL	
FOCALEZ	Particle coordinates and vertical beam dimension at distance XL	
GASCAT	Gas scattering	207
HISTO	1-D histogram	208

IMAGE	Localization and size of horizontal waist	. 209
IMAGES	Localization and size of horizontal waists	. 209
IMAGESZ	Localization and size of vertical waists	. 209
IMAGEZ	Localization and size of vertical waist	. 209
MAP2D	2-D Cartesian uniform mesh field map - arbitrary magnetic field	. 210
MAP2D-E	2-D Cartesian uniform mesh field map - arbitrary electric field	. 211
MARKER	Marker	. 212
TRANSMAT	Matrix transfer	. 248
MATRIX	Calculation of transfer coefficients, periodic parameters	. 213
MCDESINT	Monte-Carlo simulation of in-flight decay	. 214
MCOBJET	Monte-Carlo generation of a 6-D object	. 215
MULTIPOL	Magnetic multipole	.218
OBJET	Generation of an object	. 219
OBJETA	Object from Monte-Carlo simulation of decay reaction	. 221
OCTUPOLE	Octupole magnet	. 222
OPTICS	Write out optical functions	. 223
ORDRE	Taylor expansions order	. 224
PARTICUL	Particle characteristics	. 225
PICKUPS	Beam centroid path; closed orbit	. 226
PLOTDATA	Intermediate output for the PLOTDATA graphic software	. 227
POISSON	Read magnetic field data from POISSON output	. 228
POLARMES	2-D polar mesh magnetic field map	. 229
PS170	Simulation of a round shape dipole magnet	. 230
QUADISEX	Sharp edge magnetic multipoles	. 231
QUADRUPO	Quadrupole magnet	.232
REBELOTE	'Do it again'	. 234
RESET	Reset counters and flags	
SCALING	Time scaling of power supplies and R.F	. 236
SEPARA	Wien Filter - analytical simulation	. 237
SEXQUAD	Sharp edge magnetic multipole	. 238
SEXTUPOL	Sextupole magnet	. 239
SOLENOID	Solenoid	.240
SPNPRNL	Store spin coordinates into file FNAME	. 241
SPNSTORE	Store spin coordinates every <i>IP</i> other pass at labeled elements	.241
SPNPRT	Print spin coordinates	. 241
SPNTRK	Spin tracking	. 242
SRLOSS	Synchrotron radiation loss	.243
SRPRNT	Print SR loss statistics	. 244
SYNRAD	Synchrotron radiation spectral-angular densities	. 245
TARGET	Generate a secondary beam from target interaction; see CIBLE	. 180
TOSCA	2-D and 3-D Cartesian or cylindrical mesh field map	
TRAROT	Translation-Rotation of the reference frame	
TWISS	Calculation of optical parameters; periodic parameters	
UNDULATOR	Undulator magnet	
UNIPOT	Unipotential cylindrical electrostatic lens	
VENUS	Simulation of a rectangular shape dipole magnet	
WIENFILT	Wien filter	
YMY	Reverse signs of Y and Z reference axes	

Optical elements versus keywords

This glossary gives a list of keywords suitable for the simulation of common optical elements. These are classified in three categories: magnetic, electric and combined electro-magnetic elements.

Field map procedures are also listed; they provide a means for ray-tracing through measured or simulated electric and/or magnetic fields.

MAGNETIC ELEMENTS

AGS main magnet AGSMM

Decapole DECAPOLE, MULTIPOL

Dipole[s] AIMANT, BEND, DIPOLE[S], DIPOLE-M, MULTIPOL, QUADISEX

Dodecapole DODECAPO, MULTIPOL

FFAG magnets DIPOLES, FFAG, FFAG-SPI, MULTIPOL, EMMA

Helical dipole HELIX

Multipole MULTIPOL, QUADISEX, SEXQUAD

Octupole OCTUPOLE, MULTIPOL, QUADISEX, SEXQUAD

Quadrupole QUADRUPO, MULTIPOL, SEXQUAD

Sextupole SEXTUPOL, MULTIPOL, QUADISEX, SEXQUAD

Skew multipoles MULTIPOL
Solenoid SOLENOID
Undulator UNDULATOR

Using field maps

1-D, cylindrical symmetry BREVOL

2-D, mid-plane symmetry CARTEMES, POISSON, TOSCA

2-D, no symmetry MAP2D
2-D, polar mesh, mid-plane symmetry POLARMES
3-D, no symmetry TOSCA

ELECTRIC ELEMENTS

2-tube (bipotential) lens **EL2TUB** 3-tube (unipotential) lens **UNIPOT** Decapole **ELMULT** Dipole **ELMULT** Dodecapole **ELMULT** Multipole **ELMULT** N-electrode mirror/lens, straight slits **ELMIR** N-electrode mirror/lens, circular slits **ELMIRC** Octupole **ELMULT** Quadrupole **ELMULT** R.F. (kick) cavity **CAVITE** Sextupole **ELMULT** Skew multipoles **ELMULT**

Using field maps

1D, cylindrical symmetry ELREVOL

2-D, no symmetry MAP2D-E

ELECTRO-MAGNETIC ELEMENTS

Decapole **EBMULT** Dipole **EBMULT** Dodecapole **EBMULT** Multipole **EBMULT** Octupole EBMULT Quadrupole **EBMULT** Sextupole **EBMULT** Skew multipoles **EBMULT**

Wien filter SEPARA, WIENFILT

INTRODUCTION

Here after is given a detailed description of input data formatting and units. All available keywords appear in alphabetical order.

Keywords are read from the input data file by an unformatted *FORTRAN READ* statement. They may therefore need be enclosed between quotes (*e.g.*, '*DIPOLE*').

Text string data such as comments or file names, are read by formatted READ statements. Therefore no quotes are needed. Numerical variables and indices are read by unformatted READ. It may therefore be necessary that integer variables be assigned an integer value.

In the following tables

- the first column states the input numerical variables, indices and text strings,
- the second column gives brief explanations,
- the third column gives the units or ranges of the input variables and indices,
- the fourth column indicates whether the inputs are integers (I), reals (E) or text strings (A). For example, 'I, 3*E' means that one integer followed by 3 reals must be entered. 'A80' means that a text string of maximum 80 characters must be entered.

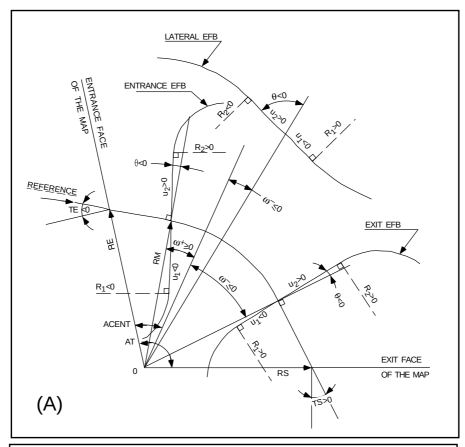
AGSMM	AGS main magnet		
IL	IL = 1, 2: print field and coordinates along trajectories	0-2	I
$XL, R_0, B1, B2,, B10,$	Length of element; radius at pole tip; field at pole tip for dipole, quadrupole,, dodecapole components	2*cm,10*kG	12*E
$X_E, \lambda_E, E_2,, E_{10}$	Entrance face Integration zone ; fringe field extent : dipole fringe field extent = λ_E ; quadrupole fringe field extent = $\lambda_E * E_2$; 20-pole fringe field extent = $\lambda_E * E_{10}$ (sharp edge if field extent is zero)	2*cm,9*no dim.	11*E
NCE , $C_0 - C_5$	same as QUADRUPO	0-6, 6*no dim.	I, 6*E
$X_S, \lambda_S, S_2,, S_{10}$ $NCS, C_0 - C_5$	Exit face Integration zone; as for entrance	2*cm, 9*no dim. 0-6, 6*no dim.	11*E I, 6*E
R1, R2, R3,, R10	Skew angles of field components	10*rad	10*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	$KPOS=1$: element aligned, 2: misaligned; shifts, tilt (unused if $KPOS=1$) for $QUADRUPO$. $KPOS=3$: effective only if $B1 \neq 0$: entrance and exit frames are shifted by YCE and tilted wrt . the magnet by an angle of \bullet either ALE if $ALE\neq 0$ \bullet or $2 Arcsin(B1 XL / 2BORO)$ if $ALE=0$	1-2, 2*cm, rad	I, 3*E

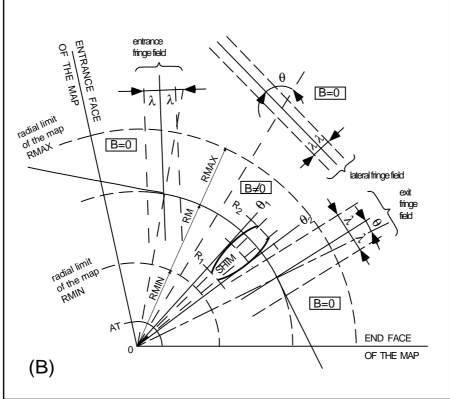
AGSQUAD	AGS quadrupole		
IL	$I\!L=1,2$: print field and coordinates along trajectories	0-2	I
$XL, R_0, B1, B2,, B10,$	Length of element; radius at pole tip; field at pole tip for dipole, quadrupole,, dodecapole components	2*cm,10*kG	12*E
$X_E, \lambda_E, E_2,, E_{10}$	Entrance face Integration zone ; fringe field extent : dipole fringe field extent = λ_E ; quadrupole fringe field extent = $\lambda_E * E_2$;	2*cm,9*no dim.	11*E
	20-pole fringe field extent = $\lambda_E * E_{10}$ (sharp edge if field extent is zero)		
NCE , $C_0 - C_5$	same as QUADRUPO	0-6, 6*no dim.	I, 6*E
$X_S, \lambda_S, S_2,, S_{10}$	Exit face Integration zone; as for entrance	2*cm, 9*no dim.	11*E
NCS , $C_0 - C_5$		0-6, 6*no dim.	I, 6*E
R1, R2, R3,, R10	Skew angles of field components	10*rad	10*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	$KPOS=1$: element aligned, 2: misaligned; shifts, tilt (unused if $KPOS=1$) for $QUADRUPO$. $KPOS=3$: effective only if $B1 \neq 0$: entrance and exit frames are shifted by YCE and tilted wrt . the magnet by an angle of • either ALE if $ALE\neq 0$ or $2 Arcsin(B1\ XL\ /\ 2BORO)$ if $ALE=0$	1-2, 2*cm, rad	I, 3*E

AIMANT	Generation of dipole mid-plane 2-D map, polar frame $B_Z = \mathcal{F}B_0 \left(1 - N\left(\frac{R-RM}{RM}\right) + B\left(\frac{R-RM}{RM}\right)^2 + G\left(\frac{R-RM}{RM}\right)^3\right)$)	
NFACE, IC, IL	Number of field boundaries $IC = 1, 2$: print field map $IL = 1, 2$: print field and coordinates on trajectories	2-3, 0-2, 0-2	3*I
IAMAX, IRMAX	Azimuthal and radial number of nodes of the mesh	$\leq 400, \leq 10^4$	2*I
B_0 , N , B , G	Field and field indices	kG, 3*no dim.	4*E
AT, ACENT, RM, RMIN, RMAX	Mesh parameters: total angle of the map; azimuth for EFBs positioning; reference radius; minimum and maximum radii	2*deg, 3*cm	5*E
	ENTRANCE FIELD BOUNDARY		
λ, ξ	Fringe field extent (normally \simeq gap size); flag: - if $\xi \geq 0$: second order type fringe field with linear variation over distance ξ - if $\xi = -1$: exponential type fringe field: $F = (1 + \exp(P(s)))^{-1}$ $P(s) = C_0 + C_1(\frac{s}{\lambda}) + C_2(\frac{s}{\lambda})^2 + + C_5(\frac{s}{\lambda})^5$	cm, (cm)	2*E
NC , $C_0 - C_5$, shift	NC = 1 + degree of $P(s)$; C_0 to C_5 : see above; EFB shift (ineffective if $\xi \ge 0$)	0-6, 6*no dim., cm	I, 7*E
$\omega^+, \theta, R_1, U_1, U_2, R_2$	Azimuth of entrance EFB with respect to ACENT; wedge angle of EFB; radii and linear extents of EFB (use $\mid U_{1,2}\mid = \infty$ when $R_{1,2}=\infty$)	2*deg, 4*cm	6*E
	(Note : $\lambda = 0$, $\omega^+ = ACENT$ and $\theta = 0$ for $\underline{sharp\ edge}$)		
	EXIT FIELD BOUNDARY (See ENTRANCE FIELD BOUNDARY)		
λ, ξ NC, $C_0 - C_5$, shift	Fringe field parameters	cm, (cm)	2*E
$\omega^{-}, \theta, R_1, U_1, U_2, R_2$	Positioning and shape of the exit EFB	0-6, 6*no dim., cm 2*deg, 4*cm	1, 7*E 6*E
	(Note : $\lambda=0,\omega^-=$ -AT+ACENT and $\theta=0$ for		

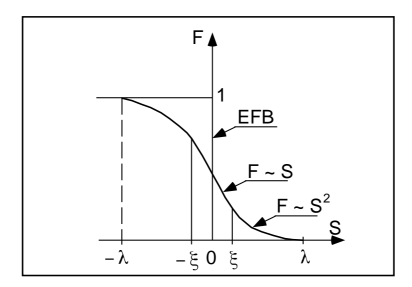
sharp edge)

if NFACE = 3 λ, ξ $NC, C_0 - C_5, \text{ shift}$ $\omega^-, \theta, R_1, U_1, U_2, R_2,$	LATERAL FIELD BOUNDARY (See ENTRANCE FIELD BOUNDARY) Next 3 records <i>only</i> if <i>NFACE</i> = 3 Fringe field parameters Positioning and shape of the lateral EFB;	cm, (cm) 0-6, 6*no dim., cm 2*deg, 5*cm	2*E I, 7*E 7*E
RM3	RM3 is the radial position on azimut ACENT		
NBS	Option index for perturbations to the field map	normally 0	I
if $NBS = 0$	Normal value. No other record required		
if $NBS = -2$	The map is modified as follows:		
$R_0, \Delta B/B_0$	B transforms to $B*\left(1+rac{\Delta B}{B_0}rac{R-R_0}{RMAX-RMIN} ight)$	cm, no dim.	2*E
if $NBS = -1$	the map is modified as follows:		
$\theta_0, \Delta B/B_0$	B transforms to $B*\left(1+rac{\Delta B}{B_0}rac{ heta- heta_0}{AT} ight)$	deg, no dim.	2*E
if NBS ≥ 1	Introduction of NBS shims		
For $I = 1$, NBS	The following 2 records must be repeated NBS times		
$R_1, R_2, \theta_1, \theta_2, \lambda$	Radial and angular limits of the shim ; λ is unused	2*cm, 2*deg, cm	5*E
$\gamma, \alpha, \mu, \beta$	geometrical parameters of the shim	2*deg, 2*no dim.	4*E
IORDRE	Degree of interpolation polynomial: 2 = second degree, 9-point grid 25 = second degree, 25-point grid 4 = fourth degree, 25-point grid	2, 4 or 25	I
XPAS	Integration step	cm	E
KPOS	Positioning of the map, normally 2. Two options :	1-2	I
if KPOS = 2 RE, TE, RS, TS	Positioning as follows: Radius and angle of reference, respectively, at entrance and exit of the map.	cm, rad, cm, rad	4*E
if KPOS = 1 <i>DP</i>	Automatic positioning of the map, by means of reference relative momentum	no dim.	E

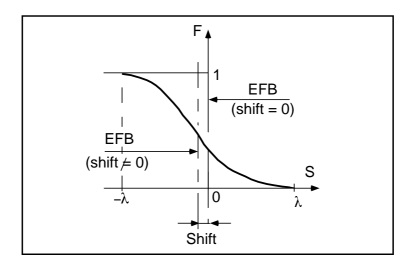




 $A: Parameters \ used \ to \ define \ the \ field \ map \ and \ geometric \ boundaries.$ $B: Parameters \ used \ to \ define \ the \ field \ map \ and \ fringe \ fields.$



Second order type fringe field.



Exponential type fringe field.

AUTOREF Automatic transformation to a new reference frame

I : Equivalent to CHANGREF (XCE = 0, YCE = Y(1), ALE = T(1)) 1-2

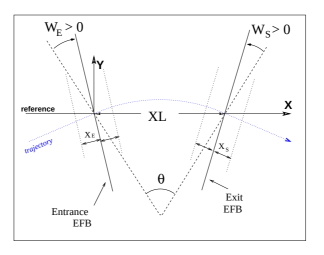
- 2: Equivalent to CHANGREF (XW, YW, T(1)), with (XW, YW) being the location of the intersection (waist) of particles 1, 4 and 5 (useful with MATRIX, for automatic positionning of the first order focus)
- 3: Equivalent to CHANGREF (XW, YW, T(I1)), with (XW, YW) being the location of the intersection (waist) of particles I1, I2 and I3 (for instance: I1 = central trajectory, I2 and I3 = paraxial trajectories that intersect at the first order focus)

 $\begin{array}{ll} \mbox{if } I=3 & \mbox{Next record only if } I=3 \\ I1, I2, I3 & \mbox{Three particle numbers} & 3*(1-10^4) & 3*I \\ \end{array}$

BEND

IL	$I\!L=1,2$: print field and coordinates along trajectories (otherwise $I\!L=0$)	0-2	I
XL, Sk, B1	Length; skew angle; field	cm, rad, kG	3*E
$X_{\mathrm{E}},\lambda_{\mathrm{E}},W_{\mathrm{E}}$	Entrance face: Integration zone extent; fringe field extent (normally \simeq gap height; zero for sharp edge); wedge angle	cm, cm, rad	3*E
$N, C_0 – C_5$	Unused ; fringe field coefficients : $B(s)=B1$ $F(s)$ with $F(s)=1/(1+\exp(P(s))$ and $P(s)=\sum_{i=0}^5 C_i(s/\lambda)^i$	unused, 6*no dim.	I, 6*E
	Exit face :		
X_S, λ_S, W_S	See entrance face	cm, cm, rad	3*E
$N, C_0 – C_5$		unused, 6*no dim.	I, 6*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	$KPOS=1$: element aligned, 2: misaligned; shifts, tilt (unused if $KPOS=1$) $KPOS=3$: entrance and exit frames are shifted by YCE and tilted wrt . the magnet by an angle of • either ALE if $ALE\neq 0$ • or $2 Arcsin(B1XL/2BORO)$ if $ALE=0$	1-2, 2*cm, rad	I, 3*E

Bending magnet, Cartesian frame



Geometry and parameters of $BE\!N\!D$ in its Cartesian frame : $X\!L=$ length, $\theta=$ deviation, $W_E,~W_S$ are the entrance and exit wedge angles.

BINARY	BINARY/FORMATTED data converter	
NF, NCol, NHDR	Number of files to convert, of data columns, of header lines. $3* \le 9$	3*I1
The next NF lines: FNAME	Name of the file to be converted. File content is assumed binary <i>iff</i> name begins with "B_" or "b_", assumed formatted otherwise.	A80

BREVOL	1-D uniform mesh magnetic field map X-axis cylindrical symmetry is assumed		
$I\!C,I\!L$	$I\!C=1,2$: print the map $I\!L=1,2$: print field and coordinates along trajectories	0-2, 0-2	2*I
BNORM, XN	Field and X-coordinate normalization coeff.	2*no dim.	2*E
TITL	Title. Start with "FLIP" to get field map X-flipped.		A80
IX	Number of longitudinal nodes of the map	≤ 400	I
FNAME [, SUM] $^{1,\;2}$	File name		A80
ID, A, B, C [, A', B', C' , B'' , etc., if $ID \ge 2$]	Integration boundary. Ineffective when $ID=0$. $ID=$ -1, 1 or ≥ 2 : as for <i>CARTEMES</i>	≥ -1 , 2*no dim., cm [,2*no dim., cm, etc.]	I,3*E [,3*E,etc.]
IORDRE	unused	2, 4 or 25	I
XPAS	Integration step	cm	Е
KPOS, XCE, YCE, ALE	<pre>KPOS=1 : element aligned, 2 : misaligned ; shifts, tilt (unused if KPOS=1)</pre>	1-2, 2*cm, rad	I, 3*E

```
OPEN (UNIT = NL, FILE = FNAME, STATUS = 'OLD' [,FORM='UNFORMATTED']) DO 1 I = 1, IX IF (BINARY) THEN READ(NL) X(I), BX(I) ELSE READ(NL,*) X(I), BX(I) ENDIF 1 CONTINUE
```

where X(I) and BX(I) are the longitudinal coordinate and field component at node (I) of the mesh. Binary file names must begin with FNAME 'B_' or 'b_'. 'Binary' will then automatically be set to '.TRUE.'.

```
myMapFile1 SUM
myMapFile2 SUM
myMapFile3
```

(all maps must all have their mesh defined in identical coordinate frame).

¹ FNAME (e.g., solenoid.map) contains the field data. These must be formatted according to the following FORTRAN sequence:

² Sumperimposing (summing) field maps is possible. To do so, pile up file names with 'SUM' following each name but the last one. *e.g.*, in the following example, 3 field maps are read and summed:

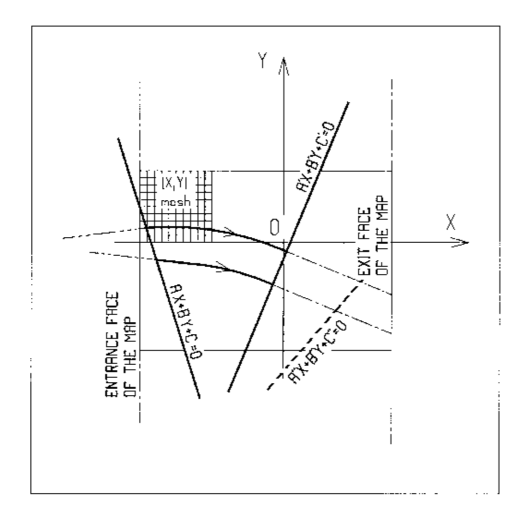
CARTEMES	2-D Cartesian uniform mesh magnetic field map mid-plane symmetry is assumed		
IC, IL	$I\!C=1,2$: print the map $I\!L=1,2$: print field and coordinates along trajectories	0-2, 0-2	2*I
BNORM, XN,YN	Field and X-,Y-coordinate normalization coeffs.	3*no dim.	3*E
TITL	Title. Start with "FLIP" to get field map X-flipped.		A80
IX,JY	Number of longitudinal (IX) and transverse (JY) nodes of the map	$\leq 400, \leq 200$	2*I
FNAME ¹	File name		A80
ID, A, B, C [, A', B', C', A'', B'' , etc., if $ID \ge 2$]	Integration boundary. Normally $ID=0$. $ID=-1$: integration in the map begins at entrance boundary defined by $AX+BY+C=0$. $ID=1$: integration in the map is terminated at exit boundary defined by $AX+BY+C=0$. $ID\geq 2$: entrance (A,B,C) and up to $ID-1$ exit $(A',B',C',A'',B'',etc.)$ boundaries	$\geq -1.2*$ no dim., cm [,2*no dim., cm, etc.]	I, 3*E [3*E,etc.]
IORDRE	Degree of interpolation polynomial (see <i>DIPOLE-M</i>)	2, 4 or 25	I
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	<pre>KPOS=1 : element aligned, 2 : misaligned ; shifts, tilt (unused if KPOS=1)</pre>	1-2, 2*cm, rad	I, 3*E

```
OPEN (UNIT = NL, FILE = FNAME, STATUS = 'OLD' [,FORM='UNFORMATTED']) IF (BINARY) THEN READ(NL) (Y(J), J=1, JY) ELSE READ(NL,100) (Y(J), J=1, JY) ENDIF 100 FORMAT(10 F8.2) DO 1 I=1,IX IF (BINARY) THEN READ(NL) X(I), (BMES(I,J), J=1, JY) ELSE READ(NL,101) X(I), (BMES(I,J), J=1, JY) 101 FORMAT(10 F8.2) ENDIF 1 CONTINUE
```

where X(I) and Y(J) are the longitudinal and transverse coordinates and BMES is the Z field component at a node (I,J) of the mesh. For binary files, FNAME must begin with 'B_' or 'b_'.

 $^{^2}$ FNAME (e.g., spes2.map) contains the field data. These must be formatted according to the following FORTRAN sequence :

^{&#}x27;Binary' will then automatically be set to '.TRUE.'



OXY is the coordinate system of the mesh. Integration zone limits may be defined, using $ID \neq 0$: particle coordinates are extrapolated linearly from the entrance face of the map, into the plane A'X + B'Y + C' = 0; after ray-tracing inside the map and terminating on the integration boundary AX + BY + C = 0, coordinates are extrapolated linearly to the exit face of the map.

CAVITE ¹	Accelerating cavity $\Delta W = qV sin(2\pi h f \Delta t + \varphi_s)$		
IOPT[.i]	Option. $i=1$ causes info output into zgoubi.CAVITE.out	0-3	I
If IOPT=0	Element inactive		
X, X	unused		
If IOPT=1 ²	f_{RF} follows the timing law given by SCALING		
\mathcal{L}, h	Reference closed orbit length; harmonic number	m, no dim.	2*E
$\hat{V},~X$	R.F. peak voltage; unused	V, unused	2*E
If IOPT=2 \mathcal{L} , h	f_{RF} follows $\Delta W_s = q \hat{V} sin \phi_s$ Reference closed orbit length ; harmonic number	m, no dim.	2*E
\hat{V},ϕ_s	R.F. peak voltage; synchronous phase	V, rad	2*E
If IOPT=3	No synchrotron motion : $\Delta W = q \hat{V} sin \phi_s$		
X, X	unused; unused	2*unused	2*E
\hat{V} , ϕ_s	R.F. peak voltage; synchronous phase	V, rad	2*E

 $^{^1}$ Use PARTICUL to declare mass and charge. 2 For ramping the R.F. frequency following $B\rho(t)$, use SCALING, with family CAVITE.

CHAMBR	Long transverse aperture limitation ¹		
IA	0 : element inactive1 : (re)definition of the aperture2 : stop testing and reset counters, print information on stopped particles.	0-2	I
IFORM[.J], C1, C2, C3, C4	IFORM = 1 : rectangular aperture ; IFORM = 2 : elliptical aperture. $J=0$, default : opening is $\pm YL=\pm C1$, $\pm ZL=\pm C2$, centered at $YC=C3$, $ZC=C4$. J=1 : opening is, in Y : $[C1,C2]$, in Z : $[C3,C4]$	1-2[.0-1]	I[.I], 4*E

Any particle out of limits is stopped. When used with an optical element defined in polar coordinates (e.g., DIPOLE) YL is the radius and YC stands for the reference radius (normally, $YC \simeq RM$).

CHANGREF Transformation to a new reference frame

"Old Style" (Figure below):

XCE, YCE, ALE Longitudinal and transverse shifts, followed by Z-axis rotation

2*cm, deg

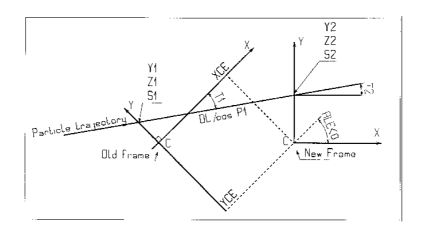
3*E

"New Style" (example below). In an arbitrary order, up to 9 occurences of :

XS 'val', YS 'val', ZS 'val', XR 'val', YR 'val', ZR 'val'

cm or deg

up to 9*(A2,E)



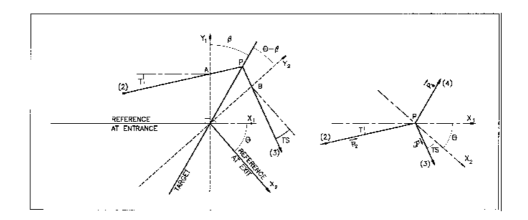
Parameters in the CHANGREF procedure.

Zgoubi data file:

```
Using CHANGREF "New Style
'OBJET'
51.71103865921708 Electron, Ekin=15MeV.
2
1 1 One particle, with
2. 0. 0.0 0.0 0.0 1. 'R' Y_0=2 cm, other coordinates zero.
1 1 1 1 1 1
'MARREF' BEG .plt -> list into zgoubi.plt.
'DRIFT' 10 cm drift.
10.
'CHANGREF'
ZR -6.34165 YS 1. First half Z-rotate, Next Y-shift.
'CHANGREF'
0. 1. 0.
'MULTIPOL' Combined function multipole, dipole + quadrupole.
2 -> list into zgoubi.plt.
5 10. 2.064995867082342 2. 0. 0. 0. 0. 0. 0. 0. 0.
0 0 5. 1.1 1.00 1.00 1.00 1.00 1.00 1. 1. 1. 1.
4 .1455 2.2670 -6395 1.1558 0. 0. 0.
0 0 0 0 0 0 0 0 0 0 0
1 step size
1 0. 0. 0.
'CHANGREF'
YS -1. ZR -6.341 First Y-shift back, next half Z-rotate.
'DRIFT' 10 cm drift.
10.
'FAISCEAU'
'END'
'END'
'END'
'END'
'END'
'STATSCEAU'
'END'
```

CIBLE, TARGET	Generate a secondary beam from target interaction
---------------	---

M_1, M_2, M_3, Q T_2, θ, β	Target, incident and scattered particle masses; Q of the reaction; incident particle kinetic energy; scattering angle; angle of the target	$5*\frac{MeV}{c^2}$, $2*\deg$	7*E
NT, NP	Number of samples in T and P coordinates after \emph{CIBLE}		2*I
TS, PS, DT	Sample step sizes ; tilt angle	3*mrad	3*E
BORO	New reference rigidity after CIBLE	kG.cm	E



Scheme of the principles of CIBLE (TARGET)

A,T= position, angle of incoming particle 2 in the entrance reference frame P= position of the interaction

B,T= position, angle of the secondary particle in the exit reference frame

 θ = angle between entrance and exit frames

 β = tilt angle of the target

COLLIMA Collimator ¹

IA 0: element inactive

1 : element active 0-2 I

2: element active and print information on stopped

particles

Physical-space collimation

IFORM[.J], C1, C2, IFORM = 1 : rectangular aperture; 1-2[.0-1] I[.I], 4*E

C3, C4 IFORM = 2 : elliptical aperture.

J=0, default : opening is $\pm YL=\pm C1, \pm ZL=\pm C2$,

centered at YC = C3, ZC = C4.

J = 1: opening is, in Y: [C1, C2], in Z: [C3, C4]

Longitudinal collimation

IFORM.J, H_{min} , H_{max} , IFORM = 6 or 7 for horizontal variable resp^{ly} S or Time, 2*cm or 2*s, I, 4*E

 V_{min} , V_{max} J=1 or 2 for vertical variable resp^{ly} 1+dp/p, kinetic-E (MeV); 2*no.dim or 2*MeV

horizontal and vertical limits

Phase-space collimation

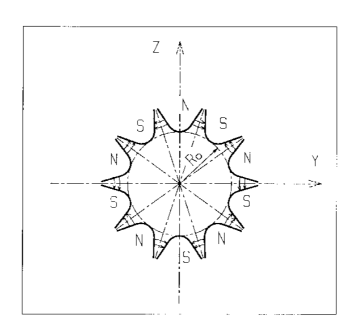
IFORM, α , β , ϵ/π , N_{σ} IFORM = 11, 14: horizontal collimation; horizontal 11-16, no.dim, I, 4*E ellipse parameters (unused if 14), emittance, cut-off 2*m, no.dim

ellipse parameters (unused if 14), emittance, cut-off IFORM = 12, 15: vertical collimation; vertical ellipse parameters (unused if 15), emittance, cut-off IFORM = 13, 16: longitudinal collimation; to be

implemented

Any particle out of limits is stopped.

DECAPOLE	Decapole magnet		
IL	$I\!L=1,2$: print field and coordinates along trajectories	0-2	I
XL, R_0, B_0	Length; radius and field at pole tip	2*cm, kG	3*E
X_E, λ_E	Entrance face : Integration zone extent ; fringe field extent ($\lesssim 2R_0$, $\lambda_E=0$ for sharp edge)	2*cm	2*E
NCE , $C_0 - C_5$	NCE = unused C_0-C_5 = Fringe field coefficients such that $G(s)=G_0/(1+\exp P(s))$, with $G_0=B_0/R_0^4$ and $P(s)=\sum_{i=0}^5 C_i(s/\lambda)^i$	unused, 6*no dim.	I, 6*E
X_S, λ_S $NCS, C_0 - C_5$	Exit face : see entrance face	2*cm 0-6, 6*no dim.	2*E I, 6*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	KPOS=1: element aligned, 2: misaligned; shifts, tilt (unused if KPOS=1)	1-2, 2*cm, rad	I, 3*E



DIPOLE	Dipole magnet, polar frame $B_Z = \mathcal{F}B_0 \left(1 + N\left(\frac{R-RM}{RM}\right) + B\left(\frac{R-RM}{RM}\right)^2 + G\left(\frac{R-RM}{RM}\right)^3\right)$)	
IL	$I\!L=1,2$: print field and coordinates along trajectories	0 - 2	I
AT,RM	Total angular extent of the dipole; reference radius	deg, cm	2*E
$ACENT, B_0, N, B, G$	Azimuth for positioning of EFBs; field and field indices	deg., kG, 3*no dim.	5*E
	ENTRANCE FIELD BOUNDARY		
λ, ξ	Fringe field extent (normally \simeq gap size); unused. Exponential type fringe field $F=1/(1+\exp(P(s)))$ with $P(s)=C_0+C_1(\frac{s}{\lambda})+C_2(\frac{s}{\lambda})^2++C_5(\frac{s}{\lambda})^5$	cm, unused	2*E
NC , $C_0 - C_5$, shift	unused ; C_0 to C_5 : see above ; EFB shift	0-6, 6*no dim., cm	I,7*E
$\omega^+, \theta, R_1, U_1, U_2, R_2$	Azimuth of entrance EFB with respect to $ACENT$; wedge angle of EFB; radii and linear extents of EFB (use $\mid U_{1,2} \mid = \infty$ when $R_{1,2} = \infty$)	2*deg, 4*cm	6*E
	EXIT FIELD BOUNDARY (See ENTRANCE FIELD BOUNDARY)		
λ, ξ $NC, C_0 - C_5$, shift	Fringe field parameters	cm, unused 0-6, 6*no dim., cm	2*E 1, 7*E
$\omega^-, \theta, R_1, U_1, U_2, R_2$	Positioning and shape of the exit EFB	2*deg, 4*cm	6*E
	LATERAL FIELD BOUNDARY (See ENTRANCE FIELD BOUNDARY)		
λ, ξ $NC, C_0 - C_5$, shift	LATERAL EFB is inhibited if $\xi = 0$	cm, unused 0-6, 6*no dim., cm	2*E 1, 7*E
$\omega^{-}, \theta, R_1, U_1, U_2, R_2,$ $RM3$	Positioning and shape of the exit EFB	2*deg, 5*cm	7*E
IORDRE, Resol	Degree of interpolation polynomial: 2 = second degree, 9-point grid 25 = second degree, 25-point grid 4 = fourth degree, 25-point grid; resolution of flying mesh is XPAS/Resol	2, 4 or 25; no dim.	I, E
XPAS	Integration step	cm	E
KPOS	Positioning of the map, normally 2. Two options :	1-2	I
if KPOS = 2 RE, TE, RS, TS	Positioning as follows: Radius and angle of reference, respectively,	cm, rad, cm, rad	4*E

at entrance and exit of the map.

if KPOS = 1 Automatic positioning of the map, by means of DP reference relative momentum

no dim.

E

DIPOLE-M	Generation of dipole mid-plane 2-D map, polar frame $B_Z = \mathcal{F}B_0 \left(1 + N\left(\frac{R-RM}{RM}\right) + B\left(\frac{R-RM}{RM}\right)^2 + G\left(\frac{R-RM}{RM}\right)^3 \right)$)	
NFACE, IC, IL	Number of field boundaries $IC=1,2$: print field map $IL=1,2$: print field and coordinates on trajectories	2-3, 0-2, 0-2	3*I
IAMAX, IRMAX	Azimuthal and radial number of nodes of the mesh	$\leq 400, \leq 200$	2*I
B_0, N, B, G	Field and field indices	kG, 3*no dim.	4*E
AT, ACENT, RM, RMIN, RMAX	Mesh parameters: total angle of the map; azimuth for positioning of EFBs; reference radius; minimum and maximum radii	2*deg, 3*cm	5*E
	ENTRANCE FIELD BOUNDARY		
λ, ξ	Fringe field extent (normally \simeq gap size); unused. Exponential type fringe field $F=1/(1+\exp(P(s)))$ with $P(s)=C_0+C_1(\frac{s}{\lambda})+C_2(\frac{s}{\lambda})^2++C_5(\frac{s}{\lambda})^5$	cm, unused	2*E
NC , $C_0 - C_5$, shift	unused ; C_0 to C_5 : see above ; EFB shift	0-6, 6*no dim., cm	I,7*E
$\omega^+, \theta, R_1, U_1, U_2, R_2$	Azimuth of entrance EFB with respect to <i>ACENT</i> ; wedge angle of EFB; radii and linear extents of EFB (use $\mid U_{1,2} \mid = \infty$ when $R_{1,2} = \infty$)	2*deg, 4*cm	6*E
	(Note : $\lambda = 0$, $\omega^+ = ACENT$ and $\theta = 0$ for sharp edge)		
	EXIT FIELD BOUNDARY (See ENTRANCE FIELD BOUNDARY)		
λ, ξ $NC, C_0 - C_5$, shift	Fringe field parameters	cm, unused 0-6, 6*nodim., cm	2*E 1, 7*E
$\omega^-, \theta, R_1, U_1, U_2, R_2$	Positioning and shape of the exit EFB	2*deg, 4*cm	6*E
	(Note : $\lambda=0,\omega^-=-AT+ACENT$ and $\theta=0$ for sharp edge)		
if NFACE = 3	LATERAL FIELD BOUNDARY (See ENTRANCE FIELD BOUNDARY) Next 3 records <i>only</i> if <i>NFACE</i> = 3	4	04F
λ, ξ	Fringe field parameters	cm, (cm)	2*E
NC, $C_0 - C_5$, shift $\omega^-, \theta, R_1, U_1, U_2, R_2, RM3$	Positioning and shape of the lateral EFB; RM3 is the radial position on azimut <i>ACENT</i>	0-6, 6*no dim., cm 2*deg, 5*cm	I, 7*E 7*E
NBS	Option index for perturbations to the field map	normally 0	I
if $NBS = 0$	Normal value. No other record required		

if $NBS = -2$	The map is modified as follows:		
$R_0, \Delta B/B_0$	B transforms to $B*\left(1+\frac{\Delta B}{B_0}\frac{R-R_0}{RMAX-RMIN}\right)$	cm, no dim.	2*E
if NBS = -1	the map is modified as follows:		
$\theta_0, \Delta B/B_0$	B transforms to $B*\left(1+rac{\Delta B}{B_0}rac{ heta- heta_0}{AT} ight)$	deg, no dim.	2*E
if NBS ≥ 1	Introduction of NBS shims		
For I = 1, NBS	The following 2 records must be repeated NBS times		
$R_1, R_2, \theta_1, \theta_2, \lambda$	Radial and angular limits of the shim ; λ is unused	2*cm, 2*deg, cm	5*E
$\gamma, \alpha, \mu, \beta$	geometrical parameters of the shim	2*deg, 2*no dim.	4*E
IORDRE	Degree of interpolation polynomial: 2 = second degree, 9-point grid 25 = second degree, 25-point grid 4 = fourth degree, 25-point grid	2, 4 or 25	I
XPAS	Integration step	cm	E
KPOS	Positioning of the map, normally 2. Two options :	1-2	I
if KPOS = 2 RE, TE, RS, TS	Positioning as follows: Radius and angle of reference, respectively, at entrance and exit of the map.	cm, rad, cm, rad	4*E
if KPOS = 1 <i>DP</i>	Automatic positioning of the map, by means of reference relative momentum	no dim.	E

2*deg, 5*cm

7*E

DIPOLES Dipole magnet N-tuple, polar frame (i) $B_Z = \sum_{i=1}^N B_{z0,i} \mathcal{F}_i(R,\theta) \left(1 + b_{1_i}(R - RM_i)/RM_i + b_{2_i}(R - RM_i)^2/RM_i^2 + \ldots\right)$ (ii) $B_Z = B_{z0,i} + \sum_{i=1}^N \mathcal{F}_i(R,\theta) \left(b_{1_i}(R - RM_i) + b_{2_i}(R - RM_i)^2 + \ldots\right)$ 0 - 2ILIL = 1, 2: print field and coordinates along trajectories I N, AT, RMI, 2*E Number of magnets in the N-tuple; no dim., total angular extent of the dipole; reference radius deg, cm Repeat N times the following sequence _ ACN, δRM^{-1} , B_0 , Positioning of EFBs : azimuth, $RM_i = RM + \delta RM$; field; 3*E, I, ind*E deg., cm, kG, $ind, b_i, (i = 1, ind)$ number of, and field coefficients (ind + 1)*no dim.ENTRANCE FIELD BOUNDARY Fringe field extent $(g = g_0 (RM/R)^{\kappa})$ cm, no dim. 2*E g_0, κ Exponential type fringe field $F = 1 / (1 + \exp(P(s)))$ with $P(s) = C_0 + C_1(\frac{s}{a}) + C_2(\frac{s}{a})^2 + \dots + C_5(\frac{s}{a})^5$ NC, $C_0 - C_5$, shift unused; C_0 to C_5 : see above; EFB shift 0-6, 6*no dim., cm I,7*E $\omega^+, \theta, R_1, U_1, U_2, R_2$ Azimuth of entrance EFB with respect to ACN; 2*deg, 4*cm 6*E wedge angle of EFB; radii and linear extents of EFB (use $\mid U_{1,2} \mid = \infty$ when $R_{1,2} = \infty$) (Note: $g_0 = 0$, $\omega^+ = ACENT$, $\theta = 0$ and KIRD=0 for sharp edge) EXIT FIELD BOUNDARY (See ENTRANCE FIELD BOUNDARY) 2*E cm, no dim. g_0, κ NC, $C_0 - C_5$, shift 0 - 6, 6*no dim., cm 1, 7*E $\omega^{-}, \theta, R_1, U_1, U_2, R_2$ 2*deg, 4*cm 6*E (Note: $g_0 = 0$, $\omega^- = -AT + ACENT$, $\theta = 0$ and KIRD=0 for sharp edge) LATERAL FIELD BOUNDARY to be implemented - following data not used 2*E cm, no dim. g_0, κ NC, $C_0 - C_5$, shift 0-6, 6*no dim., cm 1, 7*E

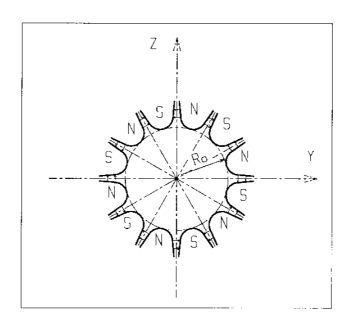
 $\omega^-, \theta, R_1, U_1, U_2, R_2, R_3$

End of repeat _

¹ Non-zero δRM requires KIRD= 2, 4 or 25.

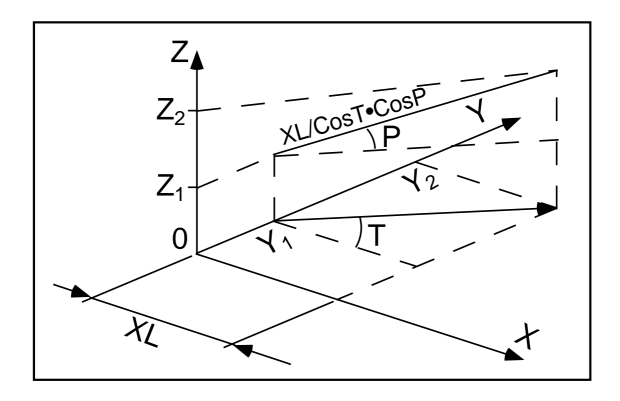
KIRD, Resol	KIRD=0: analytical computation of field derivatives; Resol = 2/4 for 2nd/4th order field derivatives computation KIRD2, 4 or 25: numerical interpolation of field derivatives; size of flying interpolation mesh is XPAS/Resol KIRD=2 or 25: second degree, 9- or 25-point grid KIRD=4: fourth degree, 25-point grid	0, 2, 4 or 25 ; no di	m.I, E
XPAS	Integration step	cm	E
KPOS	Positioning of the magnet, normally 2. Two options :	1-2	I
if KPOS = 2 RE, TE, RS, TS if KPOS = 1	Positioning as follows: Radius and angle of reference, respectively, at entrance and exit of the magnet Automatic positioning of the magnet, by means of	cm, rad, cm, rad	4*E
DP	reference relative momentum	no dim.	E

DODECAPO	Dodecapole magnet		
ΙL	$I\!L=1,2$: print field and coordinates along trajectories	0-2	I
XL, R_0, B_0	Length; radius and field at pole tip	2*cm, kG	3*E
X_E, λ_E	Entrance face : Integration zone extent ; fringe field extent ($\lesssim 2R_0, \lambda_E = 0$ for sharp edge)	2*cm	2*E
NCE , $C_0 - C_5$	NCE = unused C_0-C_5 = Fringe field coefficients such that $G(s)=G_0/(1+\exp P(s))$, with $G_0=B_0/R_0^5$ and $P(s)=\sum_{i=0}^5 C_i(s/\lambda)^i$	unused, 6*no dim.	I, 6*E
X_S, λ_S $NCS, C_0 - C_5$	Exit face : see entrance face	2*cm 0-6, 6*no dim.	2*E I, 6*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	<pre>KPOS=1 : element aligned, 2 : misaligned ; shifts, tilt (unused if KPOS=1)</pre>	1-2, 2*cm, rad	I, 3*E



DRIFT, ESL Field free drift space

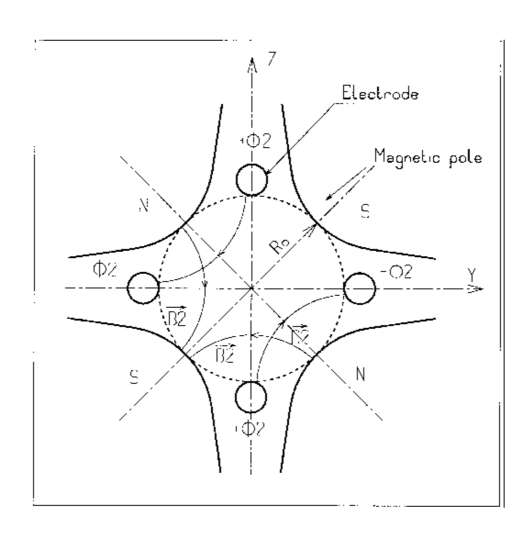
XL length cm ${\sf E}$



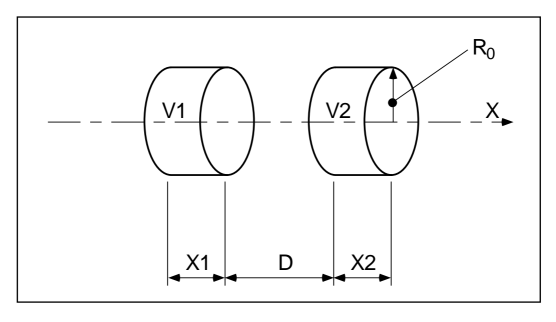
EBMULT ¹	Electro-magnetic multipole		
IL	$I\!L=1,2$: print field and coordinates along trajectories	0-2	I
$XL, R_0, E1, E2,, E10$	Electric poles Length of element; radius at pole tip; field at pole tip for dipole, quadrupole,, 20-pole electric components	2*cm, 10*V/m	12*E
$X_{E}, \lambda_{E}, E_{2},, E_{10}$	Entrance face Integration zone; fringe field extent: dipole fringe field extent = λ_E ; quadrupole fringe field extent = $\lambda_E * E_2$; 20-pole fringe field extent = $\lambda_E * E_{10}$ (for any component: sharp edge if field extent is zero)	2*cm, 9*no dim.	11*E
$NCE, C_0 - C_5$	same as QUADRUPO	0-6, 6*no dim.	I,6*E
$X_{S}, \lambda_{S}, S_{2},, S_{10}$ $NCS, C_{0} - C_{5}$	Exit face Integration zone; as for entrance	2*cm, 9*no dim. 0-6, 6*no dim.	11*E I, 6*E
R1, R2, R3,, R10	Skew angles of electric field components	10*rad	10*E
$XL, R_0, B1, B2,, B10$	Magnetic poles Length of element; radius at pole tip; field at pole tip for dipole, quadrupole,, 20-pole magnetic components	2*cm, 10*kG	12*E
$X_E, \lambda_E, E_2,, E_{10}$	Entrance face Integration zone; fringe field extent: dipole fringe field extent = λ_E ; quadrupole fringe field extent = $\lambda_E * E_2$;	2*cm, 9*no dim.	11*E
	20-pole fringe field extent = $\lambda_E * E_{10}$ (for any component : sharp edge if field extent is zero)		
$NCE, C_0 - C_5$	same as QUADRUPO	0-6, 6*no dim.	I,6*E

¹ Use PARTICUL to declare mass and charge.

$X_S, \lambda_S, S_2,, S_{10}$	Exit face Integration zone; as for entrance	2*cm, 9*no dim.	11*E
NCS , $C_0 - C_5$		0-6, 6*no dim.	I, 6*E
R1, R2, R3,, R10	Skew angles of magnetic field components	10*rad	10*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	<i>KPOS</i> =1 : element aligned, 2 : misaligned; shifts, tilt (unused if <i>KPOS</i> =1)	1-2, 2*cm, rad	I, 3*E



EL2TUB ¹	Two-tube electrostatic lens		
IL	$I\!L=1,2$: print field and coordinates along trajectories	0-2	I
X_1, D, X_2, R_0	Length of first tube; distance between tubes; length of second tube; inner radius	3*m	4*E
V_1,V_2	Potentials	2*V	2*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	<pre>KPOS=1 : element aligned, 2 : misaligned; shifts, tilt (unused if KPOS=1)</pre>	1-2, 2*cm, rad	I, 3*E

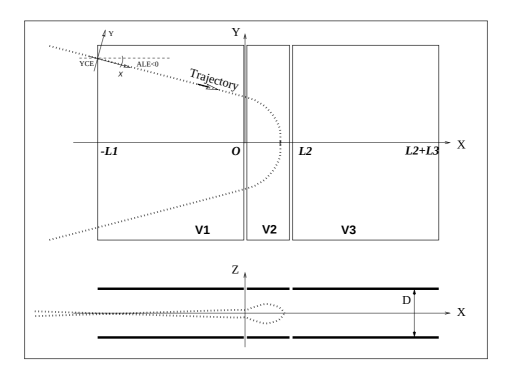


Two-electrode cylindrical electric lens.

 $^{^{1}\,}$ Use PARTICUL to declare mass and charge.

ELECTION ELECTION AND THE TOTAL STRAIGHT SH	ELMIR	Electrostatic N-electrode	e mirror/lens, straight slits
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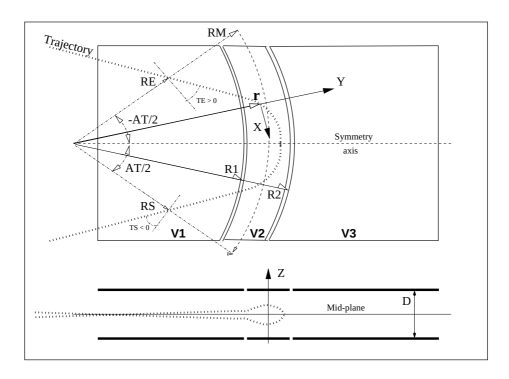
IL	IL = 1, 2: print field and coordinates along trajectories	0-2	I
N,L1,, LN, D, MT	Number of electrodes; electrode lengths; gap; mode (11/H-mir, 12/V-mir, 21/V-lens, 22/H-lens)	2-7, N*m, m	I, N*E, E, I
V1,,VN	Electrode potentials (normally $V1 = 0$)	N*V	N*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	KPOS=1: element aligned; 2: misaligned; shifts, tilt (unused if $KPOS=1$); 3: automatic positioning, $YCE=$ pitch, $ALE=$ half-deviation	1-2, 2*cm, rad	I, 3*E



Electrostatic N-electrode mirror/lens, straight slits, in the case N=3, in horizontal mirror mode (MT=11). Possible non-zero entrance quantities YCE, ALE should be specified using CHANGREF, or using KPOS=3 with YCE=pitch, ALE=half-deviation.

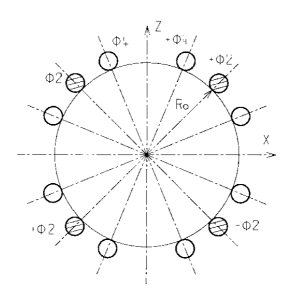
ELMIRC	Electrostatic N-electrode mirror/lens, circular slits	
IL	IL = 1, 2: print field and coordinates	0-2

IL.	along trajectories $E = 1, 2$. Print field and coordinates	0-2	1
R1, R2, AT, D	Radius of first and second slits; total deviation angle; gap	4*m 2*m, rad, m	4*E 4*E
V - VA, VB - V	Potential difference	2*V	2*E
XPAS	Integration step	cm	E
KPOS RE, TE, RS, TS	Normally $KPOS=2$ for positioning; Radius and angle at respectively entrance and exit.	1-2 cm, rad, cm, rad	I 4*E



Electrostatic N-electrode mirror/lens, circular slits, in the case ${\cal N}=3$, in horizontal mirror mode.

ELMULT ¹	Electric multipole		
IL	IL = 1, 2: print field and coordinates along trajectories	0-2	I
$XL, R_0, E1, E2,, E10$	Length of element; radius at pole tip; field at pole tip for dipole, quadrupole,, dodecapole components	2*cm, 10*V/m	12*E
$X_E, \lambda_E, E_2,, E_{10}$	Entrance face Integration zone ; fringe field extent : dipole fringe field extent = λ_E ; quadrupole fringe field extent = $\lambda_E * E_2$; 20-pole fringe field extent = $\lambda_E * E_{10}$ (sharp edge if field extent is zero)	2*cm, 9*no dim.	11*E
$NCE, C_0 - C_5$	same as QUADRUPO	0-6, 6*no dim.	I, 6*E
$X_{S}, \lambda_{S}, S_{2},, S_{10}$ $NCS, C_{0} - C_{5}$	Exit face Integration zone ; as for entrance	2*cm, 9*no dim. 0-6, 6*no dim.	11*E I, 6*E
R1, R2, R3,, R10	Skew angles of field components	10*rad	10*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	<pre>KPOS=1 : element aligned, 2 : misaligned ; shifts, tilt (unused if KPOS=1)</pre>	1-2, 2*cm, rad	I, 3*E



 $^{^{1}\,}$ Use PARTICUL to declare mass and charge.

ELREVOL ¹	1-D uniform mesh electric field map X-axis cylindrical symmetry is assumed		
IC, IL	$I\!C=1,2$: print the map $I\!L=1,2$: print field and coordinates along trajectories	0-2, 0-2	2*I
ENORM, X-NORM	Field and X-coordinate normalization coeff.	2*no dim.	2*E
TITL	Title. Start with "FLIP" to get field map X-flipped.		A80
IX	Number of longitudinal nodes of the map	≤ 400	I
FNAME ²	File name		A80
ID, A, B, C [, A', B', C', B'' , etc., if $ID \ge 2$]	Integration boundary. Ineffective when $ID=0$. $ID=$ -1, 1 or ≥ 2 : as for <i>CARTEMES</i>	≥ -1 , 2*no dim., cm [,2*no dim., cm, etc.]	I,3*E [,3*E,etc.]
IORDRE	unused	2, 4 or 25	I
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	<pre>KPOS=1 : element aligned, 2 : misaligned; shifts, tilt (unused if KPOS=1)</pre>	1-2, 2*cm, rad	I, 3*E

where X(I) and EX(I) are the longitudinal coordinate and field component at node (I) of the mesh. Binary file names FNAME must begin with 'B_' or 'b_'. 'Binary' will then automatically be set to '.TRUE.'

 $^{^{1}\,}$ Use PARTICUL to declare mass and charge.

 $^{^2}$ FNAME (e.g., e-lens.map) contains the field data. These must be formatted according to the following FORTRAN sequence :

EMMA 2-D Cartesian or cylindrical mesh field map for EMMA FFAG

IC, IL	see CARTEMES	0-2, 0-2	2*I
BNORM, XN, YN, ZN	Field and X-,Y-,Z-coordinate normalization coefficients	4*no dim.	4*E
TITL	Title. Start with "FLIP" to get field map X-flipped		A80
IX, IY, IZ, MOD[.i]	Number of nodes of the mesh in the X,Y and Z directions, $IZ=1$ for single 2-D map; MOD : operational and map $FORMAT$ reading mode 1 $MOD \leq 19$: Cartesian mesh; $MOD \geq 20$: cylindrical mesh; $.i$, optional, tells the reading $FORMAT$, default is '*'.	$ \leq 400, \leq 200, \\ 1, \geq 0[.1-9] $	3*I
FNAME 1 $(K = 1, NF)$	Names of the NF files that contain the 2-D maps, ordered from $Z(1)$ to $Z(NF)$. If $MOD=0$: a single map, superimposition of QF and QD one If $MOD=1$: a single map, interpolated from QF[x_F] and QD[If $MOD=22$: a single map, superimposition of QF and QD on If $MOD=24$: field at particle is interpolated from a (QF,QD) properties (x_F,x_D) value, taken from of set of (QF,QD) pairs region.	$[x_D]$ ones, is built for es, is built for tracking pair of maps, closest to	tracking. g.
ID, A, B, C [, A', B', C', B'' , etc., if $ID \ge 2$]	Integration boundary. Ineffective when $ID=0$. $ID=$ -1, 1 or ≥ 2 : as for <i>CARTEMES</i>	≥ -1 , 2*no dim., cm [,2*no dim., cm, etc.]	I,3*E [,3*E,etc.]
IORDRE	If $IZ = 1$: as in <i>CARTEMES</i> If $IZ \neq 1$: unused	2, 4 or 25	I
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	<pre>KPOS=1 : element aligned, 2 : misaligned ; shifts, tilt (unused if KPOS=1)</pre>	1-2, 2*cm, rad	I, 3*E

 $^{^1}$ FNAME normally contains the field map data. If MOD=24 FNAME(K) contains the names of the QF maps and QD maps, as well as the QF-QD distance attached to each one of these pairs.

FAISCEAU Print particle coordinates

Print particle coordinates at the location where the keyword is introduced in the structure.

FAISCNL Store particle coordinates in file FNAME

 $FNAME^1$ Name of storage file A80

(e.g., zgoubi.fai, or b_zgoubi.fai for binary storage).

Store coordinates every IP other pass [, at elements with appropriate label] **FAISTORE**

FNAME $^{1, 2}$ Name of storage file (e.g. zgoubi.fai) [; label(s) of the element(s) at the exit A80, [, 10*A10] [, LABEL(s)] ³ of which the store occurs (10 labels maximum)]. If either FNAME or first LABEL

is 'none' then store only occurs at location of FAISTORE. Store occurs at

all elements if first LABEL is 'all'

IPStore every IP other pass (when using REBELOTE I

with $NPASS \ge IP - 1$).

¹ Stored data can be read again using OBJET, KOBJ = 3.

FNAME = 'none' will inhibit printing.
 If first LABEL = 'none' then printing will be inhibited.

FFAG	FFAG magnet, N -tuple UNDER DEVELOPEMENT $B_Z = \sum_{i=1}^N B_{z0,i} \mathcal{F}_i(R,\theta) \ (R/R_{M,i})^{K_i}$		
IL	$I\!L=1,2$: print field and coordinates along trajectories	0 - 2	I
N, AT, RM	Number of dipoles in the FFAG N -tuple ; total angular extent of the dipole ; reference radius	no dim., deg, cm	I, 2*E
Repeat N times the follow	wing sequence		
ACN , δRM , B_{z_0} , K	Azimuth for dipole positionning ; $R_{M,i}=R\!M+\delta R\!M$; field at $R_{M,i}$; index	deg, cm, kG, no dim.	4*E
	ENTRANCE FIELD BOUNDARY		
g_0, κ $NC, C_0 - C_5$, shift $\omega^+, \theta, R_1, U_1, U_2, R_2$	Fringe field extent $(g=g_0 (RM/R)^\kappa)$ unused; C_0 to C_5 : fringe field coefficients; EFB shift Azimuth of entrance EFB with respect to ACN ; wedge angle of EFB; radii and linear extents of EFB (use $ U_{1,2} =\infty$ when $R_{1,2}=\infty$)	cm, no dim. 0-6, 6*no dim, cm 2*deg, 4*cm	2*E I,7*E 6*E
	(Note : $g_0 = 0$, $\omega^+ = ACENT$, $\theta = 0$ and KIRD=0 for sharp eq	lge)	
	EXIT FIELD BOUNDARY (See ENTRANCE FIELD BOUNDARY)		
g_0, κ $NC, C_0 - C_5$, shift $\omega^-, \theta, R_1, U_1, U_2, R_2$		cm, no dim 0-6, 6*no dim, cm 2*deg, 4*cm	2*E 1, 7*E 6*E
	(Note : $g_0=0,\omega^-=-AT+$ ACENT, $\theta=0$ and KIRD=0 for	sharp edge)	
	LATERAL FIELD BOUNDARY to be implemented - following data not used		
g_0, κ $NC, C_0 - C_5$, shift $\omega^-, \theta, R_1, U_1, U_2, R_2$		cm, no dim 0-6, 6*no dim, cm 2*deg, 4*cm	2*E 1, 7*E 6*E
End of repeat			
KIRD, Resol	KIRD=0: analytical computation of field derivatives; Resol = 2/4 for 2nd/4th order field derivatives computation KIRD2, 4 or 25: numerical interpolation of field derivatives; size of flying interpolation mesh is XPAS/Resol KIRD=2 or 25: second degree, 9- or 25-point grid KIRD=4: fourth degree, 25-point grid	0, 2, 4 or 25 ; no dir	n.I, E
XPAS	Integration step	cm	E
KPOS			
	Positioning of the magnet, normally 2. Two options :	1-2	I

at entrance and exit of the magnet Automatic positioning of the magnet, by means of if KPOS = 1

DPreference relative momentum no dim. Е

FFAG-SPI	Spiral FFAG magnet, N -tuple UNDER DEVELOPEMENT $B_Z = \sum_{i=1}^{N} B_{20,i} \mathcal{F}_i(R,\theta) (R/R_{M,i})^{K_i}$		
IL	IL = 1, 2: print field and coordinates along trajectories	0 - 2	I
N, AT, RM	Number of dipoles in the FFAG N -tuple ; total angular extent of the dipole ; reference radius	no dim., deg, cm	I, 2*E
Repeat N times the follow	ving sequence		
ACN , δRM , B_{z_0} , K	Azimuth for dipole positionning ; $R_{M,i}=R\!M+\delta R\!M$; field at $R_{M,i}$; index	deg, cm, kG, no dim.	4*E
	ENTRANCE FIELD BOUNDARY		
g_0, κ $NC, C_0 - C_5$, shift $\omega^+, \xi, 4$ dummies	Fringe field extent $(g=g_0(RM/R)^\kappa)$ unused; C_0 to C_5 : fringe field coefficients; EFB shift Azimuth of entrance EFB with respect to ACN ; spiral angle; $4\times$ unused	cm, no dim. 0-6, 6*no dim, cm 2*deg, 4*unsued	2*E I,7*E 6*E
	EXIT FIELD BOUNDARY (See ENTRANCE FIELD BOUNDARY)		
g_0, κ $NC, C_0 - C_5$, shift $\omega^-, \xi, 4$ dummies		cm, no dim 0-6, 6*no dim, cm 2*deg, 4*unused	2*E 1, 7*E 6*E
	LATERAL FIELD BOUNDARY to be implemented - following data not used		
g_0, κ $NC, C_0 - C_5$, shift $\omega^-, \theta, R_1, U_1, U_2, R_2$		cm, no dim 0-6, 6*no dim, cm 2*deg, 4*cm	2*E 1, 7*E 6*E
End of repeat			
KIRD, Resol	KIRD=0: analytical computation of field derivatives; Resol = 2/4 for 2nd/4th order field derivatives computation KIRD2, 4 or 25: numerical interpolation of field derivatives; size of flying interpolation mesh is <i>XPAS/Resol</i> KIRD=2 or 25: second degree, 9- or 25-point grid KIRD=4: fourth degree, 25-point grid	0, 2, 4 or 25; no dir	n.I, E
XPAS	Integration step	cm	E
KPOS	Positioning of the magnet, normally 2. Two options :	1-2	I
if KPOS = 2 RE, TE, RS, TS	Positioning as follows: Radius and angle of reference, respectively, at entrance and exit of the magnet	cm, rad, cm, rad	4*E
if KPOS = 1 DP	Automatic positioning of the magnet, by means of reference relative momentum	no dim.	E

FIN, END End of input data list

Any information following these keywords will be ignored

FIT, FIT2	Fitting procedure		
NV	Number of physical parameters to be varied	≤ 20	I
For $I = 1$, NV	repeat NV times the following sequence		
either:			
IR, IP, XC, DV	Number of the element in the structure; number of the physical parameter in the element; coupling switch (off = 0); variation range (\pm)		2*I, 2*E
or:			
IR, IP, XC, [Vmin, Vm]	[aax]	\leq MXL, \leq MXD,	2*I, 3*E
NC [, penalty 3]	Number of constraints [, penalty].	$\leq 20 [, \sim 10^{-n}]$	I [, E]
For $I = 1$, NC	repeat NC times the following sequence		
$IC, I, J, IR, V^4, WV, NP [, p_i(i = 1, NP)]$	IC , I and J define the type of constraint (see table below); IR : number of the element after which the constraint applies; V : value; W : weight (the stronger the lower WV) NP : number of parameters; if $NP \geq 1$, $p_i(i=1,NP)$: parameter values.	0-5, 3*(>0), current unit, 2*no dim., curr. unit	4*I, 2*E, I, <i>NP</i> *E s

 $[\]begin{array}{l} ^{1} \quad \text{MXL is set in include file } \textit{MXLD.H.} \\ ^{2} \quad \text{MXD is set in include file } \textit{MXLD.H.} \ \text{Data is of the form "integer.iii" with i a 1-digit integer.} \\ ^{3} \quad \text{FIT[2]} \quad \text{will stop when the sum of the squared residuals gets } < \textit{penalty.} \\ ^{4} \quad \textit{V} \ \text{is in current } \textbf{zgoubi} \ \text{units.} \\ \end{array}$

Type of	Parameters defining the constraints					Object definition			
constraint	I C	I	J	Constraint	#	Parameter(s) values		` '	
σ -matrix	0	1 - 6	1 - 6	$\sigma_{IJ} \ (\sigma_{11} = \beta_Y, \sigma_{12} = \sigma_{21} = \alpha_Y, \text{etc.})$					OBJET/KOBJ=5,6
Beam matrix (N=1-9 for MATRIX block 1-9))	0.N	1 - 6 7 8 9 10	1 - 6 any any any any	$\sigma_{IJ} (\sigma_{11} = \cos \mu_Y + \alpha_Y \sin \mu_Y, \text{ etc.})$ $Y \text{-tune} = \mu_Y / 2\pi$ $Z \text{-tune} = \mu_Z / 2\pi$ $\cos(\mu_Y)$ $\cos(\mu_Z)$					OBJET/KOBJ=5,6
First order parameters	1	$ \begin{array}{c c} 1 - 6 \\ 7 \\ 8 \end{array} $	1-6 i j	Transport coeff. R_{IJ} $i \neq 8$: YY-determinant; i=8: YZ-det. $j \neq 7$: ZZ-determinant; j=7: ZY-det.					OBJET/KOBJ=5
Second order parameters	2	1-6	11 – 66	Transport coeff. $T_{I,j,k}$ $(j = [J/10], k = J - 10[J/10])$					OBJET/KOBJ=6
Trajectory coordinates	3.1 3.2 3.3 3.4	1 - MAX -1 -2 -3 1 - MAX 1 - MAX 1 - MAX 1 - MAX	$ \begin{array}{r} 1 - 7 \\ 1 - 7 \\ 1 - 7 \\ 1 - 7 \\ 1 - 7 \\ 1 - 7 \\ 1 - 7 \\ 1 - 7 \\ 1 - 7 \\ \end{array} $	$F(J,I) < F(J,i) >_{i=1,\text{MAX}} \\ Sup(F(J,i))_{i=1,\text{MAX}} \\ Dist F(J,I) _{i=11,I2,dI} \\ F(J,I) - FO(J,I) \\ F(J,I) + FO(J,I) \\ \text{min. (1) or max. (2) value of } F(J,I) \\ F(J,I) - F(J,K) (K=1-\text{MAX})$	3	11 1-2 K	I2	dI	[MCJOBJET
Matched ellipse parameters	4	1 - 6	1 - 6	σ_{IJ} ($\sigma_{11}=eta_Y,\sigma_{12}=\sigma_{21}=lpha_Y,$ etc.)					OBJET/KOBJ=8; MCOBJET/KOBJ
Number of particles	5	$ \begin{array}{r} -1 \\ 1 - 3 \\ 4 - 6 \end{array} $	any any any	$N_{survived}/ extbf{MAX} \ N_{in~\epsilon_{Y,Z,X}}/N_{survived} \ N_{in~best~\epsilon_{Y,Z,X,rms}}/N_{survived}$	1	ϵ/π ϵ/π			OBJET MCOBJET MCOBJET
Spin	10 10.1	1 - IMAX 1 - IMAX	$1 - 4 \\ 1 - 3$	$S_{X,Y,Z}(I), \vec{S}(I) \ S_{X,Y,Z}(I) - SO_{X,Y,Z}(I) $					[MC]OBJET +SPNTRK

FOCALE	Particle coordinates and horizontal beam dimension at distance XL			
XL	Distance from the location of the keyword	cm	E	
FOCALEZ	Particle coordinates and vertical beam dimension at distance	e XL		
XL	Distance from the location of the keyword	cm	E	

GASCAT	Gas scattering		
KGA	Off/On switch	0, 1	I
AI,DEN	Atomic number; density		2*E

1-24, 2*

HISTO

1-D histogram

 $J, X_{\min}, X_{\max},$ NBK, NH J =type of coordinate to be histogramed ;

the following are available : current units,
• current coordinates : < 120, 1-5

1(D), 2(Y), 3(T), 4(Z), 5(P), 6(S),

• initial coordinates : $11(D_0)$, $12(Y_0)$, $13(T_0)$, $14(Z_0)$, $15(P_0)$, $16(S_0)$,

• spin: $21(S_X), 22(S_Y), 23(S_Z), 24(< S >) \; ; \\ X_{\min}, X_{\max} = \text{limits of the histogram, in units} \\ \text{of the coordinate of concern} \; ; NBK = \text{number of channels} \; ; NH = \text{number of the histogram (for independency of histograms of the same coordinate)} \; ...$

NBL, KAR, NORM, TYP Number of lines (= vertical amplitude); alphanumeric character; normalization if NORM = 1, otherwise NORM = 0; TYP = 'P': primary particles are histogramed, or 'S': secondary, or Q: all particles - for use with MCDESINT

normally 10-40, I, A1, I, A1 char., 1-2, P-S-Q

I, 2*E, 2*I

IMAGE Localization and size of horizontal waist

IMAGES Localization and size of horizontal waists

For each momentum group, as classified by means of OBJET, KOBJ = 1, 2 or 4

IMAGESZ Localization and size of vertical waists

For each momentum group, as classified by means of OBJET, KOBJ = 1, 2 or 4

IMAGEZ Localization and size of vertical waist

MAP2D	2-D Cartesian uniform mesh field map - arbitrary magnetic field					
IC, IL	$I\!C=1,2$: print the field map $I\!L=1,2$: print field and coordinates along trajectories	0-2, 0-2	2*I			
BNORM, XN,YN	Field and X-,Y-coordinate normalization coeffs.	3*no dim.	3*E			
TITL	Title. Start with "FLIP" to get field map X-flipped.		A80			
IX, JY	Number of longitudinal and horizontal-transverse nodes of the mesh (the Z elevation is arbitrary)	$\leq 400, \leq 200$	2*I			
FNAME ¹	File name		A80			
ID, A, B, C [, A', B', C', B'' , etc., if $ID \ge 2$]	Integration boundary. Ineffective when $ID=0$. $ID=$ -1, 1 or ≥ 2 : as for <i>CARTEMES</i>	≥ -1 , 2*no dim., cm [,2*no dim., cm, etc.]	I,3*E [,3*E,etc.]			
IORDRE	Degree of polynomial interpolation	2, 4	I			
XPAS	Integration step	cm	Е			
KPOS, XCE, YCE, ALE	<pre>KPOS=1 : element aligned, 2 : misaligned ; shifts, tilt (unused if KPOS=1)</pre>	1-2, 2*cm, rad	I, 3*E			

These must be formatted according to the following FORTRAN read sequence (normally compatible with TOSCA code OUTPUTS - details and possible updates are to be found in the source file 'fmapw.f'):

```
\begin{aligned} & \text{OPEN (UNIT = NL, FILE = FNAME, STATUS = 'OLD')} \\ & \text{DO 1 J = 1, JY} \\ & \text{DO 1 I = 1, IX} \\ & \text{IF (BINARY) THEN} \\ & \text{READ(NL) Y(J), Z(1), X(I), BY(I,J), BZ(I,J), BX(I,J)} \\ & \text{ELSE} \\ & \text{READ(NL,100) Y(J), Z(1), X(I), BY(I,J), BZ(I,J), BX(I,J)} \\ & \text{100} & \text{FORMAT (1X, 6E11.4)} \\ & \text{ENDIF} \\ & \text{1} & \text{CONTINUE} \end{aligned}
```

where X(I), Y(J) are the longitudinal, horizontal coordinates in the at nodes (I,J) of the mesh, Z(I) is the vertical elevation of the map, and BX, BY, BZ are the components of the field.

For binary files, FNAME must begin with 'B_' or 'b_'~; a logical flag 'Binary' will then automatically be set to '.TRUE.'

¹ FNAME (e.g., magnet.map) contains the field map data.

MAP2D-E	2-D Cartesian uniform mesh field map - arbitrary electric field					
IC, IL	$I\!C=1,2$: print the field map $I\!L=1,2$: print field and coordinates along trajectories	0-2, 0-2	2*I			
ENORM, X-,Y-NORM	Field and X-,Y-coordinate normalization coeffs.	2*no dim.	2*E			
TITL	Title. Start with "FLIP" to get field map X-flipped.		A80			
IX,JY	Number of longitudinal and horizontal-transverse nodes of the mesh (the Z elevation is arbitrary)	$\leq 400, \leq 200$	2*I			
FNAME ¹	File name		A80			
ID, A, B, C [, A', B', C', B'' , etc., if $ID \ge 2$]	Integration boundary. Ineffective when $ID=0$. $ID=$ -1, 1 or ≥ 2 : as for <i>CARTEMES</i>	≥ -1 , 2*no dim., cm [,2*no dim., cm, etc.]	I,3*E [,3*E,etc.]			
IORDRE	Degree of polynomial interpolation, 2nd or 4th order.	2, 4	I			
XPAS	Integration step	cm	E			
KPOS, XCE, YCE, ALE	<i>KPOS</i> =1 : element aligned, 2 : misaligned; shifts, tilt (unused if <i>KPOS</i> =1)	1-2, 2*cm, rad	I, 3*E			

```
\begin{aligned} & \text{OPEN (UNIT = NL, FILE = FNAME, STATUS = 'OLD')} \\ & \text{DO 1 J = 1, JY} \\ & \text{DO 1 I = 1, IX} \\ & \text{IF (BINARY) THEN} \\ & \text{READ(NL) Y(J), Z(1), X(I), EY(I,J), EZ(I,J), EX(I,J)} \\ & \text{ELSE} \\ & \text{READ(NL,100) Y(J), Z(1), X(I), EY(I,J), EZ(I,J), EX(I,J)} \\ & \text{100} & \text{FORMAT (1X, 6E11.4)} \\ & \text{ENDIF} \\ & \text{1} & \text{CONTINUE} \end{aligned}
```

where X(I), Y(J) are the longitudinal, horizontal coordinates in the at nodes (I,J) of the mesh, \$Z(1)\$ is the vertical elevation of the map, and EX, EY, EZ are the components of the field.

For binary files, FNAME must begin with 'B_' or 'b_'"; a logical flag 'Binary' will then automatically be set to '.TRUE.'

 $^{^1}$ FNAME (e.g., ''mirror.map'') contains the field map data. These must be formatted according to the following FORTRAN read sequence - details and possible updates are to be found in the source file 'fmapw.f':

MARKER Marker

Just a marker. No data

^{&#}x27;.plt' as a second LABEL will cause storage of current coordinates into zgoubi.plt

MATRIX Calcu

Calculation of transfer coefficients, periodic parameters

IORD, IFOC

Options : 0-2, 0-1 or > 10 2*I [,A]

[, zgoubi.MATRIX.out] IORD = 0 : Same effect as FAISCEAU

IORD = 1 (normally using OBJET, KOBJ = 5): First order transfer matrix; beam matrix, phase advance if using OBJET, KOBJ = 5.01;

if IFOC > 10: periodic beam matrix, tune numbers

IORD=2 (normally using OBJET, KOBJ=6): First order transfer matrix $\left[R_{ij}\right]$, second order array $\left[T_{ijk}\right]$ and higher order transfer

coefficients ; if $\mathit{IFOC} > 10$: periodic parameters,

IFOC = 0: matrix at actual location,

reference \equiv particle # 1

IFOC = 1 : matrix at the closest first order horizontal focus,

reference \equiv particle # 1

IFOC = 10 + NPER: same as IFOC = 0, and also calculates

the twiss parameters, tune numbers, etc.

(assuming that the DATA file describes one period of a

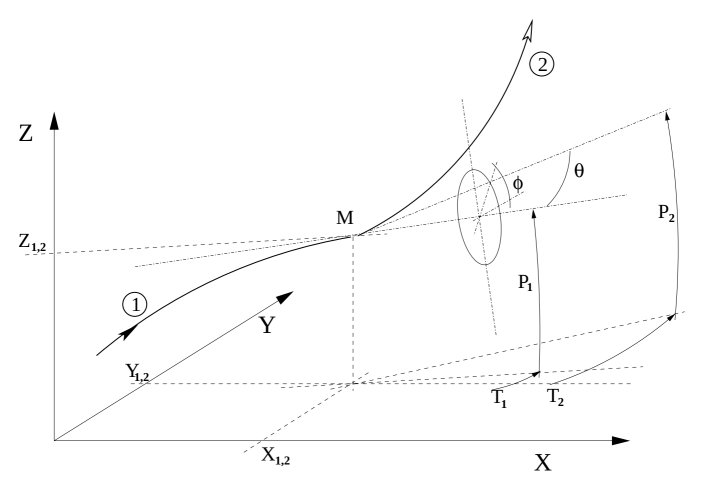
NPER-period structure).

Including 'zgoubi.MATRIX.out' will cause printout to zgoubi.MATRIX.out file

MCDESINT 1	Monte-Carlo simulation of in-flight decay
	$M1 \rightarrow M2 + M3$

[INFO,] 2 $M2$, $M3$, $\tau 2$	[Switch,]; masses of the two decay products;	[-,] $2*MeV/c^2$, s	[A4,] 3*E
	COM lifetime of particle 2		
I1, I2, I3	Seeds for random number generators	$3* \simeq 10^6$	3*I

Note that τ^2 can be left blank, in which case the lifetime of particle 2 is set to zero (it decays immediately).



Particle 1 decays into 2 and 3; **zgoubi** then calculates trajectory of 2, while 3 is discarded. θ and ϕ are the scattering angles of particle 2 relative to the direction of the incoming particle 1. They transform to T_2 and P_2 in Zgoubi frame.

MCDESINT must be preceded by PARTICUL, for the definition of the mass and lifetime of the incoming particle M1.
 Presence of 'INFO' will cause more info on decay kinematics parameters to be printed into zgoubi.res at each decay.

MCOBJET	Monte-Carlo generation of a 6-D object		
BORO	Reference rigidity	kG.cm	E
KOBJ	Type of support of the random distribution $KOBJ = 1$: window $KOBJ = 2$: grid $KOBJ = 3$: phase-space ellipses	1-3	I
I MAX	Number of particles to be generated	$\leq 10^4$	I
KY, KT, KZ, KP, KX, KD^{-1}	Type of probability density	6*(1-3)	6*I
$Y_0, T_0, Z_0, P_0, X_0, D_0$	Mean value of coordinates ($D_0 = B\rho/BORO$)	m, rad, m, rad, m, no dim.	6*E
if KOBJ = 1	In a window		
$\begin{array}{l} \delta Y, \delta T, \delta Z, \delta P, \\ \delta X, \delta D \end{array}$	Distribution widths, depending on KY , KT etc. ¹	m, rad, m, rad, m, no dim.	6*E
$N_{\delta Y}, N_{\delta T}, N_{\delta Z}, N_{\delta P}, N_{\delta X}, N_{\delta D}$	Sorting cut-offs (used only for Gaussian density)	units of σ_Y , σ_T , etc.	6*E
N_0, C_0, C_1, C_2, C_3	Parameters involved in calculation of P(D)	no dim.	5*E
IR1, IR2, IR3	Random sequence seeds	$3* \simeq 10^6$	3*I

KD can take the values

1: uniform, $p(D) = 1/2\delta D$ if $-\delta D \le x \le \delta D$

2: exponential, $p(D)=\mbox{No }\exp(C_0+C_1l+C_2l^2+C_3l^3)$ if $-\delta D\leq x\leq \delta D$

3 : kinematic, $D = \delta D * T$

 $^{^{1}}$ Let x=Y,T,Z,P or X. KY,KT,KZ,KP and KX can take the values

^{1 :} uniform, $p(x) = 1/2\delta x$ if $-\delta x \le x \le \delta x$

^{2:} Gaussian, $p(x) = \exp(-x^2/2\delta x^2)/\delta x \sqrt{2\pi}$

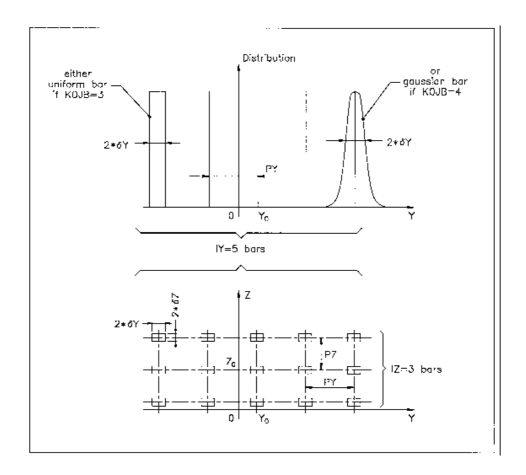
^{3:} parabolic, $p(x) = 3(1-x^2/\delta x^2)/4\delta x$ if $-\delta x \le x \le \delta x$

If $KOBJ = 2$	On a grid		
IY, IT, IZ, IP, IX, ID	Number of bars of the grid		6*I
PY, PT, PZ, PP, PX, PD	Distances between bars	m, rad, m rad, m, no dim.	6*E
$\begin{array}{l} \delta Y, \delta T, \delta Z, \delta P, \\ \delta X, \delta D \end{array}$	Width of the bars (\pm) if uniform, Sigma value if Gaussian distribution	ibidem	6*E
$N_{\delta Y}, N_{\delta T}, N_{\delta Z}, N_{\delta P}, N_{\delta X}, N_{\delta D}$	Sorting cut-offs (used only for Gaussian density)	units of σ_Y , σ_T , etc.	6*E
N_0, C_0, C_1, C_2, C_3	Parameters involved in calculation of $P(D)$	no dim.	5*E
IR1, IR2, IR3	Random sequence seeds	$3*\simeq 10^6$	3*I
if $KOBJ = 3$	On a phase-space ellipse ¹		
$\begin{array}{l} \alpha_Y, \beta_Y, \varepsilon_Y/\pi, N_{\sigma_{\epsilon_Y}} \\ [, N'_{\sigma_{\epsilon_Y}} \text{ if } N_{\sigma_{\epsilon_Y}} < 0] \end{array}^2 \\$	Ellipse parameters and emittance, Y-T phase-space; cut-off	no dim., m/rad, m, units of $\sigma(\varepsilon_Y)$	4*E [,E]
$\begin{array}{l} \alpha_Z,\beta_Z,\varepsilon_Z/\pi,N_{\sigma_{\epsilon_Z}}\\ [,N_{\sigma_{\epsilon_Z}}' \text{ if } N_{\sigma_{\epsilon_Z}}<0]^{2} \end{array}$	Ellipse parameters and emittance, Z-P phase-space; cut-off	no dim., m/rad, m, units of $\sigma(\varepsilon_Z)$	4*E [,E]
$\begin{array}{l} \alpha_X, \beta_X, \varepsilon_X/\pi, N_{\sigma_{\epsilon_X}} \\ [, N'_{\sigma_{\epsilon_X}} \text{ if } N_{\sigma_{\epsilon_X}} < 0]^2 \end{array}$	Ellipse parameters and emittance, X-D phase-space; cut-off	no dim., m/rad, m, units of $\sigma(\varepsilon_X)$	4*E [,E]
IR1, IR2, IR3	Random sequence seeds	$3* \simeq 10^6$	3*I

tier
$$\frac{1+\sigma_Y^2}{\beta_Y^2}Y^2+2\alpha_YYT+\beta_YT^2=\frac{\varepsilon_Y}{\pi}$$
 if $N_{\sigma_{\epsilon_Y}}>0$, or, if $N_{\sigma_{\epsilon_Y}}<0$ sorting within the ring

$$[\,|N_{\sigma_{\epsilon_Y}}\,|,N'_{\sigma_{\epsilon_Y}}\,]$$

 $^{^1}$ Similar possibilities, non-random, are offered with OBJET, KOBJ=8 (p. 220) 2 Works with Gaussian density type only : sorting within the ellipse fron-



Scheme of the input parameters to MCOBJET when KOBJ= 3, 4

 $\label{eq:A:A} {\bf A}: {\bf A} \mbox{ distribution of the } Y \mbox{ coordinate } \\ {\bf B}: 2\mbox{-D grid in } (Y,Z) \mbox{ space}.$

MULTIPOL	Magnetic Multipole		
IL	$I\!L=1,2$: print field and coordinates along trajectories	0-2	I
$XL, R_0, B1, B2,, B10,$	Length of element; radius at pole tip; field at pole tip for dipole, quadrupole,, dodecapole components	2*cm,10*kG	12*E
$X_E, \lambda_E, E_2,, E_{10}$	Entrance face Integration zone ; fringe field extent : dipole fringe field extent = λ_E ; quadrupole fringe field extent = $\lambda_E * E_2$; 20-pole fringe field extent = $\lambda_E * E_{10}$ (sharp edge if field extent is zero)	2*cm,9*no dim.	11*E
$NCE, C_0 - C_5$	same as QUADRUPO	0-6, 6*no dim.	I, 6*E
$X_S, \lambda_S, S_2,, S_{10}$ $NCS, C_0 - C_5$	Exit face Integration zone; as for entrance	2*cm, 9*no dim. 0-6, 6*no dim.	11*E I, 6*E
R1, R2, R3,, R10	Skew angles of field components	10*rad	10*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	$KPOS=1$: element aligned, 2: misaligned; shifts, tilt (unused if $KPOS=1$) for $QUADRUPO$. $KPOS=3$: effective only if $B1 \neq 0$: entrance and exit frames are shifted by YCE and tilted wrt . the magnet by an angle of \bullet either ALE if ALE $\neq 0$ \bullet or $2 \operatorname{Arcsin}(B1 \ XL \ / \ 2BORO)$ if ALE=0	1-2, 2*cm, rad	I, 3*E

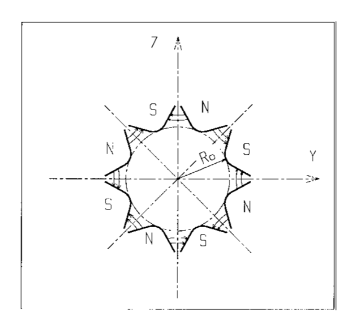
OBJET	Generation of an object		
BORO	Reference rigidity	kG.cm	E
KOBJ	Option index	1-6	I
if $KOBJ = 1[.01]$	[Non-] Symmetric object		
IY, IT, IZ, IP, IX, ID	Ray-Tracing assumes mid-plane symmetry Total number of points in $\pm Y$, $\pm T$, $\pm Z$, $\pm P$ [$+Z$, $+P$ with KOBJ = 1.01], $\pm X$. and $\pm D$ coordinates ($IY \leq 20,,ID \leq 20$)	$\text{IY*IT*IZ*IP*IX*ID} \leq 10^4$	6*1
PY, PT, PZ, PP, PX, PD	Step size in Y , T , Z , P , X and momentum ($PD = \delta B \rho / BORO$)	2(cm,mrad), cm, no dim.	6*E
YR, TR, ZR, PR, XR, DR	Reference ($DR = B\rho/BORO$)	2(cm,mrad), cm, no dim.	6*E
if $KOBJ = 2$	All the initial coordinates must be entered explicitly		
IMAX, IDMAX	total number of particles ; number of distinct momenta (if $IDMAX>1$, group particles of same momentum)	$MAX \le 10^4$	2*I
For $I = 1$, $IMAX$	Repeat <i>MAX</i> times the following line		
Y, T, Z, P, X, D, LET	Coordinates and tagging character of the $M\!A\!X$ particles ($D=B\rho/BORO$)	2(cm,mrad), cm, no dim., char	6*E, A1
$\mathbb{E}X(I=1,\mathbb{M}AX)$	MAX times 1 or -2. If $IEX(I) = 1$, trajectory number I is calculated. If $IEX(I) = -9$, it is not calculated	1 or -9	I MAXI
If KOBJ=3[.NN, NN=0003]	Reads coordinates from a storage file NN=00 (default): [b_]zgoubi.fai like data file FORMAT NN=01: read FORMAT is ``READ(NL,*) Y,T,Z,P,S,DP'' NN=02: read FORMAT is ``READ(NL,*) X,Y,Z,PX,PY,PZ NN=03: read FORMAT is ``READ(NL,*) DP,Y,T,Z,P,S,T		
IT1, IT2, ITStep	Read particles numbered IT1 to IT2, step ITStep (For more than 10^4 particles stored in <i>FNAME</i> , use ' <i>REBELOTE</i> ')	$\geq 1, \geq IT1, \geq 1$	3*I
IP1, IP2, IPStep	Read particles that belong in pass numbered IP1 to IP2, step IPStep	$\geq 1, \geq IP1, \geq 1$	3*I
YF, TF, ZF, PF, XF, DF, TF, TAG	Scaling factor. TAG-ing letter: no effect if '*', otherwise only particles with TAG=LET are retained.	7*no.dim, char.	7*E, A1
YR, TR, ZR, PR, XR, DR, TR	Reference. Given the previous line of data, all coordinate C is transformed to C*CF+CR	2(cm, mrad), cm, no dim., μ s	7*E
InitC	0 : set $new \ \vec{R}_0 = old \ \vec{R}_0$, $new \ \vec{R} = old \ \vec{R}$;	0-1	I

	1: set $new \ \vec{R}_0 = old \ \vec{R}, new \ \vec{R} = old \ \vec{R}$; 2: save $old \ \vec{R}$ in $new \ \vec{R}_0$, set $new \ \vec{R} = old \ \vec{R}_0$.		
FNAME	File name (e.g., zgoubi.fai) (NN in KOBJ=3.NN determines storage FORMAT)		A80
If KOBJ = 5[.NN, NN=01,99]	Generation of 11 particles, or 11*NN if $I \ge 2$ (for use with M	ATRIX, IORD = 1)	
PY, PT, PZ, PP, PX, PD	Step sizes in Y, T, Z, P, X and D	2(cm,mrad), cm, no dim.	6*E
YR, TR, ZR, PR, XR, DR	Reference trajectory ($DR = B\rho/BORO$)	2(cm,mrad), cm, no dim.	6*E
$If KOBJ = 5.01$ $\alpha_Y, \beta_Y, \alpha_Z, \beta_Z, \alpha_X, \beta_X,$ D_Y, D_Y', D_Z, D_Z'	additional data line : Initial beam ellipse parameters ¹	2(no dim.,m), ?, ?, 2(m,rad)	6*E, 4*E
If KOBJ = 5.NN, NN=02-99 YR, TR, ZR, PR, XR, DR	i = 1 to 98 (if, resp ly , NN=02 to 99) additional data lines : Reference trajectory # i ($DR=B\rho/BORO$)	2(cm,mrad), cm, no dim.	6*E
If $KOBJ = 6$	Generation of 61 particles (for use with $MATRIX$, $IORD = 2$	()	
PY, PT, PZ, PP, PX, PD	Step sizes in Y , T , Z , P , X and D	2(cm,mrad), cm, no dim.	6*E
YR, TR, ZR, PR, XR, DR	Reference trajectory ; $DR = B\rho/BORO$	2(cm,mrad), cm, no dim.	6*E
If KOBJ = 7	Object with kinematics		
IY, IT, IZ, IP, IX, ID	Number of points in $\pm Y$, $\pm T$, $\pm Z$, $\pm P$, $\pm X$; ID is not used	$\textit{IY*IT*IZ*IP*IX*ID} \leq 10^4$	6*I
PY, PT, PZ, PP, PX, PD	Step sizes in Y , T , Z , P and X ; PD = kinematic coefficient, such that $D(T) = DR + PD * T$	2(cm,mrad), cm, mrad ⁻¹	6*E
YR, TR, ZR, PR, XR, DR	Reference ($DR = B\rho/BORO$)	2(cm,mrad), cm, no dim.	6*E
If $KOBJ = 8$	Generation of phase-space coordinates on ellipses 2		
IY, IZ, IX	Number of samples in each 2-D phase-space; if zero the central value (below) is assigned	$\begin{split} 0 &\leq IX, IY, IZ \leq IMAX, \\ 1 &\leq IX * IY * IZ \leq IMAX \end{split}$	3*I
$Y_0, T_0, Z_0, P_0, X_0, D_0$	Central values ($D_0 = B\rho/BORO$)	m, rad, m, rad, m, no dim.	6*E
$ \alpha_Y, \beta_Y, \varepsilon_Y/\pi $ $ \alpha_Z, \beta_Z, \varepsilon_Z/\pi $ $ \alpha_X, \beta_X, \varepsilon_X/\pi $	ellipse parameters and emittances	no dim., m, m no dim., m, m no dim., m, m	3*E 3*E 3*E

 $^{^1~}$ They can be transported by using MATRIX $^2~$ Similar possibilities, random, are offered with MCOBJET, KOBJ=3 (p. 216)

OBJETA	Object from Monte-Carlo simulation of decay reaction		
	$M1+M2\longrightarrow M3+M4$ and $M4\longrightarrow M5+M6$		
BORO	Reference rigidity	kG.cm	E
IBODY, KOBJ	Body to be tracked: $M3$ ($IBODY=1$), $M5$ ($IBODY=2$) $M6$ ($IBODY=3$); type of distribution for Y_0 and Z_0 : uniform ($KOBJ=1$) or Gaussian ($KOBJ=2$)	1-3,1-2	2*I
I MAX	Number of particles to be generated (use 'REBELOTE' for more)	$\leq 10^4$	I
M_1-M_6	Rest masses of the bodies	$6*GeV/c^2$	6*E
T_1	Kinetic energy of incident body	GeV	E
Y_0, T_0, Z_0, P_0, D_0	Only those particles in the range $Y_0 - \delta Y \leq Y \leq Y_0 + \delta Y$ $D_0 - \delta D \leq D \leq D_0 + \delta D$ will be retained	2(cm,mrad), no dim.	5*E
$\delta Y, \delta T, \delta Z, \delta P, \delta D$		2(cm,mrad), no dim.	5*E
XL	Half length of object : $-XL \le X_0 \le XL$ (uniform random distribution)	cm	E
IR1, IR2	Random sequence seeds	$2^* \simeq 0^6$	2*I

OCTUPOLE	Octupole magnet		
IL	$I\!L=1,2$: print field and coordinates along trajectories	0-2	I
XL, R_0, B_0	Length; radius and field at pole tip of the element	2*cm, kG	3*E
X_E, λ_E	Entrance face : Integration zone ; Fringe field extent ($\lambda_E=0$ for sharp edge)	2*cm	2*E
$NCE, C_0 - C_5$	NCE = unused C_0-C_5 = fringe field coefficients such that : $G(s)=G_0/(1+\exp\ P(s))$, with $G_0=B_0/R_0^3$ and $P(s)=\sum_{i=0}^5 C_i(s/\lambda)^i$	any, 6*no dim.	I, 6*E
X_S, λ_S	Exit face: Parameters for the exit fringe field; see entrance	2*cm	2*E
NCS , $C_0 - C_5$		0-6, 6*no dim.	I, 6*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	<pre>KPOS=1 : element aligned, 2 : misaligned ; shifts, tilt (unused if KPOS=1)</pre>	1-2, 2*cm, rad	I, 3*E



Octupole magnet

OPTICS Write out optical functions

IOPT, label, IMP IOPT = 0/1: Off/On. Transport the beam matrix; 0-1, string, 0-1 I, A, I

'label' : Can be 'all' or existing 'LABEL_1(NOEL)';

IMP=1 causes storage of optical functions in zgoubi.OPTICS.out.

ORDRE	Taylor expansions order		
IO	Taylor expansions of \vec{R} and \vec{u} up to $\vec{u}^{(IO)}$ (default is $IO=4$)	2-5	I

PARTICUL Particle characteristics

M,Q,G, au,X Mass ; charge ; gyromagnetic factor ; MeV/c², C, no dim., s 5*E COM life-time ; unusued

If M is of the form $\{M1 \ M2\}$, then when masses are assigned to particles from a previously defined object, the first half of the particles are given the mass M1, and the second half are given the mass M2.

If Q is zero, the reference charge is left unchanged.

NOTE : Only the parameters of concern need their value be specified (for instance M, Q when electric lenses are used); others can be set to zero.

PICKUPS	Beam centroid path; closed orbit		
N	0 : inactive ≥ 1 : total number of <i>LABEL</i> 's at which beam centroid is to be recorded	≥ 0	I
For $I = 1, N$	A list of N records follows		
<i>LABEL</i> 's	N labels at which beam centroid is to be recorded	strings	N*A8

PLOTDATA Intermediate output for the PLOTDATA graphic software [34]

To be documented.

POISSON	Read magnetic field data from POISSON output		
IC, IL	$I\!C=1,2$: print the field map $I\!L=1,2$: print field and coordinates along trajectories	0-2, 0-2	2*I
BNORM, XN,YN	Field and X-,Y-coordinate normalization coeffs.	3*no dim.	3*E
TITL	Title. Start with "FLIP" to get field map X-flipped		A80
IX,IY	Number of longitudinal and transverse nodes of the uniform mesh	$\leq 400, \leq 200$	2*I
FNAME ¹	File name		A80
ID, A, B, C [, A', B', C', B'' , etc., if $ID \ge 2$]	Integration boundary. Ineffective when $ID=0$. $ID=$ -1, 1 or ≥ 2 : as for <i>CARTEMES</i>	≥ -1 , 2*no dim., cm [,2*no dim., cm, etc.]	
IORDRE	Degree of interpolation polynomial as for <i>DIPOLE-M</i>	2, 4 or 25	I
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	<pre>KPOS=1 : element aligned, 2 : misaligned ; shifts, tilt (unused if KPOS=1)</pre>	1-2, 2*cm, rad	I, 3*E

```
I = 0 \\ 11 \ CONTINUE \\ I = I+1 \\ READ(LUN,101,ERR=99,END=10) \ K, \ K, \ K, \ R, \ X(I), \ R, \ R, \ B(I) \\ 101 \ FORMAT(I1, I3, I4, E15.6, 2F11.5, 2F12.3) \\ GOTO \ II \\ 10 \ CONTINUE
```

where X(I) is the longitudinal coordinate, and B(I) is the Z component of the field at a node (I) of the mesh. K's and R's are variables appearing in the POISSON output file outpoi.lis, not used here.

 $^{^1}$ FNAME (e.g., ''outpoi.lis'') contains the field map data. These must be formatted according to the following FORTRAN read sequence - details and possible updates are to be found in the source file 'fmapw.f':

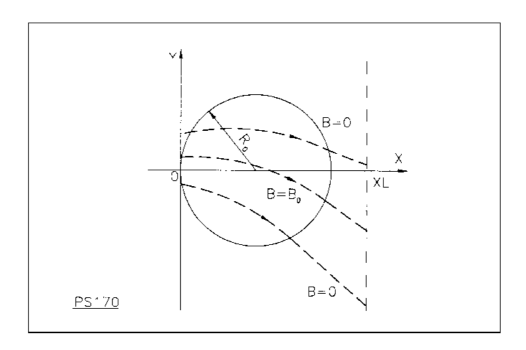
POLARMES	2-D polar mesh magnetic field map mid-plane symmetry is assumed		
IC, IL	$I\!C=1,2$: print the map $I\!L=1,2$: print field and coordinates along trajectories	0-2, 0-2	2*I
BNORM, AN,RN	Field and A-,R-coordinate normalization coeffs.	3*no dim.	3*E
TITL	Title. Start with "FLIP" to get field map X-flipped		A80
IA,JR	Number of angular and radial nodes of the mesh	$\leq 400, \leq 200$	2*I
FNAME ¹	File name		A80
ID, A, B, C [, A', B', C', B'' , etc., if $ID \ge 2$]	Integration boundary. Ineffective when $ID=0$. $ID=$ -1, 1 or ≥ 2 : as for <i>CARTEMES</i>	≥ -1 , 2*no dim., cm [,2*no dim., cm, etc.]	I,3*E [,3*E,etc.]
IORDRE	Degree of interpolation polynomial (see <i>DIPOLE-M</i>)	2, 4 or 25	I
XPAS	Integration step	cm	E
KPOS	as for DIPOLE-M. Normally 2.	1-2	I
If KPOS = 2 RE, TE, RS, TS		cm, rad, cm, rad	4*E
If KPOS = 1 <i>DP</i>		no dim.	E

```
OPEN (UNIT = NL, FILE = FNAME, STATUS = 'OLD' [,FORM='UNFORMATTED']) IF (BINARY) THEN READ(NL) (Y(J), J=1, JY) ELSE READ(NL,100) (Y(J), J=1, JY) ENDIF 100 FORMAT(10 F8.2) DO 1 I = 1,IX IF (BINARY) THEN READ (NL) X(I), (BMES(I,J), J=1, JY) ELSE READ(NL,101) X(I), (BMES(I,J), J=1, JY) ELSE READ(NL,101) X(I), (BMES(I,J), J=1, JY) EDIF 1 CONTINUE
```

where X(I) and Y(J) are the longitudinal and transverse coordinates and BMES is the Z field component at a node (I,J) of the mesh. For binary files, FNAME must begin with 'B_' or 'b_'. 'Binary' will then automatically be set to '.TRUE.'

 $^{^1}$ FNAME (e.g., spes2.map) contains the field data. These must be formatted according to the following FORTRAN read sequence - details and possible updates are to be found in the source file 'fmapw.f':

PS170	Simulation of a round shape dipole magnet		
IL	$I\!L=1,2$: print field and coordinates along trajectories	0-2	I
XL, R_0, B_0	Length of the element, radius of the circular dipole, field	2*cm, kG	3*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	<pre>KPOS=1 : element aligned, 2 : misaligned ; shifts, tilt (unused if KPOS=1)</pre>	1-2, 2*cm, rad	I, 3*E



Scheme of the PS170 magnet simulation.

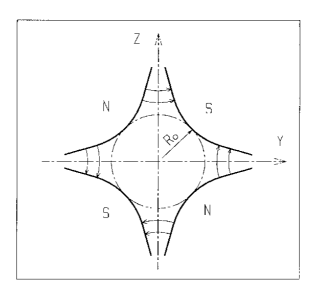
QUADISEX Sharp edge magnetic multipoles $B_Z \mid_{Z=0} = B_0 \left(1 + \frac{N}{R_0} Y + \frac{B}{R_0^2} Y^2 + \frac{G}{R_0^3} Y^3 \right)$ I ILIL = 1, 2: print field and coordinates along trajectories 0-2 XL, R_0, B_0 Length of the element; normalization distance; field 2*cm, kG 3*E N, EB1, EB2, EG1, EG2 Coefficients for the calculation of B. 5*no dim. 5*E if Y > 0: B = EB1 and G = EG1; if Y < 0: B = EB2 and G = EG2.

XPAS Integration step cm E

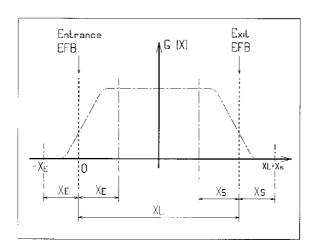
KPOS, XCE, KPOS=1: element aligned; 1-2, 2*cm, rad I, 3*E

YCE, ALE shifts, tilt (unused if KPOS=1)

QUADRUPO	Quadrupole magnet		
IL	$I\!L=1,2$: print field and coordinates along trajectories	0-2	I
XL, R_0, B_0	Length; radius and field at pole tip	2*cm, kG	3*E
X_E, λ_E	Entrance face : Integration zone extent ; fringe field extent ($\simeq 2R_0$, $\lambda_E=0$ for sharp edge)	2*cm	2*E
$NCE, C_0 - C_5$	NCE = unused C_0-C_5 = Fringe field coefficients such that $G(s)=G_0/(1+\exp P(s))$, with $G_0=B_0/R_0$ and $P(s)=\sum_{i=0}^5 C_i(s/\lambda)^i$	any, 6*no dim.	I, 6*E
	Exit face		
X_S, λ_S NCS, $C_0 - C_5$	See entrance face	2*cm 0-6, 6*no dim.	2*E I, 6*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	<pre>KPOS=1 : element aligned, 2 : misaligned ; shifts, tilt (unused if KPOS=1)</pre>	1-2, 2*cm, rad	I, 3*E



Quadrupole magnet



Scheme of the elements QUADRUPO, SEXTUPOL, OCTUPOLE, DECAPOLE, DODECAPO and MULTIPOL

(OX) is the longitudinal axis of the reference frame (0, X, Y, Z) of **zgoubi**.

The length of the element is XL, but trajectories are calculated from $-X_E$ to $XL + X_S$, by means of automatic prior and further X_E and X_S translations.

Ι

REBELOTE Jump to the beginning of zgoubiinput data file

NPASS, KWRIT, K[.n], [, Label1 [, Label2]]

NPASS: Number of runs; KWRIT = 1.1 (resp. 0.0) switches arbitrary; 3*I (inhibits) $FORTRAN\ WRITE$ s to .res and to screen; 0-1; 0, 22, 99 2A8

K option :

K=0: initial conditions (coordinates and spins) are generated following the regular functioning of object definitions. If random generators are used (e.g. in MCOBJET) their seeds will not be reset. K=22: next run will account for new parameters in

zgoubi.dat data list.

K=99: coordinates at end of previous pass are used as initial coordinates for the next pass; idem for spin components. K=99.1: Label1 is expected, subsequent passes will start from Label1 wat down to REBELOTE and so forth;

K=99.2: Label1 and Label2 are expected; last pass (# NPASS+1) will end at Label2 whereupon execution will jump to the keyword next to *REBELOTE* and will be carried out down to *'END'*.

if $K = 22^{-1}$

NPRM Number of parameters to be changed for next runs

Repeat NPR times the following sequence (tells parameters concerned, and for each its successive values):

LMNT, PRM, NV*Val Keyword # in zgoubi.dat list; parameter # under that see 'FIT' 2*I, NV*E

LMNT, PRM, NV*Val Keyword # in zgoubi.dat list; parameter # under that see 'FIT' Keyword; NV successive values (if NV < NPASS then keyword.

Reyword; NV successive values (if NV < NPASS then keyword

last value is maintained over remaining passes).

¹ K=22 is compatible with use of the FIT procedure: e.g., allows successive FITs in a run, with successive sets of optical parameters.

RESET Reset counters and flags

Resets counters involved in CHAMBR, COLLIMA HISTO and INTEG procedures

Switches off CHAMBR, MCDESINT, SCALING and SPNTRK options

SCALING	Time scaling of power supplies and R.F.		
IOPT, NFAM	<pre>IOPT = 0 (inactive) or 1 (active) ; NFAM = number of families to be scaled</pre>	0-1; 1-9	2*I
For NF=1, NFAM:	repeat NFAM times the following sequence:		
NAMEF [, Lbl [, Lbl]]	Name of family (i.e., keyword of concern), up to 2 labels		A8 [,A8[,A8]]
NT	$NT>0$: number of timings $NT=-1$: field scaling factor updated by $\it CAVITE$ $NT=-2$: $\it CAVITE$ RF law read from external data file	-2, -1 or 1-10	I
SCL(I),I=1,NT	Scaling values (a single one if $NT = -1$).	relative	NT*E
TIM(I), I = 1, NT	Corresponding timings, in units of turns (1 if $NT = -1$).	turn number	NT*I

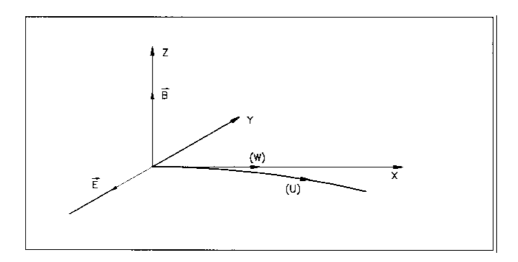
SEPARA 1 Wien Filter - analytical simulation

IA, XL, E, B,0-2, m, I, 3*E IA = 0: element inactive V/m, T

IA = 1: horizontal separation

IA = 2: vertical separation;

Length of the separator; electric field; magnetic field.



Horizontal separation between a wanted particle, (W), and an unwanted particle, (U). $\left(W\right)$ undergoes a linear motion while $\left(U\right)$ undergoes a cycloidal motion.

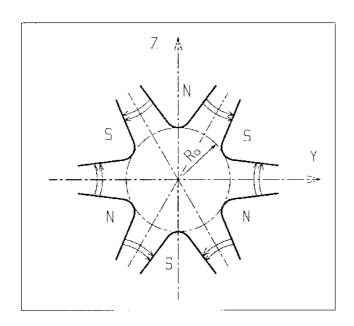
¹ SEPARA must be preceded by PARTICUL for the definition of mass and charge of the particles.

YCE, ALE

SEXQUAD	Sharp edge magnetic multipole $B_Z\mid_{Z=0}=B_0\left(rac{N}{R_0}Y+rac{B}{R_0^2}Y^2+rac{G}{R_0^3}Y^3 ight)$		
IL	IL=1,2 : print field and coordinates along trajectories	0-2	I
XL, R_0, B_0	Length of the element; normalization distance; field	2*cm, kG	3*E
N, EB1, EB2, EG1, EG2	Coefficients for the calculation of B. if $Y > 0$: $B = EB1$ and $G = EG1$; if $Y < 0$: $B = EB2$ and $G = EG2$.	5*no dim.	5*E
XPAS	Integration step	cm	E
KPOS, XCE,	KPOS=1: element aligned, 2: misaligned;	1-2, 2*cm, rad	I, 3*E

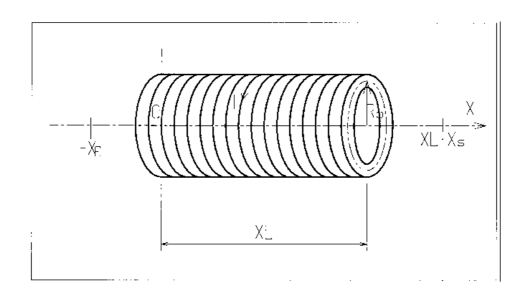
shifts, tilt (unused if *KPOS*=1)

SEXTUPOL	Sextupole Magnet		
IL	$I\!L=1,2$: print field and coordinates along trajectories	0-2	I
XL, R_0, B_0	Length; radius and field at pole tip of the element	2*cm, kG	3*E
X_E, λ_E	Entrance face : Integration zone ; fringe field extent ($\lambda_E=0$ for sharp edge)	2*cm	2*E
$NCE, C_0 - C_5$	NCE = unused C_0-C_5 = Fringe field coefficients such that $G(s)=G_0/(1+\exp P(s))$, with $G_0=B_0/R_0^2$ and $P(s)=\sum_{i=0}^5 C_i(s/\lambda)^i$	any, 6* no dim.	I, 6*E
X_S, λ_S	Exit face: Parameters for the exit fringe field; see entrance	2*cm	2*E
NCS , $C_0 - C_5$		0-6, 6*no dim.	I, 6*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	<pre>KPOS=1 : element aligned, 2 : misaligned ; shifts, tilt (unused if KPOS=1)</pre>	1-2, 2*cm, rad	I, 3*E



Sextupole magnet

SOLENOID	Solenoid		
IL	$I\!L=1,2$: print field and coordinates along trajectories	0-2	I
XL, R_0, B_0	Length ; radius ; asymptotic field (= $\mu_0 NI/XL$)	2*cm, kG	3*E
X_E, X_S	Entrance and exit integration zones	2*cm	2*E
XPAS	Integration step	cm	Е
KPOS, XCE, YCE, ALE	<i>KPOS</i> =1 : element aligned, 2 : misaligned; shifts, tilt (unused if <i>KPOS</i> =1)	1-2, 2*cm, rad	I, 3*E



SPNPRNL Store spin coordinates in file FNAME

FNAME 1 A80 Name of storage file (e.g., zgoubi.spn)

SPNSTORE Store spin coordinates every ${\it IP}$ other pass

FNAME 1 Name of storage file (e.g., zgoubi.spn) [; label(s) of the element(s) A80 $[, LABEL(s)]^{-2}$ at the exit of which the store occurs (10 labels maximum)]. [, 10*A10] IPStore every IP other pass (when using REBELOTEI

with $NPASS \ge IP - 1$).

SPNPRT Print spin coordinates

Print spin coordinates at the location where this keyword is introduced in the structure.

 $^{^1}$ FNAME = 'none' will inhibit printing. 2 If first $\it LABEL$ = 'none' then printing will be inhibited.

SPNTRK ¹	Spin tracking		
KSO	Initial conditions options	1-5	I
If KSO = 1 – 3	KSO = 1 (respectively 2, 3): all particles have their spin automatically set to $(1,0,0)$ – longitudinal [respectively $(0,1,0)$ – horizontal and $(0,0,1)$ – vertical]		
If KSO = 4	Repeat IMAX times (corresponding to the IMAX particles in 'OBJET') the following sequence:		
S_X, S_Y, S_Z	X,Y and Z initial components of the initial spin.	3*no dim.	3*E
If $KSO = 4.1$			
S_X, S_Y, S_Z	X,Y and Z components of the initial spins. These will be assigned to all particles.	3*no dim.	3*E
If KSO = 5 $TO, PO, A, \delta A$	Random distribution in a cone (see figure) Enter the following two sequences: Angles of average polarization: $A = \text{angle of the cone}$; $\delta A = \text{standard deviation}$	4*rad	4*E
IR	of distribution around A Random sequence seed	$\lesssim 10^6$	I
	Times and seed	≥ 10	-

 $^{^{1}\,}$ SPNTRK must be preceded by PARTICUL for the definition of G and mass.

SRLOSS	Synchrotron radiation loss	
KSR[.i]	Switch $i=1$ causes info output into <code>zgoubi.SRLOSS.out</code> $0-1$	2*I
STR1, STR2	Options: STR1 = 'ALL' or a particular magnet KEYWORD; 2*A STR2 = 'scale'	
Option, seed	1 : loss entails dp only $1-3,,>10^5$ 1 : loss entails dp and kick angle	I

SRPRNT Print SR loss statistics into zgoubi.res

SYNRAD	Synchrotron radiation spectral-angular densities		
KSR	Switch 0: inhibit SR calculations 1: start 2: stop	0-2	I
If $KSR = 0$			
D1, D2, D3	Dummies		3*E
If KSR = 1			
X0, Y0, Z0	Observer position in frame of magnet next to SYNRAD	3*m	3*E
If $KSR = 2$			
ν_1, ν_2, N	Frequency range and sampling	2*eV, no dim.	2*E, I

If KPOS = 2RE,TE,RS,TS

cm, rad, cm, rad

4*E

TOSCA	2-D and 3-D Cartesian or cylindrical mesh field map		
IC, IL	see CARTEMES	0-2, 0-2	2*I
BNORM, XN,YN, ZN	Field and X-,Y-,Z-coordinate normalization coefficients	4*no dim.	4*E
TITL	Title. Start with "FLIP" to get field map X-flipped		A80
IX, IY, IZ, MOD[.i]	Number of nodes of the mesh in the X,Y and Z directions, $IZ=1$ for single 2-D map; $MOD:$ operational and map $FORMAT$ reading mode $^1MOD \le 19:$ Cartesian mesh; $MOD \ge 20:$ cylindrical mesh; $.i.$, optional, tells the reading $FORMAT$, default is '*'.	$\leq 400, \leq 200,$ $\geq 1, \geq 0[.1-9]$	3*I
FNAME 1 $(K = 1, NF)$	Names of the NF files that contain the 2-D maps, ordered from $Z(1)$ to $Z(NF)$. If $MOD=0:NF=1+[IZ/2]$, the NF 2-D maps are for $0 \le$ they are symmetrized with respect to the $Z(1)=0$ plane. If $MOD=1:NF=IZ$, no symmetry assumed ; $Z(1)=Z_{max}$ $Z(1+[IZ/2])=0$ and $Z(NF)=-Z_{max}$. If $MOD=12:$ a single $FNAME$ file contains the all 3-D volume If $MOD=20-22:$ other symmetry options, see $toscap.f$ routine.	e.	A80
ID, A, B, C [, A', B', C', B'' , etc., if $ID \ge 2$]	Integration boundary. Ineffective when $ID=0$. $ID=$ -1, 1 or ≥ 2 : as for <i>CARTEMES</i>	≥ -1 , 2*no dim., cm [,2*no dim., cm, etc.]	I,3*E [,3*E,etc.]
IORDRE	If $IZ = 1$: as in <i>CARTEMES</i> If $IZ \neq 1$: unused	2, 4 or 25	I
XPAS	Integration step	cm	E
If Cartesian mesh (see M KPOS, XCE, YCE, ALE If polar mesh:	<pre>KPOS=1 : element aligned, 2 : misaligned ; shifts, tilt (unused if KPOS=1)</pre>	1-2, 2*cm, rad	I, 3*E
KPOS	as for <i>POLARMES</i> . Normally 2.	1-2	I

 1 Each file $\mathit{FNAME}(K)$ contains the field specific to elevation Z(K) and must be formatted according to the following $\mathit{FORTRAN}$ read sequence (that usually fits TOSCA code $\mathit{OUTPUTS}$ - details

and possible updates are to be found in the source file 'fmapw.f'):

```
\begin{array}{l} DO\ K=1, NF \\ OPEN\ (UNIT=NL, FILE=FNAME(K), STATUS='OLD'\ [,FORM='UNFORMATTED']) \\ DO\ J=1,\ JY \\ DO\ I=1,\ IX \\ IF\ (BINARY)\ THEN \\ READ(NL)\ Y(J),\ Z(K),\ X(I),\ BY(J,K,I),\ BX(J,K,I) \\ ELSE \\ READ(NL,*)\ Y(J),\ Z(K),\ X(I),\ BY(J,K,I),\ BX(J,K,I) \\ ENDIF \\ ENDDO \\ ENDDO \\ NL=NL+1 \\ ENDDO \end{array}
```

Note: for 2-D maps BX and BY are assumed zero at all nodes of the 2-D mesh, regardless of BX(J,1,I), BY(J,1,I) values. For binary files, FNAME must begin with 'B_' or 'b_'. 'Binary' will then automatically be set to '.TRUE.'

TRANSMAT	Matrix transfer		
IORDRE	Transfer matrix order	1-2	I
XL	Length (ineffective, for updating)	m	E
For $IA = 1, 6$:			
R(IA, IB), IB = 1, 6	First order matrix	m, rad	6 lines 6*E each
If IORDRE = 2	Following records <i>only</i> if <i>IORDRE</i> = 2		o"E each
T(IA, IB, IC),	Second order matrix, six 6*6 blocks	m, rad	36 lines 6*E each

TRAROT	Translation-Rotation		
TX, TY, TZ, RX, RY, RZ	Translations, rotations	3*m, 3*rad	6*E

TWISS Calculation of optical parameters; periodic parameters

KTW, FacD, FacA KTW = 0/1: Off/On; 0-1, any, any I,2*E

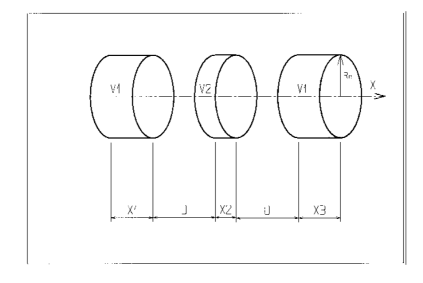
 $FacD \times D = \delta p/p$ applied, with D the momentum sampling in OBJET;

FacA: unused.

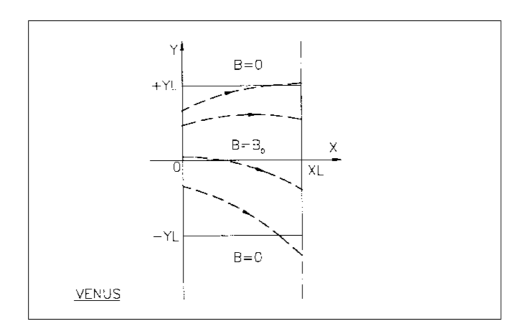
UNDULATOR Undulator magnet

To be documented

UNIPOT	Unipotential electrostatic lens		
IL	$I\!L=1,2$: print field and coordinates along trajectories	0-2	I
X_1, D, X_2, X_3, R_0	Length of first tube; distance between tubes; length of second and third tubes; radius	5*m	5*E
V_1,V_2	Potentials	2*V	2*E
XPAS	Integration step	cm	Е
KPOS, XCE, YCE, ALE	<i>KPOS</i> =1 : element aligned, 2 : misaligned; shifts, tilt (unused if <i>KPOS</i> =1)	1-2, 2*cm, rad	I, 3*E



VENUS	Simulation of a rectangular dipole magnet					
IL	$I\!L=1,2$: print field and coordinates on trajectories	0-2	I			
XL, YL, B_0	Length; width = $\pm YL$; field	2*cm, kG	3*E			
XPAS	Integration step	cm	E			
KPOS, XCE, YCE, ALE	<i>KPOS</i> =1 : element aligned, 2 : misaligned; shifts, tilt (unused if <i>KPOS</i> =1)	1-2, 2*cm, rad	I, 3*E			



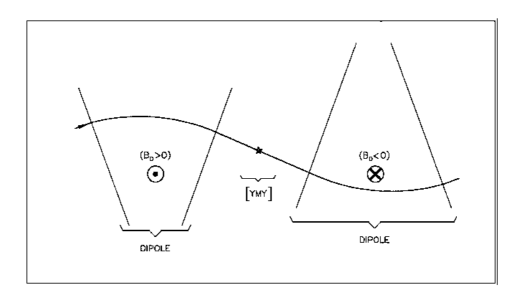
Scheme of VENUS rectangular dipole.

WIENFILT 1	Wien filter		
ΙL	$I\!L=1,2$: print field and coordinates along trajectories (otherwise $I\!L=0$)	0-2	Ι
XL, E, B, HV	Length; electric field; magnetic field; option: element inactive ($HV=0$) horizontal ($HV=1$) or vertical ($HV=2$) separation	m, V/m, T, 0-2	3*E, I
$X_{ extsf{E}}, \lambda_{E_E}, \lambda_{B_E}$	Entrance face: Integration zone extent; fringe field extent (\simeq gap height)	3*cm	3*E
C_{E0} – C_{E5} C_{B0} – C_{B5}	Fringe field coefficients for E Fringe field coefficients for B	6*no dim. 6*no dim.	6*E 6*E
	Exit face :		
$X_S, \lambda_{E_S}, \lambda_{B_S}$ C_{E0} – C_{E5} C_{B0} – C_{B5}	See entrance face	3*cm 6*no dim. 6*no dim.	3*E 6*E 6*E
XPAS	Integration step	cm	E
KPOS, XCE, YCE, ALE	<i>KPOS</i> =1 : element aligned, 2 : misaligned; shifts, tilt (unused if <i>KPOS</i> =1)	1-2, 2*cm, rad	I, 3*E

¹ Use *PARTICUL* to declare mass and charge.

YMY Reverse signs of Y and Z axes

Equivalent to a 180° rotation with respect to X-axis



The use of YMY in a sequence of two identical dipoles of opposite signs.

PART C

Examples of input data files and output result files

Examples 259

INTRODUCTION

Several examples of the use of **zgoubi** are given here. They show the contents of the input and output data files, and are also intended to help understanding some subtleties of the data definition.

Example 1: checks the resolution of the QDD spectrometer SPES 2 of SATURNE Laboratory [38], by means of a *Monte Carlo initial object* and an *analysis of images* at the focal plane with histograms. The *measured field maps* of the spectrometer are used for that purpose. The design of SPES 2 is given in Fig. 47.

Example 2: calculates the *first and second order transfer matrices* of an 800 MeV/c kaon beam line [39] at each of its four foci: at the end of the first separation stage (vertical focus), at the intermediate momentum slit (horizontal focus), at the end of the second separation stage (vertical focus), and at the end of the line (double focusing). The first bending is represented by its *3-D map* previously calculated with the TOSCA magnet code. The second bending is simulated with *DIPOLE*. The design of the line is given in Fig. 48.

Example 3: illustrates the use of MCDESINT and REBELOTE with a simulation of the in-flight decay

$$K \longrightarrow \mu + \nu$$

in the SATURNE Laboratory spectrometer SPES 3 [20]. The angular acceptance of SPES 3 is ± 50 mrd horizontally and ± 50 mrd vertically; its momentum acceptance is $\pm 40\%$. The bending magnet is simulated with *DIPOLE*. The design of SPES 3 is given in Fig. 49.

Example 4: illustrates the functioning of *the fitting procedure*: a quadrupole triplet is tuned from -0.7/0.3 T to field values leading to transfer coefficients R12=16.6 and R34=-.88 at the end of the beam line. Other example can be found in [40].

Example 5: shows the use of the *spin and multiturn tracking procedures*, applied to the case of the SATURNE 3 GeV synchrotron [7, 10, 36]. Protons with initial vertical spin $(\vec{S} \equiv \vec{S}_Z)$ are accelerated through the $\gamma G = 7 - \nu_Z$ depolarizing resonance. For easier understanding, some results are summarized in Figs. 51, 52 (obtained with the graphic post-processor, see Part D).

Example 6: shows *ray-tracing through a micro-beam line* that involves *electro-magnetic quadrupoles* for the suppression of second order (chromatic) aberrations [6]. The extremely small beam spot sizes involved (less than 1 micrometer) reveal the high accuracy of the ray-tracing (Figs. 53).

260 Examples

1 MONTE CARLO IMAGES IN SPES 2

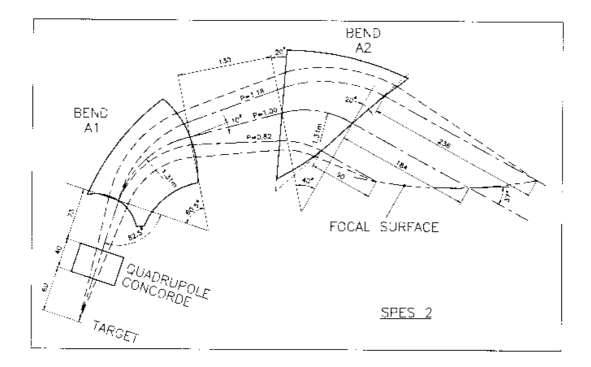


Figure 47: Design of SPES 2.

zgoubidata file.

$\label{eq:correction} \textbf{Excerpt from zgoubioutput: histograms of initial beam } \textbf{coordinates.}$

SPES2 QDD SPECTROMETER, USING FIELD MAPS; MONTE-CARLO OBJECT WITH MOMENTUM GRID.			coordinates.			
'MCOBJET'	DD MAPS/ MONTE CARDO OBOECT WITH MOMENTON	1				
2335.	REFERENCE RIGIDITY.			*****************	******	
2	DISTRIBUTION IN GRID.			HISTO		
10000	NUMGER OF PARTICLES.					
1 1 1 1 1	UNIFORM DISTRIBUTIONS			HISTOGRAMME DE LA COORDONNEE D		
. 0. 0. 0. 1.	CENTRAL VALUES OF BARS.			PARTICULES PRIMAIRES ET SECONDAIRES		
. 1 1 1 1 5	NUMBER OF BARS IN MOMENTUM.			DANS LA FENETRE: 0.9970 / 1.003		
0. 0. 0. 0001	SPACE BETWEEN MOMENTUM BARS.			NORMALISE		
. 50.e-3 0. 50.e-3 0. 0.	WIDTH OF BARS.					
. 1. 1. 1. 1. 1.	SORTING CUT-OFFS (UNUSED)		20			
9. 9. 9. 9.	FOR P(D) (UNUSED)		19			
86387 548728 472874	SEEDS.		18			
'HISTO'		2	17	D D D	D	
1 .997 1.003 80 1	HISTO OF D.		16	D D D	D	
20 'D' 1 'Q'			15	D D D	D	
'HISTO'		3	14	D D D	D	
3 -60. 60. 80 1	HISTO OF THETAO.		13	D D D	D	
20 'T' 1 'Q'			12	D D D	D	
'HISTO'		4	11	D D D	D	
5 -60. 60. 80 1	HISTO OF PHIO.		10	0 0 0	0 D	
20 'P' 1 'Q'			9	D D D		
DRIFT'		5	8 7	D D D	D D	
1.5				D D D		
CARTEMES'	QUADRUPOLE MAP.	6	6	D D D	D	
0 0	IC IL.		5	D D D	D	
96136E-3 1. 1.	BNORM, XNorm, YNorm		4	D D D D	D D	
+++ CONCORDE ++++			2		D	
39 23	IX IY.		1		D	
concord.map	field map file name, quadrupole		1		-	
0 0 0 0	NO LIMIT PLANE.			12345678901234567890123456789012345678901234567890 2 3 4 5 6 7		
	IORDRE.			2 3 4 5 0 /	o	
.5	XPAS.					
0 0 0	KPOS.			TOTAL COMPTAGE : 10000 SUR 10000		
DRIFT'		7		NUMERO DU CANAL MOYEN : 51		
1.8				COMPTAGE AU " : 2038		
CHANGREF'	POSITIONING OF THE	8		VAL. PHYS. AU " : 1.000		
. 32.5 -35.6	1-ST BENDING.			RESOLUTION PAR CANAL : 7.500E-0		
CARTEMES'		9		PARAMETRES PHYSIQUES DE LA DISTRIBUTION :		
0 0				COMPTAGE = 10000 PARTICULES		
1.04279E-3 1. 1.				MIN = 0.9980 , MAX = 1.002 , MAX-MIN = 4	00006-03	
+++ A1 ++++				MOYENNE = 1.000	.0000	
117 52				SIGMA = 1.4108E-03		
al.map	field map file name, first dipole			51GFA = 1.4100E 05		
0 0 0 0			TRA.T	IEX,D,Y,T,Z,P,S,time: 1 0.9980 0.000 -30.24 0.000 44.63	0 0000 0 0000	
_				,-,-,-,-,		
5			*****	************	******	
0 0 0				HISTO		
CHANGREF'	POSITIONING OF THE	10				
28.65 -27.6137	EXIT FRAME.			HISTOGRAMME DE LA COORDONNEE THETA		
DRIFT'		11		PARTICULES PRIMAIRES ET SECONDAIRES		
3.15	DOCUMENT OF THE	12			RD)	
CHANGREF'	POSITIONING OF THE	12		NORMALISE		
. 27.5 -19.88 CARTEMES'	2-ND BENDING.	13				
0 0		13	20			
1.05778E-3 1. 1.			19			
+++ A2 ++++			18			
132 80			17		T	
a2.map	field map file name, second dipole		16	T T	T	
0 0 0 0	ricia map rire name, become arpore		15	TT T TT T T T T T T T	TT TT	
0 0 0 0			14	TT T TT T TTTT T TTTTTT T	T T TTT TTT	
.5			13	TTTTTT TTTTTTT TTTT TTT TT TTT T TTTTTT	TTT TTTTTTT	
0 0 0			12	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	TTTTTTTTTTTTT	
CHANGREF'	POSITIONING OF THE	14	11	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	TTTTTTTTTTTTT	
18121.945	EXIT FRAME.		10	000000000000000000000000000000000000000	0000000000000	
DRIFT'	:== -:=::::::::::::::::::::::::::::::::	15	9	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT		
.55			8	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT		
'HISTO'	HISTO OF Y:	16	7	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT		
25 2. 80 1	SHOWS THE RESOLUTION		6	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT		
20 'Y' 1 'Q'	OF THE SPECTROMETER.		5	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT		
END'		17	4	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT		
			3	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT		
			2	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT		
			1	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	TTTTTTTTTTTTT	
				123456789012345678901234567890123456789012345678901234567890	10045550000000000	
				2 3 4 5 6 7		
				TOTAL COMPTAGE : 10000 SUR 10000		
				NUMERO DU CANAL MOYEN : 51		
				COMPTAGE AU " : 128		
				VAL. PHYS. AU " : 3.331E-15 (MRD)		
				RESOLUTION PAR CANAL : 1.50 (MRD)		
				PARAMETRES PHYSIQUES DE LA DISTRIBUTION :		
				COMPTAGE = 10000 PARTICULES MIN = -49.99 , MAX = 50.00 , MAX-MIN =	99.98 (MRD)	
				MOYENNE = 0.3320 (MRD)	(PIRD)	
				SIGMA = 29.04 (MRD)		
				(AMD)		
				IEX,D,Y,T,Z,P,S,time : 1 0.9980 0.000 -30.24 0.000 44.63		
			*****	*****************	******	

```
# HISTO

# HISTOGRAMME DE LA COORDONNEE PHI
PARTICULES PRIMATRES ET SECONDAIRES
DANS LA FENETRE : -60.00 / 60.00 (MRD)
NORMALISE

**DANS LA FENETRE : -60.00 / 60.00 (MRD)
NORMALISE

**DANS LA FENETRE : -60.00 / 60.00 (MRD)
NORMALISE

**DANS LA FENETRE : -60.00 / 60.00 (MRD)

**DANS LA FENETRE : -60.00 / MRD LA FENETRE EN FEN
```

Excerpt from zgoubioutput: the final momentum resolution histogram at the spectrometer focal surface.

```
16 HISTO
                            HISTO
                                          OF
                              HISTOGRAMME DE LA COORDONNEE Y PARTICULES PRIMAIRES ET SECONDAIRES DANS LA FENETRE : -0.5000 / 2 NORMALISE
                                                                                   2.000
    19
18
17
16
15
14
13
12
11
10
9
8
7
6
                                                      YYY
                                                                     YY
                                                                                     YY
                                                                                                 YYYY
                                      YY Y
YY YY
                                                      YYY
                                                                                                 YYYY
                12345678901234567890123456789012345678901234567890123456789012345678901
          TOTAL COMPTAGE

NUMERO DU CANAL MOYEN

COMPTAGE AU " "

VAL. PHYS. AU " "

RESOLUTION PAR CANAL
                                                      : 10000 SUR 10000
: 51
: 246
: 0.750 (CM)
: 3.125E-02 (CM)
          PARAMETRES PHYSIQUES DE LA DISTRIBUTION:

COMPTAGE = 10000 PARTICULES

MIN = -0.1486 , MAX = 1.652

MOVENNE = 0.7576 (CM)

SIGMA = 0.4621 (CM)
                                                                              , MAX-MIN = 1.800
                                                                                                                  (CM)
TRAJ 1 IEX,D,Y,T,Z,P,S,time : 1 0.9980 0.2475 74.43 -6.2488E-03 -6.929 697.41 0.0000
```

2 TRANSFER MATRICES ALONG A TWO-STAGE SEPARATION KAON BEAM LINE

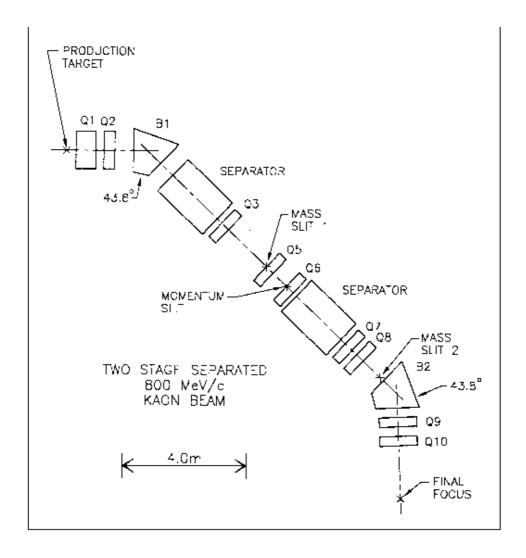


Figure 48: Design of 800 MeV/c kaon beam line.

oubidata file.	'COLLIMA' FIRST VERTICAL FOCUS, 2 MASS SLIT	
800 MeV/c KAON BEAM LINE. CALCULATION OF TRANSFER COEFFICIENTS.	2 14.6 .15E10 0. 0.	
OBJET' 1 668.5100 AUTOMATIC GENERATION OF	'DRIFT' 20.0	
AN OBJECT FOR CALCULATION 1 .1 .1 0001 OF THE FIRST ORDER TRANSFER	'QUADRUPO' Q5 0	
. 0. 0. 0. 1. COEFFICIENTS WITH 'MATRIX'	45.72 15.24 10.93 30. 30.	
PARTICUL' 2 493.646 1.60217733E-19 0. 0. 0. KAON M & Q, FOR USE IN WIEN FILTER	4 0.2490 5.3630 -2.4100 0.9870 0. 0.	
DRIFT' 3 35.00000	30. 30. 4 0.2490 5.3630 -2.4100 0.9870 0. 0.	
QUADRUPO' Q1 4	1.1 1 0.0.0.	
76.2 15.24 13.6 30. 30.	'DRIFT' 10.0	
4 0.2490 5.3630 -2.4100 0.9870 0. 0.	'MULTIPOL' COMPENSATION OF O ABERRATIONS AT VF2	
30. 30. 4	10. 15.24 0. 0. 0. 1. 0. 0. 0. 0. 0.	
1.1 1 0.0.0.	0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 4 0.2490 5.3630 -2.4100 0.9870 0. 0.	
DRIFT' 5 25.00000	0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 4 0.2490 5.3630 -2.4100 0.9870 0. 0.	
QUADRUPO' Q2 6	0. 0. 0. 0. 0. 0. 0. 0. 0. 0. .4	
5.72 15.24 -11.357	1 0.0.0.	
30. 30. 4	'DRIFT' 10.0	
30. 30. 4	'QUADRUPO' Q6	
1.1	45.72 15.24 -11.18 30. 30.	
1 0.0.0. DRIFT' 7	4 0.2490 5.3630 -2.4100 0.9870 0. 0.	
-1.898 TOSCA' 3-D MAP THE OF FIRST 8	30. 30. 4 0.2490 5.3630 -2.4100 0.9870 0. 0.	
0 0 BENDING MAGNET .0313E-3 1. 1. 1. B, X, Y, Z normalization coefficients	1.1 1 0.0.0.	
D map at z=0, from TOSCA	'DRIFT'	
9 39 1 w6_0.map	'WIENFILT' SECOND VERTICAL WIEN FILTER	
0. 0. 0.	0 2.16 -55.E5 .0215576 2	
1.1	20. 10. 10. 0.2401 1.8639 -0.5572 0.3904 0. 0.	
0 0 0 CHANGREF' 9	0.2401 1.8639 -0.5572 0.3904 0. 0.	
70.78 -43.8 AISCEAU' 10	20. 10. 10. 0.2401	
'DRIFT' 11 -49.38	0.2401 1.8639 -0.5572 0.3904 0. 0. 1.	
OCTUPOLE' 12	1. 0. 0. 0. 'DRIFT'	
10. 15.24 .6	30.0	
). 0. 4 0.2490 5.3630 -2.4100 0.9870 0. 0.	'QUADRUPO' Q7	
0. 0.	45.72 15.24 -6.44 30. 30.	
4 0.2490 5.3630 -2.4100 0.9870 0. 0. .4	4 0.2490 5.3630 -2.4100 0.9870 0. 0.	
1 0. 0. 0. SEXTUPOL' SX1, COMPENSATION 13	30. 30. 4 0.2490 5.3630 -2.4100 0.9870 0. 0.	
OF THE Theta.Phi ABERRATION 0. 15.24 2.4 AT VF1	1.1 1 0.0.0.	
0. 0. 0. 0. 4 0.2490 5.3630 -2.4100 0.9870 0. 0.	'DRIFT' 25.00000	
0. 0. 0. 0.	'QUADRUPO' Q8	
4 0.2490 5.3630 -2.4100 0.9870 0. 0. .4	45.72 15.24 8.085	
1 0. 0. 0. DRIFT' 14	30. 30. 4 0.2490 5.3630 -2.4100 0.9870 0. 0.	
50.0 'WIENFILT' FIRST VERTICAL WIEN FILTER 15	30. 30. 4 0.2490 5.3630 -2.4100 0.9870 0. 0.	
0	1.1	
.16 55.E50215576 2 0. 10. 10.	1 0.0.0. 'DRIFT'	
.2401 1.8639 -0.5572 0.3904 0. 0. .2401 1.8639 -0.5572 0.3904 0. 0.	40.0 'COLLIMA' SECOND VERTICAL FOCUS,	3
0. 10. 10. .2401 1.8639 -0.5572 0.3904 0. 0.	2 MASS SLIT 1 172E10 0.0.	
.2401 1.8639 -0.5572 0.3904 0. 0.	'MATRIX' TRANSFER COEFFICIENTS	
0. 0. 0.	2 0 'DRIFT'	
DRIFT' 16 30.	-25.0 'DIPOLE' SIMULATION OF THE MAP	
QUADRUPO' Q3 17	2 OF THE SECOND BENDING MAGNE 150 60 (upgraded version of keyword 'AIMANT')	
45.72 15.24 -6.34	79.3329 17.7656 140.4480 110. 170.	
30. 30. 4 0.2490 5.3630 -2.4100 0.9870 0. 0.	151. 4 .1455 2.26706395 1.1558 0. 0. 0.	
30. 30. 4	0.00 21.90 1.E6 -1.E6 1.E6 1.E6 151.	
1.1	4 .1455 2.26706395 1.1558 0. 0. 0.	
1 0.0.0. DRIFT' 18	-43.80 -21.90 -1.E6 -1.E6 1.E6 -1.E6 0. 0.	
10.0 MULTIPOL' SX2 + OCTU, COMPENSATION 19	4 .1455 2.26706395 1.1558 0. 0. 0. -43.80 -21.90 -1.E6 -1.E6 1.E6 -1.E6 1E6	
OF THE D.Phi AND D2.Phi	2 10.0 2.5	
0. 15.24 0. 08. 1.2 0. 0. 0. 0. 0. 0. ABERRATIONS AT VF1 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0.	2	
4 0.2490 5.3630 -2.4100 0.9870 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0. 0.	147.48099 -0.31007 147.48099 0.31007 'DRIFT'	
4 0.2490 5.3630 -2.4100 0.9870 0. 0. . 0. 0. 0. 0. 0. 0. 0. 0.	-15.00000 'QUADRUPO' Q9	
.4	0	
1 0. 0. 0. DRIFT' 20	35.56 12.7 -13.69 -13.91 30. 25.4	
90.0 MATRIX' TRANSFER COEFFICIENTS 21	4 0.2490 5.3630 -2.4100 0.9870 0. 0. 30. 25.4	
INMODER COEFFICIENTS ZI	4 0.2490 5.3630 -2.4100 0.9870 0. 0.	
2 0	.5	

```
'DRIFT'
                                                                              42
 25.00000
'QUADRUPO'
                                         010
                                                                              43
 35.56 12.7 11.97
30. 25.4
4 0.2490 5.3630 -2.4100 0.9870 0. 0.
30. 25.4
     0.2490 5.3630 -2.4100 0.9870 0. 0.
 1.1
1 0.0.0.
'DRIFT'
                                                                              44
200.0
'MATRIX'
                                         TRANSFER COEFFICIENTS
AT THE FINAL FOCUS
                                                                              45
'END'
                                                                              46
```

Excerpt of zgoubioutput: first and second order transfer matrices and higher order coefficients at the end of the line

```
FIRST ORDER COEFFICIENTS ( MKSA ):
                                -4.453265E-02 -3.049728E-04 -1.165832E-04
0.270335 4.700517E-05 1.763910E-05
05 -8.687757E-07 -3.60817 -1.731805E-02
            0.00000
                                                                                                                                                     -9.561918E-02
-7.815367E-02
                                                                                                                             0.00000
                                                                                                                                                      -3.983392E-02
                                                                                                                             1.00000
                                                                                                                                                       0.374917
                                                                      0.00000
                                                                                                                             0.00000
                                                                                                                                                          1.00000
            DetY-1 =
                                 -0.1170246601, DetZ-1 =
            R12=0 at 0.1647 m,
                                                                R34=0 at -0.6034E-01 m
     First order sympletic conditions (expected values = 0) : -0.1170 \qquad \qquad 3.4614\text{E-}06 \qquad -1.8207\text{E-}04 \qquad 3.0973\text{E-}05
                                                                               3.0973E-05 4.6007E-04 -8.0561E-05
                   SECOND ORDER COEFFICIENTS ( MKSA ):
1 11 7.34 1 21 -1.78 1 31 1.399E-02
1 12 -1.78 1 22 -530. 1 32 -1.308E-03
1 13 1.399E-02 1 23 -1.308E-03 1 33 -0.611
1 14 1.456E-02 1 24 -1.743E-03 1 34 -0.522
1 15 0.00 1 25 0.00 1 35 0.00
1 16 36.3 1 26 12.3 1 36 -2.771E-02
                                                                                                                                                           1 61 36.3
1 62 12.3
1 63 -2.771E-02
1 64 -2.211E-02
1 65 0.00
1 66 2.88
                                                              1 31 1.399E-02 1 41 1.456E-02 1 51
1 32 -1.308E-03 1 42 -1.743E-03 1 52
1 33 -0.611 1 43 -0.522 1 53
                                                                                                                            1 52
1 53
                                                                                                                                          0.00
                                                                                              1 44 0.163
1 45 0.00
                                                                                                                                         0.00
                                                                                              1 46 -2.211E-02
2 11 -303. 2 21 3.81 2 31 3.684E-02 2 12 3.81 2 23 2-5.821E-04 2 13 3.684E-02 2 23 -5.821E-04 2 33 1.05 2 14 3.581E-02 2 24 -1.638E-04 2 34 1.94 2 25 0.00 2 35 0.00
                                                              2 31 3.684E-02 2 41 3.581E-02 2 51
2 32 -5.821E-04 2 42 -1.638E-04 2 52
                                                                                                                                         0.00
                                                                                                                                                            2 62 -0.759
                                                                                              2 43 1.94
2 44 6.70
                                                                                                                             2 53
                                                                                                                                         0.00
                                                                                                                                                            2 63 -1.031E-02
2 64 -4.285E-02
                                                                                                                             2 54
                                                                                                                                          0.00
                                                              2 35 0.00
2 36 -1.031E-02
                                                                                             2 45 0.00
2 46 -4.285E-02
                             3 21 2.158E-02 3 31 20.6
3 22 64.6 3 32 1.61
3 23 1.61 3 33 0.710
3 24 0.496 3 34 0.128
3 25 0.00 3 35 0.00
3 26 8.793E-02 3 36 39.1
3 11 -0.145
                                                                                              3 41 86.0
3 42 0.496
3 12 2.158E-02
                                                                                                                             3 52
                                                                                                                                          0.00
                                                                                                                                                            3 62 8.793E-02
3 13 20.6
3 14 86.0
3 15 0.00
                                                                                                                             3 53
3 54
3 55
3 56
                                                                                              3 43 0.128
3 44 64.8
                                                                                                                                          0.00
                                                                                                                                                             3 63
                                                                                                                                                                        39.1
7.17
                                                                                                                                         0.00
                             4 21 1.146E-02 4 31
4 22 33.0 4 32
4 23 0.787 4 33
4 24 0.157 4 34
4 25 0.00 4 35
4 26 3.566E-02 4 36
4 11 -8.254E-02
                                                                           10.7
                                                                                              4 41
                                                                                                                             4 51
                                                                                                                                          0.00
4 11 -6.254E-02
4 12 1.146E-02
4 13 10.7
4 14 47.3
4 15 0.00
4 16 -0.127
                                                              4 32 0.787
4 33 0.365
4 34 6.774E-02
4 35 0.00
4 36 17.5
                                                                                             4 41 47.3
4 42 0.157
4 43 6.774
4 44 33.1
4 45 0.00
4 46 1.05
                                                                                                                             4 52
4 53
4 54
4 55
4 56
                                                                                                                                         0.00
                                                                                                                                                             4 62 3.566E-02
4 63 17.5
                                                                                                        6.774E-02
                                                                                                                                          0.00
                                                                                                          33.1
0.00
1.05
                                                                                                                                                             4 63 17.5
4 64 1.05
4 65 0.00
4 66 0.715
                                                                                                                                          0.00
5 41 -5.682E-02
5 42 6.947E-04
                                                                                                                             5 52
                                                                                                                                          0.00
                                                                                                                                                            5 62
                                                                                              5 43
5 44
5 45
5 46
                                                                                                                             5 53
5 54
5 55
5 56
                                                                                                                                         0.00
0.00
0.00
0.00
                                                                                                                                                            5 63 0.215
5 64 0.129
5 65 0.00
5 66 112.
                                                                                                        10.2
                   HIGHER ORDER COEFFICIENTS ( MKSA ):
              Y/Y3
                                          5784.8
                                      9.40037E+05
0.70673
0.42104
              Y/T3
                                        -18607.
1.04607E+05
               T/T3
              T/Z3
                                     -0.10234
              T/P3
                                       5.25793E-02
                                       32.161
18.425
-872.50
-785.20
              Z/Y3
              Z/P3
              P/Y3
                                        15.460
                                        7.5264
-409.98
-389.15
```

3 IN-FLIGHT DECAY IN SPES 3

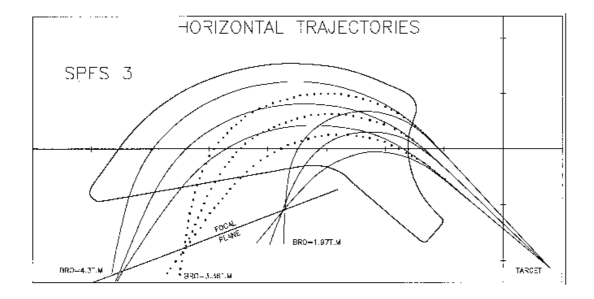


Figure 49: Design of SPES 3.

zgoubidata file

```
SIMULATION OF PION IN-FLIGHT DECAY IN SPES3
'MCOBJET'
                                                                1
3360.
                                 REFERENCE RIGIDITY (PION).
1
                                 DISTRIBUTION IN WINDOW.
                                BUNCHES OF 200 PARTICLES.
200
1 1 1 1 1 1 UNIFORM DISTRIBUTION 0. 0. 0. 0. 1. CENTRAL VALUES OF BARS.
.5e-2 50.e-3 .5e-2 50.e-3 0. 0.4 WIDTH OF BARS.
1 1 1 1 1 1 CUT-OFFS (UNUSED) 9 9. 9. 9. 9. . UNUSED.
186387 548728 472874
                                 SEEDS.
'PARTICUL'
139.6000 0. 0. 26.03E-9 0.
                                PION MASS AND LIFE TIME
'MCDESINT'
105.66 0.
                                 PION -> MUON + NEUTRINODECAY
136928 768370 548375
'ESL'
77.3627
'CHAMBR'
                                 STOPS ABERRANT MUONS.
1
1 100. 10. 245. 0.
'DIPOLE'
                                                                 6
2
180 130
80. 33. 208.5 140. 350.
46. -1.
4. .14552 5.21405 -3.38307 14.0629 0. 0. 0.
15. 0. -65. 0. 0. -65.
46. -1.
4. .14552 5.21405 -3.38307 14.0629 0. 0. 0.
-15. 69. 85. 0. 1.E6 1.E6
0.0.
4. .14552 5.21405 -3.38307 14.0629 0. 0. 0.
-15. 69. 85. 0. 1.E6 1.E6 1E6
2 10.0
4.
2
164.755 .479966 233.554 -.057963
'CHAMBR'
                                                                7
2
1 100.10.245.0.
                                TILT ANGLE OF
'CHANGREF'
                                                                8
0. 0. -49.
                                FOCAL PLANE.
'HISTO'
                                TOTAL SPECTRUM (PION + MUON).
2 -170. 130. 60 1
20 'Y' 1 'Q'
                               PION SPATIAL SPECTRUM
'HISTO'
                                                         10
2 -170. 130. 60 2
                                AT FOCAL PLANE.
20 'P' 1 'P'
                               MUON SPATIAL SPECTRUM
'HISTO'
                                                              11
2 -170. 130. 60 3
                                AT FOCAL PLANE.
20 'y' 1 'S'
'HISTO'
                                MUON MOMENTUM SPECTRUM
                                                              12
1 .2 1.7 60 3
20 'd' 1 'S'
                                AT FOCAL PLANE.
'REBELOTE'
                                (49+1) RUNS = CALCULATION OF 13
49 0.1 0
                                (49+1) * 200 TRAJECTORIES.
'END'
                                                               14
```

Excerpt of zgoubioutput: histograms of primary and secondary particles at focal surface of SPES3.

```
TOTAL
      9 HISTO
                             SPECTRUM
                               HISTOGRAMME DE LA COORDONNEE Y
PARTICULES PRIMAIRES ET SECONDAIRES
DANS LA FENETRE : -1.7000E+02 / 1.3000E+02 (CM)
NORMALISE
                                                                                          11 HISTO
                                                                                                         MUON
                                                                                                                    SPATIAL
                                                                                                                     HISTOGRAMME DE LA COORDONNEE Y
PARTICULES SECONDAIRES
DANS LA FENETRE: -1.7000E+02 / 1.3000E+02 (CM)
NORMALISE
   19
18
17
16
15
14
13
12
11
10
9
                                                19
                                                                                                                                     16
15
14
13
12
                                                YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY
                                                YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY
                                                YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY
                                              YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY
                                                                                                                         1234567890123456789012345678901234567890123456789012345678901
                TOTAL COMPTAGE
NUMERO DU CANAL MOYEN
COMPTAGE AU " "
VAL. PHYS. AU " "
                                                    9887
55
281
                                                                                                      TOTAL COMPTAGE
                                                                                                                                             605 SUR 10000
                                                                                                       NUMERO DU CANAL MOYEN
COMPTAGE AU " "
VAL. PHYS. AU " "
RESOLUTION PAR CANAL
                                                                                                      NUMERO
                                                                                                                                               50
                                                 : 0.000E+00 (CM)
                                                 : 5.000E+00 (CM)
                                                                                                                                      : -2.500E+01 (CM)
: 5.000E+00 (CM)
                PARAMETRES PHYSIQUES DE LA DISTRIBUTION:

COMPTAGE = 9887 PARTICULES

MIN = -1.6687E+02, MAX = 9.4131E+01, MAX-MIN = 2.6100E+02(CM)

MOYENNE = -9.2496E+01 (CM)

SIGMA = 5.3583E+01 (CM)
                                                                                                      PARAMETRES PHYSIQUES DE LA DISTRIBUTION:

COMPTAGE = 605 PARTICULES

MIN = -1.6687E+02, MAX = 9.4131E+01, MAX-MIN = 2.6100E+02 (CM)

MOYENDE = -2.2782E+01 (CM)

SIGMA = 5.4452E+01 (CM)
10 HISTO
                   PION
                              SPATIAL
                                                                                     12 HISTO
                              HISTOGRAMME DE LA COORDONNEE Y
PARTICULES PRIMATRES
DANS LA FENETRE: -1.7000E+02 / 1.3000E+02 (CM)
NORMALISE
                                                                                                                     HISTOGRAMME DE LA COORDONNEE D
PARTICULES SECONDAIRES
DANS LA FENETRE: 2.0000E-01 / 1.7000E+00
NORMALISE
   19
18
17
16
15
14
13
12
11
10
9
8
7
6
5
                                                d
d
d
                                                                                                                                           d
dd
                                                                                                                                       dd
                                                                                                                                          dd
                                                dd dd
ddd d dd
ddd d dd
ddd d dd
ddd d ddd d
0000 0 000 0
                                               10
                                                                                                                                                   00
                                                                                                                                    ddddd ddddd dd dd
                                                                                                                                   ddddddddddddd ddddd
                                                12345678901234567890123456789012345678901234567890123456^{\frac{1}{7}}8901
                                                                                                                             ddddddddddddddddddddddddddddddddddddd
                                                                                                                     123456789012345678901234567890123456789012345678901
                TOTAL COMPTAGE

NUMERO DU CANAL MOYEN
COMPTAGE AU " "

VAL. PHYS. AU "

RESOLUTION PAR CANAL
                                                 9282 SUR 10000
: 55
: 264
: 0.000E+00 (CM)
: 5.000E+00 (CM)
                                                                                                       TOTAL COMPTAGE
                                                                                                                                              605 SUR 10000
                                                                                                      NUMERO DU CANAL MOYEN
COMPTAGE AU " "
VAL. PHYS. AU " "
RESOLUTION PAR CANAL
                PARAMETRES PHYSIQUES DE LA DISTRIBUTION:

COMPTAGE = 9282 PARTICULES

MIN= -9.5838E+01, MAX = 9.3504E+01, MAX-MIN = 1.8934E+02 (CM)

MOYENDE = 4.9971E-01 (CM)

SIGMA = 5.3215E+01 (CM)
                                                                                                      PARAMETRES PHYSIQUES DE LA DISTRIBUTION:

COMPTAGE = 605 PARTICULES

MIN = 3.7184E-01, MAX = 1.3837E+00, MAX-MIN = 1.0119E+00

MOVENNE = 8.1639E-01

SIGMA = 2.2849E-01
```

4 USE OF THE FITTING PROCEDURE

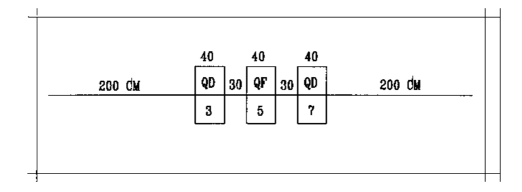


Figure 50: Vary B in all quadrupoles, for fitting of the transfer coefficients R_{12} and R_{34} at the end of the line. The first and last quadrupoles are coupled so as to present the same value of B.

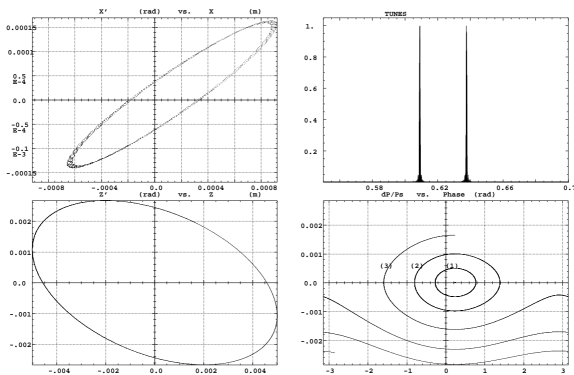
zgoubidata file.

```
MATCHING A SYMMETRIC QUADRUPOLE TRIPLET 'OBJET' 2501.73 750MeV/c
                                               750MeV/c PROTONS
11 PARTICLES FOR USE OF MATRIX
2. 2. 2. 2. 0. .001
0. 0. 0. 0. 0. 1.
 'ESL
'ESL '
200.
'QUADRUPO' 3
0
40. 15. -7.
0. 0.
6 .1122 6.2671 -1.4982 3.5882 -2.1209 1.723
0. 0.
6 .1122 6.2671 -1.4982 3.5882 -2.1209 1.723
1 0. 0. 0.
'ESL'
30.
'QUADRUPO'
0
40. 15. 3.
0. 0.
6 .1122 6.2671 -1.4982 3.5882 -2.1209 1.723
0. 0.
6 .1122 6.2671 -1.4982 3.5882 -2.1209 1.723
5.
1 0. 0. 0.
'ESL'
30.
'QUADRUPO'
0
40. 15. -7.
0. 0.
6 .1122 6.2671 -1.4982 3.5882 -2.1209 1.723
 0. 0.
6 .1122 6.2671 -1.4982 3.5882 -2.1209 1.723
5.
1 0. 0. 0.
'ESL'
200.
'MATRIX'
1 0
'FIT'
                             VARY B IN QUADS FOR FIT OF R12 AND R34
                                                                                                          10
2
3 12 7.12 2.
5 12 0. 2.
                            FIRST ORDER TRANSFER COEFFICIENTS
SYMMETRIC TRIPLET => QUADS #1 AND #3 ARE COUPLED
PARAMETER #12 OF ELEMENTS #3, 5 AND 7 IS FIELD VALUE
2 1 1 2 8 16.6 1. FIRST CONSTRAINT: R12=16.6, AFTER ELEMENT #8 (LAST DRIF 1 3 4 8 -.88 1. SECOND CONSTRAINT: R34=-.88
'END' 11
```

Excerpt of zgoubioutput: first order transfer matrices prior to and after fitting.

```
******************************
TRANSFER MATRIX WITH STARTING CONDITIONS :
               MATRICE DE TRANSFERT ORDRE 1 ( MKSA )
       STATE OF VARIABLES AFTER MATCHING:
        VARIABLE ELEMENT 3, PRMTR #12:
             COUPLED WITH ELEMENT 7, PRMTR #12
 STATUS OF VARIABLES
LMNT VAR PARAM MINIMUM
  MNT VAR PARAM MINIMUM INITIAL FINAL MAXIMUM STEP
3 1 12 -8.384E+00 -6.986E+00 -6.98648097E+00 -5.590E+00 2.424E-16
5 2 12 2.585E+00 3.230E+00 3.22956371E+00 3.877E+00 1.208E-16
                          INITIAL
                                       FINAL
                                                   MAXIMUM
 STATUS OF CONSTRAINTS
TYPE I J LMNT#
1 1 2 8
1 3 4 8 -
                     DESTRED
                                   WEIGHT
                                               REACHED
                                                             KT2
                               1.0000E+00 1.6600000E+01 8.2185E-02
1.0000E+00 -8.8000000E-01 9.1781E-01
                    1.6600E+01
                  -8.8000E-01
FINAL RUN, WITH NEW VARIABLES :
     9 MATRIX
         Frame for MATRIX calculation moved by :
          XC = 0.000 \text{ CM}, YC = 0.000 \text{ CM}, Path length of particle #1 : 580.00
                                              A = 0.00000 DEG ( = 0.000000 RD )
                                    580.0000 m
               MATRICE DE TRANSFERT ORDRE 1 ( MKSA )
       DetY-1 = -.0000011112

DetZ-1 = -.0000000156
     Determinants :
      R12=0 at
                   -3.1484 meters
      R34=0 at
                   -0.7073 meters
     First order sympletic conditions (expected values = 0):
-1.1112E-06 -1.5616E-08 0.0000E+00 0.0000E+00 0.0000E+00 0.0000E+00
```



5 MULTITURN SPIN TRACKING IN SATURNE 3 GeV SYNCHROTRON

Figure 51: Tracking over 3000 turns. These simulations exhibit the first order parameters and motions as produced by the multiturn ray-tracing.

(A) Horizontal phase-space: the particle has been launched near to the closed orbit (the fine structure is due to Y-Z coupling induced by bends fringe fields, also responsible of the off-centering of the local closed orbit - at ellipse center).

(B) Vertical phase-space: the particle has been launched with $Z_0 = 4.58 \ 10^{-3}$ m, $Z_0' = 0$. A least-square fit by $\gamma_Z Z^2 + 2\alpha_Z Z Z' + \beta_Z Z'^2 = \varepsilon_Z/\pi$ yields $\beta_Z = 2.055$ m, $\alpha_Z = 0.444$, $\gamma_Z = 0.582$ m⁻¹, $\varepsilon_Z/\pi = 12 \ 10^{-6}$ m.rad in agreement with matrix calculations. (C) Fractional tune numbers obtained by Fourier analysis for $\varepsilon_Y/\pi = \varepsilon_Z/\pi \simeq 12 \ 10^{-6}$ m.rad: $\nu_Y = 0.63795$, $\nu_Z = 0.60912$ (the integer part is 3 for both).

(**D**) Longitudinal phase-space ("(DP, phase)" in Zgoubi notations): particles with initial momentum dispersion of 5 10^{-4} (1), 10^{-3} (2), 1.65 10^{-3} (3) (out of acceptance), are accelerated at 1405 eV/turn ($\dot{B}=2.1$ T/s); analytical calculations give accordingly momentum acceptance of 1.65 10^{-3} .

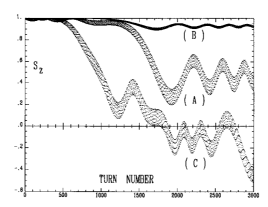


Figure 52: Crossing of $\gamma G = 7 - \nu_Z$, at $\dot{B} = 2.1$ T/s.

(A) $\varepsilon_Z/\pi=12.2\ 10^{-6}$ m.rad. The strength of the resonance is $|\varepsilon|=3.3\ 10^{-4}$. As expected from the Froissart-Stora formula the asymptotic polarization is about 0.44.

(B) The emittance is now $\varepsilon_Z/\pi = 1.2 \, 10^{-6}$ m.rad; comparison with (A) shows that $|\varepsilon|$ is proportional to $\sqrt{\varepsilon_Z}$.

(C) Crossing of this resonance for a particle having a momentum dispersion of 10^{-3} .

ubidata file (begi	-NUz, NUz=3.60877(pertur	bed)	71.6256	DTD 2 4 2	
JET'		34.04 MeV, proton	'BEND' 0 247.30039 0. 1.57	DIP 3 4 3	
1 -02 6.5E-02 .458		psilonY/pi ~ 0. (Closed orbit)	20. 8042760566 4 .2401 1.8639	5572 .3904 0. 0. 0.	
6 0.379 .458	0. 0. 1.0005 '1'	parioni/pr 0. (crosed orbit)	4 .2401 1.8639	5572 .3904 0. 0. 0.	
4 1.09 .458	0. 0. 1.001 '2' 0. 0. 1.0016 '3'		#30 120 30 bend 'ESL'	3 0. 01963495408 SD 2	
1 1 LING'			71.6256 'QUADRUPO'	QP 1	
LTIPOL			0 46.723 10763695		
15.388E-3 5034.391E-3		maG=7-Nuz+/-14E, E=3.3E-4 IN 3442 MACHINE TURNS,	0. 0. 6 .1122 6.2671 -1.4	982 3.5882 -2.1209 1.723	
3442 JADRUPO	FROM 834.041	TO 838.877 MeV	0. 0.	1982 3.5882 -2.1209 1.723	
15.388E-3 5034.391E-3			#30 50 30 Quad	3.3002 -2.1203 1.723	
3442			1 0. 0. 0. 'ESL'		
			392.148 'MULTIPOL'	QP 5	
15.388E-3 5034.391E-3 3442			0 48.6273 10. 0	765533 0. 0. 0. 0. 0. 0. 0. 0.	
VITE			0. 0. 0. 0. 0. 0.		
1.00378894 3442	RELATIVE CHA	NGE OF SYNCRHONOUS RIGIDITY	0. 0. 0. 0. 0. 0.		
TICUL' .2723 1.6021892E-19 1	.7928474 0. 0.		0. 0. 0. 0. 0. 0. #30 50 30 Quad		
TRK'			1 0. 0. 0.		
DRUPO'	QP 1	5	'ESL' 392.148		
723 10763695		763695 = FIELD FOR BORO=1 T.m	'QUADRUPO'	QP 1	
0. .1122 6.2671 -1.4982 3.	5882 -2.1209 1.723		46.723 10763695 0. 0.		
0. .1122 6.2671 -1.4982 3.	5882 -2.1209 1.723			982 3.5882 -2.1209 1.723	
50 30 Quad . 0. 0.			6 .1122 6.2671 -1.4	982 3.5882 -2.1209 1.723	
L' 6256	SD 2	6	#30 50 30 Quad 1 0. 0. 0.		
END'	DIP 3 4 3	7	'ESL' 71.6256	SD 2	
7.30039 0. 1.57776			'BEND'	DIP 3 4 3	
20. 804276056667 .2401 1.86395572	.3904 0. 0. 0.		247.30039 0. 1.57 20. 8042760566		
20. 804276056667 .2401 1.86395572	20. 8. .3904 0. 0. 0.			5572 .3904 0. 0. 0.	
0 120 30 bend 30. L'		8	4 .2401 1.8639	5572 .3904 0. 0. 0.	
6256	QP 5	9	'ESL'	3 0. 0. 01963495408 SD 2	
			71.6256 'MULTIPOL'	QP 5	
6273 10. 077319 0. 0. 0. 0. 00 .0 0.	0. 0. 0.	77319=765533+QUAD DEF FOR EXCITING THE DEPOLARI	ECT 0 ZING.6273 10. 07	65533 0. 0. 0. 0. 0. 0. 0. 0.	
.1122 6.2671 -1.4982 3. 0. 0. 0. 0. 0. 0. 0	. 0. 0. 0.	RESONNANCE.	0. 0. 0. 0. 0. 0.	0. 0. 0. 0. 0. 982 3.5882 -2.1209 1.723	
$.1122\ 6.2671\ -1.4982\ 3. \\ 0.\ 0.\ 0.\ 0.\ 0.\ 0.\ 0.$			0. 0. 0. 0. 0. 0.		
50 30 Quad			0. 0. 0. 0. 0. 0. #30 50 30 Quad		
	SD 2	10	1 0. 0. 0.		
END'	DIP 3 4 3	11	'ESL' 71.6256	SD 2	
7.30039 0. 1.57776			'BEND'	DIP 3 4 3	
20. 804276056667 .2401 1.86395572	.3904 0. 0. 0.		247.30039 0. 1.57 20. 8042760566		
20. 804276056667 .2401 1.86395572	20. 8. .3904 0. 0. 0.			5572 .3904 0. 0. 0.	
0 120 30 bend 30. L'		12	4 .2401 1.8639	5572 .3904 0. 0. 0. 3 0. 0. 01963495408	
6256 JADRUPO'	QP 1	13			
723 10763695	x- ±	13		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	
0.			'ESL' 392.148		
.1122 6.2671 -1.4982 3. 0.			'CAVITE'		
.1122 6.2671 -1.4982 3. 50 30 Quad . 0. 0.	5882 -2.1209 1.723		105.5556848673 3. 6000. 0.	SIN(phis) = .234162, dE=1.404	197 keV/
L'	SD 2	14	'FAISCNL' b_zgoubi.fai		
6256 END'	DIP 3 4 3	15	'SPNPRNL' zgoubi.spn		
7.30039 0. 1.57776			'SPNPRT' 'REBELOTE'		
20. 804276056667 .2401 1.86395572 20. 804276056667	.3904 0. 0. 0.		2999 0.1 99 'END'	TOTAL NUMBER OF TURNS = 3000	J
.2401 1.86395572 0 120 30 bend 3 0.	.3904 0. 0. 0.				
L'	SD 2	16			
6256 LTIPOL'	QP 5	17			
	0 0 0 0 0 0	0.			
6273 10. 0765533					
0. 0. 0. 0. 0. 0.	0. 0. 0. 0.				
	0. 0. 0. 0. 5882 -2.1209 1.723 0. 0. 0. 0.				

6 MICRO-BEAM FOCUSING WITH ELECTROMAGNETIC QUADRUPOLES

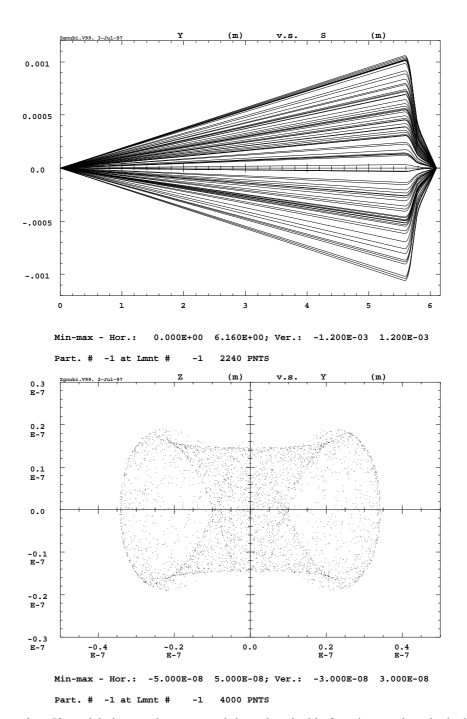


Figure 53: Upper plot: 50-particle beam tube ray-traced through a double focusing quadrupole doublet typical of the front end design of micro-beam lines. Initial conditions are : $Y_0 = Z_0 = 0$, angles T_0 and P_0 random uniform within ± 0.2 mrad, and momentum dispersion $\delta p/p$ uniform in $\pm 3\,10^{-4}$.

Lower plot: (**D**) sub-micronic cross-section at the image plane of a 4000-particle beam with initial conditions as above, obtained thanks to the second-order achromatic electro-magnetic quadrupole doublet (the inage size would be $\Delta Y \approx \Delta Z \approx \pm 50 \mu m$ with regular magnetic quadrupoles, due to the momentum dispersion). Note the high resolution of the ray-tracing which still reveals image structure of nanometric size.

zgoubidata file.

```
MICROBEAM LINE, WITH AN ELECTROMAGNETIC QUADRUPOLE DOUBLET.
                                                            RANDOM OBJECT DEFINITION
RIGIDITY (20keV PROTONS).
DISTRIBUTION IN WINDOW.
NUMBER OF PARTICLES.
 20.435
 200
                                                               NUMBER OF PARTICLES.
UNIFORM DISTRIBUTION.
CENTRAL VALUE, AND
HALF WIDTH OF DISTRIBUTION.
CUT-OFFS (UNUSED).
FOR P(D) - UNUSED.
200

1 1 1 1 1 1 1 1

0. 0. 0. 0. 0. 1.

0. .2e-3 0. .2e-3 0. 0.0003

10. 10. 10. 10. 10. 10. 10.

186387 548728 472874
                                                            SEEDS.
PARTICLE MASS AND CHARGE
  'PARTICUL'
 938.2723 1.60217733E-19 0. 0. 0. 'DRIFT'
                                                            FOR INTEGRATION IN E-FIELD.
DRIFT. 3
 59.
'EBMULT'
FIRST ELECTROMAGNETIC
.8
1 0. 0. 0.
'DRIFT'
                                                            DRIFT.
 4.9
'EBMULT'
                                                            SECOND ELECTROMAGNETIC
                                                                                   OUADRUPOLE.
.8
1 0. 0. 0.
'DRIFT'
25.
'HISTO'
                                                            DRIFT.
                                                            HISTOGRAM
OF THE Y COORDINATE.
 2 -5E-6 5E-6
20 'Y' 1 'Q'
'HISTO'
                        60 2
                                                            HISTOGRAM
 4 -5E-6 5E-6
20 'Z' 1 'O'
                                                                                            10
                        60 2
                                                               OF THE Z COORDINATE.
 20 'Z' 1
'FAISCNL'
                                                            RAYS ARE STORED IN RAYS 11
                                                            FOR FURTHER PLOTTING.
RUN AGAIN, FOR RAY-TRACING 12
TOTAL OF 200*(19+1) PARTICLES.
 19 0.1 0
'END'
```

zgoubioutput file.

```
LE PASSAGE SUIVANT EST LE
                      20-EME (ET DERNIER) PASSAGE DANS LA STRUCTURE
MCOBJET RANDOM OBJECT
Reference magnetic rigidity =
                                20 435 KG+CM
         Object built up of 200 particles Distribution in a Window
         Central values (MKSA units):
       Yo, To, Zo, Po, Xo, BR/BORO
                              0.000E+00 0.000E+00 0.000E+00 0.000E+00 0.000E+00 1.0000E+00
      Width ( +/- , MKSA units ) : DY, DT, DZ, DP, DX, DBR/BORO :
                              0.000E+00 2.000E-04 0.000E+00 2.000E-04 0.000E+00 3.0000E-04
      Cut-offs ( * +/-Width ) : NY, NT, NZ, NP, NX, NBR/BORO :
                                0.0
                                      0.0
                                            0.0
                                                 0.0
                                                        0.0
                                                              0.0
.....
  2 PARTICUL PARTICLE MASS
PARTICLE PROPERTIES :
     Masse = 938.2723000000 MeV/c2
Charge = 1.6021773300000D-19 C
```

```
3 DRIFT
                                         DRIFT.
                                                                        ESPACE LIBRE = 500.00000 CM
   TRAJ #1 D,Y,T,Z,P,S,IEX: 1.0002E+00 1.7062E-02 3.4124E-02 -2.6802E-02 -5.3603E-02
              4 DRIFT
                                             DRIFT.
                                                                        ESPACE LIBRE = 59.00000 CM
   TRAJ #1 D,Y,T,Z,P,S,IEX: 1.0002E+00 1.9075E-02 3.4124E-02 -2.9964E-02 -5.3603E-02
                                                                                                                                                                                                                                 5.59000E+02 1
......
              5 EBMULT
                                   ULTIPOLE :

LONGUEUR DE L'ELEMENT : 10.200 CM
RAYON DE GORGE RO = 10.00 CM
V-DIPOLE = 0.000000E+00 V
V-QUADRUPOLE = 0.000000E+00 V
V-SEXTUPOLE = 0.000000E+00 V
V-DECAPOLE = 0.000000E+00 V
V-DECAPOLE = 0.00000E+00 V
V-14-POLE = 0.00000E+00 V
V-18-POLE = 0.00000E+00 V
V-18-POLE = 0.00000E+00 V
LENTILLE A GRADIENT CRENEAU
              ---- MULTIPOLE
                                  | NULTIPOLE : | LONGUEUR DE L'ELEMENT : 10.200 CM | RAYON DE GORGE RO = 10.00 CM | DEPLIE | RAYON DE GORGE RO = 10.00 CM | DEPLIE | RAYON DE GORGE RO = 1.894930E+00 KG | DEPLIE | RAYON DE CONTROLE | DEPLIE |
              ---- MULTIPOLE
                                               Integration step :
                                                                                                   0.80 cm
DRIFT
                                                                         ESPACE LIBRE =
                                                                                                                        4.90000 CM
   TRAJ #1 D,Y,T,Z,P,S,IEX: 1.0002E+00 1.1032E-02 -8.0508E-01 -4.5922E-02 -1.6008E+00
                                                                                                                                                                                                                                 5.74100E+02 1
7 EBMULT SECOND
                                   ULTIPOLE :

LONGUEUR DE L'ELEMENT : 10.200 CM
RAYON DE GORGE RO = 10.00 CM
V-DIPOLE = 0.0000008+00 V
V-GUARDUPOLE = 1.3779908+04 V
V-SEXTUPOLE = 0.0000008+00 V
V-DECAPOLE = 0.0000008+00 V
V-DECAPOLE = 0.0000008+00 V
V-DECAPOLE = 0.0000008+00 V
V-14-POLE = 0.0000008+00 V
V-14-POLE = 0.0000008+00 V
V-14-POLE = 0.0000008+00 V
V-18-POLE = 0.0000008+00 V
V-18-POLE = 0.0000008+00 V
V-20-POLE = 0.0000008+00 V
                                    V-20-POLE
                                                                              0.000000E+00 V
                                       LENTILLE A GRADIENT CRENEAU
               MULTIPOLE :
LONGUEUR DE L'ELEMENT : 10.200 CM
RAYON DE GORGE RO = 10.00 CM
B-DIPOLE = 0.000000E+00 kG
B-QUADRUPOLE = 0.000000E+00 kG
B-SEXTUPOLE = 0.000000E+00 kG
B-DECAPOLE = 0.000000E+00 kG
B-DECAPOLE = 0.000000E+00 kG
B-DECAPOLE = 0.000000E+00 kG
B-14-POLE = 0.000000E+00 kG
B-14-POLE = 0.000000E+00 kG
B-18-POLE = 0.000000E+00 kG
B-18-POLE = 0.000000E+00 kG
                                       LENTILLE A GRADIENT CRENEAU
                                                 Integration step :
......
             8 DRIFT
                                         DRIFT.
                                                                         ESPACE LIBRE = 25.00000 CM
   TRAJ #1 D,Y,T,Z,P,S,IEX: 1.0002E+00 9.0257E-07 -2.3996E-01 -1.0770E-06 1.7947E+00
```

```
9 HISTO
                  HISTOGRA
                                HISTOGRAMME DE LA COORDONNEE Y
PARTICULES PRIMAIRES ET SECONDAIRES
DANS LA FENETRE : -5.0000E-06 / 5
NORMALISE
                                                                         5.0000E-06 (CM)
   19
18
17
16
15
14
13
12
11
10
9
8
7
6
5
                                                        1234567890123456789012345678901234567890123456789012345678901
                 TOTAL COMPTAGE
                                                        4000 SUR 4000
                 NUMERO DU CANAL MOYEN COMPTAGE AU " "
                                                       51
109
                 COMPTAGE AU " "
VAL. PHYS. AU " "
RESOLUTION PAR CANAL
                                                    : 0.000E+00 (CM)
: 1.667E-07 (CM)
                 PARAMETRES PHYSIQUES DE LA DISTRIBUTION:

COMPTAGE = 4000 PARTICULES

MIN = -3.4326E-06, MAX = 3.4347E-06, MAX-MIN = 6.8674E-06 (CM)

MOYENDE = -2.8531E-08 (CM)

SIGMA = 1.8619E-06 (CM)
TRAJ #1 D,Y,T,Z,P,S,IEX: 1.0002E+00 9.0257E-07 -2.3996E-01 -1.0770E-06 1.7947E+00 6.09300E+02 1

10 HISTO HISTOGRA
                                HISTOGRAMME DE LA COORDONNEE Z
PARTICULES PRIMAIRES ET SECONDAIRES
DANS LA FENETRE: -5.0000E-06 / 5
NORMALISE
                                                                         5.0000E-06 (CM)
   20
19
18
17
16
15
14
13
12
11
10
9
8
7
6
5
4
                                                                          Z
Z
Z
Z
ZZ
                                                                          ZZ
                                                       zz
                                                       zz
                                                                         ZZZ
                                                       00
                                                                         000
                                                       ZZZZ
                                                                         ZZZ
                                                      ZZZZZZZZZZZZZZZZZZZZZZZZ
                                                     ZZZZZZZZZZZZZZZZZZZZZZZZ
                                                     ZZZZZZZZZZZZZZZZZZZZZZZZ
                                \begin{smallmatrix} 1234567890123456789012345678901234567890123456789012345678901\\ 3 & 4 & 5 & 6 & 7 & 8 \end{smallmatrix}
                 TOTAL COMPTAGE
NUMERO DU CA
                 NUMBERO DU CANAL MOYEN
COMPTAGE AU " "
VAL. PHYS. AU " "
RESOLUTION PAR CANAL
                                                           51
                                                       169
                                                    : 0.000E+00 (CM)
                                                    : 1.667E-07 (CM)
                 PARAMETRES PHYSIQUES DE LA DISTRIBUTION:

COMPTAGE = 4000 PARTICULES

MIN = -1.9150E-06, MAX = 1.9110E-06, MAX-MIN = 3.8260E-06 (CM)

MOYENNE = -3.8539E-09 (CM)

SIGMA = 1.1232E-06 (CM)
TRAJ #1 D,Y,T,Z,P,S,IEX: 1.0002E+00 9.0257E-07 -2.3996E-01 -1.0770E-06 1.7947E+00 6.09300E+02 1

11 FAISCNL RAYS ARE
Print[s] occur at
12 REBELOTE RUN AGAIN,
     **** FIN D'EFFET DE 'REBELOTE' ****

IL Y A EU 20 PASSAGES DANS LA STRUCTURE
# PARTICULES ENVOYEES : 4000
PGM PRINCIPAL : ARRET SUR CLE REBELOTE
```

PART D

Running zgoubi and its post-processor/graphic interface zpop

INTRODUCTION

The basic **zgoubi** *FORTRAN* package is transportable; it has been compiled, linked and executed on several types of computers (e.g. CDC, CRAY, IBM, DEC, HP, SUN, VAX).

An additional *FORTRAN*code, **zpop**, allows the post-processing and graphic treatment of **zgoubi** output files. **zpop** is routinely used on DEC, HP and SUN stations.

1 GETTING TO RUN zgoubi AND zpop

1.1 Making the executable files zgoubi and zpop

1.1.1 The transportable package zgoubi

Compile and link the FORTRAN source file zgoubi.f, to create the executable zgoubi.

zgoubi.f is written in standard *FORTRAN*, therefore it is not necessary to link with any Library, except maybe a local math. lib.

1.1.2 The post-processor and graphic interface package zpop

Compile the FORTRAN source files zpop*.f.

Link **zpop** with the graphic library, libminigraf.a [35]. This will create the executable **zpop**, that can run on xterm type window.

1.2 Running zgoubi

The principles are the following:

- fill zgoubi.dat with the input data that describe the problem (see examples, Part C).
- Run zgoubi.
- Results of the execution will be printed into zgoubi.res and, upon options appearing in zgoubi.dat, into several other outputs files (see section 2 below).

1.3 Running zpop

- Run **zpop** on an xterm window. This will open a graphic window.
- Select options displayed on the menu.
- To access the graphic sub-menu, select option 7.
- An on-line Help provides all necessary informations on the post-processors (Fourier transform, elliptical fit, synchrotron radiation, field map contours, etc.).

2 STORAGE FILES

When explicitly requested by means of the adequate keywords, options, or dedicated *LABEL*'s, extra storage files are opened by **zgoubi** (*FORTRAN "OPEN*" statement) and filled.

Their content can be afterwards post-processed using the interactive program **zpop** and its dedicated graphic and analysis procedures.

282 2 STORAGE FILES

The **zgoubi** procedures that create and fill these extra output files are the following (refer to Part A and Part B of the guide):

- Keywords FAISCNL, FAISTORE: fill a '.fai' type file (normally named [b_]zgoubi.fai) with particle coordinates and other informations.
- Keywords *SPNPRNL*, *SPNSTORE*: fill a '.spn' type file (normally named [b_]zgoubi.spn) with spin coordinates and other informations.
- Option IC = 2, with field map keywords (e.g. *CARTEMES*, *TOSCA*): fill zgoubi.map with 2-D field map.
- Option IL = 2, with magnetic and electric element keywords: fill zgoubi.plt with the particle coordinates, and experienced field, step after step, all along the optical element.
- Using the keyword MARKER with '.plt' as a second LABEL will cause storage of current coordinates into zgoubi.plt.

Typical examples of graphics that one can expect from the post-processing of these files by **zpop** are the following (see examples, Part C):

• '.fai' type files

Phase-space plots (transverse and longitudinal), aberration curves, at the position where *FAISCNL* appears in the optical structure. Histograms of coordinates. Fourier analysis (e.g. tune numbers in multiturn tracking), calculation of Twiss parameters from phase-space ellipse matching.

• zgoubi.map

Isomagnetic field lines of 2-D map. Superimposing trajectories read from zgoubi.plt is possible.

• zgoubi.plt

Trajectories inside magnets and other lenses (these can be superimposed over field lines obtained from zgoubi.map). Fields experienced by the particles at the traversal of optical elements. Synchrotron radiation.

• zgoubi.spn

Spin coordinates and histograms, at the position where *SPNPRNL* appears in the structure. Resonance crossing when performing multiturn tracking.

284 REFERENCES

References

- [1] F. Méot et S. Valéro, Manuel d'utilisation de Zgoubi, report IRF/LNS/88-13, CEA Saclay, 1988.
- [2] F. Méot and S. Valéro, Zgoubi users' guide, Int. Rep. CEA/DSM/LNS/GT/90-05, CEA Saclay (1990) & TRIUMF report TRI/CD/90-02 (1990).
- [3] F. Méot and S. Valéro, Zgoubi users' guide Version 3, Int. Rep. DSM/LNS/GT/93-12, CEA Saclay (1993).
- [4] F. Méot and S. Valero, *Zgoubi users' guide Version 4*, FNAL Tech. Rep. FERMILAB-TM-2010 (Aug. 1997), & Int. Rep. CEA DSM DAPNIA/SEA-97-13, Saclay (Oct. 1997).
- [5] http://zgoubi.sourceforge.net/
- [6] F. Méot, The electrification of Zgoubi, Saturne report DSM/LNS/GT/93-09, CEA Saclay (1993); F. Méot, Generalization of the Zgoubi method for ray-tracing to include electric fields, NIM A 340 (1994) 594-604.
- [7] D. Carvounas, Suivi numérique de particules chargées dans un solénoïde, report de stage, CEA/LNS/GT-1991.
- [8] F. Méot, Raytracing in 3-D field maps with Zgoubi, report DSM/LNS/GT/90-01, CEA Saclay, 1990.
- [9] G. Leleux, Compléments sur la physique des accélérateurs, cours de DEA Univ. Paris-VI, report IRF/LNS/86-101, CEA Saclay, March 1986.
- [10] F. Méot, A numerical method for combined spin tracking and raytracing of charged particles, NIM **A313** (1992) 492, and proc. EPAC (1992) p.747.
- [11] V. Bargmann, L. Michel, V.L. Telegdi, *Precession of the polarization of particles moving in a homogeneous electromagnetic field*, Phys. Rev. Lett. 2 (1959) 435.
- [12] F. Méot and J. Payet, Numerical tools for the simulation of synchrotron radiation loss and induced dynamical effects in high energy transport lines, Report DSM/DAPNIA/SEA-00-01, CEA Saclay (2000).
- [13] F. Méot, Synchrotron radiation interferences at the LEP miniwiggler, Report CERN SL/94-22 (AP), 1994.
- [14] L. Ponce, R. Jung, F. Méot, LHC proton beam diagnostics using synchrotron radiation, Yellow Report CERN-2004-007.
- [15] J.D. Jackson, Classical electrodynamics, 2nd Ed., J. Wiley and Sons, New York, 1975.
- [16] F. Méot, A theory of low frequency far-field synchrotron radiation, Particle Accelerators Vol 62, pp. 215-239 (1999).
- [17] B. Mayer, personal communication, CEA Saclay, 1990.
- [18] Installed by J. S. Berg, BNL (2007). *Cf. Detection and remediation of stagnation in the Nelder-Mead algorithm using a sufficient decrease condition*, C. T. Kelley, Siam J. Optim., Vol. 10, No. 1, pp. 43-55.
- [19] L. Farvacque et al., Beta user's guide, Note ESRF-COMP-87-01, 1987; J. Payet, IRF/LNS, CEA Saclay, private communication; see also J.M. Lagniel, Recherche d'un optimum, Note IRF/LNS/SM 87/48, CEA Saclay 1987.
- [20] F. Méot and N. Willis, Raytrace computation with Monte Carlo simulation of particle decay, internal report CEA/LNS/88-18 CEA Saclay, 1988.
- [21] H.A. Enge, *Deflecting magnets*, in **Focusing of Charged Particles**, ed. A. Septier, **Vol. II**, pp 203-264, Academic Press Inc., 1967.
- [22] P. Birien et S. Valéro, *Projet de spectromètre magnétique à haute résolution pour ions lourds*, **Section IV** p.62, Note CEA-N-2215, CEA Saclay, mai 1981.
- [23] V. M. Kel'man and S. Ya. Yavor, Achromatic quadrupole electron lenses, Soviet Physics Technical Physics, vol. 6, No 12, June 1962;
 S. Ya. Yavor et als., Achromatic quadrupole lenses, NIM 26 (1964) 13-17.
- [24] A. Septier et J. van Acker, Les lentilles quadrupolaires électriques, NIM 13 (1961) 335-355; Y. Fujita and H. Matsuda, Third order transfer matrices for an electrostatic quadrupole lens, NIM 123 (1975) 495-504.
- [25] A. Septier, Cours du DEA de physique des particules, optique corpusculaire, Université d'Orsay, 1966-67, pp. 38-39.

REFERENCES 285

[26] S. P. Karetskaya et als., Mirror-bank energy analyzers, in Advances in electronics and electron physics, Vol. 89, Acad. Press (1994) 391-491.

- [27] F. Lemuet, F. Méot, Developements in the ray-tracing code Zgoubi for 6-D multiturn tracking in FFAG rings, NIM A 547 (2005) 638-651.
- [28] F. Méot, 6-D beam dynamics simulations in FFAGs using the ray-tracing code Zgoubi, ICFA Beam Dyn.Newslett.43:44-50 (2007).
- [29] J. Fourrier, F. Martinache, F. Méot, J. Pasternak, Spiral FFAG lattice design tools, application to 6-D tracking in a proton-therapy class lattice, NIM A 589 (2008).
- [30] P. Akishin, JINR, Dubna, 1992.
- [31] M.W. Garrett, Calculation of fields [...] by elliptic integrals, J. Appl. Phys., 34, 9, sept. 1963.
- [32] P.F. Byrd and M.D. Friedman, *Handbook of elliptic integrals for engineers and scientists*, pp. 282-283, Springer-Verlag, Berlin, 1954.
- [33] A. Tkatchenko, Computer program UNIPOT, SATURNE, CEA Saclay, 1982.
- [34] J.L. Chuma, PLOTDATA, TRIUMF Design Note TRI-CO-87-03a.
- [35] J.L. Hamel, mini graphic library LIBMINIGRAF, CEA-DSM, Saclay, 1996.
- [36] E. Grorud, J.L. Laclare, G. Leleux, *Résonances de dépolarisation dans SATURNE 2*, Int. report GOC-GERMA 75-48/TP-28, CEA Saclay (1975), and, Home Computer Codes POLAR and POPOL, IRF/LNS/GT, CEA Saclay (1975).
- [37] M. Froissart et R. Stora, Dépolarisation d'un faisceau de protons polarisés dans un synchrotron, NIM 7 (1960) 297-305.
- [38] J. Thirion et P. Birien, Le spectromètre II, Internal Report DPh-N/ME, CEA Saclay, 23 Déc. 1975; H. Catz, Le spectromètre SPES II, Internal Report DPh-N/ME, CEA Saclay, 1980.
- [39] P. Pile, I-H. Chiang, K. K. Li, C. J. Kost, J. Doornbos, F. Méot et als., A two-stage separated 800-MeV/c Kaon beamline, TRIUMF and BNL Preprint (1997).
- [40] F. Méot, *The raytracing code Zgoubi*, CERN SL/94-82 (AP) (1994), 3rd Intern. Workshop on Optimization and Inverse Problems in Electromagnetism, CERN, Geneva, Switzerland, 19-21 Sept. 1994.
- [41] W. H. Press et als., Numerical recipes, Cambridge Univ. Press (1987).
- [42] V. O. Kostroun, Simple numerical evaluation of modified Bessel functions and integrals [...], NIM 172 (1980) 371-374.

Index

acceleration, 69, 71, 89, 177, 234, 236 AGSMM, **77**, 153, **165** AGSQUAD, **78**, **166** AIMANT, 58, **79**, 99, 167 *ALE*, 153 AUTOREF, **84**, 171

backward ray-tracing, 152 BEND, 25, **85**, 172 BINARY, **55**, 173 BORO, **44**, **47**, **52**, 89, 180, **215**, **219**, **221** BREVOL, 24, **86**, 174

CARTEMES, 24, 86, **87**, 112, 113, 118, 119, 124, 125, 133, 152, 175, 198, 246, 282
CAVITE, 71, **89**, 90, 153, 177
CHAMBR, 69, 91, **91**, 178, 235
CHANGREF, 65, 84, 91, **92**, 94, **179**CHANGREF "New Style", 92, 179
CHANGREF "Old Style", 92, 179
checking field, 152
checking trajectories, 152
chromaticity, **145**, 146, 151

COLLIMA, 69, **95**, 181, 235 constraint (FIT, FIT2), 57, **60**, **204**

DECAPOLE, 67, **96**, 182, 233 DIPOLE, 58, 91, **97**, 101, 153, 183, 259 DIPOLE-M, 79, 81, **99**, 125, 156, 185, 228 DIPOLES, **101**, 187

DODECAPO, 26, 67, 104, 189, 233

DRIFT, 105, 190

CIBLE, 64, 94, 180

EBMULT, 25-27, 58, 67, 106, 191

EL2TUB, **107**, 193 ELMIR, **108**, 194 ELMIRC, **109**, 195

ELMULT, 27, 58, 67, 106, **110**, 196

ELREVOL, 24, **112**, 152, 197

EMMA, 113, 198 END, 54, **56**, 203 ESL, 65, **105**, 190

FAISCEAU, **141**, 199
FAISCNL, **141**, 153, 199, 282
FAISTORE, **141**, 153, 199, 282
FFAG magnet, radial, 101, **114**, 200
FFAG magnet, spiral, **116**, 202
FIN, **56**, 203
FIT, 50, 54, 57, **57**, 58, 60, 204, 234
FIT2, **57**, 204
FOCALE, **143**, 206

FOCALEZ, **143**, 206 fringe fields overlapping, 101

GASCAT, 63, 207

HISTO, 45, 65, 95, 144, **144**, 156, 208, 235

IC, 152
ID, **87**, **175**IDMAX, 48, **48**IEX, 30, **48**, 49, 76, 87, 91, 95, 141, 149, **156**IL, 152

IMAGE, 84, 143, **143**, 209 IMAGES, 47, 48, 143, **143**, 209

IMAGESZ, **143**, 209 IMAGEZ, **143**, 209

MAX, **44**, **48**, 49, 52, 65, 69, 73, 89, 90, 141, 149, 153, 156

INTEG, 235

integration step size, 156 coded, 156

negative, 65, 152 IORDRE, 25, 28, 67, 83, 87, 100, 113, 119, 133

KPOS. 153

LABEL, 71, 120, 141, 147, 149, **153**, 199, 212, 226, 241, 282

MAP2D, 25, **118**, 133, **210**MAP2D-E, 25, **119**, **211**maps, summing, 174
MARKER, **120**, 212, 282
MATRIX, 50, 57, 62, 84, 121, **145**, 146, 171, **213**, 220
MCDESINT, 45, 47, 54, 64, 65, 69, 144, 214, 235, 259
MCOBJET, 44, 45, 62, 69, 70, 156, 215, 234
Misalignement, **153**Monte Carlo, 44, 52, 64, 69, 156, 215, 221
multiparticle, 69, 156, 234
MULTIPOL, 25, 26, 58, 67, 71, 106, 110, 122, 218, 233
multiturn, 43, 71, 89, 153, 234, 259, 272, 282, 287
multiturn tracking, 69

negative charge, 44, 47, 156, 215, 219 negative momentum, 44, 47, 156, 215, 219 negative rigidity, 156 NPASS, 69, 89, 141, 144, 149, 153, 156, 199, 234, 241

OBJET, 47, 60, 62, 64, 70, 84, 89, 91, 143, 151, 153, 156, 219
OBJETA, 52, 69, 221
OCTUPOLE, 67, 123, 222, 233
OPTICS, **66**, 223

288 INDEX

ORDRE, 67, 224 outpoi.lis, 124

PARTICUL, 53, 64, 68, 73, 75, 76, 95, 106–110, 112, 130, XPAS, negative, 152 136, 138, 214, 225

PICKUPS, **147**, 153, 226 PLOTDATA, 148, 227 POISSON, 124, 228

POLARMES, 125, 156, 229

Positioning, **153** PS170, 126, 230

QUADISEX, 24, 25, 127, 231 QUADRUPO, 25, 26, 58, 59, 67, 91, 96, 110, 122, 123, 128, 131, 153, 232, 233

REBELOTE, 43, 44, 49, 54, 57, 69, 71, 89, 141, 144, 147, 149, 153, 156, 199, 219, 234, 241, 259

RESET, 70, 235

SCALING, 59, 71, 89, 90, 153, 177, 235, 236

SEPARA, 130, 237

SEXQUAD, 24, 127, 238

SEXTUPOL, 25, 67, 131, 233, 239

SOLENOID, 132, 240

spin tracking, 35, 69, 73, 92, 134, 140, 144, 149, 156, 208, 234, 242, 259

SPNPRNL, 149, 241, 282

SPNPRT, 149, 241

SPNSTORE, 149, 241, 282

SPNTRK, 54, 69, 73, 156, 235, 242

SRLOSS, 75, 243, 244

SRPRNT, 150, 244

stopped particles, 69, 70, 91, 95, 141, 156, 178, 181

storage files, 281

synchrotron motion, 69, 71, 89, 177, 234, 236

synchrotron radiation, 37, 282

synchrotron radiation loss, 75, 150, 243

synchrotron radiation spectra, 76, 245

SYNRAD, 76, 245

TARGET, 94, 180

TOSCA, 24, 55, 59, 67, 133, 153, 246, 259, 282

TRANSMAT, 121, 248

TRAROT, 134, 249

TWISS, 50, 151, 250

UNDULATOR, 135, 251

UNIPOT, 136, 252

variable (FIT, FIT2), 57, 57, 204

VENUS, 24, 25, 137, 253

WIENFILT, 25, 138, 254

XCE, 153

XPAS

negative, 65 XPAS, coded, 156 XPAS, negative, 152

YCE, 153 YMY, 139, 255

zgoubi, 281

zgoubi.dat, 142, 149, 281

zgoubi.f, 281

zgoubi.fai, 141, 149, 199, 220, 282

zgoubi.map, 152, 282

zgoubi.plt, 120, 152, 153, 212, 282

zgoubi.res, 152, 281

zgoubi.spn, 149, 241, 282

zgoubi.sre, 76

zpop, 39, 141, 149, 152, 281