# **Introduction to datreat**

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## **Datreat Purpose and Philosophy**

Datreat is a command-based multi purpose system that allows to deal with collections of (x,y)-data (data-records) each of which has an individual block of parameters. Data may stem from any field ranging e.g. from SANS diffraction results, NSE-S(Q,t) curves, magnetic field values along a path, BSS-spectra etc.. Datreat allows to plot, manipulate and fit these data-records. Theoretical models may be assembled from existing "theories" or new theories may be formulated (which, however, requires some Fortran programming). Simultaneous fitting of collections of data-records that are distinguished by one or several parameters is supported as well as coupling of fit parameters.

Some automation of data-evaluation is possible with the help of "makros". These are files that contain a collection of genuine commands including conditional and unconditional jumps that allow also the construction of loops.

Interaction with the program is effected by commands. Each command line consists of a *command-name* and a number of *items* that are *separated by blanks*.

**Command line items** may be **names**, **numbers** or **expressions** that evaluate to a number. In general **expressions** may be used at any place in **command lines** and **input files** where a **number** would be appropriate.

The system classifies all items that start with one of the following characters as *expressions*:

(+-.0123456789

any item that would start with one of these characters and would not lead to a valid *expression* leads to an error.

Despite some preliminary mechanisms to specify a *path* the recommended use of Datreat is to call the program from the current data-subdirectory and keep the specific makros and other auxiliary files there too.

RB: But it is usefull to discriminate between raw data and evaluated data plots ... See **path** at the end to set datapath and savepath

To exit the program:  $\rightarrow$  q

**Hint1:** Running datreat from an Emacs-shell offers a number of advantages, in particular the availability of command-line history (use ALT-p, ALT-n) to retrieve previous/next command inputs. In addition all editor functions are available on the whole output produced by the current session.

A simple history (20 commands) exists. Accessible by "\_xxx" the last command starting with xxx ist executed again.

**Hint2**: Command hierarchy: Commands are first used as datreat commands than as makro name. After both failed (eg no makro with name pwd) command is given to a new shell instance and executed. This means you can use any shell command as you are used to it, as long as no makro has the same name. (see history for a nice example how to use this feature)

Here we will write always MAKRO with k because the makros are detected by the German "makro" in the first line.

**Hint3:** If a makro "**initdatr**" exists within datreat/makro or in your local path this will be executed right after start of datreat. Its a normal makro and you can define your own path, standard data treatment plotting ....... whatever you can imagine.

#### **Data Structure**

#### **Files**

**Data files:** contain a number of data records each of which consists of a header, a parameter section and a list of x,y values with an optional third column of y-errors.

**Theory definition file**: always has the value **lastth.** To save previous versions copy it onto another name.

**Makro files**: contain a collection of command lines their names follow the same rules as the data files. To qualify as a makro the name must also be different from any genuine datreat-command and the first line must start with the keyword: **makro**.

Shell commands are allowed. Lines are limited to 1024 characters.

#### **Internal Representation**

Data and theory definitions are stored internally (when loaded) in common blocks with fixed (may be changed by recompiling the program with other limits) maximum capacity of 40. The data on these structures are largely accessible by predefined system functions.

# **Datreat Commands**

# Input/Output

# Input of Data / Display of List of Loaded Data

•	Take 7 Display of List of Loaded Data
in /input	filename
	reads data (x, y, y-error) from <filename>.</filename>
	<pre><filename> may not contain one of the fllowing characters: '+-0123456789.('.</filename></pre>
	<pre><filename> can be a complete path prepended by variable <datapath>. If</datapath></filename></pre>
	<pre><filename> starts with ./ or / (./path/name or /path/name ) <datapath> is NOT</datapath></filename></pre>
	prepended.
	<b>Hint:</b> if data stem form a different computer system, make sure that they do not contain tab's and extra CR-characters etc., these cause occasionally observed errors during reading.
	The last read data record is automatically selected. All previous selections are removed.
	NEW Data format
	Simpler format rules: parameters and comment ahead of Data
	parameter: any line starting with a name followed by numbers
	(parameter name; first number is set as value; others ignored)
	data: line with minimum 2 values (x,y); 3 <sup>rd</sup> is used as error; others are ignored everything not data or parameter; first comment line is stored;
	comment: everything not data or parameter; first comment line is stored; other comment lines are ignored
	ignored lines: start with "#" as first character; "values" and "parameter" also ignored if
	# is 1. char.
	new set:initiated by non Data after Datablock (parameters, comment or empty line) or
	"#nxt"
	OLD Data format for input files: q= 0.161 dpeo25pmma75 no argon :comment
	q= 0.161 dpeo25pmmâ75 no argon j04n01 ncounts vs omeg/Ghz 10308 :comment :name xname vs yname <numor></numor>
	parameter :keyword=parameter normal
	lambda
	temp 0.39580000E+03
	d_energ 0.17000000E+02
	104n019   ncounts   vs   omeg/Ghz   10308   name xname vs yname <numor>   parameter   normal</numor>
	-0.25345777E+02  0.64695009E-02  0.17290469E-02 : <x> <y> <y-error></y-error></y></x>
	-0.25129146E+02 0.63254744E-02 0.19072023E-02
	0.24479254E+02 0.88531187E-02 0.18874913E-02 0.24695885E+02 0.78431373E-02 0.16721625E-02
	: blank line #nxt :keyword=#nxt
	g= 0.002 dpeo25pmma75 no argon j04n019 ncounts vs omeg/Ghz 10308
	parameter 0.10000000E+01
	V_5V 0.50000000E+01
	values
	-0.26212299E+02
	-0.25779038E+02 0.16228497E-02 0.72576046E-03
	0.23829363E+02 0.74321813E-03 0.52553458E-03 0.24045993E+02 0.79744817E-03 0.56388101E-03
	0.24262624E+02
	#eod :keyword=#eod

# **Output of Data**

•	
save	[ <isel>] to filename</isel>
	saves data from selected respectively <isel> internal record to <filename> . If <filename> starts not with "." or "/" savepath is prepended. Current <b>restriction</b>: <filename> may not begin with one of the fllowing characters: '+-0123456789('.  <b>Hint:</b> behind the <b>data</b> the saved file will contain the theory setting definition which was active at the time</filename></filename></filename></isel>
	when the save command was issued. Note that this will only be useful if save is performed before a different theory setting which does not represent the situation of the saved data is prepared!
	Hint2: To restore the theory including parameters use :
	> csplit <saved_filename> %theory% -f <theory_filename></theory_filename></saved_filename>
	resulting in a file < theory_filename00 > which you can cp to lastth, activate it by > acl and recalculate theory by > thc
msave	filename
	saves all selected data records to <filename>. If <filename> starts not with "." or "/" savepath is prepended. Current <b>restriction</b>: <filename> may not begin with one of the fllowing characters: '+- 0123456789('.  <b>Hint:</b> behind the <b>data</b> the saved file will contain the theory setting definition which was active at the time</filename></filename></filename>
	when the <b>save</b> command was issued. Note that this will only be useful if <b>save</b> is performed before a different theory setting which does not represent the situation of the saved data is prepared!
clip	[[from <n1>] [to <n2>]] [last <n>] [errmax <val>] [rel]</val></n></n2></n1>
•	removes points from a loaded data record. Either by specifying the range of data point numbers <n1><n2>, resp. last <n> or by setting a limit to the maximum error. rel modifies the error condition to hold for relative errors.  Note: the action only affects the loaded data record and not the data in the input file.</n></n2></n1>
edit	<n> [sc <numor>]</numor></n>
	allows to edit data of data-record <n> or data-record with <numor>. The contents is written to a file <i>datbuf</i> which is laded to an editor and reread after the editor is left.  <b>Note:</b> the action only affects the loaded data record and not the data in the input file</numor></n>
putpar	f_parname <value></value>
<u> </u>	adds a parameter or changes a parameter value of the selected data-record.
dir	[clength <displayed-comment-length>] [f_parnam1] [f_parnam2]</displayed-comment-length>
	displays a list of the loaded data records. Selected records are marked by a '!'.  The clength parameter allows to control the number of characters that are dsplayed from a comment.
	If f_parname is given only data records that contain a parameter with that name are listed and the
	parameter value is displayed. It is possible to specify more than one f_parname.
Z	
	zeroes data-record counter,. logically clears data-record list.
	  2-11

## **Associated variables:**

nbuf	evaluates to the number of loaded items, i.e. points to the end of the dir-list.
num(j)	evaluates to the "numor" of the j-th loaded data record.
nv(j)	evaluates to the length (number of points) of the j-th loaded data record.
xv(i,j)	evaluates to the x-value of the i-th point of the j-th loaded data record.
yv(i,j)	evaluates to the y-value of the i-th point of the j-th loaded data record.
ye(i,j)	evaluates to the y-value error of the i-th point of the j-th loaded data record.
sumx(i1,i2,j)	evaluates to the sum of x-values from the i1-th point to the i2-th point of the j-th loaded data record.
sumy(i1,i2,j)	evaluates to the sum of y-values from the i1-th point to the i2-th point of the j-th loaded data record.

## **Selecting dataset**

In general actions triggered by datreat commands are performed on all data records that are selected.

In special cases a certain sequence of selections assigns data records to different roles in command performance (see command descriptions).

sel	<n1> <n2><nx> [add]</nx></n2></n1>
	selects exactly those data records with (internal) numbers <n1>, <n2><nx>.  The option add causes that the previous selctions are kept and the new ones are added to the list.</nx></n2></n1>
sel	<n1> -<nx> [add]</nx></n1>
	selects all data records with (internal) numbers <n1><nx></nx></n1>
sel	all f_parnam <value> band <range> [add]</range></value>
	selects all data records which have a parameter with name <b>f_parnam</b> which has a value within the range: <pre><value>± <range>.</range></value></pre>
sel	next f_parnam <value> band <range> [add]</range></value>
	selects next data record (after actually selected) which hase a parameter with name <b>f_parnam</b> which has a value within the range: <value>± <range>.</range></value>
sel	[fits] [fit+]
	selects fitted data, or fits associated to selected data (fit+).
sel	
	removes all selections.
dsl	
	displays a list of the current selections. (see also dir)
purge	<n1> <n2><nx> [all]</nx></n2></n1>
	removes data records <n1><nx> or all data records from the dir-list.  <b>Note</b>: all selection will be removed!</nx></n1>

#### **Associated variables:**

sel	evaluates to the last selected data record number.
nsel	evaluates to the number of selected data records (length of selected list).
isel(i)	evaluates to the data record number of the i-th selected list item.

# Plotting

We use xmgrace as plotting interface. The obsolete gr is working but not supported.

<pre></pre> <pre></pre> <pre></pre> <pre></pre>			
gp/gplot	<k0><com><gr><jp><sv><so><cleanio></cleanio></so></sv></jp></gr></com></k0>		
	gplot,gp to plot selected files in connected xmgrace		
gp plots into same grace until connection is closed			
	without open connection a new grace process is created		
	common parameter were extracted and a legend is shown build up from uncommon params'		
	in grace: C-t write text		
	C-d delete text;		
	C-M moves objekt		
	C-L moves legend		
	all parameters were transferred to set comments in grace to identify datasets		
	save your plots in the file menu as grace file (?.agr or ?.xgr ) if you want to modify later		
	Print it in menu File/Print Setup (Device setup opens)		
	choose format and size ( press apply )		
	(for .eps choose Tight box in device options)		
	printed in menu point File/Print		
Options:	stored		
help, h, ?	for this helptext		
switches on/off'			
fits, fi	hide fits (but transferred )		
fitpara, fp	write last fit parameters		
error, er	no error bars (but transferred)		
legend, le	hide legend; if hidden and parameter names were found they build up legend;		
	eg "gplot nl q" hides parameters in legend exept q values		
	or "gp le" set to no legend; afterwards "gp q T" creates legend im q and T are given		
noplot, np	dont plot , just set switches		
jointpar,jp	write joint parameters'		
close, cl	close connection to grace after plotting; with "save name cl" in a script a new plot is saved		
	and then closed		
clearplo,clear			
autoscal, as	autoscale switched off/on, default on		

normal:		
closenow, cn	closes connection without plotting of new data	
kill, ki		
command, com	rest of input line as direct command to grace	
Command, com	see xmgrace help or grace files as example'	
graph, gr	plot data in graph i, a new graph inside grace is build,' number i is stored for next dataset	
grapii, gr	arrange them in EDIT/ARRANGE Graphs'	
start	The rest of the line will be sent directly to grace'; used to customise the startup behaviour of	
Start	xmgrace. Its a semicolon separated list of xmgrace commands e.g. (see saved *.agr file )	
	##> gp start title "NSE data";subtitle "\xt\N is a fouriertime";xaxis label "\xt\N / ns"	
save, sv	save graceplot with following name (.agr is added)'; if name is "on" updated plots are saved	
Save, sv	always to "lastgrace.agr"; without name its saved to "lastgrace.agr" and switches save off.	
setoffset, so	offset for numbering new data, default 0	
Cotonioot, co	data with smaller numbers are preserved'	
cleanio	deletes iofile.tmpg?s? if exists'	
o o o o o o o o o o o o o o o o o o o	adiotec following for it extends	
	hints to grace	
gp com <grcom></grcom>	grcom is a direct command to grace	
	some usefull examples:	
	page size 400, 300 resizes page to 400, 300 pixels (try what you like most)	
	xaxes scale Normal scale set to lin	
	title "this is the main title" obvious	
	yaxis label "S(q,\xt) / S(q,0)" obvious, see xmgrace help for text	
	yaxes scale Logarithmic scale set to log	
	redraw plot needed sometimes do see the effect of previous	
	commands	
	the full command in datreat is "gp com redraw"	
to install the .grace directory with default configuration files of xmgrace config.grace to your home directory as .grace	to install the .grace directory with default configuration files of xmgrace copy the directory	
	config.grace to your home directory as .grace	
	To change the default startup layout of your gracewindow open	
	.grace/templates/Default.agr	
	and save it again after you changed whatever you wanted.	
	To change the default behaviour look in grace.user to use something like this input filter for .dat files.	
	accept only lines starting with +,- or a number for .dat files	
gp start xxxx	DEFINE IFILTER "egrep '^ *([+,-,0-9,\] \$)' %s" PATTERN "*.dat"  xxx is the rest of the line after start. It is a semicolon separated list of	
gp start xxxx	xmgrace commands like above and will be used for all of your following	
	new xmgrace windows in this session.	
	To have a permanent behaviouzr use initdatr initialization file in your makro	
	path.	
	puu.	

#### Plotting (OLD)

obsolete but working new => gplot

plot	[xmin <xmin>] [xmax <xmax>] [ymin <ymin>] [ymax <ymax>]</ymax></ymin></xmax></xmin>
	plots selected data records (including fit/thc results if present) using the (optionally) specified range.
	Default is symbols for data and lines for fit/thc results.
plot	[option1][ option_n]

	OPTIONS:	
	log_x	: log scale on x-axis (range given by xmin, xmax must be > 0)
	lin_x	: resets the default linear x-scaling
	log_y	: log scale on y-axis (range given by ymin, ymax must be > 0)
	lin_y	: resets the default linear y-scaling
		sn>] : assigns symbols to plotted curves
		cn>]: assigns colors to plotted curves
	errplo	: plot with error bars
	noerrplo	: plot without error bars (default)
	parplo	: plot with written parameters at the side (default)
	noparplo [f_pa	ırnam1] [[f_parnam_n]]
		: plot without parameters at the side except those listed: [f_parnam1
	framx <fx></fx>	
	framy <fy></fy>	
	frlux <ux></ux>	
	frluy <uz></uz>	: changes size and shape of plot area (axes) on the screen <fx> <fy> dimension</fy></fx>
		: of plot window, <ux> <uy> location of lower left (values in cm relative A4)</uy></ux>
	legsize <ls></ls>	
	legx <lx></lx>	
	legy <ly></ly>	: text size and relative location of block of written parameters.
	txsize <ts></ts>	
	font <n></n>	: text size <ts> (cm) and font (by number <n>)</n></ts>
	text	
	notext	: allow or suppress text plotting completely.
	o <option></option>	: GR-software option specifier string (graxs-subprogram). the format is e.g.
	0	: ox x=1 {shows x-axes below your graph}{2 puts the axes above} and {3 on
	both sides1/-2/-3 s	shows a grid}
p0		
	sets/changes options	only without producing a plot.
tit	<title-string></title-string>	
	sets a title string tha	t occurs as title on the plot.
rename		p>] [yaxis <new-string>] [name <new-id>]</new-id></new-string>
	sets new names (lab	els) for xaxis, yaxis or ident of the selected data record.

The plot command is based on the call of the FZJ/ZAM GR-software package. It asks for an output device number to poduce the plot. For X-terminals (and that is the standard now for all workplaces) the number to be given is 211.

#### How to get a printed output?

If the plot shall be printed or stored as eps-file, the output device number is **62.** This causes the plot command to write a file named *gliXX.eps* with *XX* the number of the plot invocation during the current session. This file may be copied to another name and/or printed using the UNIX **lpr** command.

# Fitting

# **Setting-up Theories and Parameters**

	Incomes and Farameters	
theos	<string></string>	
	lists names of all available theory types. <string> If string is given only theos starting with string are listed.</string>	
	The always available pseudo-theory <b>eval</b> is not quoted. It is intended to allow for the ad-hoc fitting of	
	simple expressions that have to be enterd using the <b>yfitformel</b> command.	
ac	theoryname [multiply] [range <datparname> min <p1> max <p2> ]</p2></p1></datparname>	
	activates one instance of theory theoryname which is appended to the current list. <b>Options:</b>	
	<b>multiply</b> if given the value from this theory instance is multiplied to the result of the previous list,	
	range if given the name of the data parameter datparname and min and max values are to be	
	specified. The theory will only be evaluated if the value of <datparname> as parameter of a dataset under</datparname>	
	consideration is present and its value is in the interval between <p1> and <p2>. Hint: the range option is intended to be used for fits of datasets of a series with different values of an</p2></p1>	
	experimental parameter (like an increasing Temperature stored as <datparname>). A theory instance may</datparname>	
	be activated for each dataset and assigned to it by appropriate specification of the range interval. Those theory parameters that shall be common to the whole data sets may be coupled (see below). Any	
	individual scalings etc. may simultaneously be effected by keeping the corresponding parameters as	
	individual fit parameters,	
acl	activates last theory stored in file lastth	
	<b>HINT:</b> To reuse a complete theory including parameters (how to restore a theory setting	
	from saved data see <b>save</b> ) copy the lastth to a different file name and copy it back to file name lastth if needed like this:	
	>ac strexpo : activates theory strexpo	
	>emacs lastth : opens emacs to modify parameters close it.or use <b>chgthpar</b>	
	>cp lastth lasth_strexpo : copy to new filename (backup of your settings)	
	to reuse it:	
	>cp lasth_strexpo lastth >acl	
dac		
	desactivates all theory instances. After <b>dac</b> the theory definition is completely cleared.	
al		
	lists current theory definition .on console and into file <i>lastth</i> .	
chgthpar	theoryname [ <instance>] t_parname par <p1> scale <s1></s1></p1></instance>	
21.94.194.	sets parameter value <p1> and scale <s1> for parameter _t_parname of the <instance>-th instance of</instance></s1></p1>	
	theory theoryname. If scale <s1>=0 the parameter is fixed during fits, otherwise <s1> should set to a value of the same order of magnitude as the expected parameter value.</s1></s1>	
label		
IdDel	theoryname [ <instance>] t_parname label assigns label label to parameter _t_parname of the <instance>-th instance of theory theoryname. The</instance></instance>	
	label may contain of a maximum of 4 characters. The labels are used to establish parameter couplings	
	between different theory instances.	
couple	theoryname [ <instance>] t_parname label <factor></factor></instance>	
	installs a coupling of a labelled parameter onto a parameter as quoted in this command, The coupling factor is <a href="factor">factor</a> . All changes with respect to the start values of the labelled parameter are multiplied by	
	facor <factor> and added to the parameter quoted in this command. Coupling to several lablelled</factor>	
	parameters is possible. Parameters with couplings will not be fitted, the corresponding scale is automatically set to zero.	
yfitform;	$(xx^*p(1)+p(2)^*p(3))$	
	definition of the special theory <b>eval</b> . xx means the actual x-value, p(1)p(9) are parameters of the	
	theory. Note the necessary ';' between yfitformel and the expression in brackets. f_parnams may be	
	used.	

cth	obsolete but useful; change theory parameter
	you can use a command like "emacs lastth" or kwrite lastth"
	its the same without extra makro (if you append "&" datreat doesn't wait until the
	process finishes. But remember to save your changes before you continue and to
	revert to eventual changes if you proceed)
	Makro: allows editing of the complete theory definition list.
	Data format of theory list:
	theory zimm 1. instance of theory zimm
	a1 intensit 0.9765E+00 0.1E+00 0.39E-02
	eta_solv 0.8220E-03 0.1E-02 0.18E-04
	epsilon 0.1000E-06 0.0E+00 0.00E+00
	temp 0.3000E+03 0.0E+00 0.00E+00
	com_diff 0.0000E+00 0.0E+00 0.00E+00
	theory zimm  2. instance of theory zimm
	intensit 0.9765E+00 0.0E+00 0.39E-02 <mark>a1 -1</mark>
	eta_solv
	epsilon 0.1000E-06 0.0E+00 0.00E+00
	temp 0.3000E+03 0.0E+00 0.00E+00 com diff 0.0000E+00 0.0E+00 0.00E+00
	theory rouse range q min 0.1 max 0.3  1. instance of theory rouse intensit 0.0000E+00 0.0E+00
	xi_frict 0.1000E+09 0.0E+00 0.00E+00
	b_segmnt 0.5000E+01 0.0E+00 0.00E+00
	epsilon 0.0000E+00 0.0E+00
	temp 0.3000E+03 0.0E+00 0.00E+00
	com_diff 0.0000E+00 0.0E+00 0.00E+00
	q_width 0.0000E+00 0.0E+00 0.00E+00
	end
	label par scale couplings
	All numeric entries may also be expresssions which will be converted to their numerical values upon
	reading. Remember: expressions must be given in brackets () or begin by +.

# Theory Computations and Fittings

thc	[n <num> x1 <x1> x2 <x1>] [convolute xc1 <xc1> xc2 <xc2>] computes theory values for the selected data using the activated theory list. Default is theory evaluation at the location of data points only. If n is specified the values are not computed at the data point positions but on <num> equidistant points in the interval between x1 and x2. If the value <num> is negative the interval is divided in equidistant steps on a logarithmic scale.  The option convolute applies for SANS data only and requires special parameters in the data sets (see SANS).</num></num></xc2></xc1></x1></x1></num>		
fit	[x1 <x1> x2 <x1> go]</x1></x1>		
	uns a fit of all selected data records with the list of activated theories. By <x1>, <x2> a (sub) interval for data comparison may be specified. If any parameters are give go is needed to get an immediate start of fit, otherwise a second call of fit without parameters starts the execution.  Further options: writes one line into the opened parameter file according to the actual selection and/or theory status. fit  [sc scan1 [scan2]]  [x1 startvalue] [x2 endvalue] and other the opts. [auto] [maxit max_number_of_iterations] [ngood est_no. of valid digits in the theory] [maxstep max_step_between_to_values] [trustreg trustregion] [maxfn] max_fun_calls] [relerr] [abserr] (default) [wrtfit] writes fitted curve data (x,y,yth) at each step onto file fitdat.tmp [nowrtfit] (default) [errors] [map mgrid] [div ndiv] go [go]</x2></x1>		

Wth the **fit** -command the parameters of the activated theories are fitted to your selected data (more than one data-record may be selected!) x1 specifies the lower x-value of the fit-interval, x2 the upper value. If  $\frac{1}{1}$  auto is given x1 and x2 values are ignored and all data are respected in the fit, the next time x1 or x2 is given again a corresponding limit is again established. See also the command **clip** . With **maxfn** you can set the maximum number of function calls that take place during the fit-procedure. The **go**-option must be explicitely stated every time the **fit** command is used, if not given, **fit** will only find the parameters for the activated theory, the curve-fitting can then be done by once more entering: **fit** without any parameters. Convolution with a resolution function (SANS) may be performed; for details see **fit**! Another way to deal with resolution convolution is decribed in the **BSS** part. map option causes fit to writes files map.xy and map.ssq that contain the landscape of ssq vs paramaters (max. no of free pars.=3) The parameters varied are those with nonzero fitscale. The stepwidth used is fitscale/div (div may be given as parameter in the command line, default is 20). map must be followed by the *number of grid points* in each direction. Use *ssq.qli* to display the resulting files (2D).If map is given **NO fitting** will be performed! The comman dline must be finished by the parameter: **go** Using parameter values for output or in makros is possible by taking advantage of one of several mechanisms: 1: fit and the add a number of paramters to the resulting curve buffers these may be addressed in other functions or viewed in the plot. These parameters describe the theory parameters and errors. In the moment we are still restricted to 8 characters for the names therefore the theory parameters are coded as follows: first 2 characters = first two charactres of theory name 3rd character = one digit (1..9) telling the number of the theory instance in the activation list. 4..8 character = 1..5 charcater of the th-parameter name The corresponding names for errors are build by putting 'e' in front of the coded parameter name and truncating the result to 8 chars; those parameters may be used in expressions. See **plot noparplo** to get a plot with a non crowded parameter part 2. The last instance of a parameter name is used to create a user defined variable of that name and the value of the parameter. The corresponding error has the same name with a trailing 'e.'. This mechanism is only viable if one or a few instances of different theories are defined. 3. The internal vectors that hold the parameters are accessible via **+th\_par(i,j)**, **+th\_err(i,j)** are evaluated to the value of the i-th parameter (parameter-error) of the j-th instance of any theory. not yet implemented th init initialises the theories if they were changed during execution of datreat After changes in a theory or adding a new theory you have to compile it and linkit by invoking "make pur"in the main directory of datreat. this will look for new theories and compile the shared library libtheospur.so in datreat/lib which is used for the theories. All this is done automatically with the comand th\_make below compiles the theos library afterwards you need to reinitialise the theories with th\_init

## Saving Parameters and Fit Results

#### **Output/Edit Theory Parameters**

al	
activist	

lists current theory definition .on console and into file *lastth*.

# Write a List of Any Parameters/Variables

print	filename expr1 [expr2] [expr3][expr9]		
	<b>appends</b> one line of numbers to file <filename> expr19 denote expressions that compute to a number.</filename>		
	They may parameter names, user-defined variables, automatic variables or expressions containing numbers		
	and/or these variables. <b>Remember:</b> only strings that start with one of the following characters: '+-		
	0123456789.(' will compute to a number, an open bracket '(' requires a closing counterpart ')', blanks are		
	not allowed within an expression. Refer to section <i>PredefinedVariables/Functions</i> for a list of useful links		
	to internal information.		

## **Cast Parameters into a Datreat Input File**

open	filename parnam1 [no.theo1] parnam2 [no.theo2]		
	will prepare a file named file <filename> in datreat readable input format for output. This file will be filled with the values of the parameters with names <parameters (1st="" <no.theo="" <parameters="" activated="" and="" as="" associated="" be="" by="" case="" currently="" file="" from="" giving="" in="" it="" latter="" list)="" may="" number="" of="" or="" parameter-block="" parameters="" selected="" selection="" setting,="" specified,="" taken="" the="" theory="" with="" x-values="" y-values.="">, that the parameter is to be taken from the <no.theo>- th instance of the theory's parameter name in the activated list. Open write a header.</no.theo></parameters></filename>		
write			
	writes one line into the opened parameter file according to the actual selection and/or theory status.		
	<b>Tip:</b> to use any other values make them a parameter of the selected file by the command putpar.		
close			
	closes the open parameter collecting file and writes #eod mark.		

# Data Manipulation

## **General Purpose**

arit   f1 < f2 < f2 > to <numor> [mult] [div] [norm nonorm]    </numor>	1			
assigned <numor>. Default is addition, the options mult or div select multiplication or division of y norm or nonorm relate to monitor normalization, this is only effective if the data-records contain a corresponding parameter monitor.  Combine raster <xstart> <dx> <n> to <numor> [norm nonorm]  combines contents of selected data records and interpolates them to a common grid (raster).  norm or nonorm relate to monitor normalization, this is only effective if the data-records contain a corresponding parameter monitor.  function [<value>]  applies a function to x or y-values f selected data records. See also: xformel and yformel as alternated function:  x : x&gt; x  log(x) : ln(x)&gt; x  exp(x) : exp(x)&gt; x  x**2 : x²&gt; x  x**2 : x²&gt; x  x+ <s1> : x* f1&gt; x  x+ <s1> : x* s1&gt; x  sqrt(x) : sqrt(x)&gt; x  rouse : q²*√(w14*x)&gt; x (q from parameter, w14 from command or parameter)  zimm : (q**3*kT/(6*pi*eta)*x)**(2/3)&gt; x (q, temp, eta_solv from parameter, or command)</s1></s1></value></numor></n></dx></xstart></numor>	ı			
norm or nonorm relate to monitor normalization, this is only effective if the data-records contain a corresponding parameter monitor.  combine raster <xstart> <dx> <n> to <numor> [norm nonorm] combines contents of selected data records and interpolates them to a common grid (raster). norm or nonorm relate to monitor normalization, this is only effective if the data-records contain a corresponding parameter monitor.  function [<value>]  applies a function to x or y-values f selected data records. See also: xformel and yformel as alternated function:  x : x&gt; x log(x) : ln(x)&gt; x exp(x) : exp(x)&gt; x x**2 : x²&gt; x x**4 : x* f1&gt; x x**51 : x* f1&gt; x x+ <s1> : x* f1&gt; x sqrt(x) : sqrt(x)&gt; x rouse : q²*√(w14*x)&gt; x (q from parameter, w14 from command of parameter) zimm : (q**3*kT/(6*pi*eta)*x)**(2/3)&gt; x (q, temp, eta_solv fparameter, or command)</s1></value></numor></n></dx></xstart>	ı			
combine raster < xstart> < dx> < n> to < numor> [norm nonorm]  combine contents of selected data records and interpolates them to a common grid (raster).  norm or nonorm relate to monitor normalization, this is only effective if the data-records contain a corresponding parameter monitor.  fun function [< value>]  applies a function to x or y-values f selected data records. See also: xformel and yformel as alternated function:  x : x> x log(x) : ln(x)> x exp(x) : exp(x)> x x**2 : x²> x x**4 < f1> : x* f1> x x+ < s1> : x+ s1> x sqrt(x) : sqrt(x)> x rouse : q²*√(w14*x)> x (q from parameter, w14 from command or parameter)  zimm : (q**3*kT/(6*pi*eta)*x)**(2/3)> x (q, temp, eta_solv for parameter, or command)				
combines contents of selected data records and interpolates them to a common grid (raster).  norm or nonorm relate to monitor normalization, this is only effective if the data-records contain a corresponding parameter monitor.  fun function [ <value>]  applies a function to x or y-values f selected data records. See also: xformel and yformel as alternate function:  x : x&gt; x log(x) : ln(x)&gt; x exp(x) : exp(x)&gt; x x**2 : x²&gt; x x**4 + <s1> : x* f1&gt; x sqrt(x) : sqrt(x)&gt; x rouse : q²*√(w14*x)&gt; x (q from parameter, w14 from command parameter) zimm : (q**3*kT/(6*pi*eta)*x)**(2/3)&gt; x (q, temp, eta_solv five parameter, or command)</s1></value>				
norm or nonorm relate to monitor normalization, this is only effective if the data-records contain a corresponding parameter monitor.  function [ <value>]  applies a function to x or y-values f selected data records. See also: xformel and yformel as alternate function:  x : x&gt; x log(x) : ln(x)&gt; x exp(x) : exp(x)&gt; x x**2 : <math>x^2</math>&gt; x  <math>x^*</math> ** ** ** ** ** ** ** ** ** ** ** ** *</value>				
corresponding parameter <code>monitor</code> .  fun function [ <value>]  applies a function to x or y-values f selected data records. See also: <code>xformel</code> and <code>yformel</code> as alternate function:</value>				
function [ <value>]  applies a function to x or y-values f selected data records. See also: xformel and yformel as alternate function:  x : x&gt; x log(x) : ln(x)&gt; x exp(x) : exp(x)&gt; x x**2 : <math>x^2</math>&gt; x x**61&gt; : <math>x^2</math> f1&gt; x x+ <s1> : <math>x^2</math> sqrt(x)&gt; x rouse : <math>q^{2*}\sqrt{(wl4*x)}</math>&gt; x (q from parameter, wl4 from command parameter) zimm : <math>(q^*3*kT/(6*pi*eta)*x)**(2/3)</math>&gt; x (q, temp, eta_solv f parameter, or command)</s1></value>	ives.			
function: $ x \qquad : x \qquad> x \\ log(x) \qquad : ln(x) \qquad> x \\ exp(x) \qquad : exp(x) \qquad> x \\ x^*2 \qquad : x^2 \qquad> x \\ x^* \qquad : x^* f1 \qquad> x \\ x+  \qquad : x+ s1 \qquad> x \\ sqrt(x) \qquad : sqrt(x)> x \\ rouse \qquad : q^2*\sqrt{(wl4*x)}> x \qquad (q from parameter, wl4 from command exparameter) \\ zimm \qquad : (q^*3*kT/(6*pi*eta)*x)**(2/3)> x (q, temp, eta_solv final parameter, or command) $	ives.			
$\begin{array}{llllllllllllllllllllllllllllllllllll$				
$\begin{array}{llllllllllllllllllllllllllllllllllll$				
exp(x) : exp(x)> x x**2 : x²> x x* <f1> : x* f1&gt; x x+ <s1> : x+ s1&gt; x sqrt(x) : sqrt(x)&gt; x rouse : q²*√(w14*x)&gt; x (q from parameter, w14 from command of parameter) zimm : (q**3*kT/(6*pi*eta)*x)**(2/3)&gt; x (q, temp, eta_solv for parameter, or command)</s1></f1>				
$x+ < s1>$ : $x+ s1$ > $x$ $sqrt(x)$ : $sqrt(x)> x$ rouse : $q^2*\sqrt{(w14*x)}$ > $x$ (q from parameter, w14 from command parameter) zimm : $(q**3*kT/(6*pi*eta)*x)**(2/3)$ > $x$ (q, temp, eta_solv f parameter, or command)				
$x+ < s1>$ : $x+ s1$ > $x$ $sqrt(x)$ : $sqrt(x)> x$ rouse : $q^2*\sqrt{(w14*x)}$ > $x$ (q from parameter, w14 from command parameter) zimm : $(q**3*kT/(6*pi*eta)*x)**(2/3)$ > $x$ (q, temp, eta_solv f parameter, or command)				
rouse : q²*√(wl4*x)> x (q from parameter, wl4 from command of parameter)  zimm : (q**3*kT/(6*pi*eta)*x)**(2/3)> x (q, temp, eta_solv f parameter, or command)	x+ <s1> : x+ s1&gt; x</s1>			
<pre>parameter) zimm : (q**3*kT/(6*pi*eta)*x)**(2/3)&gt; x (q, temp, eta_solv f parameter, or command)</pre>	sqrt(x)   : sqrt(x)> x rouse : $q^2*\sqrt{(wl4*x)}> x$ (q from parameter, wl4 from command or			
parameter, or command)	parameter)			
v : v> v				
log(y) : ln(y)> y exp(y) : exp(y)> y				
y**2 : y <sup>2</sup> > y				
y* [f1] : y* f1> y y+ [s1] : y+ s1> y				
sqrt(y) : sqrt(y)> y				
deff : y/x²> y				
with appropriate calculation of errors  The results are written to newly created data records.				
<b>xformel</b> ;(xx*3+q)				
<b>yformel</b> ;(sin(yy)*xx^3)				
allows to specify expression by which the x and y-values (denoted xx and yy within the expressions)	of the			
selected data record are transformed. The action is initiated by the command <mark>funfun</mark> .  Note the semicolon; which is necessary here. The expression is given within brackets, no blanks.				
funfun [immediate] [op]				
treats selected data-records according to the x- and y formulae which are either taken from internal st	_			
as specified by the <b>xformel</b> and <b>yformel</b> commands if the option <mark>immediate</mark> is given <b>OR</b> are taken consecutive lines (max length 132) in the file <b>formdat</b> .	as			
If the option op is given the data transformation id performed "on place" otherwise new data records	are			
created.				
Errors are treated appropriately.				
creates an error column for the selected data-records by using the expression specified by the command				
yformel.	ıd			

#### **SANS Related**

invers	[bkgr <b>]</b>			
	creates a data-record with $x \rightarrow x^2$ and $y \rightarrow 1/(y-b)$ .			
mirror	<k0> <k1> <k2></k2></k1></k0>			
	generates a mirror-image of a selected data-record <mark>k0</mark> is the estimated x-center <mark>k1</mark> and <mark>k2</mark> specify the range that shall be used to determine the center.			
qc				
	converts X-values (channels) of the first selected data-record to q-values. For that purpose the data-record has to have the following f_parameters: xk0, lambda, detdis, bklen ; i.e. channel of zero angle, wavelength, detector distance, size of one detector channel.			
spline	[iequal <i>] [nneu <n>] [auto noauto smpar <s>] spline interpolation of the fiirst selected data-record. nneu specifies the number of points of the splined data. auto selects automatic parameter determination, noauto allows to specify smpar <s> to control the amount of smoothing.</s></s></n></i>			
des	[nneu <n>] [qmax <qm>] [errabs <ea>] [errrel <er>]</er></ea></qm></n>			
	infinite slit height desmearing. <mark>nneu</mark> specifies the number of points of the treated data. Needs spline immediately before beeing called. Needs <mark>f_parameter: delqv</mark>			
mux/	trans  thick <t> xmax <xmax> nfft <n></n></xmax></t>			
dmx				
	SANS multiple scattering computation / removal.			
	c yi < i0 * sigma(x) * [unit sample thickness:d] = c lim[d>0] (i-measured(x,d) / d) c the output data on y will the be scaled such that they represent c yo < i0 * i-measured(x,d) / d (note: no limit !!) c c i.e. the scattering intensity including all orders of multiple c elastic small angle scattering will be calculated for a sample c of unit thickness assuming the same primary intensity factor,i0, c as for the input data.			
	c input variables: c t: transmission reduction factor due to small angle sc. c xmax: largest scattering angle (or q) to be considered c dx: increment of x c nx: no. of points to be generated c nfft: fft no. of points (optimal choice nfft=2**m) c output variables: c x(1nfft/2): output x-values c y(1nfft/2): output y-values			

# **Obsoletes**

out-gli	<filename></filename>	
	writes data in simple x y column form.	
inscn	<filename></filename>	

# makro Language

#### Call

A makro is a file (name max. 8 characters) which contains a header line and a collection of command lines.

The filename must be different from the names of genuine commands.

At call a number of values may be given after the name that replace the parameter-variables in the makro.

I.e. call:

#### makname 1.23 op3 (expression1) .xx.

invokes a makro with name maknam the first parameter evaluates to 0.123000e+00 the second is a string op3 the third gets the value of (expression1) and the last one is again a string XX. Also shell commands are involved, so you can also change file names etc.

Nested calls up to a nesting level of 10 are supported. However, all variables have global scope!

It is possible to use normal shell commands in a makro eg to copy a fit result or rename a file. To load a list of files with unknown filenames (perhaps in a makro to load all b\_\_\_???? files inside a directory ( with a line "q 0.05" use grep ))

Some commands to generate this makro (do it inside of datreat):

echo makro > newmakro :writes newmakro with line "makro" ls b\_\_\_41\* | sed 's#\#in #' >> newmakro :sed appends "in " in front of filenames echo sel 1 2 3 4 select some data

echo gp > newmakro : gp will plot selected data

start it with newmakro

think about the possibilities

#### **Header and Parameters**

The first line of a makro must start with the keyword: **makro**.

Then a number of arguments **arg1** ...may follow which generally give the heading line the following structure:

```
makro _a1_ _a2_ _this _ another
```

in the makro body the arguments are *string replaced* at any place where they occur by the values given when the makro is called. Expressions are first evaluated.

#### **Expressions**

Expressions evaluate to a (real) number and may be used throughout **in commands and input** *files* wherever a number is expected. The system identifies an expression by the occurrence of one of the following characters at the beginning of a blank-separated item:

```
+-.0123456789(
```

they may contain the following binary operators:



they may use the **system specific**, user-defined or automatically defined variables and *functions* as well as the following standard mathematical functions:

sqrt(x), sin(x), cos(x), tan(x), asin(x), acos(x), atan(x), ln(x), exp(x), abs(x), int(x)

Expression may *not contain any blanks*, since these are considered as item separators throughout the system.

## Loops

Loops must be formulated explicitly using **if** and **goto**.

## Commands that relate to the makro language

set	v_name <(value expression)>		
	creates uservariable v_name with value or value of expression		
vars?			
	lists all defined variables and values		
??	(expression)		
	displays the value of an expression, internal parnames from datasets can be used  (value from first of multiple selected sets used)		
	?? +q evaluates the internal data variable q to its value		
	expression to be evaluated are indicated by brackets () or +- or contain +*/- like a+b*q		
	set q2 q^2 to calculate the square of q and access it latter as uservariable q2		
_			
clr	v_name		
	removes a variable. <b>Hint:</b> a variable of the same name will mask the visibility of am f_parameter. To access the f_parameter the variable must be cleared. Vice versa creation of a corresponding variable may deliberately be used to perform this making.		
if	(expression1) < > = <= >= (expression2) then cmd goto :		
	realizes an if condition by comparing the values of two expressions.		
goto	:label		
	jumps to label. The label has to start with a colon ':' and must be the only item of a line.		
cms	<system~command> obsolete non commands or makros used as shell</system~command>		
	commands		
	transmits the given system command to the shell. All blanks within the system command may be replaced by '~'. By using this mechanism errors may be avoided that would occur if the command contains items beginning with '+','-' or '.' which normally would be considered as expressions to be evaluated by the datreat input-communication module.		

## **Predefined Variables/Function**

nbuf	evaluates to the number of loaded items, i.e. points to the end of the dir-list.		
num(j)	evaluates to the "numor" of the j-th loaded data record.		
nv(j)	evaluates to the length (number of points) of the j-th loaded data record.		
xv(i,j)	evaluates to the x-value of the i-th point of the j-th loaded data record.		
yv(i,j)	evaluates to the y-value of the i-th point of the j-th loaded data record.		
ye(i,j)	evaluates to the y-value error of the i-th point of the j-th loaded data record.		
XX	evaluates to the currently appropriate x(i,j)-value (in xyformel expressions).		
уу	evaluates to the currently appropriate y(i,j)-value (in xyformel expressions).		
ye	evaluates to the currently appropriate y(i,j)-error (in xyformel expressions).		
maxx / minx	evaluates to the maximum/minimum (x) of the current data-record.		
maxy / miny	evaluates to the maximum/minimum (y) of the current data-record.		
sumx(i1,i2,j)	evaluates to the sum of x-values from the i1-th point to the i2-th point of the j-th loaded data record.		
sumy(i1,i2,j)	evaluates to the sum of y-values from the i1-th point to the i2-th point of the j-th loaded data record.		
sel	evaluates to the last selected data record number.		
nsel	evaluates to the number of selected data records (length of selected list).		
isel(i)	evaluates to the data record number of the i-th selected list item.		
sc(i)	evaluates to the data-record address (number in list) for data with numor=i. (inverse of num(j))		

th_par(i,j)	evaluates to the value of the i-th parameter of the j-th activated theory instance.	
th_err(i,j)	evaluates to the error of the i-th parameter of the j-th activated theory instance.	
f_parnam(i)	evaluates to the value of parameter f_parnam of the i-th data-record.	
f_parnam	evaluates to the parameter of the currently considered data record (e.g. in xyformel expressions)	

## **Automatically Created Variables**

#### Main

ssq0	chi-squared from last fit / thc.	
t_parnam (fit) parameter of theory only unique if there is just one theory instance with this parameter		
e.t_parnam	evaluates to the error of t_parnam.	

**Theories** create a number of user-variables and/or f\_parameters, see there to get the corresponding descriptions.

## Interaction with the system

#### History

history	shows history of last 20 commands	
hist	writes them to file history	
<b>"_???"</b>	if the command has an underscore as first character the last command starting with the same characters is	
underscore	redone immediately.	
hist clear	clears internal history list	
	to create a makro from history:	
	echo makro > newmakro : create	s a file newmakro with first line "makro"
	histo clear :clear h	istory
	: do son	nething
	histo : shows	history and writes it without line numbers to file history
	cat history >> newmakro :append	s history to newmakro
	newmakro : use nev	wmakro whenever you need

#### Path

You can give presets in the "initdatr" makro somewhere in your datreat makro path.

path	shows path definitions		
macrpath	<newpath> change to newpath</newpath>	defailt "./"	additional path to search fpr makros
datapath	<newpath> change to newpath</newpath>	defailt "./"	path to search for input datafiles
savepath	<newpath> change to newpath</newpath>	defailt "./"	path where something is stored

#### **Active working directory**

cd	<path></path>	changes active working directory. "" is working

# Collection of Standard and Example makros

#### cth obsolete but there

Editing of theory parameters:

makro

al

acl

## **Programming**

#### Installation

#### **Prerequesities**

gfortran

**LAPACK** -- Linear Algebra PACKage

from http://www.netlib.org/lapack/

BLAS (Basic Linear Algebra Subprograms)

from http://www.netlib.org/blas/

should be installed on your system

(if you have problems with blas (missing functions) ask your system administrator to install from the original package.

In lapack is a subset of blas included but it is only a subset)

Graphical output

xmgrace should be installed

http://plasma-gate.weizmann.ac.il/Grace/

other libraries are included in datreat

#### Compile: (tested on CENTOS 64 bit and Ubuntu 32bit)

- Datreat is a local operating program -> userspace you dont need root for datreat itself.
- uncompress the tar.gz or your source file
- In datreat main folder go to "src" where you find the "Makefile", type in a shell >>make
- If you have a \$HOME/bin directory a link to the executable in datreat/bin will be placed there, which calls datreat from the correct place, if bin is in your PATH variable you can call it from everywhere.
- Afterwards you can reduce the used space on disc with "make clean".
   To delete all compiled files use "make cleanall" This is necessary after release of a new version or if you have problems with changes of names
- **Test:** After compile start with datreat change dir with "cd test" start macro "testmac" as example macro

make <Options>"

Options:" builds datreat

clean deletes object files executables were not touched, libraries untouched cleanall deletes everything including libraries exept source, for a new start

help shows this "

distribution make a tar.gz to distribute to others with pure source files and documentation

## Linking theories

The theories section is or should be a **standalone library** which can , in near future , be used without datreat. The corresponding theos Makefile is within src/theos . theories are linked automatically with a limitation to 40 theories. Please select your needed theories and move the unneeded theories to another folder. The old version theories in f77 are still working. Please use only free available non-commercial libraries.

Naming convention is: th\_give\_a\_usefull\_descriptive\_name.f90

Additional helper function: without\_a\_th\_in\_front\_and\_a\_descriptive\_name.f90 It is planned to make it possible to use also C instead of Fortran90. Perhaps at some point we will use Python as command interpreter with the old Fortran theories. But till now this is only my plan --Ralf

## **Creating New Theories**

```
Example
```

```
function th_hohdiff(x,pa,thnam,parnam,npar, ini, nopar ,params,napar,mbuf)
                        Water dynamics for TOF vibrational+rotational+translational motion RANDOM JUMP - DIFFUSION SEARS
                        J. Teixeira
                         Experimental determination of the nature of diffusive motions of water molecules at low temperatures
Phys. Rev. A 31, 1913–1917 (1985)
! implicit none real th_hohdiff character*8 thnam,parnam(20) ! theory and parameternames real pa(20)
 integer
integer
integer
                               npar,ini,ier
mbuf
                                                                 ! Anzahl der Parameter data , intent(inout) : name des parameters n , intent(inout) :: value des parameters n , intent(inout) ::
                                                                                                                                       , intent(inout) ::
intent(inout) ::
                                nopar
 character*80 napar(mbuf)
real params(mbuf)
real result
 integer nj
real amplitu
real u quadra
real self dif
real tau0_dif
                               nparx
 real a_rot
real d_rot
real shiftt
real elas_dw
real linbgr
 real q logical found_q double precision pi,h,w,elastic,Dw,gamma,Transdif,j0_a_q,j1_a_q,Rotdiff,I_q_omega
 real e_mev ! meV in data! internal val. name of independent varable
if(ini.eq.0) then ! initialization of theories
  thnam = 'hohdiff' ! name in datreat
  nparx = 7 ! number of parameters
  if(npar.lt.nparx) then
    write(6,"(' theory: ',a8,' no of parametrs=',i8,' exceeds current max. = ',i8)")thnam,nparx,npar
    th_hohdiff = 0
    return
           return
endif
npar = nparx
           Please give the Units and meaning here! name of theory parameters parnam(1) = 'amplitu' ! overall amplitude parnam(2) = 'u_quadra' ! A^2 Debye-Waller-Factor <u^2> parnam(3) = 'self dif' ! A^2/ns Self Diffusion coefficient Random Jump diffusion parnam(4) = 'tau0_dif' ! ns residence time Random Jump diffusion parnam(5) = 'a rot' ! A rotation radius (distance OH in h2O' = 0.98A) parnam(6) = 'd_rot' ! ns^-1 water rotational diffusion parnam(7) = 'shiftt' ! as pa(1) shift elatisc bgr to 0 e_mev+shiftt*q tetum
 return
endif
e_mev = x ! r

amplitu = pa(1)

u_quadra= pa(2)

self_dif = pa(3)

tau0_dif = pa(4)

a_rot = pa(5)

d_rot = pa(5)

d_rot = pa(7)

q = 0.1 ! A^-1

! get parameters
                                         ! meV
                                                                 X
                                                    Q waveyector
   get parameters from the data call getpar ('q ',q,nopar,param
                                             ',q,nopar,params,napar,mbuf,ier) !A^-1 scatteringvector
  pi=3.1415926535897
!h=4.13567E-15 eVs= 4.13567E-6 eVns
h = 4.13567E-6
! convert meV to ns^-1
w= ((e mev+shiftt*q)*0.001)/(h/2*pi)
if (w.eq.0) then
                                              w = 1e - 20
 w=1e-20
endif
! Debye Waller factor
DW=exp(-1*u_quadra*q**2)
!Translation Diffusion
gamma=self_dif*q**2/(1+self_dif*q**2*tau0_dif)
Transdif=(1/pi)*(gamma)/(gamma**2+w**2)
! Rotational fiffusion
```

Please give blue lines in given format. We will extract a description from these lines All comment with! Will be a description including a paper.

The parnam section will be a description of parameters and Units.

getpar searches for data parameters with name ''(8 char long))

setpar will store these params in the dataset.

## **Creating New Commands**

????? If you want??

## **Available Theories**

Use theos more is coming soon

# OLD datreat help \*\*\*\*\* datreat information \*\*\*\*\* symbols: <,> are used to mark a datreat-command (usually the short form) [,] mark optional parameters commands available: <command/synonym/synonym..> symbols: com

<aclast acl=""> :</aclast>	reactivates the last theory used
<activ1st al=""> :</activ1st>	lists active theories & current parameters
	change single theory parameter value or
scale	
<label> :</label>	supply a theory parameter with a label
<pre><couple> :</couple></pre>	install a coupling of a
	theory parameter with a label
	allows arithmetic operations on your data
<pre><combine> :</combine></pre>	interpolates and combines data from
	several datasets onto one raster
	clears list of selected datarecords
	executes a cms command (xedit,list,)
<pre><desactiv dac=""> :</desactiv></pre>	deactivates all theories
	lists all loaded and generated datarecords
	lists all selected datarecords
<pre><edit> <clip> :</clip></edit></pre>	allows editing of a specified datarecord
	remove points (by numer or error-limit) allows saving of a specified datarecord
	multiple data-record save (all
selected+fits)	multiple data-record save (all
	starts a fit of the selected data with the
1120	activated theory
<yfitform> :</yfitform>	enters a fitformula for eval-theory !
	opens a file for the collection of
	<pre>parameters(fitparametres) vs parameters(f.</pre>
<pre><write> :</write></pre>	writes the actual parameter(fitp.) into
	the opened file
<pre><close> :</close></pre>	close the opened parameter collection file
	gives infos
•	reads a datafile
	converts i vs q-data to i-1 vs q**2
<iout> :</iout>	sets output-level
<mirror m=""> :</mirror>	mirrors the data
<numorpls> :</numorpls>	sets the increment of scan-numbers when
l. + /	the data is treated. (default 10000)
	plots selected datarecords
	allows to set plot-parameters. no plot!
	erases specified datarecords from buffer converts channel-numbers to q
<fun></fun>	treat x and/or y -values by some functions
	x-formula and y-formula from file fromdat
<fft> :</fft>	experimental small angle multiple scatt.
<quit q=""> :</quit>	leaves the program
<sel> :</sel>	selects specified datarecords
<sym> :</sym>	makes data symmetric
<pre><spline> :</spline></pre>	generates spline coefficients smooth data
	infinte slit height desmearing
	computes data according to activated theo-
	ries (no fit !)

<theos> : lists all theories available <title/tit> : defines a title for the plots : renames axis of selected items <rename>

: puts a parameter into the list of selected <putpar>

: clears databuffer <zero/z>

<set> <clr> : set/create a user-defined variable : remove user-defined variables

: if-construct for makros <if> : goto construct for makros: display all userdefined variables <goto>

<vars?>

: reference to internal data. <ref>

#### adressing a specific datarecord:

there are two ways to access a specific set of data: either use the linenumber of the databuffer-directory or use the 'scan'-code of your data, which is defined in your datafile or generated by some operations of the program. nearly all of the commands require the keyword 'sc', when you want to use the scan-code.

makro-facilities: you can define a makro containing a series of commands which may use parameters. it is possible to call makros recursively!!

to generate a makro use the <cms> xedit-command.

the first line must be:

makro [par1] [par2] [...] now you can simply type your commands - one per line! to use the parameters simply type the parameter-name.

example: <plot> sc par1 xmax par2 ymax 60 makros allow for the use of a few basic like !!!> programming features as set, if and goto. any numerical parameters may be a formula. to identify a parameter as number or formula the parameter expression has to begin with

one of the following characters: '(+-.0..9'.

there are two standard makros, which, of course, can be altered to suit your purposes: doit fname scannr and cth. the first makro requires a filename and its scancode as inputs and will read the file, find the mirror-axes, symmetrize and convert channel-numbers to q, the second makro calls xedit with the last activated theories and parameters. this allows easy theory-parameter-changes.

----- general remarks -----

to execute datreat simply type : datreat after a while you will see a list of all available theories and then the request-prompt: '---->'. always when this symbol appears, you can type a command. datreat will try to execute the command or - if there is no such command - it will look for a makro with that name. if there is no command and no makro with that name, no action will take place.

a command line has the following syntax:

```
<cmd> parname1 value1 [value2] [..] [parname2] [value1] [...]
```

cmd is the command-name, parameters are adressed by names and seperated with a blank. with certain parameters you can set more than one value. these have to be seperated by blanks, too.

usually every parameter once defined is kept until it is explicitly changed. one important exeption from this rule is the list of selected datarecords. <m>,<sym> and <qc> allow only one selected item. if there is more than one in the list, only the first datarecord is used!

the maximum no of names and of numerical values that may be given at once is 20, the maximum length of an input line is 80 characters. one input line may contain several commands seperated by ';'.

once you activated a theory and fitted your data, the fit-curve will automatically be plotted. you can recognize a fit-curve in the data-buffer (use <dir> to see) by a negative scan-number. if you don't want the fit further displayed, use <purge> to erase it.

```
----- additional remarks -----
```

new theoretical curves for fitting may be programmed by use of one of the subroutines thx1 ... thxn as templates (near the end of the listing). if you look at the predefined theories, you will immediately see, what you have to do.

to save a generated datarecord, use the <edit>-command. when you enter the fullscreen-editor, simply type 'file file xxxx' into the command-line, where xxxx is the name, you want to give your file.

\*

\* ----- detailed explanations ----ac <ac> theoryname [par1] [scale1] [par2] [scale2] [...]
or <ac> theoryname parname par1 [scale1] [parname...]

the theorynames can be listed with the <al>-command.
the number of parameters to be passed depends on the number you
defined for this theory (see general remarks).
the scale is used in the fit-procedure, for sensible results it

or <ac> theoryname multiply parname par1 [scale1] [parname...]

the scale is used in the fit-procedure. for sensible results it should be ten to the power of the corresponding parameter.

you can activate more than one theory by calling <ac> several times. all activated theories will be added or multiplied respectively. i.e. if the multiply flag is given the corresponding theory result is multiplied with the result from the preceding theories (sum or product) instead of beeing added.

acl <acl>

theories that were activated before are reactivated with the parameters as stored in file lasth. this file is rewritten every times when <al> (list activated theories) is executed (this takes also place during the fit-procedure). you can edit file lasth using the <cms> x-command. typing of multiply (blank separated) behind a theory name flags this as multiplicative theory, the result of which is multiplied with the result from the previous theories

to yield the new (intermediate) result.

the command sequence al, <edit>, acl is combined in the cth makro, which thereby provides a tool for easy change of theory(fit)-parameters.

al <al>

all activated theories and their actual parameters are listed. this command rewrites the file lastth.

chgthpar <chgthpar> theoryname [number] parametername [par new\_value]
[scale new\_value]

changes the parameter given by parametername in the specified theory to

a new value and/or the scale of the parameter.

par and scale may be specified both or individually.

If the theoryname is not unique (e.g. the same theory is active several times)

 $$\operatorname{the}$$  optional <number> specifies the (sequence) number of it in the list of

activated theories.

label <label> theoryname [which occurence] parametername label

supplies a theory parameter with a label. the label may contain up to 4 characters, it may not be a number nor may its first character be a number. the label may be referenced if linear couplings of parameters are to be employed during a fit. see command couple.

cth <cth>

is a standard makro to faciliate the changing of theory-(fit) parameters. it loads the actual parameter set into the editor, thereby allows for editing and rereads the (changed) parameters after filing of the editor input.

installs a coupling of a labeled parameter onto a parameter quoted in this command. labels may be assigned to parameters by the label command. the whole procedure may also be done by editing the fitparameter-file. see command label.

arit <arit> [normflag] [typ] [sc scan1 [scan2 ...]] [f1 factor1] [f2 factor2] [to scan]

this command requires two selected scans; these can be selected in the command with the sc-parameter.

If sc is not give arit expects two curves to be selected, they are used as source in the sequence they were selected.

normflag must have one of the values 'norm' or 'nonorm' - if not specified, norm is defaulted.

typ may be div or mult (if not give add is assumed)

the command performs arithmetic operations with your data: factor1 \* scan1-data + factor2 \* scan2-data is written into the destination-scan datarecord. if not specified by to, the destination is defaulted.

the norm-option normalises by the monitor-parameter given in your data-file. if this parameter is not specified, the program will give a message.

errors are preserved

this command requires two or more selected scans; these may be selected in the command with the sc-parameter. the data are collected on a new scan, which has x-values that are created by the raster directive. y-values are generated by interpolation from the input data sets. if the input data sets are provided with a monitor parameter, data and monitor values are summed for each overlapping channel. finally all data are normalized to the summed monitor values. if no monitor-data are given, 1 is assumed. errors are evaluated using the 1/sqrt(n) ansatz. x-points that are not present in the set of input points are set to zero.

cs <cs>

this command clears the list of selected data-records.

cms <cms> cms-command

with <cms> it is possible to use most of the cms-commands, especially the xeditor. when a cms-command is executed, it will return to the program.

dac <dac> [number1] [number2 ...]

the theories are deactivated according to their number of activation. if you use <dac> without a parameter, all theories are deactivated.

you can reactivate these theories by <acl>.

dir <dir> [<clength> value]

this command shows you all the datarecords presently loaded. it is recommended to press the pa2-button before executing this command.

The optional parameter <clength> followed by a value=1..80 set the number of characters of the comment to be displayed.

dsl <dsl>

lists all selected datarecords.

edit <edit> [number] [sc scan]

with <edit> you can change the datarecords that are loaded or generated by the program. this command will call the xeditor.

<edit> also gives you the possibility to save these records on
your disk: you only have to specifiy a filename when leaving the
xeditor by file or ffile. if you only type file, your datarecord
is stored in file datbuf and your original file (if existing)
will not be affected.

save <save> to filename SEE ALSO: msave <save> dirnum to filename <save> n name to filename <save> sc numor to filename <save>

store a datarecord on disk permanently with the name: file filename a. if no item is specified the first datarecord in the selection table will be saved. by giving dirnum the dirnum-th record as shown in the dir-list may be saved. by specifying n name a file with the internal name <name> will be save and by sc <numor> a file with the corresponding numor will be saved. if no destination <filename> is given the data will be saved onto file lastsave.

msave <msave> filename

store a datarecords on disk permanently with the name : file filename a.

Records that are selected are stored (associated fit results are also stored).

<open>

<open> filename parnam1 [no.theo1] parnam2 [no.theo2]
will prepare a file named file <filename> a for output.
this file will be filled with the values of the parameters
with names <parnam1> as x-values and <parnam2> as y-values.
the parameters may be taken from the parameter-block associated
with the currently selected file (1st file of selection list)
or from the parameters associated with the currently activated
theory setting, in the latter case it may be specified
by giving <no.theo> that the parameter is to be taken from
the <no.theo>-th theory that is activated.

<write>

write one line into the opened parameter file according to the actual selection and/or theory status.

<close> <close>

close the open parameter collecting file.

treat all selected files by applying some functions to the x and/or y-values.

option : op or np : item ---> item or item ---> new item : x ---> X optionx: Х : ln(x) ---> xlog(x) $: exp(x) \longrightarrow x$ exp(x) x\*\*2 ´ : x\*\*2 ---> X x\* [f1] : x\* f1 ---> X x+ [s1] : x+ s1 ---> x  $\operatorname{sqrt}(x)$  :  $\operatorname{sqrt}(x)$  ---> x

```
: q^{**}2^{*}sqrt(w14^{*}x) --> x (q from
                    rouse
parameter, wl4 from command or parameter)
                                : (q^**3*kT/(6*pi*eta)*x)**(2/3) --> x (q,
                    zimm
temp, eta_solv from parameter)
                                                                       (optio
nal eta from command)
         optionx:
                               : ln(y) ---> y
                    log(y)
                               : exp(y) ---> y
                    exp(y)
                               : y**2
                                        ---> y
                    y* [f1]
                               : y* f1 ---> y
                               : y+ s1 ---> y
                    y+ [s1]
                               : sqrt(y)---> y
                    sqrt(y)
                    deff
                               : y/(x^**2) -> y
         with appropriate errors
         x-values and y-values of all selected items are treated
funfun
         by the formula lines that are given in file formdat a
         as first and second line. the lines may not contain
         any blanks within the formula. they may be written
         as usual formulas, the x-values are to be referred
         by xx, the y-values by yy.
         exponentiation is indicated by ^ eg. xx^2, xx may
         not be negative!
         log is ln.
         functions available: sin, cos, tan, asin, acos, atan, ln, exp,
                              sqrt, int, abs
         the trig. function operate with rad or degree, which may
         be switched by the setrad or setdeg command.
         the output may be reduced by iout -5.
fit
            <fit> [sc scan1 [scan2 ...]]
                  [x1 startvalue] [x2 endvalue] .. and other the opts.
                  [auto]
                  [maxit max._number_of_iterations]
                  [ngood est_no. of valid digits in the theory]
                  [maxstep max._step_between_to_values]
                  [trustreg trustregion]
                  [maxfn max_fun_calls]
                  [relerr]
                  [abserr]
                              (default)
                  [wrtfit]
                  [nowrtfit] (default)
                  [errors]
                  [map mgrid] [div ndiv] go
                  [go]
         with the <fit>-command the parameters of the activated theories
         are fitted to your selected data (more than one data-record may
         be selected !)
         x1 specifies the lower x-value of the fit-intervall, x2 the
         upper value.
         if auto is given x1 and x2 values are ignored and all data are
         respected in the fit. the next time x1 or x2 is given again
         a corresponding limit is again established.
         with maxfn you can set the maximum number of function calls that
         take place during the fit-procedure.
```

the (volatile) errors option causes the computation of statictical errors for the fit-parameters, provided errors for the data are given. the errors option must be combined with the go option.

the go-option must be explicitely stated every time the <fit>command is used. if not given, <fit> will only find the parameters for the activated theory. the curve-fitting can then be done by once more typing <fit> without any parameters.

convolution with a resolution function may be performed. for details see thc !

fit and the add a number of paramters to the resulting curve buffers

these may be addressed in other functions or viewed in the plot. These parameters describe the theory parameters and errors. In the moment we are still restricted to 8 characters for the names therefore the theory parameters are coded as follows:

first 2 characters = first two charactres of theory name

3rd character = one digit (1..9) telling the number of the theory in the activation list.

4..8 character = 1..5 charcter of the th-parameter name

the corresponding errors are build by puuting 'e' in front of the coded parameter name and truncating the result to 8 chars.

See --> plot noparplo to get a plot with a noncrowded parameter part

those parameters may be used in expressions.

OPTION: map

if --> fit map writes files map.xy and map.ssq that contain the landscape of ssq vs paramateters (max. no of free pars.=3) The parameters varied are those with nonzero fitscale. The stepwidth used is fitscale/div (div may be given as

parameter

in the command line, default is 20).

Map must be followed by the number of grid points in each direction.

Use ssq.gli to display the resulting files (2D). If map is given NO fitting will be performed ! The commandline must be finished by the parameter: go

yfitform enter a formula that may be used as fitting function by invoking eval as theory. the formula is entered as a pseudo-resline after a;
yfitform ;p(1)+xx\*p(2)+xx\*xx\*p(3)
where xx is the independent variable and p(i) are the parameters as given with eval.
! this type of function evaluation is quite inefficient!
however it may be used for a quick check of simple ideas without changing the program code.

help with the <help>-command you get these information or, if you use <help> <command>, you will get specific information about one datreat-command.

in <in> filename

```
must be stored in the following format:
         (the uppercaes words are keywords!)
         line 1: arbitrary comments
         line 2: filename y-label vs x-label
                                                   scancode
         line 3:
                                                 {must be empty !}
         line 4: parameters
         line 5: par1 value1
         line x: parxy valuexy {10 is the maximum number of parameters}
       {let's call it line 10}
         line 11:
                                                 {must be empty !}
         line 12: values
         line 13ff.: [x]
                           x-value [y]
                                        y-value
                                                    [e] error
             or
                         x-value1 x-value2 x-value3 ...
                     У
                         y-value1 y-value2 y-value3 ...
                     е
                        error1
                                  error2
                                           error3 ....
       {always one line x-values and one line y-values !}
         line y:
                                                 {must be empty !}
         line y+1: #eod
         there are two files that can be linked to your data by the
         get-command of the xeditor: file kopf must be linked to the
         beginning of your data and provides the header, file eod gives
         an empty line and the #eod-keyword.
invers
            <invers> [bkgr background]
         this command converts the first selected datarecord to
         i-1 vs q**2 - format. the original data must be i vs q.
         with bkgr you can subtract a constant background-level
         before inverting.
            <iout> [number]
iout
         with <iout> you can set your output-level.
            <m> est_mid x1 x2
m
         the <m>-command generates a mirror-image of a selected data-
         record (it must be selected before !).
         in est_mid you estimate the center-x.
         with x1 and x2 you specifiv the range that shall be used to
         determine the center.
numorplos
           <numorpls> offset
         this command should only be used in the beginning of a session,
         it sets the offset, that is added to the scan-code, when any
         operation is performed by the program.
         the default-value is 10000. this means, that usually you can
         identify your data as follows:
               00xxx: raw-data
                                        i(ntensity) vs kanal
               10xxx :
                       mirror-data
                                                          {see <m>}
                                        dto.
                       symmetric-data sym-i vs kanal
               20xxx :
                                                          {see <sym>}
                        converted-data i vs q
               30xxx :
                                                          {see <qc>}
               40xxx: "inverted"-data i-1 vs q**2
                                                          {see <invers>}
              -xxxxx : fit-curve
                                                          {see <fit>}
         to keap your head clear of garbage, you should use the cycle-
```

this command allows you to read a datarecord from disk. the data

numbers of your experiment as scan-numbers. to distinguish "horizontal"-data from "vertical"-data use 1xxx for horizontal and 0xxx for vertical data. if this definition becomes popular, it will be easier to exchange data with other users.

```
 [sc scan1 [scan2 ...]]
   *[fsc fitscan1 [fitscan2 ...]]*
    [xmin minimum_x_value]
    [xmax maximum_x_value]
    [ymin min_y] [ymax max_y]
    [framx length_of_x_axes] [framy length_of_y_axes]
    [frlux beginning_of_x_axes] [frluy beg_of_y_axes]
    [symb symbol1 [symbol2 ...]]
    [icolo color1 [color2 ...]]
    [o... option-specifier]
    [fitflag]
    [textflag]
    [show_parameter_flag]
    [txsize textsize]
    [font fontnumber]
    [legsize legend_size]
    [legx relative_x_position_of_legend]
    [legy rel_y_pos_of_legend]
    [errplo / noerrplo]
    [parplo / noparplo]
    [log_x / lin_x]
    [log_y / lin_y]
```

the <plot>-command is one of the most powerful commands of this program. once you know how to use the parameters, you can do nearly every plot you like. after execution you are asked, whether you want to print the plot elsewhere.

with sc you can select certain data-records as usually.

fsc may be used to select several fit-curves. if you performed a fitting, this fit is automatically selected. after selecting other curves by sc or a <sel>-command, you have to specify the fit-curves that shall be plotted. this is especially useful, when you want to have different fits for different intervals. attention: you must give the corresponding positive scan-number to select a fit! this allows it to select fit-curves automatically in a makro.

choose the borders of your plot with xmin, xmax, ymin and ymax. the frxxx-parameters can be used to size and locate your plot. symb chooses the symbols that are used to represent a data-point in the plot. the first number will be used for the first selected item, the second for the second, ... for detailed information see the gr-software-handbook or (if you read a print of this guide to datreat) the appendix.

icolo chooses the colors for your plot. only selected datarecords can be colored. the codes are:

- 0 black (default for all curves)
- 1 red

p

- 2 blue
- 3 green
- 4 purple
- 5 yellow
- 6 zyan.

o... specifies the options for your axes. for detailed information see the gr-software-handbook (graxs-subprogram). the format

is (example): ox x=1  $\{\text{shows x-axes below your graph}\}$ .  $\{2 \text{ puts the axes above and 3 on both sides. } -1/-2/-3 \text{ shows a grid}\}$ 

fitflag can be fits or nofits and says if fit-curves are to be presented.

textflag is text or notext and is used to show title and legend or not.

show\_parameter\_flag is parplo or noparplo and determines whether (when text is set) all parameters are shown or only the legend. txsize determines the size of the text on the axes and the title.

with font you can choose a graphic font. detailed information is again given in the gr-software-handbook or in the appendix. legsize sets the size in which the legend-text and parameters are plotted.

legx and legy can be used to move the legend elsewhere, when it would cover your data (or if you prefer a different place).

{if you want to set plot-parameters only, use <plot>>} errplo / noerrplo activates/desactivates plotting of errorbars

parplo / noparplo may be use to switch (on/off) the writing of the parameters associated to each file at the right border of the plot. If noparplo is in effect one may select a number of parameters that is plotted anyway by specifying the name of those parameters in the plot command (volatile).

log\_x, log\_y yields log-scaling of the x/y-axes. Use lin\_x, lin\_y to go back to linear scaling.

```
p0 <p0> [... see p(lot) ... ]
```

<plot0> sets parameters for following plots, but does not plot
anything. this command is especially useful for defining an
initial setting according to your taste. the parameters are
described in <p(lot)>.

this command is used to delete datarecords from your directory. use the <dir>-command before to get the right numbers. all selections will be removed.

```
qc <qc>
```

<qc> converts the symmetric i vs kanal -data into i vs q. this
command operates on the first selected item.

```
q p
```

leaves the program. if you send plots to external printing-media it may be useful to quit datreat once a while, because the plot is only started when the program is quitted.

the <sel>-command can be used to select certain datarecords for

further operations. The same effect is achieved when using sc as keyword in several commands.

If direct addressing is used, the OPTION add allows to keep

NEW

the previous selction and to add further recods to the selection. when an operation generates a new datarecord this new record is automatically selected.

Option: fit+ adds the corresponding fits to the selection

NEW

given as list of sequence numbers

NEW

parameter sc selects scans according to their numors sc+ adds numor selections to present list only a number list selects entries according to their sequence number in the dir-list OPTION fits searches for old fitted items and selects them as fits of the selected items if the numors do match.

sym <sym>

this command calculates the mean-values of the data on the left and right side of the center and creates a new datarecord with only one side.

spline

<spline> [auto]/[noauto] [nneu nneu] [smpar smpar]
spline approximation of scattered data on the selected scan.
the smoothing parameter smpar is automatically determined if
the option auto is given. otherwise (noauto) smpar must be
specified (by try and error, look at the plot!). smpar will
influence the degree of smoothing. nneu specifies the number
of points (density) the smoothed synthetic new scan should
get within the x-range of the original scan.
the spline polynomial coefficients of the last spline call
are stored in an internal common block for subsequent use,
i.e. in des or fft.

des

des [qmax qmax] [nneu nneu] infinite slit height desmearing up to a q-value of qmax. the new dataset will contain nneu points. the data to be desmeared must be splined immediately before des is invoked.

thc  $\langle thc \rangle [n] n [x[c]1 x1] [x[c]2 x2] [auto] [convolute] [off]$ 

with <thc> you can recalculate fit-data. n is the number of points that are used for calculation; 0 means: take value from last fit.

NEGATIVE n will cause an even distribution x-values on a logarithmic scale, if x1=0 or x2=0 or x1>=x2 this option is ignored.

if auto is given x1 and x2 values are ignored and all data are respected in the calcul. the next time x1 or x2 is given again a corresponding limit is again established.

if convolute the external routine datconv is taken to convolute the calculated data before leaving thc. the number of points and range before convolution is specified by n, xc1 and xc2 ( instead of n, x1 and x2 ! ). the result gets the x-values of the selected template(s). the x-ranges before and after convolution may be different depending on the transformation that is implicit in the

convolution kernel (e.g. q-values ---> scattering angles). usually x belongs is a physical variable before and an experiment variable after convolution. xc1/2 refer to the physical variable range. the selected item has to be in experiment space.

all convolution settings are also valid during fit. convolution is valid until convolution off is specified. any parameters needed to specify the convolution kernel are to be given as parameters in the selected data-item, they are extracted by datconv using the getpar routine.

theos <theos>

if you want to know what theories are available in your version of datreat, simply type theos and you will see.

tit <tit> titlestring

with the <title>-command you can define a title that is displayed with your plots if the text-option is set.

putpar <putpar> <parname> <value>
 a parameter with name <parname> will be created and set
 to the value <value> in all selected items. if the
 parameter already exists, its value is updated.

z <z>

this command is used to clear (<zero>) the buffer. no datarecords can be selected afterwards. the <dir>-command will show an empty list.

makro {see general information also}

there are some standard-makros which simplify the use of this program:

doit filename scancode loads the specified data-record and does everything up to the

q-conversion automatically.

cth calls the xeditor to edit the last activated theorie-para-

meters and then activates this

theorie.

plnorm sets standard values for plot

i vs q.

plzimm sets standard values for plot

i-1 vs q\*\*2.

set set/create a user-defined variable.
usage is set <name> <value> [ <name> <value> ...
variables may be listed by the command: vars?
some commands transfer their parameters to the
uservariable-stack, so that these variables can
be referred to.

```
clr
           remove user-defined variables
           usage clr all removes all uservariables.
           clr <name1> <name2> ... <namex> removes the
           variables with <name1> .. <namex>
           set angle-units to degree for the commandline
setdeg
           trigonometric functions.
setrad
           set angle-units to rad for the commandline
           trigonometric functions.
??
           display the value of a formula.
           ?? <expression>
           displays the evaluated value of <expression>
if
           if-construct for makros
           usage if <expression1> = <expression2> then <commandline>
                 if <expression1> > <expression2> then <commandline>
                 if <expression1> < <expression2> then <commandline>
                 if <expression1> <= <expression2> then <commandline>
                 if <expression1> >= <expression2> then <commandline>
                 if <expression1> <> <expression2> then <commandline>
goto
           goto construct for makros
           usage goto :label
           the label :label has to start with ':', the
           line containing the label may not contain any
           other commands.
           lists all userdefined variables
vars
ref
         internal data may be referenced within expressions.
                 --> current x-value (funfun)
                 --> current y-value (funfun)
         xv(i,j) --> j-th x-value of i-th databuffer
         yv(i,j) --> j-th y-value of i-th databuffer
         ye(i,j) --> j-th yerror-value of i-th databuffer
         sumx(i,j1,j2)
         sumy(i,j1,j2) --> sums
                --> first selected buffer
         sel
                 --> number of loaded buffers
         nbuf
         maxx
         maxy
         minx
                --> max and min of the first selected curve
         miny
                --> current outputlevel
         <parname> --> parametervalue of selected curve
         <parname(n)-> parametervalue curve n
         <fitpar> --> name of fitparameter, only the
                       last of equally named fit-parameters
                       is accessible
         th_par(ip,it) --> value of the ip-th parameter of the
                                                                     NFW
                           it-th activated theory
         th_err(ip,it) --> error of the ip-th parameter of the
                                                                     NEW
                           it-th activated theory
```

```
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new

sel all [parna] [val] band [val]
sel next [parna] .....
sel add +.....
theory kohl with gaussian resolution parameters for folding (makros resit ...)
```

theor

#### **BSS-Data Converter**

convert\_dat2.f

#### **NEW DEVELOPMENT: Thinktank**

create a new entry in the theory header line (same level as 'multiply') to specify a parameter range in which the theory is evaluated, outside a zero contribution is added. The parameter value stems from the parameter block of the actual data-record (iadda).

E.g.: theory XYZ [multiply] [range <parname> <val-min> <val-max>] default: full range.

Programming Interface by multiple include files/preprocessor (see McStas as example). Intro-section

Closing-section

???

Preprocessor/ vs Description