The general integral equation

\[ u(t) = f(t) + \int_0^t K(t, s, u(s))ds \]
ecomes

\[ u = f(t) + \text{int}\{K(t,t',u)\} \]

The convolution equation:

\[ v(t) = \exp(-t) + \int_0^t e^{-(t-s)^2} v(s)ds \]
would be written as:
\[ v(t) = \exp(-t) + \int \exp(-t^2) \, dv \]

If one wants to solve, say,
\[ u(t) = \exp(-t) + \int_0^t (t - t')^{-\mu} K(t, t', u(t')) \, dt' \]

the form is:
\[ u(t) = \exp(-t) + \int \mu \{ K(t, t', u) \} \]

and for convolutions, use the form:
\[ u(t) = \exp(-t) + \int \mu \{ w(t) \# u \} \]

**NETWORKS**

special zip=conv(type,npts,ncon,wgt,root)

where root is the name of a variable and wgt is a table, produces an array zip with npts:
\[
zip[i] = \sum_{j=-ncon}^{ncon} wgt[j + ncon] \cdot root[i + j]
\]

The sparse network has the syntax:

special zip=sparse(npts,ncon,wgt,index,root)

where wgt and index are tables with at least npts * ncon entries. The array index returns the indices of the offsets to which to connect and the array wgt is the coupling strength. The return is
\[
zip[i] = \sum_{j=0}^{ncon} wgt[j \cdot ncon + j] \cdot root[k]
\]

where
\[
k = index[i \cdot ncon + j]
\]

The other two types of networks allow more complicated interactions:

special zip=fconv(type,npts,ncon,wgt,root1,root2,f)

evaluates as
\[
zip[i] = \sum_{j=-ncon}^{ncon} wgt[ncon+j] \cdot f(root1[i+j], root2[i])
\]

and

special zip=fsparse(npts,ncon,wgt,index,root1,root2,f)

evaluates as
\[
zip[i] = \sum_{j=0}^{ncon} wgt[ncon+i+j] \cdot f(root1[k], root2[i])
\]

where
\[
k = index[i \cdot ncon + j]
\]

**OPTIONS**

The format for changing the options is:

@ name1=value1, name2=value2, ...

where name is one of the following and value is either an integer, floating point, or string. (All names can be upper or lower case). The first four options can only be set outside the program. They are:

- **MAXSTOR=integer** sets the total number of time steps that will be kept in memory. The default is 5000. If you want to perform very long integrations change this to some large number.
• BACK= \{Black,White\} sets the background to black or white.
• SMALL=\texttt{fontname} where \texttt{fontname} is some font available to your X-server. This sets the “small” font which is used in the Data Browser and in some other windows.
• BIG=\texttt{fontname} sets the font for all the menus and popups.
• SMC={0,...,10} sets the stable manifold color
• UMC={0,...,10} sets the unstable manifold color
• XNC={0,...,10} sets the X-nullcline color
• YNC={0,...,10} sets the Y-nullcline color

The remaining options can be set from within the program. They are

• LT=\texttt{int} sets the linetype. It should be less than 2 and greater than -6.
• SEED=\texttt{int} sets the random number generator seed.
• XP=\texttt{name} sets the name of the variable to plot on the x-axis. The default is \texttt{T}, the time-variable.
• YP=\texttt{name} sets the name of the variable on the y-axis.
• ZP=\texttt{name} sets the name of the variable on the z-axis (if the plot is 3D.)
• NMP=\texttt{int} tells XPP how many plots will be in the opening screen.
• XP2=\texttt{name},YP2=\texttt{name},ZP2=\texttt{name} tells XPP the variables on the axes of the second curve; XP8 etc are for the 8th plot. Up to 8 total plots can be specified on opening. They will be given different colors.
• AXES=\{2,3\} determine whether a 2D or 3D plot will be displayed.
• TOTAL=value sets the total amount of time to integrate the equations (default is 20).
• DT=value sets the time step for the integrator (default is 0.05).
• NPLOT=value tells XPP how frequently to output the solution to the ODE. The default is 1, which means at each integration step.
• T0=value sets the starting time (default is 0).
• TRANS=value tells XPP to integrate until \(T=\text{TRANS}\) and then start plotting solutions (default is 0.)
• NMESH=\texttt{integer} sets the mesh size for computing nullclines (default is 40).
• METH=\{ discrete,euler,modeuler,rungekutta,adams,gear,volterra, backeul, qualrk, stiff,cvode,5dp,83dp,2rb,ymp \} sets the integration method (default is Runge-Kutta.)
• ATOLER=value sets the absolute tolerance for CVODE.
• TOLER=value sets the error tolerance for the Gear, adaptive RK, and stiff integrators. It is the relative tolerance for CVODE.
• BOUND=value sets the maximum bound any plotted variable can reach in magnitude. If any plottable quantity exceeds this, the integrator will halt with a warning. The program will not stop however (default is 100.)
• DELAY=value sets the maximum delay allowed in the integration (default is 0.)
• PHI=\texttt{value},THETA=\texttt{value} set the angles for the three-dimensional plots.
• XLO=\texttt{value},YLO=\texttt{value},XHI=\texttt{value},YHI=\texttt{value} set the limits for two-dimensional plots (defaults are 0,\,-2,20,2 respectively.) Note that for three-dimensional plots, the plot is scaled to a cube with vertices that are \(\pm 1\) and this cube is rotated and projected onto the plane so setting these to \(\pm 2\) works well for 3D plots.
• XMAX=\texttt{value},XMIN=\texttt{value},YMAX=\texttt{value},YMIN=\texttt{value},ZMAX=\texttt{value},ZMIN=\texttt{value} set the scaling for three-d plots.
• OUTPUT=\texttt{filename} sets the filename to which you want to write for “silent” integration. The default is “output.dat”.
• POIMAP=\{ section,maxmin \} sets up a Poincare map for either sections of a variable or the extrema.
• POIVAR=name sets the variable name whose section you are interested in finding.
• POIPLN=value is the value of the section; it is a floating point.
- POISGN={ 1, -1, 0 } determines the direction of the section.
- POISTOP=1 means to stop the integration when the section is reached.
- RANGE=1 means that you want to run a range integration (in batch mode).
- RANGEOVER=name, RANGESTEP=number, RANGELOW=number, RANGEHIGH=number, RANGERESET=Yes,No, RANGEOOLDIC=Yes,No all correspond to the entries in the range integration option.
- TOR_PER=value, defined the period for a toroidal phasespace and tell x XPP that there will be some variables on the circle.
- FOLD=name, tells XPP that the variable name is to be considered modulo the period. You can repeat this for many variables.
- AUTO-stuff. The following AUTO-specific variables can also be set: NTST, NMAX, NPR, DSMIN, DSMAX, DS, PARMIN, PARMAX, NORMMIN, NORMMAX, AUTOVAR.

**COLOR MEANING**
- 0-Black/White; 1-Red; 2-Red Orange; 3-Orange; 4-Yellow Orange; 5-Yellow; 6-Yellow Green; 7-Green; 8-Blue Green; 9-Blue; 10-Purple.

**KEYWORDS**
You should be aware of the following keywords that should not be used in your ODE files for anything other than their meaning here.

\[
\sin \cos \tan \text{atan atan2 sinh cosh tanh exp delay ln log log10 t pi if then else asin acos heav sign ceil flr ran abs del\_shft max min normal besselj bessely erf erfc arg1 \ldots arg9 @ $ + - / * ^ ** shift | > < == >= <= != not \# int sum of i'
\]

These are mainly self-explanatory. The nonobvious ones are:

- heav(arg1) the step function, zero if arg1<0 and 1 otherwise.
- sign(arg) which is the sign of the argument (zero has sign 0)
- ran(arg) produces a uniformly distributed random number between 0 and arg.
- besselj, bessely take two arguments, \( n, x \) and return respectively, \( J_n(x) \) and \( Y_n(x) \), the Bessel functions.
- erf(x), erfc(x) are the error function and the complementary function.
- normal(arg1,arg2) produces a normally distributed random number with mean arg1 and variance arg2.
- max(arg1, arg2) produces the maximum of the two arguments and min is the minimum of them.
- if(<exp1>)then(<exp2>)else(<exp3>) evaluates <exp1> If it is nonzero it evaluates to <exp2> otherwise it is <exp3>. E.g. if(x>1)then(ln(x))else(x-1) will lead to ln(2) if x=2 and -1 if x=0.
- delay(<var>,<exp>) returns variable <var> delayed by the result of evaluating <exp>. In order to use the delay you must inform the program of the maximal possible delay so it can allocate storage.
- del\_shift(<var>,<shift>,<delay>). This operator combines the delay and the shift operators and returns the value of the variable <var> shifted by <shift> at the delayed time given by <delay>.
- ceil(arg), flr(arg) are the integer parts of <arg> returning the smallest integer greater than and the largest integer less than <arg>.
- t is the current time in the integration of the differential equation.
- pi is \( \pi \).
- arg1, \ldots, arg9 are the formal arguments for functions
- int, # concern Volterra equations.
- shift(<var>,<exp>) This operator evaluates the expression <exp> converts it to an integer and then uses this to indirectly address a variable whose address is that of <var> plus the integer value of the expression. This is a way to imitate arrays in XPP. For example if you defined the sequence of 5 variables, \( u0, u1, u2, u3, u4 \) one right after another, then shift(u0,2) would return the value of u2.
• \textit{sum(}\textless\textit{ex1},\textless\textit{ex2})\textit{of(}\textless\textit{ex3}\textgreater\textit{)} is a way of summing up things. The expressions \textless\textit{ex1},\textless\textit{ex1} are evaluated and their integer parts are used as the lower and upper limits of the sum. The index of the sum is i' so that you cannot have double sums since there is only one index. \textless\textit{ex3}\textgreater is the expression to be summed and will generally involve i'. For example \textit{sum(1,10)of(}i')\textit{) will be evaluated to 55. Another example combines the sum with the shift operator. \textit{sum(0,4)of(shift(}u0,i')\textit{)) will sum up }u0\textit{ and the next four variables that were defined after it.