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The **G Math Toolkit Reference Manual** describes the features, functions, and operation of the G Math Toolkit. With this toolkit and your LabVIEW or BridgeVIEW application, you can perform complex mathematical calculations. The G Math Toolkit is intended for use by scientists, engineers, and mathematicians, or anyone needing to solve mathematical problems in a simple, quick and efficient manner. It can also be used as an educational aid by those interested in learning and expanding their knowledge of mathematics. To use this manual effectively, you should be familiar with the G programming language and the basic theory behind the type of problem that you want to solve.

**Organization of the G Math Toolkit Reference Manual**

The **G Math Toolkit Reference Manual** is organized as follows:

- Chapter 1, *Introduction*, introduces the G Math Toolkit, presents some examples of how you can use this toolkit for your day-to-day mathematical tasks, and describes some of the basics of working with the G Math Toolkit.

- Chapter 2, *Parser VIs*, describes the VIs that act as an interface between the end user and the programming system. These VIs parse the formula, which is in the form of a string, and convert the formula string to a form that can be used for evaluating results.

- Chapter 3, *Data Visualization VIs*, describes the VIs that are used for plotting and visualizing data in several different forms. These include advanced methods such as animation, contour plots, and surface cuts.

- Chapter 4, *Ordinary Differential Equation VIs*, describes the VIs you can use to solve ordinary differential equations, both symbolically and numerically.

- Chapter 5, *Zero Finder VIs*, describes the VIs that find the zeros of 1D or *n*-dimension, linear or nonlinear functions (or system of functions).

- Chapter 6, *Optimization VIs*, describes the VIs you can use to determine local minima and maxima of real 1D or *n*-dimension
functions. You can choose between optimization algorithms based on derivatives of the function and algorithms working without these derivatives.

- Chapter 7, *1D Explorer VIs*, describes the VIs you can use to study real-valued 1D functions given in symbolic form. You can study different qualities of function graphs with and without additional parameters.

- Chapter 8, *2D Explorer VIs*, describes the VIs you can use to examine 2D functions given in symbolic form, where parameterization is allowed. You can numerically calculate extrema and partial derivatives.

- Chapter 9, *Function VIs*, describes the VIs you can use to evaluate some common mathematical functions.

- Chapter 10, *Transform VIs*, describes VIs that implement some transforms commonly used in mathematics and signal processing.


- Appendix B, *References*, section contains references to the mathematical theory or algorithm implemented in each VI.

- Appendix C, *Customer Communication*, contains forms you can use to request help from National Instruments or to comment on our products and manuals.

- The *Glossary* contains an alphabetical list and description of terms used in this manual, including abbreviations, acronyms, metric prefixes, mnemonics, and symbols.

## Conventions Used in This Manual

The following conventions are used in this manual:

- **bold**
  
  Bold text denotes a parameter, menu name, menu item, or dialog box button or option.

- **italic**
  
  Italic text denotes emphasis, a cross reference, or an introduction to a key concept.

- **bold italic**
  
  Bold italic text accompanied by the icon to the left denotes a note, which alerts you to important information.

- **monospace**
  
  Text in this font denotes text or characters that are to be literally input from the keyboard, sections of code, programming examples, and syntax examples. This font is also used for the proper names of disk
drives, paths, directories, programs, subprograms, subroutines, device
names, functions, variables, filenames, and extensions, and for
statements and comments taken from program code.

<> Angle brackets enclose the name of a key on the keyboard—for example,
<PageDown>.

- A hyphen between two or more key names enclosed in angle brackets
denotes that you should simultaneously press the named keys—for
example, <Control-Alt-Delete>.

<Keyname> Key names are capitalized.

Abbreviations, acronyms, metric prefixes, mnemonics, symbols, and
terms are listed in the Glossary.

Related Documentation

The following documents contain information that you may find helpful
as you read this manual:

- LabVIEW User Manual
- LabVIEW Analysis VI Reference Manual
- HiQ for Macintosh User Manual
- HiQ for Macintosh Function Reference Manual
- HiQ for Windows User Manual

Customer Communication

National Instruments wants to receive your comments on our products
and manuals. We are interested in the applications you develop with our
products, and we want to help if you have problems with them. To make
it easy for you to contact us, this manual contains comment and
configuration forms for you to complete. These forms are in
Appendix C, Customer Communication, at the end of this manual.
The G Math Toolkit is a new approach to solving advanced mathematics and analysis problems. With the G Math Toolkit and the G programming language underlying LabVIEW and BridgeVIEW, you can graphically assemble your problem-solving program as a block diagram. In addition, the interactive front panels introduce a new level of interactivity to math and analysis problem solving. With the G Math Toolkit, you can put the compiled speed, connectivity, open architecture, and flexibility of G to work in your applications. This chapter explains what the G Math Toolkit is, and illustrates how you can use this toolkit for your day-to-day mathematical tasks.

Some of the advantages of G programming over classical procedural programming languages such as FORTRAN, C, Pascal, and so on, include:

- Visual programming
- Data driven system design
- Virtual instrumentation
- Platform independence

The graphical user interface combined with the data driven paradigm makes G the perfect tool for complex measurement and analysis tasks because of the commonality of such tasks with mathematical operations and algorithms. You can use G as a programming system for almost all kinds of numerically and symbolically oriented mathematical routines.

The core of the G Math Toolkit is the Parser VI Library, which makes manipulating formulas on LabVIEW front panels possible. All the G Math Toolkit VI libraries are interconnected, and leverage off each other. For example, the Ordinary Differential Equation (ODE) VI library is based on the Function and the Parser VI libraries.

The strength of the complete G Math Toolkit is not just the power of individual VIs in performing mathematical calculations, but also in combining VIs to solve extremely complex problems. In addition, you can interface real-world measurements to the mathematical algorithms...
in order to obtain practical solutions. This flexibility makes the toolkit an extremely powerful analysis tool.

**G Math Toolkit Examples and Parameter Help**

Study the G Math Toolkit examples to learn some of the possible theoretical and practical applications of this package. These example libraries, installed in the `gmath` directory in your LabVIEW or BridgeVIEW Examples directory, are grouped into the following categories:

- **Graphics.llb** examples showing the graphics capabilities
- **Math.llb** examples from mathematics
- **Mechanix.llb** examples applied to the field of mechanics
- **Misc.llb** miscellaneous “real world” examples
- **Optimiz.llb** examples related to optimization
- **Sig_proc.llb** examples applied to signal processing

Also, many of the G Math Toolkit VIs have example text on the front panel of the VI showing how to input various parameters. Double-click on the VI icon to open the front panel of the VI and examine this helpful text.

**Using the Parser VIs**

The Parser VIs are a collection of VIs that directly connect an end user to the programming system. Until now, only the LabVIEW Formula Node could manage formula expressions. But the Formula Node is a pure programming tool, not directly accessible to the user from the front panel.

In many applications, being able to enter formulas directly from the front panel is extremely useful. This is possible using the G Math Toolkit. The following examples describe some typical scenarios where you might use the G Math Toolkit.
Example 1
Given a set of measurements \((x_i, y_i)\), where \(i = 0, 1, \ldots, n - 1\), fit the set of data points into a model equation, such as the following.

\[ Y = a \sin(bX) + c \cos(dX) \]

or

\[ Y = a + bX + c \exp(dX) \]

or more generally

\[ Y = f(X, a, b, \ldots, c) \]

(Where \(a, b, \ldots, c\) are the unknown model parameters)

G can calculate such optimal system parameters as

\[ \sum_{i=0}^{n-1} (y_i - f(x_i, a, b, \ldots, c))^2 = \text{min!} \]

G uses the Levenberg Marquardt method to solve this minimization problem. Before now, the model equation had to be fixed in the formula node before running the program. With the Parser VIs, you can input the model equation on the front panel.

Example 2
Another typical example consists of the following three-step process.

1. Collect a set of discrete measurements 
   \((x_i, y_i, z_i)\) for \(i = 0, 1, \ldots, n - 1\)
2. Fit this set based on a model such as 
   
   \[ Z = f(X, Y, a, b, \ldots, c) \]

   with unknown parameters \(a, b, \ldots, c\)
3. Determining those points \((x, y)\) with
   
   \[ z = f(x, y, a, b, \ldots, c) = 0 \]
   or
   \[ z = f(x, y, a, b, \ldots, c) = \text{max!} \]
   or
   \[ z = f(x, y, a, b, \ldots, c) = \text{min!} \]

An easy way to input discrete measurements on a LabVIEW front panel makes calculating the roots, minima, and maxima for a general model simple for your end user. Notice that at the beginning of the previous process, the correct formula 

\[ Z = f(X, Y, a, b, \ldots, c) \]

is not known.
Example 3

Another application for the G Math Toolkit is to help control an x-y step motor or a robot control that positions objects in 2D or 3D space during runtime. The path of the object can be calculated with the VIs found in this toolkit.

Another 2D and 3D application of the G math toolkit is for surface description. Wings of airplanes, among other parts of machines and instruments in the real technical world, can be described by mathematical curves and surfaces. In nondestructive testing (using ultrasound, eddy currents, or X-rays), a prescan of the structure under test is initially done, followed by a more accurate scan in areas where the measured and expected values differ.

Because it is virtually impossible to store the measurements of all the curves and surfaces of an entire structure, it is necessary to take a set of coarse measurements as a preparatory step. The prescan of the wing of a plane is then followed by a more accurate scan of a smaller part of this wing. Because the Parser VIs can handle formulas on the front panel, you can use them to calculate the 2D and 3D curves of the wing in an effective manner.

Example 4

The study of solutions of differential equations (especially parameter studies) is not only a question of appropriate numeric algorithms such as the Euler method, Runge Kutta method, or the Cash Karp method but also a question of formula manipulation. With the G Math Toolkit you can manipulate differential equations on the front panel. See the examples in the math.llb example collection for sample versions of this approach to solving differential equations.

The Parser VIs in More Detail

A Parser VI scans an input string and interprets this string as a formula, or a collection of formulas. Then, the Parser VI transforms the formulas into numeric calculations and outputs the results. The Parser VI routines deal only with real numbers.

There are some differences between the parser in the G Math Toolkit and the Formula Node found in the original LabVIEW package. The following table outlines these differences.

<table>
<thead>
<tr>
<th>Element</th>
<th>Formula Node</th>
<th>Parser VI Routines</th>
</tr>
</thead>
<tbody>
<tr>
<td>Variables</td>
<td>No restrictions</td>
<td>Only a, a0, ..., a9, ...</td>
</tr>
<tr>
<td></td>
<td></td>
<td>z, z0, ..., z9, are valid</td>
</tr>
<tr>
<td>Binary functions</td>
<td>max, min, mod, rem</td>
<td>Not available</td>
</tr>
</tbody>
</table>
Chapter 1  Introduction

The precedence of operators is the same for the G Math Toolkit Parser VIs as in the Formula Nodes of G. Refer to Chapter 2, Parser VIs, for more information on specific VIs.

All functions in the G Math Toolkit use the following syntax:

\[
\text{function} \ (\text{argument})
\]

The following table lists the functions you can use with the Parser VIs.

<table>
<thead>
<tr>
<th>Function</th>
<th>Corresponding G Math Function name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>abs(x)</td>
<td>Absolute Value</td>
<td>Returns the absolute value of ( x ).</td>
</tr>
<tr>
<td>acos(x)</td>
<td>Inverse Cosine</td>
<td>Computes the inverse cosine of ( x ).</td>
</tr>
<tr>
<td>acosh(x)</td>
<td>Inverse Hyperbolic Cosine</td>
<td>Computes the inverse hyperbolic cosine of ( x ) in radians.</td>
</tr>
<tr>
<td>asin(x)</td>
<td>Inverse Sine</td>
<td>Computes the inverse sine of ( x ) in radians.</td>
</tr>
<tr>
<td>asinh(x)</td>
<td>Inverse Hyperbolic Sine</td>
<td>Computes the inverse hyperbolic sine of ( x ) in radians.</td>
</tr>
<tr>
<td>atan(x)</td>
<td>Inverse Tangent</td>
<td>Computes the inverse tangent of ( x ) in radians.</td>
</tr>
<tr>
<td>atanh(x)</td>
<td>Inverse Hyperbolic Tangent</td>
<td>Computes the inverse hyperbolic tangent of ( x ) in radians.</td>
</tr>
<tr>
<td>ci(x)</td>
<td>Cosine Integral</td>
<td>Computes the cosine integral of ( x ) where ( x ) is any real number.</td>
</tr>
</tbody>
</table>
### Chapter 1  Introduction

<table>
<thead>
<tr>
<th>Function</th>
<th>Corresponding G Math Function name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ceil(x)</td>
<td>Round to +Infinity</td>
<td>Rounds $x$ to the next higher integer (smallest integer ( \geq x )).</td>
</tr>
<tr>
<td>cos(x)</td>
<td>Cosine</td>
<td>Computes the cosine of $x$ in radians.</td>
</tr>
<tr>
<td>cosh(x)</td>
<td>Hyperbolic Cosine</td>
<td>Computes the hyperbolic cosine of $x$ in radians.</td>
</tr>
<tr>
<td>cot(x)</td>
<td>Cotangent</td>
<td>Computes the cotangent of $x$ in radians (1/tan(x)).</td>
</tr>
<tr>
<td>csc(x)</td>
<td>Cosecant</td>
<td>Computes the cosecant of $x$ in radians (1/sin(x)).</td>
</tr>
<tr>
<td>exp(x)</td>
<td>Exponential</td>
<td>Computes the value of $e$ raised to the power $x$.</td>
</tr>
<tr>
<td>expm1(x)</td>
<td>Exponential(Arg)–1</td>
<td>Computes the value of $e$ raised to the power of $x - 1$ ($e^x - 1$)</td>
</tr>
<tr>
<td>floor(x)</td>
<td>Round to –Infinity</td>
<td>Truncates $x$ to the next lower integer (Largest integer ( \leq x )).</td>
</tr>
<tr>
<td>gamma(x)</td>
<td>Gamma Function</td>
<td>$\Gamma(n + 1) = n!$ for all natural numbers $n$.</td>
</tr>
<tr>
<td>getexp(x)</td>
<td>Mantissa and exponent</td>
<td>Returns the exponent of $x$.</td>
</tr>
<tr>
<td>getman(x)</td>
<td>Mantissa and exponent</td>
<td>Returns the mantissa of $x$.</td>
</tr>
<tr>
<td>int(x)</td>
<td>Round to nearest integer</td>
<td>Rounds its argument to the nearest even integer.</td>
</tr>
<tr>
<td>intrz</td>
<td>Round toward zero</td>
<td>Rounds $x$ to the nearest integer between $x$ and zero.</td>
</tr>
<tr>
<td>ln(x)</td>
<td>Natural Logarithm</td>
<td>Computes the natural logarithm of $x$ (to the base $e$).</td>
</tr>
<tr>
<td>lnpl(x)</td>
<td>Natural Logarithm (Arg + 1)</td>
<td>Computes the natural logarithm of $(x + 1)$.</td>
</tr>
<tr>
<td>Function</td>
<td>Corresponding G Math Function name</td>
<td>Description</td>
</tr>
<tr>
<td>------------</td>
<td>------------------------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>log(x)</td>
<td>Logarithm Base 10</td>
<td>Computes the logarithm of $x$ (to the base 10).</td>
</tr>
<tr>
<td>log2(x)</td>
<td>Logarithm Base 2</td>
<td>Computes the logarithm of $x$ (to the base 2).</td>
</tr>
</tbody>
</table>
| pi(x)      | Represents the value $\pi = 3.14159...$ | $\pi(x) = x^*\pi$
               | $\pi(1) = \pi$
               | $\pi(2.4) = 2.4^*\pi$ |
| rand( )    | Random Number (0–1)                | Produces a floating-point number between 0 and 1. |
| sec(x)     | Secant                             | Computes the secant of $x$ ($1/\cos(x)$). |
| si(x)      | Sine Integral                      | Computes the sine integral of $x$ where $x$ is any real number. |
| sign(x)    | Sign                               | Returns 1 if $x$ is greater than 0. Returns 0 if $x$ is equal to 0. Returns −1 if $x$ is less than 0. |
| sin(x)     | Sine                               | Computes the sine of $x$ in radians. |
| sinc(x)    | Sinc                               | Computes the sine of $x$ divided by $x$ in radians ($\sin(x)/x$). |
| sinh(x)    | Hyperbolic Sine                    | Computes the hyperbolic sine of $x$ in radians. |
| spike(x)   | Spike function                     | spike(x) returns:
               | 1 if $0 \leq x \leq 1$
               | 0 for any other value of $x$. |
| sqrt(x)    | Square Root                        | Computes the square root of $x$. |
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<table>
<thead>
<tr>
<th>Function</th>
<th>Corresponding G Math Function name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>square(x)</td>
<td>Square function</td>
<td>square (x) returns: 1 if $2n \leq x \leq (2n + 1)$ 0 if $2n + 1 \leq x \leq (2n + 2)$ where x is any real number and n is any integer.</td>
</tr>
<tr>
<td>step(x)</td>
<td>Step function</td>
<td>step(x) returns: 0 if $x &lt; 0$ 1 if any other condition obtains.</td>
</tr>
<tr>
<td>tan(x)</td>
<td>Tangent</td>
<td>Computes the tangent of $x$ in radians.</td>
</tr>
<tr>
<td>tanh(x)</td>
<td>Hyperbolic Tangent</td>
<td>Computes the hyperbolic tangent of $x$ in radians.</td>
</tr>
</tbody>
</table>

**Error Structure**

The Parser VIs use the following error handling structure. This structure consists of a Boolean status button, a signed 32-bit integer numeric code indicator, and a string source indicator. These error handler components are explained below:

- **status** is TRUE if an error occurred. If status is TRUE, this VI does not perform any operations.
- **code** is the error code number identifying the error.
- **source** is a message explaining the error in more detail.

The default status of the error in structure is TRUE (no error), indicated by an error code of 0.
With this structure, you can programmatically check the accuracy of formulas and control the data flow in case of errors. The application uses the source field of the error handling structure as a storage for a incorrect or invalid formula input. This field displays limited error descriptions if an error is detected in your program. See Appendix A, Error Codes, for the error codes and the messages of the Parser VI routines.

There are three main kinds of Parser routines:

- The VIs representing the functionality of the LabVIEW Formula Node (the Eval Formula Node, Parse Formula Node, and Eval Parsed Formula Node VIs)
- The VIs analyzing a simple string as one formula (the Eval Formula String, Parse Formula String, and Eval Parsed Formula String VIs)
- The VIs producing whole sets of function values (the Eval Single-Variable Array, Function Explorer, and Eval Multi-Variable Scalar VIs)

The first two categories of Parser VIs can be further divided into two subcategories, the direct form and the indirect form. As an example, the direct version of the Eval Formula Node VI is represented by the following block diagram.
On the other hand, the indirect forms split the VI in two subVIs, as shown in the following illustration. You can use the indirect form in larger applications, where a two-step process (parsing and then evaluating) is more efficient.

### Variables

The Parser VIs accept only the following variables:

- \(a, a0, \ldots, a9\)
- \(b, b0, \ldots, b9\)
- \(\ldots\)
- \(z, z0, \ldots, z9\)

**Note:** *For variable and function names, only lowercase letters are allowed. The toolkit interprets capital letters as errors.*

For the sake of uniqueness, all numbers in exponential notation use the 1E-1 convention (with the capital letter E). Using 1e-1 (with the lowercase letter e) results in an error message.

The following table shows some common error codes, the description of the error, and an error example.
<table>
<thead>
<tr>
<th>Code</th>
<th>Error Description</th>
<th>Error Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>No error</td>
<td>sin(x)</td>
</tr>
<tr>
<td>1</td>
<td>Bracket problem at the beginning</td>
<td>1+x)</td>
</tr>
<tr>
<td>2</td>
<td>Incomplete function expression</td>
<td>sin(x)+</td>
</tr>
<tr>
<td>3</td>
<td>Bracket problem</td>
<td>()</td>
</tr>
<tr>
<td>4</td>
<td>Bracket problem at the end</td>
<td>(1+x</td>
</tr>
<tr>
<td>5</td>
<td>Wrong decimal point</td>
<td>1.2 (US)</td>
</tr>
<tr>
<td>6</td>
<td>Wrong number format</td>
<td>1e-3 instead of 1E-3</td>
</tr>
<tr>
<td>7</td>
<td>Wrong function call</td>
<td>sin()</td>
</tr>
<tr>
<td>8</td>
<td>Not a valid function</td>
<td>sins(x)</td>
</tr>
<tr>
<td>9</td>
<td>Incomplete expression</td>
<td>x+</td>
</tr>
<tr>
<td>10</td>
<td>Wrong variable name</td>
<td>a11</td>
</tr>
<tr>
<td>11</td>
<td>Wrong letter</td>
<td>sin(X)</td>
</tr>
<tr>
<td>12</td>
<td>Too many decimal points</td>
<td>1.23.45</td>
</tr>
<tr>
<td>21</td>
<td>Contains more than one variable</td>
<td>1+x+y4</td>
</tr>
<tr>
<td>22</td>
<td>Inconsistency in variables or numbers</td>
<td>Depends on application</td>
</tr>
<tr>
<td>23</td>
<td>Contains variables</td>
<td>Depends on application</td>
</tr>
<tr>
<td>24</td>
<td>Variables output problem</td>
<td>Depends on application</td>
</tr>
</tbody>
</table>
Parser VIs

The Parser VIs act as an interface between the end user and the programming system. These VIs parse the formula, which is in the form of a string, and convert the formula string to a form that can be used for evaluating results. The Parser VIs are found in the PARSER.LLB library.

Parser VI Descriptions

Eval Formula Node

Functionally equivalent to the Formula Node in LabVIEW, but with variables that can be entered on the front panel.

- **Input Values** is an array of numbers with a one-to-one relation to **Variables Input**.
- **Variables Input** is an array of input strings, each of which represents a valid variable name.
- **Variables Output** is an array of output strings, each of which represents a valid variable name.
- **formula** is a string consisting of one or more formulas separated by semicolons. Each formula is built up by **Variables Input** on the right side of an equation and **Variables Output** on the left side.
error in (no error) describes error conditions occurring before this VI executes. If an error has already occurred, this VI returns the value of the error in cluster in error out. The VI executes normally only if no incoming error exists; otherwise it merely passes the error in value to error out. Refer to the Error Structure section of Chapter 1, Introduction, in this manual for a description of the error in cluster components.

Output Values is a 1D array of numbers corresponding to Variables Output and formula.

error out contains error information. If the error in cluster indicated an error, the error out cluster contains the same information. Otherwise, error out describes the error status of this VI.

Example

The following inputs

formula: \( y = 3x + 4z; \ p = q^2 - 5; \)

Variables Input: \([x, z, q]\)

Input Values: \([1, 2, 3]\)

Variables Output: \([y, p]\)

result in

Output Values: \([11.00, 4.00]\)

Eval Formula String

Interprets a string as a numeric calculation and determines the result.

formula is a string representing the calculation without any variables. It consists only of numbers and mathematical functions.
error in (no error) describes error conditions occurring before this VI executes. If an error has already occurred, this VI returns the value of the error in cluster in error out. The VI executes normally only if no incoming error exists; otherwise it merely passes the error in value to error out. Refer to the Error Structure section of Chapter 1, Introduction, in this manual for a description of the error in cluster components.

y value is the result of the calculation.

error out contains error information. If the error in cluster indicated an error, the error out cluster contains the same information. Otherwise, error out describes the error status of this VI.

Example

The following input

formula: \( \sin(\pi(1/2)) + 3*5 - 2 \)

results in

y value: 14.00

Eval Multi-Variable Array

Calculates the function values of a given function at an arbitrarily given set of \( n \)-dimension points by \( y_i = f(x_{1i}, x_{2i}, \ldots, x_{ni}) \), where \( f \) is an \( n \)-dimension function given by the formula, and \( (x_{1i}, x_{2i}, \ldots, x_{ni}) \) are \( n \) independent variables.

formula is a string representing the \( n \)-dimension function under investigation.

X Values is a 2D array of X Values. Each row of the array represents the fixed values of each of the Variables of the \( n \)-dimension function. The other dimension of the array marks the different \( n \)-dimension points at which the function has to be calculated.

Variables is an array of strings. Each element of the array stands for a variable name of the \( n \)-dimension independent terms.
**Eval Multi-Variable Scalar**

Calculates exactly one function value based on a given formula $y = f(x_1, x_2, ..., x_n)$.

- **formula** is a string representing the formula of $n$ independent Variables.
- **X Values** is an array of X Values corresponding to the $n$ Variables.

The following inputs:

- **formula**: $3x_1 + 4x_2 + x_3^2$
- **X Values**: 
  
  $$[1, -1, 4; 2, 1]$$
  
  ([1, -1, 2] are the values for $x_1$, $x_2$, $x_3$ in the first iteration; [0, 4, 1] for the second)
- **Variables**: $[x_1, x_2, x_3]$ (for $x_1$, $x_2$, and $x_3$)

result in

- **Y Values**: 
  
  $$[3, 17]$$
Variables is an array of strings representing the n independent variables of the given formula. There is a one-to-one relation between Variables and X Values.

error in (no error) describes error conditions occurring before this VI executes. If an error has already occurred, this VI returns the value of the error in cluster in error out. The VI executes normally only if no incoming error exists; otherwise it merely passes the error in value to error out. Refer to the Error Structure section of Chapter 1, Introduction, in this manual for a description of the error in cluster components.

y value is the value of formula (X Values).

error out contains error information. If the error in cluster indicated an error, the error out cluster contains the same information. Otherwise, error out describes the error status of this VI.

Note: This VI calculates one and only one value of a given n-dimension function. If you want to calculate a collection of function values, use the Eval Multi-Variable Array, Parse Formula String and Eval Parsed Formula String VIs.

Example

The following inputs

formula: \(3x_1 + 4x_2 + x_3^2\)

X Values: \([1, -1, 2]\)

Variables: \([x_1, x_2, x_3]\)

result in

y value: 3.00
Eval Parsed Formula Node

The Eval Parsed Formula Node VI separates the parsing process from the evaluating process of the Eval Formula Node VI and improves the runtime behavior of a program containing the Eval Formula Node VIs at different locations. This VI has a direct connection from the Parse Formula Node VI and completes an Eval Formula Node VI calculation. You can wire the Parsed Formula Node input of this VI directly from the Parsed Formula Node output of the Parse Formula Node VI.

**Input Values** is an array of numbers, each of which corresponds to the value that is given to each of the variables. These variables can be input in the Variables Input control of the Parse Formula Node VI. The coded form of these variables is available in the Variables input decode parameter of the Parsed Formula Node cluster.

**Parsed Formula Node** is a cluster consisting of:

- **Y Values** is a 2D array of numbers representing a storage of detected and analyzed numbers of formula of the Parse Formula Node VI.
- **Tables** is a 3D array with 3 columns. The first column contains a code that stands for the operator, the other two contain codes that stand for the operands.
- **Variables input decode** is the intermediate and coded state of Variables Input. See also the Parse Formula Node VI.
- **Variables output decode** is the intermediate and coded state of Variables Output. See also the Parse Formula Node VI.

You can wire the Parsed Formula Node cluster directly from the corresponding output of the Parse Formula Node VI.

**error in (no error)** describes error conditions occurring before this VI executes. If an error has already occurred, this VI returns the value of the error in cluster in error out. The VI executes normally only if no incoming error exists; otherwise it merely passes the error in value to error out. Refer to the Error Structure section of Chapter 1, Introduction, in this manual for a description of the error in cluster components.
Output Values is a 1D array of numbers corresponding to Variables Output Decode, Y Values and Tables.

error out contains error information. If the error in cluster indicated an error, the error out cluster contains the same information. Otherwise, error out describes the error status of this VI.

Eval Parsed Formula String
Takes the output of Parse Formula String VI and fixes input values to calculate function values.

X Values is an array of X Values corresponding to Variables.

Variables is a 1D array of strings representing the independent variables. There is a one-to-one relation between X Values and Variables.

Parsed Formula is a cluster consisting of:

Y Values is a 1D array of numbers representing a storage of detected and analyzed numbers of formula.

Table is a 2D array with 3 columns. The first column contains a code that stands for the operator, the other two contain codes that stand for the operands.

This input can be wired directly from the corresponding output of the Parse Formula String VI.

error in (no error) describes error conditions occurring before this VI executes. If an error has already occurred, this VI returns the value of the error in cluster in error out. The VI executes normally only if no incoming error exists; otherwise it merely passes the error in value to error out. Refer to the Error Structure section of Chapter 1, Introduction, in this manual for a description of the error in cluster components.

y value is the result of the interpretation process, such as the function value.
error out contains error information. If the error in cluster indicated an error, the error out cluster contains the same information. Otherwise, error out describes the error status of this VI.

Eval Single-Variable Array
Calculates an array of function values at given points in a given interval by \( y_i = f(x_i) \) where \( f \) is the one-dimensional (1D) function given by the user formula.

**formula** is a string representing the 1D function under investigation.

**X Values** is the given array of input values, \( x_i \).

**error in** (no error) describes error conditions occurring before this VI executes. If an error has already occurred, this VI returns the value of the error in cluster in error out. The VI executes normally only if no incoming error exists; otherwise it merely passes the error in value to error out. Refer to the Error Structure section of Chapter 1, Introduction, in this manual for a description of the error in cluster components.

**Y Values** is the 1D array of function values of formula at the given points in the array X Values.

error out contains error information. If the error in cluster indicated an error, the error out cluster contains the same information. Otherwise, error out describes the error status of this VI.

**Example**
To calculate the formula \( y = x^2 \), for \( x = 1, 2, 3, 4, \) and \( 5 \) the input arrays would be

**formula**: \( x^2 \)

**X Values**: [1, 2, 3, 4, 5]

resulting in the output array

**Y Values**: [1, 4, 9, 16, 25]
Eval Single-Variable Scalar
Calculates exactly one function value of a given 1D function, \( y = f(x) \), where \( f \) is the function specified by the user formula.

**formula** is a string representing the function under investigation. Only one variable can be integrated in this **formula**.

**x value** is the 1D point, \( x \), at which the function value has to be calculated.

**error in (no error)** describes error conditions occurring before this VI executes. If an error has already occurred, this VI returns the value of the **error in** cluster in **error out**. The VI executes normally only if no incoming error exists; otherwise it merely passes the **error in** value to **error out**. Refer to the **Error Structure** section of Chapter 1, **Introduction**, in this manual for a description of the **error in** cluster components.

**y value** is the \( y \) value evaluated by the **formula** (\( x \) value).

**error out** contains error information. If the **error in** cluster indicated an error, the **error out** cluster contains the same information. Otherwise, **error out** describes the error status of this VI.

**Note:** This VI calculates only one value of a given 1D function. It is difficult to analyze the formula in the background. Therefore, if you want to calculate a collection of Eval Single-Variable Scalar VIs, use the Eval Single-Variable Array, Parse Formula String, and Eval Parsed Formula String VIs.

**Example**
Refer to the previous Eval Single-Variable Array VI. The same example applies to this VI, except that the input **x value**, and the output **y value** are scalars.
Parse Formula Node

Analyzes the Eval Formula Node VI inputs and yields an intermediate state as an input for the Eval Parsed Formula Node VI.

Variables Input is a 1D array of input strings, each of them representing the name of a valid input variable.

Variables Output is a 1D array of output strings, each of them representing the name of a valid output variable.

Formula is a string consisting of one or more formulas separated by semicolons.

Error in (no error) describes error conditions occurring before this VI executes. If an error has already occurred, this VI returns the value of the Error in cluster in Error out. The VI executes normally only if no incoming error exists; otherwise it merely passes the Error in value to Error out. Refer to the Error Structure section of Chapter 1, Introduction, in this manual for a description of the Error in cluster components.

Parsed Formula Node is a cluster consisting of:

Y Values is a 2D array of numbers representing a storage of detected and analyzed numbers of formula.

Tables is a 3D array with 3 columns. The first column contains a code that stands for the operator, the other two contain codes that stand for the operands.

Variables Input Decode is the intermediate and coded state of Variables Input.

Variables Output Decode is the intermediate and coded state of Variables Output.

Error out contains error information. If the Error in cluster indicated an error, the Error out cluster contains the same information. Otherwise, Error out describes the error status of this VI.
Parse Formula String

Analyzes a string as a formula and produces two numeric arrays. These arrays can be used by the Eval Parsed Formula String VI to produce X Values.

- **formula** is a string representing the formula. The formula can contain any number of valid variables.
- **error in (no error)** describes error conditions occurring before this VI executes. If an error has already occurred, this VI returns the value of the **error in** cluster in **error out**. The VI executes normally only if no incoming error exists; otherwise it merely passes the **error in** value to **error out**. Refer to the **Error Structure** section of Chapter 1, **Introduction**, in this manual for a description of the **error in** cluster components.

**Parsed Formula** is a cluster consisting of:
- **Y Values** is a 1D array of numbers representing a storage of detected and analyzed numbers of **formula**.
- **Table** is a 2D array with 3 columns. The first column contains a code that stands for the operator, the other two contain codes that stand for the operands.

This output can be wired directly to the corresponding input of the Eval Parsed Formula String VI.

**error out** contains error information. If the **error in** cluster indicated an error, the **error out** cluster contains the same information. Otherwise, **error out** describes the error status of this VI.

Usually, the Parse Formula String VI is used at the beginning of the calculation of function values. The Eval Parsed Formula String VI completes the calculation. This division works well if the analyzing process of the Parse Formula String VI finishes before the calculation processes. Keep this in mind for your own programming.
Substitute Variables

Substitutes a formula string by given rules. The rules have a parameter name - parameter content structure.

original formula is a string representing the formula, where parameter names stand for formulas.

Substitution Rules is an array of clusters describing the substitution rules.

parameter name can have any length.

parameter content must have a one-to-one relation with parameter name.

error in (no error).

formula after substitution is the final formula after all substitution rules are performed.

error out is the structure of the error handler is used as the output.

Note: A parameter name that begins with a capital letter E can produce unpredictable errors, if parts of the original string represent numbers like 1E-2. Avoid terms beginning with E in such cases.

Example

The following inputs

original formula: \( \text{ALPHA} \cdot \cos(t) + \beta \)

Substitution Rules:

\( \text{ALPHA} \rightarrow \sin(t) \)

\( \beta \rightarrow 2 \cdot t \cdot \exp(t) \)

result in

substitution formula: \((\sin(t)) \cdot \cos(t) + (2 \cdot t \cdot \exp(t))\)
Data Visualization VIs

The Data Visualization VIs are used for plotting and visualizing data in several different forms. These include advanced methods such as animation, contour plots, and surface cuts. They can be found in the VISUALIZ.LLB library.

Data Visualization VI Descriptions

Animate Graphs

Shows a collection of graphs and selects one graph from the whole collection of graphs. This VI works as an interactive tool.

- **Graphs** is an input array of graphs to be animated. This array is made up of clusters each of which is composed of two arrays.
  - **x values** is an array of the x values for a particular graph
  - **y values** is an array of corresponding y values

- **step rate [sec]** is the time between two consecutive graph representations in seconds.

- **selected graph** is the selected graph. This cluster consists of two arrays.
  - **x values** is an array of the x values for the selected graph
  - **y values** is an array of corresponding y values
selected graph index is the index of the graph you select, starting with 0.

There are three possible modes of operation:

- **Automatic**—the VI sequentially displays each graph in the array, one after the other, at the speed given by step rate [sec].
- **Forward**—you can manually step through each graph to be displayed, in the order of increasing array index.
- **Backward**—you can manually step through each graph to be displayed, in the order of decreasing array index.

**Note:** The step rate [sec] can be influenced by the user during runtime of the VI.

### Common Intensity Maps

Depicts an array of real values in a common intensity plot, with the data arranged on a rectangular grid.

- **X-Y data** is the 2D array of values \( z = f(x, y) \), where \( x \) runs from 0 to \( x \) length and \( y \) runs from 0 to \( y \) length.
- **map selector** selects one item from a predefined list of Common Intensity Maps distributions. This list is shown in the table below. The value of map selector can range from 1 to 6.
- **graph** is a 2D array consisting of \( x \) and \( y \) values. You can wire this output directly to a graph for display.
- **error**. See Appendix A, Error Codes, for a list of error codes, if map selector is out of range.

The following predefined color palettes are available.

<table>
<thead>
<tr>
<th>Map Selector</th>
<th>Description</th>
<th>Number of Colors</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>256 fire</td>
<td>White, yellow, red, and blue</td>
</tr>
<tr>
<td>2</td>
<td>256 rainbow</td>
<td>Rainbow colors</td>
</tr>
<tr>
<td>3</td>
<td>256 log up</td>
<td>White and black in log up scale</td>
</tr>
</tbody>
</table>
Contour Plot
Depicts an array of real values on a contour plot, with the data arranged on a rectangular grid.

<table>
<thead>
<tr>
<th>Map Selector</th>
<th>Description</th>
<th>Number of Colors</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>256 log down</td>
<td>White and black in log down scale</td>
</tr>
<tr>
<td>5</td>
<td>32 color</td>
<td>Green, blue, and white</td>
</tr>
<tr>
<td>6</td>
<td>32 ice</td>
<td>Green and blue</td>
</tr>
</tbody>
</table>

**X-Y data** is the 2D array of values $z = f(x, y)$, where $x$ runs from 0 to $x$ length and $y$ runs from 0 to $y$ length.

**contour lines** is the number of contour lines. The default value is 5. This number can vary between 1 and 255.

**graph** is a 2D array consisting of $x$ and $y$ values. You can wire this output directly to a graph for display.

**error**. See Appendix A, Error Codes, for a list of error codes, if contour lines is out of range.

**Note:** The axis scaling of the rectangular region can be influenced by attribute nodes.
Cut nD Surface
Depicts an n-dimension function in the form of n cuts along the n axes.

The function $f(x_1, x_2) = \sin(x_1 \ast x_1 - x_2) - \cos(\sin(x_2) - x_1)$ is investigated in the interval $(-2, 2) \times (-2, 2)$ with the help of a contour plot as shown in the following figure.

- **Cut length** is the length of the n cuts.
- **Number of points** is the number of points of each graph (cut).
- **Formula** is the formula of the nD function.
- **Variables** is the n necessary variables to describe the nD function under observation.
- **Center Point** is the center point of the n different cuts. (See Figure 3-1 in the Parametric Curve 3D VI description in this chapter.)
- **Graph** is the resulting graph as combination of all cuts along the axes.
- **Values** is the same data as graph in the form of a 2D array of numbers.
- **Error**. See Appendix A, Error Codes, for a list of error codes. Fine tuning between formula, Variables and Center Point is necessary.
The following diagram explains the meaning of the center point and the cuts through the center points.

![Diagram showing center point and cuts](image)

**Draw Graph**

Produces a graph interactively with the help of mouse actions.

- **distance** is the minimal radial distance between two different points of the graph. In a neighborhood of radius **distance**, no other point of the graph can be placed.
- **graph** is a cluster consisting of two arrays holding the x and y data points drawn on the graph by the user.
- **error**. See Appendix A, *Error Codes*, for a list of error codes, if **distance** ≤ 0.

You can generate new points on the graph and delete existing points with this VI. Position the mouse cursor where you want to create or delete a point. Click the mouse and release it and the point is created or deleted. The graph stores the points in consecutive order. You can also clear all points by using the **CLEAR** button.

**Note:** The value of distance can be adjusted by the user during runtime of the VI.
The following diagram shows a generated graph. The cross hairs mark the current position of the cursor. You can add and delete points.

**Mesh 3D**
Depicts a surface under various conditions. You can completely control the position of the observer. This also lets you calculate the projections of a chosen point of the surface. This VI works as an interactive tool.

**X-Y-Z data** is the visualized 2D data material of the curve. All points of the curve are given in the form \((x, y, z)\).

**View Info** is a cluster of data that describes the viewpoint of the observer.

**Origin** is the point the observer is looking at. (See Figure 3-1 for an illustration.) The default value is \((0, 0, 0)\).

**Axis** is the proportional factor of each axis. You can stretch or compress an axis independently of the other. The default value is \((2, 2, 2)\).
psi describes the azimuth of the observer. This is the angle (in radians) of rotation about the z axis at the origin. When psi = 0, the viewpoint is along the x axis looking away from x = 0 toward the positive values of x and perpendicular to the y axis.

radius describes the distance from the origin to the actual position of the observer.

phi describes the elevation of the observer. This is the angle (in radians) made with the x-y plane by the line of sight from the observer to the origin. When phi = \pi/2, the viewpoint is directly above the z axis.

See Figure 3-1 for an explanation of psi, radius, and phi.

interactive mode. FALSE represents “not interactive,” TRUE stands for “interactive.” The default value is FALSE.

graph is an array of clusters, each consisting of to arrays containing the x and y values of the graph data determined by View Info.

Selected Point is the coordinates of exactly one selected point of the graph. This point can interactively be determined. The projections of this point on the x, y and z axis are also presented.

Final View Info is a collection of data describing the viewpoint of the observer at the end of the interactive mode. The data in this cluster has the same structure as the data in the View Info cluster.

Parametric Curve 2D

Presents a 2D curve based on the data of an array. The VI can work in an interactively working mode, where the points of the curve can exactly be determined with the help of projections.

X-Y data is the 2D array of the x and y components of the curve.

interactive mode. FALSE represents “not interactive,” TRUE stands for “interactive.” The default value is FALSE.
**graph** is the graphical presentation of the curve completed by the coordinates of a selected point on the curve.

**Selected Point** is the coordinates of the selected point of the curve.

### Parametric Curve 3D

Depicts a curve in 3D under various conditions. You can control the position of the observer. This VI also lets you calculate the projections of a chosen point of the curve. This VI works as an interactive tool.

**X-Y-Z data** is the visualized 2D array of data of the curve. All points of the curve are given in the form \((x, y, z)\).

**View Info** is a cluster of data that describes the viewpoint of the observer.

- **Origin** is the point the observer is looking at. (See Figure 3-1 for an illustration.) The default value is \((0, 0, 0)\).
- **Axis** is the proportional factor of each axis. One can stretch or compress an axis independently of the other. The default value is \((2, 2, 2)\).
- **psi** describes the azimuth of the observer. This is the angle (in radians) of rotation about the \(z\) axis at the origin. When \(psi = 0\), the viewpoint is along the \(x\) axis looking away from \(x = 0\) toward the positive values of \(x\) and perpendicular to the \(y\) axis.
- **radius** describes the distance from the origin to the actual position of the observer.
- **phi** describes the elevation of the observer. This is the angle (in radians) made with the \(x\)-\(y\) plane by the line of sight from the observer to the origin. When \(phi = \pi/2\), the viewpoint is directly above the \(z\) axis.

**interactive mode**. FALSE represents “not interactive,” TRUE stands for “interactive.” The default value is FALSE.
**graph** is an array of clusters, each consisting of two arrays containing the $x$ and $y$ values of the graph data determined by **View Info**.

**Selected Point** is the coordinates of exactly one selected point of the graph. This point can interactively be determined. The projections of this point on the $x$, $y$ and $z$ axis are also presented.

**Final View Info** is a collection of data describing the viewpoint of the observer at the end of the interactive mode. The data in this cluster has the same structure as the data in the **View Info** cluster.

Figure 3-1 illustrates the meaning of observer and origin.
The following diagram shows the 3D curve \((\sin(t), \cos(t), t)\) from the view point of an outer observer. The projections are also shown.

**Read Graphs**
Reads graphs saved with the Write Graphs VI and displays the graph data.

- **path** is the path to the file containing the graph data.
- **Graphs** is the graphical representation of the saved graph data.
- **# of Graphs** is the number of saved graphs.
- **error**. See Appendix A, Error Codes, for a list of error codes. Common file errors can occur.

**Note:** See the description of the Write Graphs VI for details of the file format.
Write Graphs
Writes a collection of graphs to a file in ASCII format.

**Graphs** is an array of clusters, each consisting of two arrays. These arrays consist of the x and y values for the graphs you are writing to a file.

**path** is the path to the location where you intend to write the file. You can provide a specific path, or wire this input to the Open/Create/Replace File VI.

**# of Graphs** is number of graphs, which is determined in the **Graphs** cluster.

**error**. See Appendix A, *Error Codes*, for a list of error codes. Common file errors can also occur.

This VI produces a readable ASCII in a table with the following structure.

<table>
<thead>
<tr>
<th>Row</th>
<th>Content</th>
<th>Discussion</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Graphs</td>
<td>Identifies the table as a collection of graphs</td>
</tr>
<tr>
<td>2</td>
<td>Tab separated set of numbers, beginning with the number of graphs being written, followed by the number of points in each graph</td>
<td>Number of graphs and number of points in each of the graphs</td>
</tr>
<tr>
<td>3</td>
<td>The tab separated components of the first graph, beginning with the x components and followed by y components</td>
<td>Data of the first graph</td>
</tr>
<tr>
<td>4</td>
<td>The tab separated components of the second graph, beginning with the x components and followed by y components</td>
<td>Data of the second graph</td>
</tr>
<tr>
<td>...</td>
<td>...</td>
<td>...</td>
</tr>
</tbody>
</table>
Ordinary Differential Equation VIs

You can use these VIs to solve ordinary differential equations, both symbolically and numerically. They can be found in the ODE.LLB library.

Ordinary Differential Equation VI Descriptions

ODE Cash Karp 5th Order
The Cash Karp method solves ordinary differential equations with start conditions. The Cash Karp method works with an adaptive step rate and is computationally more efficient than the Euler method and the Runge Kutta method.

X is an array of strings of variables.

time start is the start point of the ODE. The default value is 0.

time end is the end point of the time interval under investigation. The default value is 1.0.

h is the step rate at the beginning. The default value is 0.1.

X0 is the vector of the start condition \( x_0, \ldots, x_n \). There is a one-to-one relation between the components of X0 and X.
accuracy controls the accuracy of the solutions. The default value is 0.0, which specifies the maximum deviation of the calculated solution from the actual solution.

time is the string denoting the time variable. The default variable is $t$.

$F(X,t)$ is a 1D array of strings representing the right sides of the differential equations.

Times is a 1D array representing the time steps. The ODE Cash Karp method yields arbitrarily chosen time steps between time start and time end.

$X$ Values is a 2D array of the solution vector $x_1, \ldots, x_n$. The top index runs over the time steps, the bottom index runs over the elements of $x_1, \ldots, x_n$.

ticks is the time in milliseconds for the whole calculation.

error. See Appendix A, Error Codes, for a list of error codes. Especially, the wrong inputs $X$, $X0$, and $F(X,t)$ can produce errors.

The Cash Karp method is an embedded Runge Kutta formula and is based on a fifth order strategy (with six steps).

\[
\begin{align*}
    k_1 &= h F(X(t_n), t_n) \\
    k_2 &= h F(X(t_n) + a_2 h, t_n + b_{21} k_1) \\
            & \quad \vdots \\
    k_6 &= h F(X(t_n) + a_6 h, t_n + b_{61} k_1 + \ldots + b_{65} k_5) \\
\end{align*}
\]

and

\[
X(t_{n+1}) = X(t_n) + c_1 k_1 + \ldots + c_6 k_6
\]

\[
X^*(t_{n+1}) = X(t_n) + c_1^* k_1 + \ldots + c_6^* k_6
\]

with
\[
t_{n+1} = t_n + h
\]

The
\[
a_2, \ldots, a_6, \quad b_{21}, \ldots, b_{65}, \quad c_1, \ldots, c_6, \quad c_1^*, \ldots, c_6^*
\]

are fixed real numbers. This choice determines the quality of the method.
The actual step size $h_{new}$ can be determined with the help of the accuracy value, the old step size $h$ and the difference

$$
\Delta = \left| X(t_{n+1}) - X^*(t_{n+1}) \right|
$$

according to

$$
h_{new} = h \left[ \frac{accuracy}{\Delta} \right]^{\frac{1}{5}}
$$

---

**Note:** It may happen that the value of the last element in the Times indicator turns out to be greater than the value entered in the time end control. This is a property of the Cash Karp method. This method is very accurate, but you have no control of the step rate. In order to guarantee that the end point specified in time end is taken into consideration, the last step may turn out to be too long.

---

**Example**

The following diagram shows the solution of the following system of ordinary differential equations in a 3D representation.

\[
\begin{align*}
\frac{dx(t)}{dt} &= 10(y(t) - x(t)) \\
\frac{dy(t)}{dt} &= x(t)(28 - z(t)) - y(t) \\
\frac{dz(t)}{dt} &= x(t)y(t) - \frac{8}{3}z(t)
\end{align*}
\]

$t \in [0, 40]$

$x(0) = 0.6$

$y(0) = 0.6$

$z(0) = 0.6$
The above equations and boundary conditions are entered in the VI as:

**time start:** 0.00
**time end:** 40.00
**X0:** [0.6, 0.6, 0.6]
**F(X,t):** [10*(y – x), x*(28 – z) – y, x*y – (8/3)*z]

**ODE Euler Method**
The Euler method solves ordinary differential equations with start conditions.
Chapter 4 Ordinary Differential Equation VIs

The general form of an ordinary differential equation (ODE) is

\[
\begin{align*}
\dot{x}_1(t) &= f_1(x_1(t), \ldots, x_n(t), t) \\
& \quad \vdots \\
\dot{x}_n(t) &= f_n(x_1(t), \ldots, x_n(t), t)
\end{align*}
\]

differential equations

with

\[
\begin{align*}
\begin{bmatrix}
x_1(t_0) \\
\vdots \\
x_n(t_0)
\end{bmatrix} = \begin{bmatrix}
x_{10} \\
\vdots \\
x_{n0}
\end{bmatrix}
\end{align*}
\]

starting conditions

The functions \( f_1, \ldots, f_n \), the numbers \( x_{10}, \ldots, x_{n0} \) and the start point \( t = t_0 \) are given. With the conventions

\[
F = (f_1, \ldots, f_n), \quad X(t) = (x_1(t), \ldots, x_n(t)) \quad \text{and} \quad X_0 = (x_{10}, \ldots, x_{n0})
\]

we have

\[
\begin{align*}
\dot{X}(t) &= F(X(t), t) \\
X(t_0) &= X_0
\end{align*}
\]

You have to determine functions \( X \) fulfilling the above equations.

- \( X \) is an array of strings of variables.
- \( \text{time start} \) is the start point of the ODE. The default value is 0.
- \( \text{time end} \) is the end point of the time interval under investigation. The default value is 1.0.
- \( h \) is the fixed step rate. The default value is 0.1.
Chapter 4  Ordinary Differential Equation VIs

\[ X_0 \text{ is the vector of the start condition } x_{10}, \ldots, x_{n0}. \text{ There is a one-to-one relation between the components of } X_0 \text{ and } X. \]

\[ \text{time} \text{ is the string denoting the time variable. The default variable is } t. \]

\[ F(X,t) \text{ is a 1D array of strings representing the right sides of the differential equations.} \]

\[ \text{Times} \text{ is an array representing the time steps. The Euler method yields equidistant time steps between } \text{time start} \text{ and } \text{time end}. \]

\[ X \text{ Values} \text{ is a 2D array of the solution vector } x_1, \ldots, x_n. \text{ The top index runs over the time steps, the bottom index runs over the elements of } x_1, \ldots, x_n. \]

\[ \text{ticks} \text{ is the time in milliseconds for the whole calculation.} \]

\[ \text{error. See Appendix A, Error Codes, for a list of error codes. Especially, the wrong values of the inputs } X, X_0, \text{ and } F(X,t) \text{ can produce errors.} \]

The Euler method is the most basic and often useful strategy to solve ODEs. Beginning with \( t_0 \) and a fixed step rate \( h \) (usually relatively small) the new values

\[ X(t_0 + h) = X(t_0) + hF(X(t_0), t) \]
\[ X(t_0 + 2h) = X(t_0 + h) + hF(X(t_0 + h), t_0 + h) \]
\[ \vdots \]

are calculated. This process stops, if \( \text{time start} + nh \geq \text{time end} \), where \( \text{time end} \) is the right endpoint of the time interval under investigation.

**Example**

The following diagram shows the solution of the following ordinary differential equation.

\[ \frac{dx(t)}{dt} = \sin(tx) + \sin(t + x) + \cos(t - x) \quad t \in [0, 20] \]
\[ x(0) = 1 \]
The above equation and initial condition are entered on the front panel as:

**time start:** 0.00

**time end:** 20.00

**X0:** 1.00

**F(X,t):** \( \sin(t^*x) + \text{sinc}(t + x) + \cos(t - x) \)

**ODE Linear nth Order Numeric**
Solves an \( n \)th order homogeneous linear differential equation with constant coefficients in numeric form.

- **A** in the matrix of coefficients of the different derivatives of a function \( x(t) \), starting with the coefficient of the lowest order term. The coefficient of the highest order derivative is assumed to be equal to 1.0 and does not need to be entered.

- **X0** is a vector of the start conditions of the \( n \)th order differential equation, starting with the start condition of the lowest order derivative.
number of points is the number of equidistant time points between time start and time end. The default value is 10.

time start is the start point of the nth order linear system. The default value is 0.0.

time end is the end point of the time interval under investigation. The default value is 1.0.

Times is an array representing the time steps. The method yields equidistant time steps between time start and time end.

X is the vector of the solution x at the equidistant time points.

error. See Appendix A, Error Codes, for a list of error codes. Especially, discrepancies between the length of X0 and A can produce errors.

Consider the nth order linear homogeneous differential equation

\[ x^{(n)} + a_{n-1}x^{(n-1)} + \ldots + a_1x^{(1)} + a_0x = 0 \]

with \( x(0) = x_0 \)

\[ x^{(1)}(0) = x_{10} \]

\[ \vdots \]

\[ x^{(n-1)}(0) = x_{n-10} \]

(0 represents the more general value of time start.) There is a strong connection between the equation

\[ x^{(n)} + a_{n-1}x^{(n-1)} + \ldots + a_1x^{(1)} + a_0x = 0 \]

and the zero finding problem

\[ z^n + a_{n-1}z^{n-1} + \ldots + a_1z + a_0 = 0 \]

The n zeroes of the last equation determine the structure of the solution of the ODE. If we have n distinct complex zeroes \( \lambda_1, \ldots, \lambda_n \), the general solution of the nth order differential equation can be expressed by

\[ x(t) = \beta_1 \exp(\lambda_1 t) + \ldots + \beta_n \exp(\lambda_n t) \]
The unknowns $\beta_1, \ldots, \beta_n$ can be determined by the start condition

\[
x(0) = \beta_1 + \ldots + \beta_n \\
x^{(1)}(0) = \beta_1\lambda_1 + \ldots + \beta_n\lambda_n \\
\vdots \\
\vdots \\
x^{(n-1)}(0) = \beta_1\lambda_1^{n-1} + \ldots + \beta_n\lambda_n^{n-1}
\]

Note: The case of repeated eigenvalues $\lambda_1, \ldots, \lambda_n$ is more complex and is not treated here. An error code of -23017 is given if this happens.

Note: By convention, the value of the highest coefficient is taken as 1.0, and does not need to be entered in the $A$ control. The other coefficients are entered starting with the lowest order coefficient.

Example

To solve the differential equation

\[x'' - 3x' + 2x = 0\]

with the I.C. as with $x(0) = 2$ and $x'(0) = 3$

enter


ODE Linear nth Order Symbolic

Solve an nth order homogeneous linear differential equation with constant coefficients in symbolic form.
Chapter 4  Ordinary Differential Equation VIs

A inputs the coefficients of the consecutive derivatives of a function \( x(t) \), starting with the coefficient of the lowest order term. The coefficient of the highest order derivative is assumed to be equal to 1.0 and does not need to be entered.

**X0** is a vector of the start conditions of the nth order differential equation, starting with the coefficient of the lowest order derivative.

**formula** is the symbolic solution.

**error**. See Appendix A, *Error Codes*, for a list of error codes. Discrepancies between the length of **X0** and **A** can produce errors.

The general solution has the following form (See the ODE Linear nth Order Numeric VI description in this chapter.):

\[
x(t) = \beta_1 \exp(\lambda_1 t) + \ldots + \beta_n \exp(\lambda_n t)
\]

with complex \( \beta_1, \ldots, \beta_n \) and \( \lambda_1, \ldots, \lambda_n \). But all inputs are real, and thus the solution also has this property. As a consequence, the symbolic solution is a linear combination of \( \exp-, \sin-, \) and \( \cos-\) functions with real coefficients.

**Note:** *Only the case of pairwise different \( \lambda_1, \ldots, \lambda_n \) is treated. For the case of repeated eigenvalues, an error code of \(-23017\) is given.*

**Note:** *By convention, the value of the highest coefficient is taken as 1.0, and does not need to be entered in the \( \Lambda \) control. The other coefficients are entered starting with the lowest order coefficient.*

**Example**

To solve the differential equation

\[
x'' - 3x' + 2x = 0
\]

with the I.C. as with \( x(0) = 2 \) and \( x'(0) = 3 \)

enter

\[
A = [2, -3] \text{ and } X0 = [2, 3]
\]
ODE Linear System Numeric

Solves an $n$-dimension homogeneous linear system of differential equations with constant coefficients, for a given start condition. The solution is based on the determination of the eigenvalues and eigenvectors of the underlying matrix $A$. The solution is given in numeric form.

$A$ is the $n$ by $n$ matrix describing the linear system.

$X0$ is the $n$ vector describing the start condition.

**number of points** is the number of equidistant time points between **time start** and **time end**. The default value is 10.

**time start** is the start point of the linear system, that is, the time with $X(time \ start) = X0$. The default value is 0.0.

**time end** is the end point of the time interval under investigation. The default value is 1.0.

**Times** is an array representing the time steps. The method yields equidistant time steps between **time start** and **time end**.

**X Values** is the matrix of the solution $X$ at the equidistant time points.

**error**. See Appendix A, Error Codes, for a list of error codes. Especially, discrepancies between the dimensions of $X0$ and $A$ can produce errors.

Linear systems can be described by

$$\frac{dX(t)}{dt} = AX(t)$$

$X(0) = X0$

if **time start** = 0.

Here $X(t) = (x_0(t), \ldots, x_n(t))$ and $A$ represents a $n$ by $n$ real matrix. The linear system can be solved by the determination of the eigenvalues and eigenvectors of $A$. Let $S$ be the
set of all eigenvectors spanning the whole \( n \)-dimensional space. The transformation

\[ Y(t) = SX(t) \]

yields

\[ \frac{dY(t)}{dt} = SAS^{-1}Y(t) \]

\[ Y(0) = SX_0 \]

The matrix \( SAS^{-1} \) has diagonal form, so that the solution is obvious. The solution \( X(t) \) can be determined by back-transformation \( X(t) = S^{-1}Y(t) \).

- **Note:** *No ticks output is implemented because the essential operation is the calculation of eigenvectors and eigenvalues of the matrix \( A \). This operation is negligible for relatively small dimensions of \( A \).*

- **Note:** *This VI works fine for almost all cases of real matrices \( A \) which can have repeated eigenvalues, conjugate complex eigenvalues, and so on. The exception is the case of a singular eigenvector matrix, that is, a matrix in which the eigenvectors do not span the entire space. An error of \(-23016\) is given if the eigenvector matrix is singular.*

**Example**

The following diagram shows the four components of the solution of the linear differential equation described by the following system:

\[
\begin{align*}
\frac{dx_1(t)}{dt} &= \begin{bmatrix} -7 & -6 & 4 & -1 \end{bmatrix} x_1(t) \\
\frac{dx_2(t)}{dt} &= \begin{bmatrix} -6 & 2 & 1 & -2 \end{bmatrix} x_2(t) \\
\frac{dx_3(t)}{dt} &= \begin{bmatrix} 4 & 1 & 0 & 2 \end{bmatrix} x_3(t) \\
\frac{dx_4(t)}{dt} &= \begin{bmatrix} -1 & -2 & 2 & -7 \end{bmatrix} x_4(t)
\end{align*}
\]

with

\[
\begin{align*}
x_1(0) &= 1 \\
x_2(0) &= 2 \\
x_3(0) &= 3 \\
x_4(0) &= 4
\end{align*}
\]
Enter the equations above on the front panel as shown:

\[ A: \begin{bmatrix} -7, -6, 4, 1; -6, 2, 1, -2; 4, 1, 0, 2; -1, -2, 2, -7 \end{bmatrix} \]

\[ X0: \begin{bmatrix} 1, 2, 3, 4 \end{bmatrix} \]

\[ \text{time start: 0.00} \]

\[ \text{time end: 1.00} \]

**ODE Linear System Symbolic**

Solves an \( n \)-dimension linear system of differential equations with a given start condition. The solution is based on the determination of the eigenvalues and eigenvectors of the underlying matrix. The solution is given in symbolic form.

\[ A \]

\[ X0 \]

\[ \text{formula} \]

\[ \text{error} \]

\[ \text{formula is a string with the solution of the linear system in the standard formula notation of LabVIEW. The solution vector elements are separated by carriage return.} \]

\[ A \text{ is the } n \text{ by } n \text{ matrix describing the linear system.} \]

\[ X0 \text{ is the } n \text{ vector describing the start condition.} \]

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**error.** See Appendix A, *Error Codes*, for a list of error codes. In particular, discrepancies between the dimensions of \( X0 \) and \( A \) can produce errors.

**Note:** This VI works fine for almost all cases of real matrices \( A \) which can have repeated eigenvalues, conjugate complex eigenvalues, and so on. The exception is the case of a singular eigenvector matrix, that is, a matrix in which the eigenvectors do not span the whole space. An error of -23016 is given if the eigenvector matrix is singular.

**Example**

The linear differential equation described by the following system:

\[
\begin{align*}
\frac{dx_1(t)}{dt} & = 7 - 6 x_1(t) - 4 x_2(t) - \frac{1}{2} x_3(t) - \frac{1}{2} x_4(t) \\
\frac{dx_2(t)}{dt} & = 6 x_1(t) + 2 x_2(t) + \frac{1}{2} x_3(t) + \frac{1}{2} x_4(t) \\
\frac{dx_3(t)}{dt} & = -2 x_1(t) + 4 x_2(t) - \frac{2}{7} x_3(t) - x_4(t) \\
\frac{dx_4(t)}{dt} & = -2 x_1(t) - 6 x_2(t) + \frac{1}{2} x_3(t) + \frac{1}{2} x_4(t)
\end{align*}
\]

with

\[
\begin{align*}
x_1(0) & = 1 \\
x_2(0) & = 2 \\
x_3(0) & = 3 \\
x_4(0) & = 4
\end{align*}
\]

has the solution

\[
\begin{align*}
+1.62\exp(-12.46t) & - 1.28\exp(-6.30t) + 0.63\exp(1.34t) + 0.04\exp(5.42t) \\
+0.84\exp(-12.46t) & - 0.29\exp(-6.30t) + 1.51\exp(1.34t) - 0.06\exp(5.42t) \\
-0.73\exp(-12.46t) & + 0.01\exp(-6.30t) + 3.69\exp(1.34t) + 0.02\exp(5.42t) \\
+0.87\exp(-12.46t) & + 2.67\exp(-6.30t) + 0.45\exp(1.34t) + 0.01\exp(5.42t)
\end{align*}
\]
Enter the equations above on the front panel as follows:

\[ A: \begin{bmatrix} -7, -6, 4, 1; -6, 2, -2; 4, 1, 0, 2; -1, -2, 2, -7 \end{bmatrix} \]

\[ X0: [1, 2, 3, 4] \]

**ODE Runge Kutta 4th Order**

The Runge Kutta method solves ordinary differential equations with start conditions. The Runge Kutta method works with a fixed step rate but with a higher degree of accuracy than the common Euler method.

- **X** is an array of strings of variables.
- **time start** is the start point of the ODE. The default value is 0.0.
- **time end** is the end point of the time interval under investigation. The default value is 1.0.
- **h** is the fixed step rate. The default value is 0.1.
Chapter 4  Ordinary Differential Equation VIs

**X0** is the vector of the start condition \( x_{10}, \ldots, x_{n0} \). There is a one-to-one relation between the components of **X0** and **X**.

**time** is the string denoting the time variable. The default variable is \( t \).

**F(X,t)** is an array of strings representing the right sides of the differential equations.

**Times** is an array representing the time steps. The Runge Kutta method yields equidistant time steps between **time start** and **time end**.

**X Values** is a 2D array of the solution vector \( x_1, \ldots, x_n \). The top index runs over the time steps, the bottom index runs over the elements of \( x_1, \ldots, x_n \).

**ticks** is the time in milliseconds for the whole calculation.

**error**. See Appendix A, *Error Codes*, for a list of error codes. Especially, wrong inputs **X**, **X0** and **F(X,t)** can produce errors.

The Runge Kutta method of 4th order works as a five stage process, more precisely

\[
\begin{align*}
k_1 &= hF(X(t_n), t_n) \\
k_2 &= hF\left(X(t_n) + \frac{k_1}{2}, t_n + \frac{h}{2}\right) \\
k_3 &= hF\left(X(t_n) + \frac{k_2}{2}, t_n + \frac{h}{2}\right) \\
k_4 &= hF\left(X(t_n) + k_3, t_n + h\right)
\end{align*}
\]

and

\[
X(t_{n+1}) = X(t_n) + \frac{k_1}{6} + \frac{k_2}{3} + \frac{k_3}{3} + \frac{k_4}{6}
\]

with \( t_{n+1} = t_n + h \)

The method ends, if \( t_n \geq \text{time end} \).
Example

The following diagram shows the solution of the following system of ordinary
differential equations:

\[
\begin{align*}
\frac{dx(t)}{dt} &= 10(y(t) - x(t)) \\
\frac{dy(t)}{dt} &= x(t)(28 - z(t)) - y(t) \\
\frac{dz(t)}{dt} &= x(t)y(t) - \frac{8}{3}z(t)
\end{align*}
\]

\[t \in [0, 50]\]

\[x(0) = 1\]
\[y(0) = 1\]
\[z(0) = 1.\]

Enter these equations on the front panel as shown:

time start: 0.00

time end: 50.00

X: [x, y, z]

X0: [1, 1, 1]

F(X,t): [10*(y - x), x*(28 - z) - y, x*y - (8/3)*z]
Chapter 4 Ordinary Differential Equation VIs

Note: Even though there are actually three solutions, a first glance at the graph almost seems to show only two solutions. This is because the solutions for x and y are very similar, so they almost overlap.
Zero Finder VIs

These VIs find the zeros of 1D or n-dimensional, linear or nonlinear functions (or system of functions). They are found in the library ZERO.LLB.

Zero Finder VI Descriptions

Find All Zeros of f(x)
Determines all zeros of a 1D function in a given interval.

- **accuracy** controls the accuracy of the determined zeros. The default value is 1E-8, which specifies the maximum deviation of the calculated solution from the actual solution.
- **step type**. A value of 0 represents a fixed number of function values, a value of 1 represents an optimal step size. In general, the second value leads to more accurate zeros.
- **algorithm**. Use 0 for the Ridders method; 1 for the Newton Raphson method. The default value is 0.
- **start** is the start point of the interval under investigation. The default value is 0.0.
- **end** is the end point of the interval. The default value is 1.0.
- **formula** is a string describing the function.
- **Zeros** are the determined zeros of **formula**.
**Example**

The following diagram shows the graph and the zeros of \( \sin(\text{sinc}(\gamma(x))) \) in the interval \((-2, 2)\). These values are entered on the front panel as:

- **start:** \(-2.00\)
- **end:** \(2.00\)
- **formula:** \(\sin(\text{sinc}(\gamma(x)))\)
Newton Raphson Zero Finder

Determines a zero of a 1D function close to two points with the help of the derivative of this 1D function. The two values form a search limit for the unknown zero of the 1D function.

**accuracy** controls the accuracy of the zero determination. The default value is 1E-8, which specifies the maximum deviation of the calculated solution from the actual solution.

**h** is the delta value to calculate the derivative of the given **formula**. The default value is 1E-8.

**start** is the first point close to the **zero** that the algorithm is trying to determine. The default value is 0.0.

**end** is the second point close to the **zero** that the algorithm is trying to determine. The default value is 1.0.

**formula** is a string representing the 1D function **zero**.

**f(zero)** is the function value at the point given by **zero**.

**zero** is only a good approximation for the exact value.

**ticks** is the time in milliseconds for the calculation.

**error**. See Appendix A, *Error Codes*, for a list of error codes.

Let \( f \) be the given function. Use a method that combines the simple midpoint strategy and the Newton strategy

midpoint strategy: \( x_{\text{new}} = \frac{x_1 + x_2}{2} \)

Newton strategy: \( x_{\text{new}} = x_1 - \frac{f(x_1)}{f'(x_1)} \)
where \( x_1 \) and \( x_2 \) are given guesses with \( f(x_1) \cdot f(x_2) < 0 \).

The following figure demonstrates the Newton strategy.

**Nonlinear System Single Solution**

Determines the solutions of nonlinear systems of equations in \( n \) dimensions beginning with a starting point in \( n \) dimensions.

- **accuracy** controls the accuracy of the determined zeros. The default value is 1E-8, which specifies the maximum deviation of the calculated solution from the actual solution.

- \( h \) is a small distance to calculate derivatives. The default value is 1E-8.

- **Start** is the start point in \( n \)D.
X is an array of strings defining the independent variables.

F(X) is an array of strings defining the functions in nD.

Zero is the determined zero of F(X).

F(Zero) are the function values of the Zero. Usually, these values are very close to 0.

ticks is the time in milliseconds to analyze the formula and to produce the Zero.

error. See Appendix A, Error Codes, for a list of error codes.

Let F be the n-dimension function and let X be a given point in n-dimensions.

Furthermore, let \( f = \frac{1}{2} F \cdot F \).

The algorithm is looking for such a vector \( P \) that \( F(X + dP) \leq F(X) \) for all \( 0 \leq d \leq 1 \). In a second step, an appropriate value \( d^* \) is calculated, so that \( F(X + d^*P) \) is considerably smaller than \( F(X) \). This process is repeated until \( F(X) = 0 \) is reached. What follows is an approximation for \( F(X) = 0 \).

**Nonlinear System Solver**

Determines a set of solutions of a nonlinear system of equations in n-dimensions beginning with a randomly chosen start point in n-dimensions.

accuracy controls the accuracy of the determined zeros. The default value is 1E-8.

number of trials is the elaborate number of randomly chosen start points. The algorithm starts with these points and is looking for zeros close to these points. The default value is 5.

h is a small distance to calculate derivatives. The default value is 1E-8.
**Start** is an array describing the left corner of the nD interval. The randomly chosen start points of the zero-finding algorithm can be found in the n-dimensional rectangle spanned by **Start** and **End**.

**End** is an array describing the right corner of the n-dimensional interval. The randomly chosen start points of the zero-finding algorithm can be found in the n-dimensional rectangle spanned by **Start** and **End**.

**X** is an array of strings defining the independent variables.

**F(X)** is an array of strings defining the functions in nD.

**Zeros** are the determined zeros of **F(X)**.

**F(Zeros)** are the function values of **zeros**. Usually, these values are very close to 0.

**ticks** is the time in milliseconds to analyze the formula and to produce the **zeros**.

**error**. See Appendix A, *Error Codes*, for a list of error codes.

**Example**

This algorithm is based on Nonlinear System Single Solution VI. This VI determines that the nonlinear system

\[
\begin{align*}
2x + 3y + z^2 - 6 &= 0 \\
-4x + y^2 + 4z + 7 &= 0 \\
x^2 + y + z - 3 &= 0
\end{align*}
\]

has two solutions

(1.0000, 1.0000, 1.0000) and (-0.4050, 0.5931, 2.2429).

The above equations, with appropriate **Start** and **End** values are entered into the VI front panel as:

**Start**: \([-1, -1, -1]\)

**End**: \([4, 4, 4]\)

**X**: \([x, y, z]\)

**F(X)**: \([2x + 3y + z^2 - 6, -4x + y^2 + 4z + 7, x^2 + y + z - 3]\)
Note: Only the left side of the above equations needs to be entered into $F(X)$. The VI assumes that the right side is zero.

**Polynomial Real Zero Counter**

Calculates the number of zeros of a given real polynomial in a real interval without determining of the values of these zeros.

- **start** is the leftmost point of the interval. The default value is 0.0.
- **end** is the rightmost point of the interval. The default value is 0.0.
- **Polynomial** is an array representing the polynomial under investigation. The first element of this array relates to the constant coefficient of the Polynomial.
- **number of zeros** is the exact number of zeros of the Polynomial in the interval (start,end).
- **error**. See Appendix A, Error Codes, for a list of error codes. When start > end, the application interprets it as an error situation.

The Sturm algorithm has to deal with two situations. Let $p(x)$ be a real polynomial in $x$, and let $p'(x)$ be the derivative of $p(x)$. If $d(x)$ denotes the greatest common divisor of $p(x)$ and $p'(x)$, $d(x) = \gcd(p(x), p'(x))$, so $p(x)$ has multiple zeros, if and only if $d(x)$ is a nonconstant polynomial. In other words, the polynomial $p(x)/d(x)$ has only single zeros. Repeating this idea, you can combine the given polynomial $p(x)$ as the product of simple polynomials, each of them with single real zeros. The number of zeros of $p(x)$ is equal to the sum of all zeros of the defined simple polynomials having only single zeros.
If you need to determine the number of zeros of a real polynomial \( p(x) \) with the single zero property, use the following Euclidean Algorithm.

\[
\begin{align*}
p(x) &= q_1(x)p_1(x) - p_2(x) \\
p_1(x) &= q_2(x)p_2(x) - p_3(x) \\
&\vdots \\
p_{r-2}(x) &= q_{r-1}(x)p_{r-1}(x) - p_r(x) \\
p_{r-1}(x) &= q_r(x)p_r(x)
\end{align*}
\]

where \( p_1(x) = p'(x) \)

The Sturm’s chain \((p(x), p_1(x), \ldots, p_r(x))\) determines two values \( W(\text{start}) \) and \( W(\text{end}) \). \( W(x) \) is the number of sign changes of the chain \((p(x), p_1(x), \ldots, p_r(x))\). The number of all zeros of \( p(x) \) is exactly equal to \( W(\text{end}) - W(\text{start}) \).

**Note:** The algorithm makes extensive use of polynomial operations like multiplication, division, and greatest common divisor (gcd). You can select these subVIs and reuse them in your own projects.

**Note:** If you are interested in all real zeros of a real polynomial, the choices \( \text{start} = -(|a_0| + \ldots + |a_n|) \) and \( \text{end} = +(|a_0| + \ldots + |a_n|) \), where \( a_i \) denotes the coefficients, are sufficient to determine this number.

**Ridders Zero Finder**

Determines a zero of a 1D function in a given interval. The function has to be continuous and has to have different signs at the end points of the interval.

**accuracy** controls the accuracy of the zero determination. The default value is 1E-8.

**start** is the leftmost point of the given interval. The default value is 0.0.

**end** is the rightmost point of the given interval. The default value is 1.0.
formula is a string representing the 1D function.

zero is the determined zero of formula.

f(zero) is the function value at the point zero. Usually, the value of zero is only a good approximation for the exact value.

ticks is the time in milliseconds for the whole calculation.

error. See Appendix A, Error Codes, for a list of error codes. When start > end, the application interprets it as an error condition. The function values at the points start and end must have different signs to guarantee the existence of a zero in (start,end).

Given the function f(x) with \( f(a)f(b) < 0 \).

Ridders method determines \( c = (a + b)/2 \) and calculates the new guess

\[
c_{\text{new}} = c \pm \frac{\text{sign}(f(a) - f(b))f(c)}{\sqrt{f(c)^2 - f(a)f(b)}}
\]

The triplets start, \( c_{\text{new}} \), end are the base for the new iteration, depending on whether \( f(\text{start})f(c_{\text{new}}) < 0 \) or \( f(c_{\text{new}})f(\text{end}) < 0 \). The algorithm stops, if \( |a - b| < \text{accuracy} \).

Ridders method is very fast and reliable.
Ridders method is a generalization of the depicted estimation of a zero, as shown in the following illustration.

\[
x_0 = \frac{x_1 f(x_2) - x_2 f(x_1)}{f(x_2) - f(x_1)}
\]
Optimization VIs

Use these VIs to determine local minima and maxima of real 1D or $n$-dimension functions. You can choose between optimization algorithms based on derivatives of the function and algorithms working without these derivatives. You can also use special methods like Linear Programming, Levenberg Marquardt in symbolic form, Pade, and Chebyshev Approximation. An overview of the optimization routines is shown in the following illustration.
Brent with Derivatives 1D

Determines a local minimum of a given 1D function in a given interval.

- **accuracy** controls the accuracy of the determined minimum of **formula**. The method stops, if two consecutive approximations differ not more than the value of **accuracy**.
- **a (start)** is the left point of the bracketing interval. The default value is 0.0.
- **b (start)** is the middle point of the bracketing interval. The default value is 0.0.
- **c (start)** is the right point of the bracketing interval. The default value is 0.0.
- **formula** is a string describing the function under investigation. The Parser VIs check the syntax of this string.
- **minimum** is the determined local minimum of **formula**.
- **f(minimum)** is the function value of the determined local minimum.
- **ticks** is the time in milliseconds for the whole calculation.
- **error**. See Appendix A, *Error Codes*, for a list of error codes.

Refer to the Golden Section 1D VI description for the definition of a bracketing interval. Here the derivative of **formula** is used. This leads to a faster algorithm, because the new bracketing interval has better properties than those of the Golden Section 1D VI.
Chebyshev Approximation
Determines a given function using Chebyshev polynomials.

**number of points** is the number of equidistant points in the interval \((\text{start}, \text{end})\). The default value is 10.

**start** is the start point of the interval. The default value is 0.0.

**end** is the end point of the interval. The default value is 1.0.

**order** is the degree of the Chebyshev approximation (that is, the number of different Chebyshev polynomials \(T_0(x), T_1(x), \ldots, T_n(x)\) describing the formula). The default value is 3.

**formula** is a string describing the function under investigation. The Parser VIs check the syntax of this string.

**C** is an array of coefficients belonging to \(T_0(x), T_1(x), \ldots\).

**X** is the x values dividing \((\text{start}, \text{end})\) in equidistant subintervals.

**Y** is the y values of the Chebyshev polynomial at points X.

**error**. See Appendix A, *Error Codes*, for a list of error codes.

Let \(n\) be a given natural number. The function \(f(x)\) can approximately be represented by

\[
f(x) = c_0 T_0(x) + \ldots + c_n T_n(x)
\]

where \(T_0(x), \ldots, T_n(x)\) are the first Chebyshev polynomials.

The \(c_0, \ldots, c_n\) can be calculated as sums of the form

\[
\sum_{k=1}^{n} f(x_k) T_j(x_k)
\]

where \(x_k = \cos\left(\frac{\pi k}{n}\right)\) for \(k = 1, \ldots, n\).
Downhill Simplex nD

Determines a local minimum of a function of \( n \) independent variables with the Downhill Simplex method.

- **accuracy** controls the accuracy of the minimum. The method stops if two consecutive approximations differ no more than the value of accuracy. The default value is 1E-8.

- **Start** is an array of points at which the optimization process is starting. These points form a simplex in \( n \)-dimension.

- **X** is an array of strings representing the x variables.

- **f(X)** is the string representing the function of the \( X \) variables.

- **Minimum** is the determined local minimum in \( n \)-dimension.

- **f(Minimum)** is the function value of \( f(X) \) at the determined minimum.

- **ticks** is the time in milliseconds for the whole calculation.

- **error**. See Appendix A, Error Codes, for a list of error codes. The Parser VIs check the syntax of the inputs.

The Downhill Simplex algorithm, also called the Nelder and Mead method, works without partial derivatives. The Downhill Simplex algorithm consists of catching the minimum of the function, \( f(X) \), with the help of simple geometrical bodies, specifically a simplex. A simplex in 2D is a triangle, a simplex in 3D is a tetrahedron and so on. You must have \((n + 1)\) starting points, each of dimension \( n \), forming the initial simplex. The user must enter only one point of these \((n + 1)\). The \((n + 1)\) dimensional simplex is automatically constructed. For the example given below \((f(x,y) = x^2 + y^2)\), you must enter two numbers (describing exactly one point in 2D). The algorithm generates a new simplex by some elementary operations like reflections, expansions, and contractions. In the end, the minimum is concentrated in a very small simplex.
Example

The simplex sequence tending to the minimum (0,0) of the function \( f(x, y) = x^2 + y^2 \) is shown in the following diagram. The function is entered on the front panel as:

Start: \([3.2, 1]\)

\[ X: \begin{bmatrix} x, y \end{bmatrix} \]

\[ f(X): \begin{bmatrix} x^2 + y^2 \end{bmatrix} \]

Conjugate Gradient nD

Determines a local minimum of a function of \( n \) independent variables with the Conjugate Gradient method.

\[ \text{accuracy} \] controls the accuracy of the minimum. The method stops if two consecutive approximations differ no more than the value of \( \text{accuracy} \). The default value is 1E-8.

\[ \text{gradient method} \]. A value of 0 represents the Fletcher Reeves method, a value of 1 represents the Polak Ribiere method. The default value is 0.
line minimization. A value of 0 represents an algorithm without usage of the derivatives, a value of 1 represents an algorithm with usage of the derivatives. The default value is 0.

Start is a point in n-dimension at which the optimization process starts.

X is an array of strings representing the X variables.

f(X) is the string representing the function of the X variables.

Minimum is the determined local minimum in n-dimension.

f(Minimum) is the function value of f(X) at the determined minimum.

ticks is the time in milliseconds for the whole calculation.

error. See Appendix A, Error Codes, for a list of error codes. The Parser VIs check the syntax of the inputs.

The Fletcher Reeves and the Polak Ribiere algorithm are based on the determination of best-suited directions and 1D subminimizations.

The following diagram shows a start point and a start direction. New points and new directions are calculated by the Conjugate Gradient nD VI.
Find All Minima 1D
Determines all local minima of a given function in a given interval.

**accuracy** controls the accuracy of the values of **Minima**. The method stops if two consecutive approximations differ no more than the value of **accuracy**. The default value is 1E-8.

**step type**. A value of 0 represents a fixed number of function values, a value of 1 represents an optimal step rate. In general, the second value leads to more accurate **Minima**. The default value is 0.

**algorithm**. A value of 0 represents the Golden Section method, a value of 1 represents the Brent method. The default value is 0.0.

**start** is the start point of the interval. The default value is 0.0.

**end** is the end point of the interval. The default value is 1.0.

**formula** is a string representing the function under investigation.

**Minima** is an array of all found minima of **formula** in the interval 
(start, end).

**f(Minima)** is the function values at the points **Minima**.

**ticks** is the time in milliseconds for the whole calculation.

**error**. See Appendix A, Error Codes, for a list of error codes.

**Note:** If you want to find out the maxima of a function, you must take ~function as the inputs. The ~f(Minima) are the correct maximal function values.
Example

Find all Minima 1D of \( f(x) = \cos(x^2) \). All minima are determined. The following diagram shows the plot of \( f(x) \). The boxes on the plot are the locations of the minima. The inputs are entered on the front panel as:

- **start:** \(-1.0\)
- **end:** \(6.0\)
- **formula:** \(\cos(x^2)\)

Find All Minima nD

Determines the minima of an \( n \)-dimension function in a given \( n \)-dimension interval. The **accuracy** controls the accuracy of the minima. The default value is \(1E-8\).
algorithm. A value of 0 represents the Conjugate Gradient method, a value of 1 represents the Downhill Simplex method. The default value is 0.

gradient method. A value of 0 represents the Fletcher Reeves method, a value of 1 represents the Polak Ribiere method. The default value is 0.

line minimum. A value of 0 represents the line optimization without derivatives, a value of 1 represents the line optimization with derivatives. The default value is 0.

number of trials is the number of the randomly chosen start points of the optimization process. These points belong to the interval \( (\text{start}, \text{end}) \). The default value is 5.

Start is the start point in \( n \)-dimension.

End is the end point in \( n \)-dimension.

\( X \) is an array of strings describing the \( n \) variables.

\( F(X) \) is a string describing the \( n \)-dimension function of \( X \).

\( X \) Values is a matrix describing all determined local minima.

\( F \) Values is the function values at the points \( X \) Values.

ticks is the time in milliseconds for the whole calculation.

error. See Appendix A, Error Codes, for a list of error codes.

Fitting on a Sphere

The algorithm determines the best spherical fit on a cloud of points in 3D.

\( X \) is the \( x \) coordinates of the points of the cloud.

\( Y \) is the \( y \) coordinates of the points of the cloud.

\( Z \) is the \( z \) coordinates of the points of the cloud.

\( x_0, y_0, z_0 \) are the calculated midpoints of the given cloud.
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Chapter 6  Optimization VIs

**r** is the calculated radius of the given cloud.

**error.** See Appendix A, *Error Codes*, for a list of error codes. Most error situations result from discrepancies between the sizes of X, Y, and Z.

The min-functional is based on an uncommon idea:
find \( x_0, y_0, z_0 \) and \( r \) with

\[
\sum_{i=1}^{n} \left( (x_i - x_0)^2 + (y_i - y_0)^2 + (z_i - z_0)^2 - r^2 \right)^2 = \text{min!}
\]

This leads to a simple linear equation in \( x_0, y_0, \) and \( z_0 \).

**Golden Section 1D**
Determine a local minimum of a given 1D function with the help of a bracketing of the minimum. The Golden Section Search method is used.

**accuracy** controls the accuracy of the determined minimum of **formula.** The method stops, if two consecutive approximations differ not more than the value of **accuracy.** The default value is 1E-8.

**a (start)** is the left point of the bracketing interval. The default value is 0.0.

**b (start)** is the middle point of the bracketing interval. The default value is 0.0.

**c (start)** is the right point of the bracketing interval. The default value is 0.0.

**formula** is a string describing the function under investigation. The Parser VIs check the syntax of this string.

**minimum** is the determined local minimum of **formula.**
**f(minimum)** is the function value at the determined local minimum.

**ticks** is the time in milliseconds for the whole calculation.

**error.** See Appendix A, *Error Codes*, for a list of error codes.

A bracketing triplet \((a, b, c)\) of a 1D continuous function \(f\) is a combination of three points with \(f(a) > f(b)\) and \(f(c) > f(b)\). This guarantees the existence of a local minimum of \(f\) in the interval \((a, c)\).

The Golden Section Search method determines beginning with a bracketing triplet \((a, b, c)\) a new one with a considerably smaller expansion. Repeating this scheme often yields a good approximation of the local minimum. The new bracketing point is essentially calculated by the following equation.

\[
\frac{x - b}{c - a} = (\sqrt{5} - 2) \quad \text{(Golden Section Search Method)}
\]

The following diagram shows the relationship between \(a\), \(b\), \(c\) and \(f(a)\), \(f(b)\), \(f(c)\).
Levenberg Marquardt

Uses the Levenberg Marquardt method to determine a nonlinear set of coefficients that minimize a chi2 quantity.

- **Standard Deviation** is the standard deviation for data points. If they are equal or you do not know, leave this array empty. Internally, LabVIEW sets all data points to 1.0.

- **X** is the input array. The number of valid input points must be greater than zero and greater than the number of specified coefficients. The number of elements in X must be equal to the number of elements in Y.

- **Y** is the input array. The number of valid input points must be greater than zero and greater than the number of specified coefficients. The number of elements in Y must be equal to the number of elements in X.

- **Initial Guess Coefficient** denotes your initial-guessed solution.

- **max iteration** is the maximum executing iteration. If the VI reaches maximum iteration without finding a solution, the function returns an error. You have to increase the max iteration or adjust the Initial Guess Coefficient to get a solution. The default value is 200.

- **model description** is a cluster containing the fitting equation.
  - **model** is a string describing the model equation.
  - **Parameters** is an array of strings of the unknown parameters.
  - **x** is a string describing the independent variable.

- **Covariance** is the matrix of covariances.

- **Best Fit Coefficients** is the set of coefficients that minimize the penalty (chi-squared) function. The chi-squared function is given by
In this equation, \((x_i, y_i)\) are the input data points, and \(f(x_i; a_1, ..., a_M) = f(X, A)\) is the nonlinear function where \(a_1, ..., a_M\) are coefficients. If the measurement errors are independent and normally distributed with constant, standard deviation \(\sigma_i = \sigma\), this is also the least-square estimation.

The input arrays \(X\) and \(Y\) define the set of input data points. The VI assumes that you have prior knowledge of the nonlinear relationship between the \(x\) and \(y\) coordinates. That is, \(f = f(X, A)\) where the set of coefficients, \(A\), is determined by the Levenberg Marquardt algorithm.

Using this function successfully depends on how close your initial guess coefficients are to the solution. Therefore, it is always worth taking the time and effort to obtain good initial guess coefficients to the solution from any available resources before using the function.

**Best Fit** is the fitted data.

**mse** is the chi2 value.

**ticks** is the time in milliseconds for the whole calculation.

**error**. See Appendix A, Error Codes, for a list of error codes.

More information on the Levenberg Marquardt method can be found in the Non-Linear Lev-Mar Fit VI description in the LabVIEW Analysis VI Reference Manual.

**Linear Programming Simplex Method**

 Determines the solution of a linear programming problem.

\[
\chi^2 = \sum_{i=0}^{N-1} \left( \frac{y_i - f(x_i; a_1, ..., a_M)}{\sigma_i} \right)^2
\]

C is a vector describing the linear functional to maximize.

\(M\) is a matrix describing the different constraints.

\(B\) is a vector describing the right sides of the constraints inequalities.
maximum is the maximal value of a \( x \) under the constraints.

\( X \) is the solution vector.

ticks is the time in milliseconds for the whole calculation.

error. See Appendix A, Error Codes, for a list of error codes. The nonexistence of a solution \( x \) leads to an error.

The optimization problem

\[ cx = \text{max!} \]

with the constraints

\[ x \geq 0 \quad \text{and} \quad mx \geq b. \]

Here \( X = (x_1, \ldots, x_n) \), \( C = (c_1, \ldots, c_n) \), \( B = (b_1, \ldots, b_k) \) and \( M \) a \( k \) by \( n \) matrix. Now you must decide whether or not an optimal vector \( x \) does exist, and if so, determine this vector \( x \). The solution of a linear programming problem is a two-step process. The first step transforms the original problem into a problem in restricted normal form (essentially without inequalities in the formulation). The second step consists of the solution of this restricted normal form problem.

Note: The previous formulation seems to be special. But there are many ways to reformulate terms. For instance, \( dx \leq e \) is equivalent to \( -dx \geq -e \), and, \( dx = e \) is equivalent to the combination \( dx \geq e \) and \( -dx \geq -e \).

**Pade Approximation**

Determines the coefficients of a rational polynomial to best suit a given set of first derivatives.

\[ m \]
\[ n \]
\[ C[0..m+n] \]
\[ A[0..m] \]
\[ B[0..n] \]
\[ \text{error} \]

\( m \) is the degree of the polynomial of the numerator.

\( n \) is the degree of the polynomial of the denominator.

\( C[0..m+n] \) is the array describing the first derivatives of the given function.

\( A[0..m] \) is the polynomial of the numerator.
Let $f$ be a given function with known values $f(0), f'(0), \ldots, f^{(n+m)}(0)$. There is a unique rational polynomial ($m \geq n$)

$$R(x) = \frac{a_0 + a_1x + \ldots + a_mx^m}{1 + b_1x + \ldots + b_nx^n}$$

with $R(0) = f(0), R'(0) = f'(0), \ldots, R^{(m+n)}(0) = f^{(m+n)}(0)$.

The rational polynomial can be determined by solving a special linear equation.
1D Explorer VIs

You can use this collection of VIs to study real-valued 1D functions given in symbolic form. You can study different qualities of function graphs with and without additional parameters. You can calculate and derive characteristic properties of these curves, such as length, integration, differentiation, extrema and zeroes. You can connect the X Values and Y Values outputs of the VIs to a suitable graph indicator for a visual representation of the waveform. These VIs are available in the 1DEXPLO.LLB library.

Notice that some of the VIs are named with the words Optimal Step. For a function that has many maxima, minima, or singularities, the standard method (using equidistant points) may yield incorrect results. In such cases the Optimal Step method is better. This method starts off by taking equidistant points, but also checks the steepness of the curve between these points. In very steep portions of the graph (determined by the value of epsilon on the front panel), a new point is generated in between. As a rule, the smaller the value of epsilon, the more the points that are generated, and the better the graph.

1D Explorer VI Descriptions

Curve Length
Calculates the curve length of a 1D function between start and end.

start is the start point of the interval under investigation. The default value is 0.0.

end is the end point of the interval. The default value is 1.0.
**Chapter 7  1D Explorer VIs**

**formula** is a string describing the function under investigation. You can use any valid symbol as the variable name. See Chapter 1, *Parser VIs*, in this manual.

**X Values** is the array of all regarded points in the interval \((\text{start}, \text{end})\).

**Curve Length** is an array of the values of the curve length of **formula** between **start** and **end** at all **X Values**.

**ticks** is the time in milliseconds to analyze the formula and to produce the **X Values** array and the **Curve Length** array.

**error**. See Appendix A, *Error Codes*, for a list of error codes. The accuracy of formula is verified by the Parser VIs.

The curve length of a given function \(f(t)\) between **start** and **end** can be calculated by the following equation.

\[
L = \int_{\text{start}}^{\text{end}} \sqrt{1 + \left(\frac{df(t)}{dt}\right)^2} \, dt
\]

This calculation is realized using the Integration VI, found in the **1DEXPLO.LLB** library, and that is why the calculations are based on the Runge Kutta method.

**Differentiation**

Calculates both function values and the values of the derivative of a given 1D function defined by a formula at equidistant points in an interval.

**number of points** is the number of all calculated points. The default value is 10.

**start** is the start point of the interval under investigation. The default value is 0.0.

**end** is the end point of the interval. The default value is 1.0.
**formula** is a string describing the function under investigation. You can use any valid symbol as the variable name. See Chapter 1, *Parser VIs*, in this manual.

**X Values** is the array of all equidistantly positioned points in the interval (start, end).

**Y Values** are the values of the function.

**Derivative of Y** are the values of the derivative of the function at the points X Values.

**ticks** is the time in milliseconds to analyze the formula and to produce the X Values array, Y Values array and Derivative of Y array.

**error.** See Appendix A, *Error Codes*, for a list of error codes. The accuracy of formula is verified by the Parser VIs.

**Note:** *Even though there is a modified method (Optimal Step) for functions, the G Math Toolkit does not have one for the Differentiation VI. If you are interested in highly accurate values of Differentiation, start with the symbolic differentiation (by hand). Then use the Eval \( y = f(x) \) Optimal Step VI, where \( f \) is the derivative of the function.*

The function and the derivative of \( f(x) = \sin(\text{sinc}(x)) \) are investigated in the interval \((-20, 20)\). The following diagram shows both \( f(x) \) and \( f'(x) \).
Eval Polar to Rect
Calculates the values of a polar parametric curve in 2D.

**number of points** is the number of all calculated points. The angle variable is split into equidistant subpoints. The default value is 10.

**start** is the start point of the interval under investigation. The default value is 0.0.

**end** is the end point of the interval. The default value is 1.0.

**radius as function of angle** is a string representing the formula, \( r = r(\phi) \).

**X** is the array of the values of the first component, \( r(\phi) \cdot \cos(\phi) \).

**Y** is the array of the values of the second component, \( r(\phi) \cdot \sin(\phi) \).

**ticks** is the time in milliseconds to analyze the formula and to produce the **X** and the **Y** array.

**error**. See Appendix A, Error Codes, for a list of error codes. The accuracy of **radius as function of angle** is checked with the help of some of the Parser VIs.

Let \( \phi \) be the angle and \( r(\phi) \) the radius in polar coordinate notation. Then it is

\[
x = r(\phi) \cos \phi
\]

\[
y = r(\phi) \sin \phi
\]

where \( \phi \) runs over the specified interval.
Example

The function \( r(t) = \exp(\cos(t)) - (2\cos(4t)) + \sin(t/12)^5 \) describes a butterfly curve in the plane in polar coordinates, as shown in the following diagram. The diagram can be generated by entering the following values on the front panel:

- number of points: 1000
- start: 0
- end: 30
- radius as function of angle: \( \exp(\cos(t)) - 2\cos(4t) + \sin(t/12)^5 \)

Eval Polar to Rect Optimal Step

Operates like the Eval Polar to Rect VI, but with a significantly higher degree of accuracy.

number of points is the number of calculated points at the beginning of the execution. The default value is 10. Usually, many more points will be added.
**Chapter 7  1D Explorer VIs**

**epsilon** controls the construction of points inside the construction area. The default value is 0.05.

**start** is the start point of the interval under investigation. The default value is 0.0.

**end** is the end point of the interval. The default value is 1.0.

**radius as function of angle** is a string representing the formula $r = r(\phi)$.

$X$ is the array of the values of the first component, $r(\phi)\cos(\phi)$.

$Y$ is the array of the values of the second component, $r(\phi)\sin(\phi)$.

**ticks** is the time in milliseconds to analyze the formula and to produce the $X$ and the $Y$ array.

**error**. See Appendix A, *Error Codes*, for a list of error codes. The accuracy of **radius as function of angle** is checked with the help of some of the Parser VIs.

**Eval X-Y(a,t)**
A generalized version of the Eval X-Y(t) VI with the possibility of adding some parameters into the formula.

**number of points** is the number of all calculated points. The independent variable is split into equidistant subpoints. The default value is 10.

**start** is the start point of the interval under investigation. The default value is 0.0.

**end** is the end point of the interval. The default value is 1.0.
Parameters is an array of clusters describing the parameters.

- name of the parameter which uses the conventions of the Parser VIs.
- value of the parameter.

Formulas is an array of two strings describing the two function components. You can use any valid symbol as the variable name. See Chapter 1, Parser VIs, in this manual.

- X is the array of the values of the first component.
- Y is the array of the values of the second component.

ticks is the time in milliseconds to analyze the Formulas and to produce the X and the Y array.

error. See Appendix A, Error Codes, for a list of error codes. The accuracy of Formulas and Parameters is checked with the help of some of the Parser VIs.

Eval X-Y(t)
Calculates the values of a function \((f(t), g(t))\), where \(t\) runs over an interval. Both components are given by Formulas. The \(t\)-values are chosen equidistantly in the interval.

- number of points is the number of all calculated points. The independent variable is split into equidistant subpoints. The default value is 10.
- start is the start point of the interval under investigation. The default value is 0.0.
- end is the end point of the interval. The default value is 1.0.
- Formulas is an array of two strings describing the two formula components. You can use any valid symbol as the variable name. See Chapter 1, Parser VIs, in this manual.
- X is the array of the values of the first component.
Y is the array of the values of the second component.

ticks is the time in milliseconds to analyze the formula and to produce the X and the Y array.

error. See Appendix A, Error Codes, for a list of error codes.
The accuracy of Formulas is checked with the help of some of the Parser VIs.

Example

The functions \((t\times\sin(t),\sqrt{t}\times\cos(t))\) describe a curve in the plane, with \(t\) ranging between \((0, 12)\). The following diagram shows this curve.

Enter the data on the front panel as shown:

number of points: 100

start: 0

end: 12

Formulas: \((t\times\sin(t), \sqrt{t}\times\cos(t))\)
Eval X-Y(t) Optimal Step
Calculates the values of a more complex function \((f(t), g(t))\), where \(t\) runs over an interval. Both components are given by Formulas.

**Formulas** is an array of two strings describing the two function components. You can use any valid symbol as the variable name. See Chapter 1, Parser VIs, in this manual.

- **number of points** is the number of calculated points at the beginning of the execution. The default value is 10. Usually, many more points will be added.
- **epsilon** controls the construction of points in between. The default value is 0.05.
- **start** is the start point of the interval under investigation. The default value is 0.0.
- **end** is the end point of the interval. The default value is 1.0.
- **X** is the array of the values of the first component.
- **Y** is the array of the values of the second component.
- **ticks** is the time in milliseconds to analyze the formula and to produce the **X** and the **Y** array.
- **error**. See Appendix A, Error Codes, for a list of error codes. The accuracy of **Formulas** is checked with the help of some of the Parser VIs.
Eval $y=f(a,x)$

A generalized version of the Eval $y=f(x)$ VI, with the possibility of adding some parameters into the formula.

- **number of points** is the number of all calculated points. The independent variable is split into equidistant subpoints. The default value is 10.

- **start** is the start point of the interval under investigation. The default value is 0.0.

- **end** is the end point of the interval. The default value is 1.0.

- **Parameters** is an array of clusters describing the parameters.
  - **name** of the parameter which uses the conventions of the Parser VIs.
  - **value** of the parameter.

- **formula** is a string describing the function under investigation. You can use any valid symbol as the variable name. See Chapter 1, *Parser VIs*, in this manual.

- **X** is the array of equidistant points between **start** and **end**.

- **Y** is the function values at the points **X**.

- **ticks** is the time in milliseconds to analyze the formula and to produce the **X** and the **Y** array.

- **error**. See Appendix A, *Error Codes*, for a list of error codes. The accuracy of **formula** and **Parameters** is checked with the help of some of the Parser VIs.
**Eval y=f(x)**

Calculates the values of a 1D function given by a formula at equidistant points in an interval.

**number of points** is the number of all calculated points. The independent variable is split into equidistant subpoints. The default value is 10.

**start** is the start point of the interval under investigation. The default value is 0.0.

**end** is the end point of the interval. The default value is 1.0.

**formula** is a string describing the function under investigation. You can use any valid symbol as the variable name. See Chapter 1, *Parser VIs*, in this manual.

**X Values** is the array of equidistant points between **start** and **end**.

**Y Values** is the array of function values from the corresponding points of **X Values**.

**ticks** is the time in milliseconds to analyze the formula and to produce the **X Values** array and the **Y Values** array.

**error**. See Appendix A, *Error Codes*, for a list of error codes. The accuracy of **formula** is verified by the Parser VIs.

**Note:** *You can directly connect the (X Values, Y Values) array to a graph indicator to see the result of Eval y=f(x) VI.*
Eval \( y=f(x) \) Optimal Step

Calculates the values of a more complex 1D function given by a formula.

- **number of points** is the number of calculated points at the beginning of the execution. The default value is 10. Usually, many more points will be added.

- **epsilon** controls the construction of points in the sense of optimal steps. The default value is 0.05. The smaller **epsilon** the more values will be produced.

- **start** is the start point of the interval under investigation. The default value is 0.0.

- **end** is the end point of the interval. The default value is 1.0.

- **formula** is a string describing the function under investigation. You can use any valid symbol as the variable name. See Chapter 1, Parser VIs, in this manual.

**X Values** is the array of all points in the interval \((\text{start}, \text{end})\). In particular, all start points produced by **number of points, start** and **end** belong to the array, but many further points belong too.

**Y Values** are the function values at the points **X Values**.

**ticks** is the time in milliseconds to analyze the formula and to produce the **X Values** and the **Y Values** array.

**error**. See Appendix A, Error Codes, for a list of error codes. The accuracy of **formula** is verified by the Parser VIs.
Integration
Calculates both the function values and the integral of a 1D function between start and end.

start is the start point of the interval under investigation. The default value is 0.0.

end is the end point of the interval. The default value is 1.0.

formula is a string describing the function under investigation. You can use any valid symbol as the variable name. See Chapter 1, Parser VIs, in this manual.

X Values is the array of all regarded points in the interval (start, end).

Y Values are the values of the function.

Integral of Y are the values of the integral of formula between start and end at all X Values values.

ticks is the time in milliseconds to analyze the formula and to produce the X Values array and the Integral of Y array.

error. See Appendix A, Error Codes, for a list of error codes. The accuracy of formula is verified by the Parser VIs.

Let \( f(t) \) be the given function. The calculation of

\[
I = \int_{\text{start}}^{\text{end}} f(t) \, dt
\]

can be reformulated as an ordinary differential equation

\[
\frac{dI(s)}{ds} = f(s)
\]

\( I(\text{start}) = 0 \)
Chapter 7  1D Explorer VIs

The algorithm is based on this reformulation. The Runge Kutta method is used for accuracy. See the ODE Runge Kutta 4th Order VI description in Chapter 4, Ordinary Differential Equation VIs, for more information.

Limit
Determine the left and right limits of a 1D function at a given point.

- **point** is the point at which the limits have to be calculated. The default value is 0.0.
- **delta** is the distance to the far left and right of **point**. The default value is 1E-10.
- **formula** is a string describing the function under investigation. You can use any valid symbol as the variable name. See Chapter 1, Parser VIs, in this manual for more information.
- **left limit**. The accuracy is up to 8 decimal digits.
- **right limit**. The accuracy is up to 8 decimal digits.
- **ticks** is the time in milliseconds to analyze the formula and to produce limits. Usually, the time is negligible for the limit operations.
- **error**. See Appendix A, Error Codes, for a list of error codes. The accuracy of **formula** is verified by the Parser VIs.

The algorithm calculates only the two values \( f(\text{point} - \text{delta}) \) and \( f(\text{point} + \text{delta}) \). Furthermore, delta is internally rounded to a power of 2.

**Note:** A very small delta value can result in numerical inaccuracies. It is highly recommended that you take a value of delta = 1E-10 in all cases.

The function \( f(x) = (1 + 1/x)x \) has the famous limit \( e \) (Euler) if \( x \) tends to infinity. You can use the Limit VI to determine the Euler number if you define \( g(x) = (1 + x)^{(1/x)} \) for small positive \( x \). By entering \((1+x)^{(1/x)}\) in the formula control on the front panel, you can find the limit. The following diagram shows the convergence of \( f(x) \) to \( e \).
**Zeroes and Extrema of f(x)**

Determines all zeroes and extrema of a 1D function in a given interval.

**accuracy** controls the accuracy of the zeroes and the extrema. The default value is 1E-8.

**step type**. A value of 0 (fixed function) represents uniformly spaced function values, a value of 1 (modified function) represents the optimal step size. In general, the second value leads to more accurate zeroes and extrema. The default value is 0.

**algorithm**. A value of 0 selects the Ridders method; a value of 1 selects the Newton Raphson method. The default value is 0.

**start** is the start point of the interval under investigation. The default value is 0.0.

**end** is the end point of the interval. The default value is 1.0.
formula is a string describing the function.

Minima are the determined minimal values of formula.

f(Minima) are the function values at Minima.

Zeroes are the determined zeroes of formula.

f(Zeroes) are the function values of Zeroes. Usually, these values are very close to 0.

Maxima are the determined maximal values of formula.

f(Maxima) are the function values at Maxima.

ticks is the time in milliseconds to analyze the formula and to produce the Minima, Zeroes, and Maxima.

error. See Appendix A, Error Codes, for a list of error codes.
2D Explorer VIs

You can use the 2D Explorer VIs to examine 2D functions given in symbolic form, where parameterization is allowed. You can numerically calculate extrema and partial derivatives. These VIs are available in the 2DEXPLO.LLB library.

2D Explorer VI Descriptions

Eval X-Y-Z(a,t1,t2)
Describes a surface in 3D with two variables running over two different intervals. Additionally, you can integrate an arbitrary set of parameters, where any parameter gets its value with the help of an element of an array of parameter clusters on the front panel.

- **number of points** describes the number of grid points for both variables. The default value is 25.
- **Start** are the start points of both variables (that is, an array of length 2). The default values are (0,0).
- **End** are the end points of both variables (that is, an array of length 2). The default values are (1,1).
- **Parameters** is an array of clusters describing the parameters.
  - **name** of the parameter which uses the same conventions as the Parser VIs.
  - **value** of the parameter.
Chapter 8 2D Explorer VIs

**Formulas** is an array of three strings describing the three functions.

**Variables** is an array of two strings representing the two variables with respect to the naming conventions of the Parser VIs. The default variables are \((t_1, t_2)\).

**X** is a 1D array of the x values at the grid points. These values are calculated on a 2D grid defined by the discrete values of the **Variables** (usually \(t_1\) and \(t_2\)). The size of **X** is given by number of points.

**Y** is a 1D array of the y values at the grid points. These values are calculated on a 2D grid defined by the discrete values of the **Variables** (usually \(t_1\) and \(t_2\)). The size of **Y** is given by number of points.

**Z** is a 1D array of the z values at the grid points. These values are calculated on a 2D grid defined by the discrete values of the **Variables** (usually \(t_1\) and \(t_2\)). The size of **Z** is given by number of points.

**ticks** is the time effort for the whole calculation of the function values in milliseconds.

**error**. See Appendix A, *Error Codes*, for a list of error codes. Errors can result from incorrect function definitions and from discrepancies between the function definition and the array of variables.

**Example**

As an example, consider the ellipsoid given by

\[ \frac{x^2}{a^2} + \frac{y^2}{b^2} + \frac{z^2}{c^2} = 1. \]

Choosing the parameter values (in the **Parameters** cluster) as \(a = 1\), \(b = 2\), and \(c = 3\), you can input the following three equations in the **Formulas** control on the front panel to get the **X**, **Y** and **Z** values of the ellipsoid.

\[ a \cos(t_1) \]

\[ b \sin(t_1) \sin(t_2) \]

\[ c \sin(t_1) \cos(t_2) \]
Eval X-Y-Z(t1,t2)

Describes a surface in 3D with two variables running over two different intervals.

**number of points** describes the number of grid points for both variables. The default value is 25.

**Start** are the start points of variables (that is an array of length 2). The default value is (0,0).

**End** are the end points of both variables (that is, an array of length 2). The default value is (1,1).

**Formulas** is a 3-element array of formula strings representing a function definition of exactly two different variables. The naming conventions of the Parser VIs are valid.

Variables is an array of two strings representing the two variables with respect to the naming conventions of the Parser VIs. The default variables are \((t_1, t_2)\).

**X** is the array of the \(x\) values at the grid points defined by the parameters.

**Y** is the array of the \(y\) values at the grid points defined by the parameters.

**Z** is the array of the \(z\) values at the grid points defined by the parameters.

**ticks** is the time effort for the whole calculation of all function values in milliseconds.

**error**. See Appendix A, *Error Codes*, for a list of error codes. Errors can result from incorrect function definitions and by discrepancies between the function definition and the array of variables.
Example

The three formulas
\[ x(t1, t2) = \sin(t1) \]
\[ y(t1, t2) = \cos(t1)\sin(t2) \]
\[ z(t1, t2) = \cos(t1)\cos(t2) \]

describe a part of the unit sphere in 3D. In the **Formulas** control, only the right hand side of the above three equations needs to be entered.

**Eval y=f(a,x1,x2)**

Calculates a 2D array of function values defined on a grid. Additionally, you can integrate an arbitrary set of parameters, where any parameter gets its value with the help of an element of an array of parameter clusters on the front panel.

- **number of points** describes the number of grid points for both variables. The default value is 25.
- **Start** are the start points of both variables (that is, an array of length 2). The default value is (0,0).
- **End** are the end points of both variables (that is, an array of length 2). The default value is (1,1).
- **Parameters** is an array of clusters describing the parameters.
  - **name** of the parameter which uses the same conventions as the Parser VIs.
  - **value** of the parameter.
- **formula** is a string describing a function, where both variables and parameter variables can be used.
- **Variables** is an array of two strings representing the two variables with respect to the naming conventions of the Parser VIs. The default variables are \((x1, x2)\).
**X1 Values** is a 1D array of the used \( x_1 \) arguments.

**X2 Values** is a 1D array of the used \( x_2 \) arguments.

**Y Values** is the resulting 2D array of the function values.

**ticks** is the time effort for the whole calculation of the function values in milliseconds.

**error.** See Appendix A, Error Codes, for a list of error codes. Errors can result from incorrect function definitions and from discrepancies between the function definition and the array of variables. Additional errors may occur if the parameter variables violate the general parser rules or collide with the set of variables.

**Example**

The function \( f(a, b, x_1, x_2) = a \cdot \text{sinc}(\text{gamma}(x_1 + x_2)) - b \cdot \sin(x_1) \cdot \cos(x_2) \) with \( a = 1 \) and \( b = 2 \) is a common function in \( x_1 \) and \( x_2 \). The variables \( a \) and \( b \) stand for **Parameters.** Only the right hand side of this equation needs to be entered on the **Formula** control. Thus the values entered on the front panel are

**Parameters:**

- parameters(0) \( a \) = 1.00
- parameters(1) \( b \) = 2.00

**formula:**

\[ a \cdot \text{sinc}(\text{gamma}(x_1 + x_2)) - b \cdot \sin(x_1) \cdot \cos(x_2) \]
Eval \( y=f(x_1,x_2) \)
Calculates a 2D array of function values defined on a grid.

- **number of points** describes the number of grid points for both variables. The default value is 25.
- **Start** are the start points of both variables (that is, an array of length 2). The default value is \((0,0)\).
- **End** are the end points of both variables (that is, an array of length 2). The default value is \((1,1)\).
- **formula** is a string representing a function definition of exactly two different variables. The naming conventions of the Parser VIs are valid.
- **Variables** is an array of two strings representing the two variables with respect to the naming conventions of the Parser VIs. The default variables are \((x_1,x_2)\).
- **X1 Values** is a 1D array of the used \(x_1\) arguments.
- **X2 Values** is a 1D array of the used \(x_2\) arguments.
- **Y Values** is the resulting 2D array of the function values.
- **ticks** is the time effort for the whole calculation of all function values in milliseconds.
- **error**. See Appendix A, *Error Codes*, for a list of error codes. Errors can result from incorrect function definitions and from discrepancies between the function definition and the array of variables.

**Note:** You can use the Data Visualization VI Library to visualize the results of the Eval \( y=f(x_1,x_2) \) VI. For more details, refer to the VI descriptions for the Contour Plot, Mesh 3D, and Common Intensity Maps VIs.
The following diagram shows the visualization of the function 
\[ f(x_1, x_2) = \sin(3x_1) \cos(3x_2) \] in the interval \((-2, 2) \times (-2, 2)\).

**Extrema of \( f(x_1,x_2) \)**
The algorithm is looking for local extrema of a given function of two variables on a given rectangle.

- **Start** are the start points of both variables (that is, an array of length 2). The default values are (0,0).
- **End** are the end points of both variables (that is, an array of length 2). The default values are (1,1).
- **formula** is a string describing a function.
- **Variables** is an array of two strings representing the two variables with respect to the naming conventions of the Parser VIs. The default variables are \((x_1,x_2)\).
- **number of trials** is the number of randomly spaced starting points of the algorithm.
Minima is a 2D array of all local minima of the given function. Any local minimum is presented by two coordinates.

Maxima is a 2D array of all local maxima of the given function. Any local maximum is presented by two coordinates.

ticks is the time effort for the whole calculation of the function values in milliseconds.

error. See Appendix A, Error Codes, for a list of error codes. Errors can result from incorrect function definitions and from discrepancies between the function definition and the array of variables.

The absolute distance between two extrema must be equal or larger than 1E-6 for the two vectors to register as different from each other.

Note: While the number of trials can be very large, there is no guarantee you can find all or at least one zero, local minimum or local maximum of the given function. Moreover, such points do not exist in all cases.

Note: Though the randomly chosen start points of the extrema algorithm belong to the initially given rectangle, sometimes the determined extrema can be found outside of the rectangle. In this case, these values will also be presented.

Partial Derivatives of \( f(x_1, x_2) \)

Calculates a 2D array of the partial derivatives of a function of two independent variables.

number of points describes the number of grid points for both variables. The default value is 25.

Start are the start points of both variables (that is, an array of length 2). The default value is (0,0).

End are the end points of both variables (that is, an array of length 2). The default value is (1,1).
derivative is a value of 0 represents the partial derivative of the first variable. A value of 1 represents the partial derivative of the second variable.

formula is a string describing a function.

Variables is an array of two strings representing the two variables with respect to the naming conventions of the Parser VIs. The default value is \((x_1, x_2)\).

Partial derivative of \(f(x_1, x_2)\) is the 2D array of the fixed partial derivative at the defined grid points. For a derivative of 0, the function \(df(x_1, x_2)/dx_1\) is calculated, for a derivative of 1 the function \(df(x_1, x_2)/dx_2\) is calculated.

X2 Values is the resulting 1D array.

Y Values is the resulting 2D array.

ticks is the time effort for the whole calculation of the function values in milliseconds.

ticks is the time effort for the whole calculation of the function values in milliseconds.

error. See Appendix A, Error Codes, for a list of error codes. Errors can result from incorrect function definitions and discrepancies between the function definition and the array of variables. An additional error can occur, if the derivative is neither 0 nor 1.
Example

The $x_1$-derivative of the function $f(x_1, x_2) = \sin(x_1^2 - x_2) - \cos(\sin(x_2) - x_1)$ is investigated in the interval $(-2, 2)$ by $(-2, 2)!$ The values are obtained by entering the following on the front panel:

Start: $[-2, -2]$

End: $[2, 2]$

Formula: $\sin(x_1^2 - x_2) - \cos(\sin(x_2) - x_1)$
Function VIs

These are a group of VIs you can use to evaluate some common mathematical functions. They are available in the FUNCTION. LLB library.

Function VI Descriptions

Bessel Function Jn(x)
The Bessel function $J_n(x)$ is defined by

$$J_n(x) = \left(\frac{1}{2x}\right)^n \sum_{k=0}^{\infty} \frac{\left(-\frac{1}{4}x^2\right)^k}{k!(n+k+1)}$$

with $n = 0, 1, \ldots$

- $x$ is any real number.
- $n$ is a non-negative integer.
- $J_n(x)$ is the Bessel function of the first kind of order $n$.

Generally the power series for $J_n(x)$ is not computationally useful. More useful are recurrence relations and rational approximations of $J_n$ and $Y_n$. The recurrences are

$$J_{n+1}(x) = \frac{2n}{x} J_n(x) - J_{n-1}(x)$$

$$Y_{n+1}(x) = \frac{2n}{x} Y_n(x) - Y_{n-1}(x)$$

The treatment of the cases $x \leq n$ and $x > n$ is completely different for the sake of accuracy.
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Bessel Function \( Y_n(x) \)

The Bessel function \( Y_n(x) \) is defined by

\[
Y_n(x) = \frac{J_n(x) \cos(n\pi) - J_{-n}(x)}{\sin(n\pi)} \quad \text{with } n = 0, 1, \ldots
\]

based on the definition of the Bessel functions \( J_n(x) \).

\[
\begin{align*}
x & \quad (\text{input}) \quad \text{Yn(x)} \\
n & \quad (\text{integer}) \quad \text{Yn(x)}
\end{align*}
\]

- \( x \) is any real number.
- \( n \) is a nonnegative integer.
- \( Y_n(x) \) is the Bessel function of the second kind of order \( n \).

See the description of the Bessel Function \( J_n(x) \) VI in this chapter for more information.

Bessel Polynomial

The Bessel polynomial \( P_n(x) \) of order \( n \) is defined by a recurrence relation

\[
P_n(x) = P_{n-1}(x) + \frac{x^2}{4(n-1)^2-1} P_{n-2}(x) \quad \text{for } n = 2, 3, \ldots
\]

where \( P_0(x) = 1 \) and \( P_1(x) = 1 + x \).

\[
\begin{align*}
x & \quad (\text{input}) \quad \text{Bessel polynomial (n,x)} \\
n & \quad (\text{integer}) \quad \text{Bessel polynomial (n,x)}
\end{align*}
\]

- \( x \) is any real number.
- \( n \) is a positive integer.

Bessel polynomial \( (n, x) \) is the result of the calculation of \( P_n(x) \) for the given values of \( n \) and \( x \).

\( \square \) Note: Bessel Polynomials have strong connections to the Bessel filters. See the LabVIEW Analysis VI Reference Manual for details.
The following diagram shows the first four Bessel polynomials from bottom to top on the right side. The range of \( x \) is from \([-6, 6]\).

**Beta Function**

The beta function \( B(w, z) \) is defined by

\[
B(w, z) = \frac{\Gamma(w)\Gamma(z)}{\Gamma(w + z)}
\]

\( z \) is any real number.

\( w \) is any real number.

\( \text{beta} (w,z) \) is the result of the calculation of \( B(w, z) \) for the given values of \( w \) and \( z \).

The calculation of \( B(w, z) \) uses the Gamma Function VI.

**Note:** \( B(w, z) \) is undefined, if \( w \) or \( z \) is a non-positive integer.
Chapter 9  Function VIs

**Binomial Coefficient**

The binomial coefficient is determined by

\[
\binom{n}{k} = \frac{n!}{k!(n-k)!}
\]

- \(n\) is any non-negative integer.
- \(k\) is any non-negative integer.

**binomial coefficient** \((n,k)\) is the result of the calculation of the binomial coefficient for the given values of \(n\) and \(k\).

**error**. See Appendix A, *Error Codes*, for a list of Error Codes, if \(n < k\).

Binomial coefficients can have many digits, even in the case of relatively small numbers \(n\) and \(k\). The data type most suited for the binomial coefficient is a double real. You can directly calculate the factorial functions, \(n!\), \(k!\), and \((n-k)!\), with the Gamma Function VI.

**Chebyshev Polynomial**

The Chebyshev polynomial \(T_n(x)\) is defined by

\[
T_n(x) = \cos(n \arccos x) \text{ for } n = 0, 1, \ldots \text{ and real numbers } x.
\]

These functions form the base of the so called Chebyshev approximation. For \(i \neq j\) it is

\[
\int_{-1}^{1} \frac{T_i(x)T_j(x)}{\sqrt{1-x^2}} \, dx = 0
\]
All $T_n(x)$ form an orthogonal system over the weight function

$$\frac{1}{\sqrt{1-x^2}}$$

$x$ is the real argument.

$n$ is the order of the Chebyshev polynomial.

$T(n,x)$ is the value of the $n$th Chebyshev polynomial at the point $x$.

**Note:** The result of this definition does not look like a polynomial at first glance, but you can use trigonometric rules to show that $T_n$ is a polynomial of degree $n$ in the variable $x$.

**Note:** Chebyshev polynomials have strong connections to some very important filter types. See the LabVIEW Analysis VI Reference Manual for details.

The following diagram shows the first four Chebyshev polynomials of degrees 0, 1, 2, and 3.
Continued Fraction

The continued fraction of two sequences \((a_0, a_1, \ldots, a_n)\) and \((b_0, b_1, \ldots, b_n)\) is defined by the following term.

\[
result = \frac{a_0}{b_0 + \frac{a_1}{b_1 + \ldots}}
\]

Continued fractions are invaluable tools for calculating special functions.

\begin{align*}
A \quad \text{is the 1D array of the numerator part of the continued fraction.} \\
B \quad \text{is the 1D array of the denominator part of the continued fraction.} \\
\text{result} \quad \text{is a real value representing the result of the continued fraction.} \\
\text{error} \quad \text{See Appendix A, Error Codes, for a list of Error Codes, if the dimensions of A and B are not equal.}
\end{align*}

Cosine Integral

The cosine integral is defined by

\[
\text{ci}(x) = \gamma + \ln x + \int_0^x \frac{\cos s - 1}{s} ds \quad \text{with the Euler constant } \gamma.
\]

It is \(\text{ci}(-x) = \text{ci}(x) - i\pi\).

The algorithm accepts only nonnegative real numbers as input.
Chapter 9   Function VIs

**x** is any real non-negative number.

\( \text{ci}(x) \) is the result of the calculation of the cosine integral for the given value of \( x \).

**error.** See Appendix A, *Error Codes*, for a list of Error Codes, if \( x \leq 0 \).

The calculation of \( \text{ci}(x) \) can be done with the help of \( E(x) \). See the Sine Integral VI description in this chapter for more information.

The following diagram shows the graph of the cosine integral, in the interval (0,15).

---

**Gamma Function**

The gamma function of \( x \) is the generalization of the common factorial function \( n! \). The relation between these two functions is \( \Gamma(n + 1) = n! \) for all natural numbers \( n \). The gamma function is defined by

\[
\Gamma(x) = \int_0^x s^{x-1} \exp(-s)ds
\]

for real and complex \( x \), and has the property \( \Gamma(x + 1) = x\Gamma(x) \).
Chapter 9  Function VIs

x is any real number.

**gamma (x)** has singularities for all nonpositive integers x.

The calculation of $\Gamma(x + 1)$ is based on the Lanszos formula

$$\Gamma(x + 1) = (x + 5.5)^x + 0.5 \exp(-x - 5.5) \sqrt{2\pi} \left( c_0 + \frac{c_1}{x + 1} + \ldots + \frac{c_6}{x + 6} \right)$$

with fixed and well calculated $c_0, c_1, \ldots, c_6$.

The following diagram shows the graph of the gamma function in the interval (–4, 4).

---

**Incomplete Beta Function**

The incomplete beta function is defined by

$$I_x(a, b) = \frac{1}{B(a, b)} \int_0^x s^{a-1}(1-s)^{b-1} \, ds$$

with $a, b > 0$ and $0 \leq x \leq 1$

where $B(a, b)$ denotes the beta function of $a$ and $b$. 
**Chapter 9 Function VIs**

**Incomplete Beta Function**

- \( x \) is a real number between 0 and 1
- \( a \) is a positive real number.
- \( b \) is a positive real number.

**incomplete beta \((x, a, b)\)** is the result of the calculation of \( I_x(a, b) \) for the given values of \( x \), \( a \), and \( b \).

**error**. See Appendix A, Error Codes, for a list of Error Codes, if

\[ x \not\in [0, 1] \]
\[ a \leq 0 \]
\[ b \leq 0 \]

An efficient strategy for calculating of \( I_x(a, b) \) is based on the continued fraction

\[
I_x(a, b) = \frac{x^a (1-x)^b}{a B(a, b)} \left[ \frac{1}{d_1} \right] \quad \text{with} \quad d_i \text{ depending on } i, a, b, \text{ and } x.
\]

**Incomplete Gamma Function**

The incomplete gamma function is defined by

\[
P(a, x) = \frac{1}{\Gamma(a)} \int_0^x \exp(-s)s^{a-1} \, ds \quad \text{for } a > 0
\]

\[ x \text{ is any real number.} \]
a is a positive real number.

**Incomplete gamma** \((a,x)\) is the result of the calculation of \(P(a, x)\) for the given values of \(a\) and \(x\).

**Error.** See Appendix A, *Error Codes*, for a list of Error Codes, if \(a \leq 0\).

The integral in the definition of \(P(a, x)\) can approximately be derived by the following equation.

\[
\int_0^x \exp(-s)s^{a-1}ds = \exp(-x)x^a \sum_{n=0}^{100} \frac{\Gamma(a)}{\Gamma(a+1+n)}x^n
\]

The following diagram shows the incomplete gamma functions with \(a = 0, 1, 2, 3\) from top to bottom.
Jacobian Elliptic Function

The value of \( sn \) (Jacobian Elliptic Function) is determined by the relation

\[
u = \int_{0}^{sn} \frac{ds}{\sqrt{(1 - s^2)(1 - k^2 s^2)}}
\]

where \( u \) and \( k \) are given real numbers. The other Jacobian Elliptic functions have the following definitions.

\[
\begin{align*}
sn^2 + cn^2 &= 1 \\
k^2 sn^2 + dn^2 &= 1
\end{align*}
\]

\[
sc = \frac{sn}{cn}
\]

\[
\begin{align*}
u & \quad \text{is any real number.} \\
k & \quad \text{is a real number with } 0 \leq k \leq 1.
\end{align*}
\]

\[
\begin{align*}
\text{sn}(u, k) & \\
\text{cn}(u, k) & \\
\text{sc}(u, k) & \\
\text{dn}(u, k) & \\
\text{error} &
\end{align*}
\]

error codes. See Appendix A, Error Codes, for a list of error codes if \( k < 0 \) or \( k > 1 \).
Legendre Elliptic Integral 1st Kind

The Legendre Elliptic Integral 1st kind is defined by the following equation.

\[ F(\phi, k) = \int_{\phi}^{\infty} \frac{d\phi}{\sqrt{1 - k^2 \sin^2 \phi}} \]

\( \phi \) is any real number.

\( k \) is a real number with \( 0 \leq k \leq 1 \).

\( F(\phi, k) \) is the result of the calculation of \( F(\phi, k) \) for the given values of \( \phi \) and \( k \).

\textbf{error}. See Appendix A, Error Codes, for a list of Error Codes, if \( k < 0 \) or \( k > 1 \).

The calculation uses the relation

\[ F(\phi, k) = \sin \phi R_F(\cos^2 \phi, 1 - k^2 \sin^2 \phi, 1) \]

where \( R_F \) is the elliptic integral in the Carlson form

\[ R_F(x, y, z) = \frac{1}{2} \int_{0}^{\infty} \frac{ds}{\sqrt{(s+x)(s+y)(s+z)}} \]
Sine Integral

The sine integral is defined by

\[ \text{si}(x) = \int_0^x \frac{\sin s}{s} \, ds \]  

It is \( \text{si}(-x) = -\text{si}(x) \)

\( x \) is any real number.

\( \text{si}(x) \) is the result of the calculation of the sine integral for the given value of \( x \).

An efficient algorithm is based on a continued fraction of the combined function

\[ E(x) = -ci(x) + i\left( \text{si}(x) - \frac{\pi}{2} \right) \] more precisely on

\[ E(x) = \exp(-ix) \left( \frac{1}{1 + ix} - \frac{1^2}{3 + ix} - \frac{2^2}{5 + ix} - \cdots \right) \]

The following diagram shows the graph of the sine integral in the interval \((0, 15)\).
Spike Function

The spike function is defined by

\[
spike(x) = \begin{cases} 
1 & \text{if } 0 \leq x < 1 \\
0 & \text{else}
\end{cases}
\]

\( \text{x is any real number.} \)

\( \text{spike(x)} \) is the value of \( \text{spike(x)} \) for the given value of \( x \).

\( \text{Note: You can define more complex functions by variation and combination of the Step Function, Spike Function, and Square Function VIs, respectively.} \)

The diagram below illustrates the following example.

Example: \( \text{spike}(x) - \text{spike}(-x) \) in the interval \((-4.0, 4.0)\)
Square Function

The square function is defined by the following equation.

\[
square(x) = \begin{cases} 
1 & \text{if } 2n \leq x < 2n + 1 \quad n = \ldots, -1, 0, 1, \ldots \\
0 & \text{if } 2n + 1 \leq x < 2n + 2 \quad n = \ldots, -1, 0, 1, \ldots 
\end{cases}
\]

\(x\) is any real number.

\(\square(x)\) is the value of \(\square(x)\) for the given value of \(x\).

Note: You can define more complex functions by variation and combination of the Step Function, Spike Function and Square Function VIs, respectively.

The diagram below illustrates the following example.

Example: \(\square(x) \times \square(x - 0.8)\) in the interval \((-4.0, 4.0)\)
**Step Function**

The step function is defined by

\[
step(x) = \begin{cases} 
0 & \text{if } x < 0 \\
1 & \text{else} 
\end{cases}
\]

Note: You can define more complex functions by variation and combination of the Step Function, Spike Function, and Square Function VIs, respectively.

The diagram below illustrates the following example.

Example: \(step(x) + \text{spike}(x)\) in the interval \((-4.0, 4.0)\)
Transform VIs

These VIs implement some transforms commonly used in mathematics and signal processing. They are available in the TRANS.LLB library.

Transform VI Descriptions

Buneman Frequency Estimator

This VI estimates the frequency of a given sine wave with unknown wavelength.

$X$ is the sampled signal at consecutive times.

$\beta$ is the estimation of the frequency of the sine wave represented by $X$.

$error$ See Appendix A, Error Codes, for a list of error codes.

Sometimes, an underlying time signal is not exactly periodic with period $n$, where $n$ denotes the size of the data array. How, then, do you determine the unknown period? The Buneman algorithm calculates the unknown frequency $\beta$ by

$$\beta = b + \frac{n}{\pi} \tan^{-1} \left( \frac{\sin \left( \frac{\pi}{n} \right)}{\cos \left( \frac{\pi}{n} \right) + \frac{|F_b(X)|}{|F_{b+b}(X)|}} \right)$$

Here $F_b$ denotes the value of the Fourier transform of the signal $X$ at the frequency $b$. The value of $b$ can be determined by the greatest value of $|F_b(X)|$.

The formula for $\beta$ is exact for pure sine waves and a good estimation in all other cases.
Daubechies4 Function

The Daubechies4 function is calculated at evenly spaced sampled points.

- **number of steps**: The algorithm calculates exactly 4 times the number of steps points of the Daubechies4 function. The default value is 128.
- **iterations**: The number of iterations. The quality of the output increases with the number of iterations. The default value is 10.
- **error**: See Appendix A for error codes.

The following figure shows the Daubechies4 function. This function is the base of a well known class of wavelet transforms.
Dual Signal FFT

One step computation of the Fourier transforms of two real signals of equal length.

Signal 1 is the array of the first signal. The length of Signal 1 must equal the length of Signal 2.

Signal 2 is the array of the second signal. The length of Signal 2 must equal the length of Signal 1.

FFT 1 is the complex Fourier transform of the first signal.

FFT 2 is the complex Fourier transform of the second signal.

error. See Appendix A for error codes. Especially, the length of Signal 1 and Signal 2 have to be the same.

From two signals X and Y of the same length, an artificial signal Z = X + iY can be constructed. From the complex Fourier transform of the complex signal Z the complex Fourier transforms of the signals X and Y can be simply derived. This method is much faster than the successive application of the real Fourier transform of X and Y.

Note: This VI is especially useful, if you need to calculate a continued stream of Fourier transforms of real signals.

Fractional FFT

Realizes the fractional fast Fourier transform of complex signals with arbitrarily chosen signal lengths.

X is a given time signal. It may be complex.

alpha can be any complex number. The default value is 0.00 + 0.00i.

Fract FFT {X} is the fractional fast Fourier transform.

error. See Appendix A, Error Codes, for a list of error codes.
Chapter 10  Transform VIs

The fractional Fourier transform is a natural generalization of the classical Fourier transform. With \( X = \{x_0, x_1, \ldots, x_{n-1}\} \) it is

\[
\text{Fractional FT}(X)(j) = \sum_{k=0}^{n-1} \exp(-2\pi i j k \alpha) x_k
\]

where \( \alpha \) is a complex number.

The classical Fourier transform is a special case, namely

\[
\alpha = \frac{1}{n}
\]

There is a very efficient algorithm for calculating the fractional Fourier transform called Fractional Fast Fourier Transform (FFFT). This algorithm is based on the chirp z strategy. The starting point is the obvious algebraic relation \(2jk = j^2 + k^2 - (k-j)^2\). With this relation it is

\[
\text{Fractional FFT}(X)(j) = \exp(-\pi j^2 \alpha) \sum_{k=0}^{n-1} y_k z_{j-k}
\]

with

\[
y_k = x_k \exp(-\pi i k^2 \alpha) \quad \text{and} \quad z_k = \exp(\pi i k^2 \alpha)
\]

It is possible to extend the signals \( Y \) and \( Z \) in such a manner that the above sum is a cyclic convolution.

\[\text{Note:} \quad \text{The Laplace transform is a special case of the fractional Fourier transform with alpha being purely imaginary. The fractional Fourier transform has many other interesting applications, such as analyzing signals with noninteger periodic components, high resolution trigonometric interpolation, and detecting lines in images.}\]
The following diagram shows the fractional fast Fourier transform of the function \( f(x) = x \) with \( \alpha = 0.1 + 0.0002i \).

**Laplace Transform Real**

Realizes the real Laplace transform of a real-time signal.

The real Laplace transform of a real signal \( x(s) \) is defined by

\[
Laplace\{X\}(s) = \int_0^\infty x(t) \exp(-st) dt \quad \text{for} \quad s \geq 0, \quad \text{and} \quad s \ \text{real}.
\]

Here, \( x(t) \) is defined for all \( t \geq 0 \). The discrete version of the Laplace transform of a discretely and evenly-sampled signal is a simple generation of the above continuous version.

- \( X \) is the array describing the evenly sampled time signal. The first element of this array belongs to \( t = 0 \), the last to \( t = \text{end} \).
- \( \text{end} \) is the instant in time of the last sample. The entire sample interval is between 0 and \( \text{end} \).
Laplace \( \{X\} \) is the result of the Laplace transform as an array.

**error**. See Appendix A, *Error Codes*, for a list of error codes, if **end** is out of range.

The definition of the Laplace transform is not of much use if the time signal increases very rapidly with the time. The discrete version of the Laplace transform cannot fully detect the convergence behavior of the original definition.

The discrete version of the Laplace transform is computationally very expensive. An efficient strategy for the discrete Laplace transform is based on the Fractional Fast Fourier Transform (FFFT). The definition of the FFFT is as follows:

\[
FFFT\{X\}(t) = \int_{-\infty}^{\infty} x(s) \exp(-i\alpha st) ds
\]

with an arbitrarily chosen complex \( \alpha \). See the description of the Fractional FFT VI in this chapter for more details.

**Example**

The following diagram shows the real Laplace transform of the function \( f(t) = \sin(t) \) in the interval \((0, 6)\). This is entered on the front panel as:

**end**: 6.00

**X**: values of \( \sin(t) \) for \( 0 \leq t \leq 6 \)
Power Spectrum Fractional FFT
Yields the absolute values of the fractional Fourier transform of a complex signal.

\[ \text{Power Spectrum FFT}(x) = |\text{FFT}(x)| \]

Prime FFT
Prime FFT is a special form of the Fourier transform. Use this VI if signal length is a prime.

Determining the Prime FFT is a five step process.
1. Determination of a generator \( g \) of the group modulo \( p \), where \( p \) is the prime length of \( X \). A generator \( g \) is such a natural number that
\[ 1 < g < p \quad \text{with} \quad \{g^0, g^1, \ldots, g^{p-1}\} = \{1, 2, \ldots, p-1\} \]
Chapter 10  Transform VIs

2. Reduction \{x_0, x_1, \ldots, x_{p-1}\} \rightarrow \{x_1, x_2, \ldots, x_{p-1}\}

3. Construction of two new signals each with a power of 2 length according to the following scheme:

first signal:
\{0, 0, \ldots, x_g, x_g, \ldots, x_g, 0, 0, \ldots\}

second signal:
\{w^{g-2}, w^{g-3}, \ldots, w^1, w^0, w^{g-2}, \ldots, w^1, 0, 0, \ldots\}

where \( w = \exp\left\{\frac{-2\pi i}{p}\right\} \)

4. Convolution of these two signals with the help of the Convolution Theorem.

5. Reduction and reorganization of the convolution result.

Sparse FFT
Computes the first elements of the Fourier transform of the input signal \( X \). You can use this VI to perform an FFT on an array of complex numbers.

\[ X \] is the given time signal.

\textbf{number of FFT values} is the number of interesting Fourier transform elements. The default value is 1.

\textbf{FFT \{X\}} is the truncated Fourier transform of \( X \).

\textbf{error}. See Appendix A, Error Codes, for a list of error codes. The size of \( X \) has to be a multiple of the \textbf{number of FFT values}. 
Let \( n \) be the length of \( X \) and \( m \) the number of FFT values. Then, the Fourier transform can be calculated by

\[
FFT\{X\}(j) = \sum_{k=0}^{d-1} \exp\left(\frac{-2\pi i j k}{n}\right) \sum_{l=0}^{m-1} \exp\left(\frac{-2\pi i l k}{m}\right)x_{k+l d}
\]

where \( n = dm \) and \( j = 0, \ldots, m - 1 \). In other words, the Fourier transform \( FFT\{X\} \) can be reduced to the combination of Fourier transforms of the length \( m \). Under certain circumstances this is more efficient than the classical Fourier transform method. The efficiency depends strongly on the relation between \( m \) and \( n \).

**Sparse Signal FFT**

The VI computes the Fourier transform of a signal \( X \), a so called sparse signal.

\( X \) is the nonzero part of the signal.

**signal** length is the length of the combined (zero-padded) signal, **signal length** can not be smaller than \( m \), where \( m \) is the length of \( X \). The default value is 1.

**FFT \{X\}** is the Fourier transform of the combined signal. The size of this array is **signal length**.

**error**. See Appendix A, **Error Codes**, for a list of error codes. **signal length** must be a multiple of \( X \).

Let \( n \) be the length of the combined signal and \( m \) the length of \( X \). Then it is

\[
FFT\{X\}(j) = \sum_{k=0}^{m-1} \exp\left(\frac{-2\pi i j k}{n}\right)x_k
\]

This can efficiently be calculated using the Fractional Fourier Transform. The efficiency depends strongly on the relation between \( m \) and \( n \).
STFT Spectrogram
Computes the signal energy distribution in the joint time-frequency domain, using the Short-Time Fourier Transform (STFT) algorithm. This VI performs a sliding FFT.

\[ X \] is the time waveform.

**time increment** is the base 2 logarithm of the time spacing, in samples, between each row of the Spectrogram output. For example, if you sampled the time waveform at \( f_s \) Hz, the spacing between the rows of Spectrogram is \( \text{time increment} / f_s \) second.

Increasing **time increment** decreases the computation time and reduces memory requirements, but also reduces time-domain resolution. Decreasing **time increment** improves time-domain resolution, but increases the computation time and memory requirements.

The default value is 1.

**window length** is the actual length of the selected window. The default value is 1.

**window selector** determines the type of analysis window the VI uses to compute **STFT Spectrogram** \( \{X\} \). The **window selector** parameter can have the following values.

0: Rectangular
1: Blackman
2: Hamming
3: Hanning
4: Gaussian

The default is the Rectangular window.
STFT Spectrogram \{X\} is a 2D array that describes the time waveform energy distribution in the joint time-frequency domain.

The number of rows (time axis) in STFT Spectrogram \{X\} is equal to the number of elements in the time waveform divided by time increment, and then rounded up. The number of columns (frequency axis) in STFT Spectrogram \{X\} is equal to

\[
\text{window length} + 1
\]

error. See Appendix A, Error Codes, for a list of error codes.

Unevenly Sampled Signal Spectrum

Calculates the power spectrum of a signal that is unevenly spaced in time.

XTime is the discrete- and unevenly-spaced times.

X represents the data material at times XTime. There is a one-to-one relation between XTime and X.

Power Spectrum FFT \{X\} Frequency are the frequency points at which the power spectrum is calculated.

Power Spectrum FFT \{X\} is the power spectrum, in the sense of the Lomb normalized periodogram.

error. See Appendix A, Error Codes, for a list of error codes, if XTime and X have different lengths.

The algorithm used is based on the Lomb normalized periodogram. Let the data \(x_k\) be given at the time points \(t_k\), i.e. \(X = \{x_0, x_1, \ldots, x_{n-1}\}\) and \(XTime = \{t_0, t_1, \ldots, t_{n-1}\}\). Furthermore,

\[
\bar{x} = \frac{1}{n} \sum_{k=0}^{n-1} x_k \quad \text{and} \quad \sigma^2 = \frac{1}{n-1} \sum_{k=0}^{n-1} (x_k - \bar{x})^2
\]
Then the Lomb normalized periodogram is defined as follows:

\[
P(\omega) = \frac{1}{2\sigma^2} \left\{ \left[ \sum_{k=0}^{n-1} (x_k - \bar{x}) \cos \omega(t_k - \tau) \right]^2 + \left[ \sum_{k=0}^{n-1} (x_k - \bar{x}) \sin \omega(t_k - \tau) \right]^2 \right\}
\]

with

\[
\tau = \frac{1}{2\omega} \arctan \left( \frac{\sum_{k=0}^{n-1} \sin 2\omega t_k}{\sum_{k=0}^{n-1} \cos 2\omega t_k} \right)
\]

The following diagram shows the Fourier transform, of length 256, of a signal that has been sampled at unequal intervals of time. The signal is a combination of sine waves of frequencies 20, 40, 60, and 80 Hz. The duration of the signal is 1 sec. The sampling frequency was chosen as 256 Hz, giving the frequency resolution of 1 Hz.
**Walsh Hadamard**

Realizes the real Walsh Hadamard transform.

\[ X \] is an array of power of two length.

**Walsh Hadamard** \( \{X\} \).

**error.** See Appendix A, *Error Codes*, for a list of error codes. Especially, the signal length of \( X \) has to be a power of 2.

**Note:** The Walsh Hadamard transform has similar properties to the more well known Fourier transform, but the computational effort is considerably smaller.

The Walsh Hadamard transform is based on an orthogonal system consisting of functions of only two elements \(-1\) and \(1\). For the special case of \( n = 4 \) the Walsh Hadamard transform of the signal \( X = \{x_0, x_1, x_2, x_3\} \) can be noted in the following matrix form.

\[
WH\{X\} = \begin{bmatrix}
1 & 1 & 1 & 1 \\
1 & -1 & 1 & -1 \\
1 & 1 & -1 & -1 \\
1 & -1 & -1 & 1
\end{bmatrix} \begin{bmatrix}
x_0 \\
x_1 \\
x_2 \\
x_3
\end{bmatrix}
\]

The matrices on the right hand side have simple construction rules. If \( WH_n \) and \( WH_{n+1} \) denote the Walsh Hadamard matrices of dimension \( 2^n \) and \( 2^{n+1} \), respectively, the rule is

\[
WH_{n+1} = \begin{bmatrix}
WH_n & WH_n \\
WH_n & -WH_n
\end{bmatrix}
\]

where \(-WH_n\) is meant in the element wise sense.

**Note:** The Walsh Hadamard transform fulfills the Convolution Theorem:

\[
WH\{X \ast Y\} = WH\{X\} WH\{Y\}
\]
The following diagram shows the Walsh Hadamard transform of a pulse pattern signal of length 256, delay 32, and width 64.

### Walsh Hadamard Inverse

Realizes the inverse of the real Walsh Hadamard transform.

\[ WH^{-1}\{X\} = \frac{1}{n} WH\{X\}, \]  

where \( n \) is the length of the signal \( X \).

\( X \) is an array of power of 2 length.

**Walsh Hadamard Inverse \( \{X\} \).**

**error**. See Appendix A, *Error Codes*, for a list of error codes. Especially, the signal length of \( X \) has to be the a power of 2.

If \( WH\{X\} \) denotes the Walsh Hadamard transform of \( X \) and \( WH^{-1}\{X\} \) the inverse Walsh Hadamard transform, then it is

\[ WH\{WH\{X\}\} = WH\{WH^{-1}\{X\}\} = X \]

Furthermore, the following very simple formula is valid.

\[ WH^{-1}\{X\} = \frac{1}{n} WH\{X\}, \]  

where \( n \) is the length of the signal \( X \).
Wavelet Transform Daubechies4

Realizes the Wavelet transform based on the Daubechies4 function.

\[
\begin{bmatrix}
c_0 & c_1 & c_2 & c_3 \\
c_3 & -c_2 & c_1 & -c_0 \\
  & c_0 & c_1 & c_2 & c_3 \\
  & c_3 & -c_2 & c_1 & -c_0 \\
  & & & & \\
  & & & & \\
  & & & & \\
  & & & & \\
\end{bmatrix}
\]

\[C = \begin{bmatrix}
c_0 & c_1 & c_2 & c_3 \\
  & c_3 & -c_2 & c_1 & -c_0 \\
  & c_2 & \cdot & \cdot & \cdot & c_0 & c_1 \\
  & c_1 & \cdot & \cdot & \cdot & \cdot & c_3 & -c_2 \\
\end{bmatrix}\]

Note: Wavelet transforms form an extremely fast growing part of the general signal theory. Two important applications are compressions of data and solutions of large systems of linear equations.

The Wavelet Transform Daubechies4 transform can be defined using the transformation matrix

X are the samples of the input signal. The length of the signal has to be a power of 2, otherwise an error code is given.

\[\textbf{Wavelet Daubechies4 \{X\}}\]

\[\textbf{error}\]. See Appendix A, Error Codes, for a list of error codes.

[DBL]  
132
Here blank entries signify zeroes. The numbers $c_0, c_1, c_2,$ and $c_3$ have to fulfill certain orthogonal properties

\[
c_0^2 + c_1^2 + c_2^2 + c_3^2 = 1 \\
c_2c_0 + c_3c_1 = 0 \\
c_3 - c_2 + c_1 - c_0 = 0 \\
0c_3 - 1c_2 + 2c_1 - 3c_0 = 0
\]

with the unique solution

\[
c_0 = \frac{1 + \sqrt{3}}{4\sqrt{2}} \\
c_1 = \frac{3 + \sqrt{3}}{4\sqrt{2}} \\
c_2 = \frac{3 - \sqrt{3}}{4\sqrt{2}} \\
c_3 = \frac{1 - \sqrt{3}}{4\sqrt{2}}
\]

**Note:** You can solve the above system of nonlinear equations in $c_0, c_1, c_2,$ and $c_3$ directly with the Nonlinear System Single Solution VI of this package.

The Wavelet Daubechies4 transform of the array $X$ is defined by

\[
\text{Wavelet Daubechies4} \{X\} = C \ast X.
\]

**Wavelet Transform Daubechies4 Inverse**

Realizes the inverse of the Wavelet transform based on the Daubechies4 function.

\[
X \quad \text{Wavelet Daubechies4 Inv} \{X\} \quad \text{error}
\]

\[
\text{[DBL]} \quad \text{X are the samples of the input signal. The length of the signal has to be a power of 2, otherwise an error code is given.}
\]

\[
\text{[DBL]} \quad \text{[I32]} \quad \text{Wavelet Daubechies4 Inv} \{X\}.
\]

\[
\text{error}. \text{ See Appendix A, Error Codes, for a list of error codes.}
\]
The Wavelet Transform Daubechies4 Inverse transform can be defined with the help of the transformation matrix

\[
C = \begin{bmatrix}
    c_0 & c_3 & c_2 & c_1 \\
    c_1 & -c_2 &  &  \\
    & c_2 & c_0 & c_3 \\
    & c_3 & -c_0 & c_1 & -c_2 \\
    & & & & \\
    c_2 & c_1 & c_0 & c_3 \\
    c_3 & -c_0 & c_1 & -c_2 \\
    c_2 & c_1 & c_0 & c_3 \\
    c_3 & -c_0 & c_1 & -c_2
\end{bmatrix}
\]

Here blank entries signify zeroes. The numbers \( c_0, c_1, c_2, \) and \( c_3 \) have to fulfill certain orthogonal properties, namely

\[
\begin{align*}
c_0^2 + c_1^2 + c_2^2 + c_3^2 &= 1 \\
c_2c_0 + c_3c_1 &= 0 \\
c_3 - c_2 + c_1 - c_0 &= 0 \\
0c_3 - 1c_2 + 2c_1 - 3c_0 &= 0
\end{align*}
\]
with the unique solution

\[
\begin{align*}
    c_0 &= \frac{1 + \sqrt{3}}{4\sqrt{2}} \\
    c_1 &= \frac{3 + \sqrt{3}}{4\sqrt{2}} \\
    c_2 &= \frac{3 - \sqrt{3}}{4\sqrt{2}} \\
    c_3 &= \frac{1 - \sqrt{3}}{4\sqrt{2}}
\end{align*}
\]

The inverse Wavelet Daubechies4 transform of the array \( X \) is defined by

\[\text{Wavelet Daubechies4 Inv}\{X\} = C^{-1}X\]

It is \( CC^{-1} = C^{-1}C = I \) (Refer to the definition of the Wavelet Transform Daubechies4 VI.)

The following diagram shows the Wavelet Transform Daubechies4 Inverse VI of a function with two spikes at the points 13 and 69. The signal length is 1024.
WVD Spectrogram
Computes the signal energy distribution in the joint time-frequency domain using the Wigner-Ville distribution algorithm.

\[
\begin{align*}
&\text{\textbf{WVD Spectrogram} } \{X\} \\
&\text{time increment} \\
&\text{error}
\end{align*}
\]

\(X\) is the time waveform.

**time increment** is the base 2 logarithm of the time spacing, in samples between each row of the WVD Spectrogram \(\{X\}\) output. For example, if you sample the time waveform at \(fs\) Hz, the spacing between the rows of WVD Spectrogram \(\{X\}\) is **time increment** \(fs\) seconds. The default value is 1.

Increasing **time increment** decreases the computation time and reduces memory requirements, but also reduces time-domain resolution. Decreasing **time increment** improves time-domain resolution, but increases the computation time and memory requirements.

\(\text{WVD Spectrogram} \{X\}\) is a 2D array that describes the time waveform energy distribution in the joint time-frequency domain.

**error**. See Appendix A, *Error Codes*, for a list of error codes.
Appendix

Error Codes

The following tables contain the error codes for the G Math Toolkit.

Table A-1.  G Math Toolkit Error Codes

<table>
<thead>
<tr>
<th>Error Code Number</th>
<th>Error Code Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>No error</td>
</tr>
<tr>
<td>-23001</td>
<td>Syntax error of parser</td>
</tr>
<tr>
<td>-23002</td>
<td>Discrepancy between function, variables and coordinates</td>
</tr>
<tr>
<td>-23003</td>
<td>Number of contours out of range</td>
</tr>
<tr>
<td>-23004</td>
<td>Number of color palettes out of range</td>
</tr>
<tr>
<td>-23005</td>
<td>Negative distance</td>
</tr>
<tr>
<td>-23006</td>
<td>Not a valid path</td>
</tr>
<tr>
<td>-23007</td>
<td>Not a graphs file</td>
</tr>
<tr>
<td>-23008</td>
<td>Wrong input, Euler method</td>
</tr>
<tr>
<td>-23009</td>
<td>Wrong input, Runge Kutta method</td>
</tr>
<tr>
<td>-23010</td>
<td>Wrong input, Cash Karp method</td>
</tr>
<tr>
<td>-23011</td>
<td>Nonpositive step rate</td>
</tr>
<tr>
<td>-23012</td>
<td>Nonpositive accuracy</td>
</tr>
<tr>
<td>-23013</td>
<td>Matrix vector conflict</td>
</tr>
<tr>
<td>-23014</td>
<td>A and X0 have different dimensions</td>
</tr>
<tr>
<td>-23015</td>
<td>Empty X0</td>
</tr>
</tbody>
</table>
### Appendix A  Error Codes

#### Table A-1.  G Math Toolkit Error Codes

<table>
<thead>
<tr>
<th>Error Code Number</th>
<th>Error Code Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>–23016</td>
<td>Singular eigenvector matrix</td>
</tr>
<tr>
<td>–23017</td>
<td>Multiple roots</td>
</tr>
<tr>
<td>–23018</td>
<td>Left point is a root</td>
</tr>
<tr>
<td>–23019</td>
<td>Right point is a root</td>
</tr>
<tr>
<td>–23020</td>
<td>Left point greater than right point</td>
</tr>
<tr>
<td>–23021</td>
<td>Both function values have the same sign</td>
</tr>
<tr>
<td>–23022</td>
<td>Nonpositive accuracy or nonpositive delta x(h)</td>
</tr>
<tr>
<td>–23023</td>
<td>Wrong dimension of start</td>
</tr>
<tr>
<td>–23024</td>
<td>No root found</td>
</tr>
<tr>
<td>–23025</td>
<td>Nonvalid triplet (a,b,c)</td>
</tr>
<tr>
<td>–23026</td>
<td>No optimum found</td>
</tr>
<tr>
<td>–23027</td>
<td>Not exactly one variable</td>
</tr>
<tr>
<td>–23028</td>
<td>Wrong model equation</td>
</tr>
<tr>
<td>–23029</td>
<td>Levenberg Marquardt has failed</td>
</tr>
<tr>
<td>–23030</td>
<td>m &gt;= n &gt;= 0 is violated or the matrix of derivatives has the wrong dimension</td>
</tr>
<tr>
<td>–23031</td>
<td>No valid point</td>
</tr>
<tr>
<td>–23032</td>
<td>Maximum does not exist</td>
</tr>
<tr>
<td>–23033</td>
<td>Vectors have different dimensions or empty vectors</td>
</tr>
<tr>
<td>–23034</td>
<td>Ill conditioned system</td>
</tr>
<tr>
<td>–23035</td>
<td>Nonpositive number</td>
</tr>
</tbody>
</table>
Table A-1.  G Math Toolkit Error Codes

<table>
<thead>
<tr>
<th>Error Code Number</th>
<th>Error Code Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>–23036</td>
<td>Different parameters</td>
</tr>
<tr>
<td>–23037</td>
<td>Not exactly two functions</td>
</tr>
<tr>
<td>–23038</td>
<td>No variables in expression</td>
</tr>
<tr>
<td>–23039</td>
<td>Parameter problem</td>
</tr>
<tr>
<td>–23040</td>
<td>Derivative out of range</td>
</tr>
<tr>
<td>–23041</td>
<td>Not exactly two variables</td>
</tr>
<tr>
<td>–23042</td>
<td>Negative argument</td>
</tr>
<tr>
<td>–23043</td>
<td>Argument out of range (0,1]</td>
</tr>
<tr>
<td>–23044</td>
<td>Argument out of range [0,1]</td>
</tr>
<tr>
<td>–23045</td>
<td>n&lt;k</td>
</tr>
<tr>
<td>–23046</td>
<td>Empty array</td>
</tr>
<tr>
<td>–23047</td>
<td>Argument out of range [0,100]</td>
</tr>
<tr>
<td>–23048</td>
<td>Invalid time increment</td>
</tr>
<tr>
<td>–23049</td>
<td>Invalid window length</td>
</tr>
<tr>
<td>–23050</td>
<td>Signal length not a multiple of number</td>
</tr>
<tr>
<td>–23051</td>
<td>Signal length not a power of two</td>
</tr>
<tr>
<td>–23052</td>
<td>Signal length not a prime and $\geq 5$</td>
</tr>
<tr>
<td>–23053</td>
<td>Signal length not a power of two and $\geq 4$</td>
</tr>
<tr>
<td>–23054</td>
<td>Non–unique variables</td>
</tr>
</tbody>
</table>
The following table shows the possible parser error codes for the G Math Toolkit.

<table>
<thead>
<tr>
<th>Error Code Number</th>
<th>Error Code Description</th>
<th>Error Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>No error</td>
<td>sin(x)</td>
</tr>
<tr>
<td>1</td>
<td>Bracket problem at the beginning</td>
<td>1+x)</td>
</tr>
<tr>
<td>2</td>
<td>Incomplete function expression</td>
<td>sin(x)+</td>
</tr>
<tr>
<td>3</td>
<td>Bracket problem</td>
<td>()</td>
</tr>
<tr>
<td>4</td>
<td>Bracket problem at the end</td>
<td>(1+x</td>
</tr>
<tr>
<td>5</td>
<td>Wrong decimal point</td>
<td>1.2 (US)</td>
</tr>
<tr>
<td>6</td>
<td>Wrong number format</td>
<td>1e–3 instead of 1E–3</td>
</tr>
<tr>
<td>7</td>
<td>Wrong function call</td>
<td>sin()</td>
</tr>
<tr>
<td>8</td>
<td>Not a valid function</td>
<td>sins(x)</td>
</tr>
<tr>
<td>9</td>
<td>Incomplete expression</td>
<td>x+</td>
</tr>
<tr>
<td>10</td>
<td>Wrong variable name</td>
<td>a11</td>
</tr>
<tr>
<td>11</td>
<td>Wrong letter</td>
<td>sin(X)</td>
</tr>
<tr>
<td>12</td>
<td>Too many decimal points</td>
<td>1.23.45</td>
</tr>
<tr>
<td>21</td>
<td>Contains more than one variable</td>
<td>1+x+y4</td>
</tr>
<tr>
<td>22</td>
<td>Inconsistency in variables or numbers</td>
<td>Depends on application</td>
</tr>
<tr>
<td>23</td>
<td>Contains variables</td>
<td>Depends on application</td>
</tr>
<tr>
<td>24</td>
<td>Variables output problem</td>
<td>Depends on application</td>
</tr>
</tbody>
</table>
References

The following references may prove helpful to you as you use the VIs and examples found in the G Math Toolkit.


Appendix B  References


Customer Communication

For your convenience, this appendix contains forms to help you gather the information necessary to help us solve your technical problems and a form you can use to comment on the product documentation. When you contact us, we need the information on the Technical Support Form and the configuration form, if your manual contains one, about your system configuration to answer your questions as quickly as possible.

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- **United States**: (512) 794-5422
  - Up to 14,400 baud, 8 data bits, 1 stop bit, no parity
- **United Kingdom**: 01635 551422
  - Up to 9,600 baud, 8 data bits, 1 stop bit, no parity
- **France**: 1 48 65 15 59
  - Up to 9,600 baud, 8 data bits, 1 stop bit, no parity

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support@natinst.com

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National Instruments has branch offices all over the world. Use the list below to find the technical support number for your country. If there is no National Instruments office in your country, contact the source from which you purchased your software to obtain support.

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<thead>
<tr>
<th>Country</th>
<th>Telephone</th>
<th>Fax</th>
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<tbody>
<tr>
<td>Australia</td>
<td>03 9879 9422</td>
<td>03 9879 9179</td>
</tr>
<tr>
<td>Austria</td>
<td>0662 45 79 90 0</td>
<td>0662 45 79 90 19</td>
</tr>
<tr>
<td>Belgium</td>
<td>02 757 00 20</td>
<td>02 757 03 11</td>
</tr>
<tr>
<td>Canada (Ontario)</td>
<td>905 785 0085</td>
<td>905 785 0086</td>
</tr>
<tr>
<td>Canada (Quebec)</td>
<td>514 694 8521</td>
<td>514 694 4399</td>
</tr>
<tr>
<td>Denmark</td>
<td>45 76 26 00</td>
<td>45 76 26 02</td>
</tr>
<tr>
<td>Finland</td>
<td>09 527 2321</td>
<td>09 502 2930</td>
</tr>
<tr>
<td>France</td>
<td>01 48 14 24 24</td>
<td>01 48 14 24 14</td>
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<tr>
<td>Germany</td>
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<td>Hong Kong</td>
<td>2645 3186</td>
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<td>02 596 7455</td>
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<td>Mexico</td>
<td>5 520 2635</td>
<td>5 520 3282</td>
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<tr>
<td>Netherlands</td>
<td>0348 433466</td>
<td>0348 430673</td>
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<tr>
<td>Norway</td>
<td>32 84 84 00</td>
<td>32 84 86 00</td>
</tr>
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<td>Singapore</td>
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<td>2265887</td>
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<tr>
<td>Spain</td>
<td>91 640 0085</td>
<td>91 640 0533</td>
</tr>
<tr>
<td>Sweden</td>
<td>08 730 49 70</td>
<td>08 730 43 70</td>
</tr>
<tr>
<td>Switzerland</td>
<td>056 200 51 51</td>
<td>056 200 51 55</td>
</tr>
<tr>
<td>Taiwan</td>
<td>02 377 1200</td>
<td>02 737 4644</td>
</tr>
<tr>
<td>U.K.</td>
<td>01635 523545</td>
<td>01635 523154</td>
</tr>
</tbody>
</table>
Technical Support Form

Photocopy this form and update it each time you make changes to your software or hardware, and use the completed copy of this form as a reference for your current configuration. Completing this form accurately before contacting National Instruments for technical support helps our applications engineers answer your questions more efficiently.

If you are using any National Instruments hardware or software products related to this problem, include the configuration forms from their user manuals. Include additional pages if necessary.

Name __________________________________________________________________________
Company _______________________________________________________________________
Address ________________________________________________________________________
_______________________________________________________________________________
Fax (___)___________________ Phone (___) _________________________________________

Computer brand ___________ Model ___________ Processor___________________
Operating system (include version number) ____________________________________________
Clock speed ______MHz   RAM _____MB______________________________  Display adapter
Mouse ___yes   ___no     Other adapters installed _______________________________________

Hard disk capacity _____MB __________________________________________________Brand
Instruments used _________________________________________________________________
_______________________________________________________________________________

National Instruments hardware product model __________________________________  Revision
Configuration ___________________________________________________________________
National Instruments software product _________________________________________  Version
Configuration ___________________________________________________________________

The problem is: __________________________________________________________________
_______________________________________________________________________________
_______________________________________________________________________________
_______________________________________________________________________________
_______________________________________________________________________________

List any error messages: ___________________________________________________________
_______________________________________________________________________________
_______________________________________________________________________________

The following steps reproduce the problem:____________________________________________
_______________________________________________________________________________
_______________________________________________________________________________
_______________________________________________________________________________
_______________________________________________________________________________
_______________________________________________________________________________
G Math Hardware and Software Configuration Form

Record the settings and revisions of your hardware and software on the line to the right of each item. Complete a new copy of this form each time you revise your software or hardware configuration, and use this form as a reference for your current configuration. Completing this form accurately before contacting National Instruments for technical support helps our applications engineers answer your questions more efficiently.

National Instruments Products

DAQ hardware _______________________________________________________________
Interrupt level of hardware ____________________________________________________
DMA channels of hardware _____________________________________________________
Base I/O address of hardware __________________________________________________
Programming choice __________________________________________________________
HiQ, NI-DAQ, LabVIEW, or BridgeVIEW version __________________________________
Other boards in system _________________________________________________________
Base I/O address of other boards _______________________________________________
DMA channels of other boards _________________________________________________
Interrupt level of other boards ________________________________________________

Other Products

Computer make and model _______________________________________________________
Microprocessor ______________________________________________________________
Clock frequency or speed _______________________________________________________
Type of video board installed _________________________________________________
Operating system version _____________________________________________________
Operating system mode _________________________________________________________
Programming language _________________________________________________________
Programming language version _________________________________________________
Other boards in system _________________________________________________________
Base I/O address of other boards _______________________________________________
DMA channels of other boards _________________________________________________
Interrupt level of other boards ________________________________________________
Documentation Comment Form

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Title:  *G Math Toolkit Reference Manual*
Edition Date:  November 1996
Part Number:  321290A-01

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Glossary

Numbers/Symbols

- ° degrees
- ε epsilon
- % percent
- π pi
- 1D One-dimensional
- 2D Two-dimensional
- 3D Three-dimensional
- nD n-dimensional
- $f'$ or $\dot{f}$ derivative of $f$

A

array Ordered, indexed set of data elements of the same type.

ASCII American Standard Code for Information Interchange.
Glossary

B

Bessel filters  These filters have a maximally flat response in both magnitude and phase. The phase response in the passband, which is usually the region of interest, is nearly linear. Bessel filters can be used to reduce nonlinear phase distortion inherent in all IIR filters.

Bessel function  The Bessel function of the first kind of order \( n \), \( J_n(x) \), is defined by

\[
J_n(x) = \left( \frac{1}{\pi x} \right)^{1/2} \sum_{k=0}^{\infty} \frac{(-1)^k x^{2k}}{2^{k} k! \Gamma(n+k+1)}
\]

with \( n = 0, 1, \ldots \). The Bessel function of the second kind of order \( n \), \( Y_n(x) \), is defined by

\[
Y_n(x) = \frac{J_n(x) \cos(n\pi) - J_{-n}(x)}{\sin(n\pi)} \quad \text{with} \quad n = 0, 1, \ldots
\]

Bessel polynomial  The Bessel polynomial \( P_n(x) \) of order \( n \) is defined by a recurrence relation

\[
P_n(x) = P_{n-1}(x) + \frac{x^2}{4(n-1)^2 - 1} P_{n-2}(x) \quad \text{for} \quad n = 2, 3, \ldots
\]

where \( P_0(x) = 1 \) and \( P_1(x) = 1 + x \).

beta function  An integral defined by

\[
B(w, z) = \int_0^1 t^{w-1}(1-t)^{z-1} \, dt \quad (w > 0, z > 0)
\]

or, in terms of the gamma function

\[
B(w, z) = \frac{\Gamma(w)\Gamma(z)}{\Gamma(w + z)}
\]
binomial coefficient The binomial coefficient is given by
\[ \binom{n}{k} = \frac{n!}{k!(n-k)!} \]

block diagram Pictorial representation of a program or algorithm. In G, the block diagram, which consists of executable icons called nodes and wires that carry data between the nodes, is the source code for the VI. The block diagram resides in the block diagram window of your G development environment.

bracketing interval A bracketing triplet \((a, b, c)\) of a continuous 1D function \(f\) is a combination of three points with \(f(a) > f(b)\) and \(f(c) > f(b)\). This guarantees the existence of a local minimum of \(f\) in the interval \((a, c)\).

bracketing of a minimum A root of a continuous function is said to be bracketed by a pair of points, \(a < b\), when \(f(a)\) and \(f(b)\) are of opposite signs. A minimum is bracketed when there are three points, \(a < b < c\), with \(f(a) > f(b)\) and \(f(c) > f(b)\).

Brent Method For solving nonlinear equations, methods that converge rapidly to the solution are unreliable because convergence may not occur unless started close enough to the actual solution. On the other hand, the more reliable methods are slower. The Brent method is a hybrid method that combines both the safety of the bisection method and the rapid convergence of inverse quadratic interpolation.

Cash Karp method A numerical method for solving ordinary differential equations with start conditions. The Cash Karp method is an embedded Runge Kutta formula and is based on a fifth order strategy (with six steps). The Cash Karp method works with an adaptive step rate and is computationally more efficient than the Euler method or the Runge Kutta method.
Glossary

chart A 2D display of one or more plots, in which the display retains previous data, up to a maximum which you can define. The chart receives the data and updates the display point by point or array by array, retaining a certain number of past points in a buffer for display purposes.

Chebyshev polynomial The Chebyshev polynomial, for real numbers \( x \), is given by

\[ T_n(x) = \cos(n \arccos(x)) \]

This results in

\[ T_0 = 1, \quad T_1(x) = x, \quad T_2(x) = 2x^2 - 1, \quad T_3(x) = 4x^3 - 3x, \] and so on.

chi-squared A penalty function given by

\[ \chi^2 = \sum_{i=0}^{N-1} \left( \frac{y_i - f(x_i; a_1, ..., a_M)}{\sigma_i} \right)^2 \]

In this equation, \((x_i, y_i)\) are the input data points, and \(f(x_i; a_1, ..., a_M) = f(X,A)\) is the nonlinear function where \(a_1, ..., a_M\) are coefficients. If the measurement errors are independent and normally distributed with constant, standard deviation \(\sigma_i = \sigma\), this is also the least-square estimation.

The input arrays \(X\) and \(Y\) define the set of input data points. It is assumed that one has prior knowledge of the nonlinear relationship between the \(x\) and \(y\) coordinates. That is, \(f = f(X,A)\) where the set of coefficients, \(A\), is determined by the Levenberg-Marquardt algorithm.

common intensity map/plot A method of displaying 3 dimensions of data on a 2D plot with the use of color.

conjugate gradient method This method can be used for multidimensional unconstrained minimization. It determines the local minimum of a function of \(n\) independent variables. The direction of the gradient is modified at each iteration. This is done by forming a sequence of conjugate (the inner product being orthogonal) search directions and tends to avoid slow convergence which may result by repeated
searching in the same direction. To minimize the function $f(x)$, given an initial guess $x_0$, we first calculate $g_0 = \nabla f(x_0)$. The rest of the algorithm consists of following a sequence of steps until convergence. One of these steps consists of solving for a quantity $\beta_{k+1}$, where

$$\beta_{k+1} = \frac{(g_{k+1}^T g_{k+1})/(g_k^T g_k)}{(Fletcher & Reeves)}$$

or

$$\beta_{k+1} = \frac{((g_{k+1} - g_k)^T g_{k+1})/(g_k^T g_k)}{(Polak & Ribiere)}$$

**continued fraction** The continued fraction of two sequences $(a_0, a_1, \ldots, a_n)$ and $(b_0, b_1, \ldots, b_n)$ is defined by the following term.

$$\text{result} = \frac{a_0}{b_0 + \frac{a_1}{b_1 + \ldots}}$$

Continued fractions are valuable tools for calculating special functions.

**contour plot** A plot where contour lines are used to connect points of equal value.

**control** Front panel object for entering data to a VI interactively or to a subVI programmatically.

**cosine integral** The cosine integral is defined by:

$$\text{ci}(x) = \gamma + \ln x + \int_0^x \frac{\cos s - 1}{s} \, ds$$

with the Euler constant $\gamma$ and $x$ as any real non-negative number.

**curve in 3D** A special parametric plot $(x(t), y(t), z(t))$, where the parameter $t$ runs over a given interval.
Glossary

**D**

data flow Programming system consisting of executable nodes in which nodes execute only when they have received all required input data and produce output automatically when they have executed. G applications are dataflow systems.

Daubechies4 Function A Daubechies wavelet with 4 coefficients.

dimension Size and structure attribute of an array.

discrete Having discontinuous values of the independent variable, usually time.

Downhill Simplex Method Determines a local minimum of a function of $n$ independent variables. The Downhill Simplex algorithm, also called the Nelder and Mead method, works without partial derivatives. The algorithm consists of catching the minimum of the function, $f(X)$, with the help of simple geometrical bodies, namely with a simplex. A simplex in 2D is a triangle, a simplex in 3D is a tetrahedron and so on. You must have $n + 1$ starting points, each of dimension $n$, forming the initial simplex. The user must enter only one point of these $(n+1)$. The $(n+1)$ dimensional simplex is automatically constructed. The algorithm generates a new simplex by some elementary operations like reflections, expansions, and contractions. In the end, the minimum is concentrated in a very small simplex.

**E**

eigenvalues Values of $\lambda$ for which the matrix equation $Ax = \lambda x$ has a nontrivial solution ($x \neq 0$) are known as eigenvalues or characteristic values.

eigenvectors The solutions of $x$ for which the matrix equation $Ax = \lambda x$ has a nontrivial solution ($x \neq 0$) are known as eigenvectors or characteristic vectors.

error in The error structure entering a VI.

error out The error structure leaving a VI.

error structure The LabVIEW error structure consists of a Boolean status indicator, a numeric code indicator, and a string source indicator.
Euler method

A numerical method for solving ordinary differential equations with start conditions. This is a single-step method because it depends on information at only one point in time to advance to the next point.

extrema

Maxima and minima.

F

FFT

Fast Fourier Transform.

FFFT

Fractional Fast Fourier Transform.

Fletcher & Reeves

See Conjugate Gradient Method.

formula node

Node that executes formulas that you enter as text. This node is especially useful for lengthy formulas that would be cumbersome to build in block diagram form.

front panel

The interactive user interface of a VI. Modeled from the front panel of physical instruments, it is composed of switches, slides, meters, graphs, charts, gauges, LEDs, and other controls and indicators.

G

G

The LabVIEW graphical programming language.

gamma function

The gamma function of \( x \) is the generalization of the common factorial function \( n! \). The relation between these two functions is \( \Gamma(n+1) = n! \) for all natural numbers \( n \). The gamma function is defined by

\[
\Gamma(x) = \int_0^\infty s^{x-1} \exp(-s)ds
\]

for real and complex \( x \), and has the property \( \Gamma(x+1) = x\Gamma(x) \).
golden section search Determines a local minimum of a given 1D function with the help of a bracketing of the minimum. Consider a real valued 1D function \( f(x) \), unimodal on the interval \((a,c)\). The Golden Section Search method determines, beginning with a bracketing triplet \((a,b,c)\), a new one with a considerably smaller expansion. Repeating this scheme often yields a good approximation of the local minimum. The new bracketing point, \( x \), is essentially calculated by the following equation.

\[
\frac{x - b}{c - a} = (\sqrt{5} - 2)
\]

graph A 2D display of one or more plots. A graph receives and plots data as a block.

Hz Hertz. Cycles per second.

incomplete beta function The incomplete beta function is defined by:

\[
I_x(a,b) = \frac{1}{B(a,b)} \int_0^x s^{a-1} (1-s)^{b-1} ds
\]

with \( a, b > 0 \) and \( 0 \leq x \leq 1 \)

where \( B(a,b) \) denotes the beta function of \( a \) and \( b \).

incomplete gamma function The incomplete gamma function is defined by

\[
P(a,x) = \frac{1}{\Gamma(a)} \int_0^x \exp(-s)s^{a-1} ds
\]

for \( a > 0 \).

indicator Front panel object that displays output.

Inf Digital display value for a floating-point representation of infinity.
J

Jacobian elliptic function

The value of \( sn \) (Jacobian Elliptic Function) is determined by the relation

\[
  u = \int_{0}^{sn} \frac{ds}{\sqrt{(1-s^2)(1-k^2 s^2)}}
\]

where \( u \) and \( k \) are given real numbers. The other Jacobian Elliptic functions have the following definitions

\[
  sn^2 + cn^2 = 1, \quad k^2 sn^2 + dn^2 = 1, \quad sc = \frac{sn}{cn}
\]

L

LabVIEW

Laboratory Virtual Instrument Engineering Workbench.

Legendre elliptic integral

The Legendre Elliptic Integral of the 1st kind is defined by the following equation.

\[
  F(\varphi, k) = \int_{0}^{\varphi} \frac{d\varphi}{\sqrt{1-k^2 \sin^2 \varphi}}
\]

library

See VI Library.

linear programming

Suppose that we are given \( f \), a linear function of the variables \( x1, x2, x3, \ldots, xn \), and a set of constraints on these variables in terms of linear inequalities. Linear Programming consists of methods for maximizing or minimizing \( f \). Such problems are usually found in the areas of economics, distribution of goods, production, and approximation theory.
Glossary

M

matrix Two-dimensional array.

maxima The maximum value(s) of a function.

minima The minimum value(s) of a function.

N

NaN Digital display value for a floating-point representation of not a number, typically the result of an undefined operation, such as 0/0.

Newton Raphson method Determines a zero of a 1D function close to two points with the help of the derivative of this 1D function. The two values form a search limit for the unknown zero of the 1D function. The Newton Raphson method (or Newton method) is an iterative method for solving equations of the form \( f(x) = 0 \) where the derivative of \( f, f' \), is continuous. Given two values, \( x_1 \) and \( x_2 \), with \( f(x_1) \cdot f(x_2) < 0 \), first use the midpoint method to calculate

\[
x(0) = (x_1 + x_2)/2
\]

Then use the Newton method to calculate

\[
x(n+1) = x(n) - f(x(n))/f'(x(n))
\]

Replace \( x(n) \) by \( x(n+1) \) and repeat equation (2) till a certain termination condition is met. This termination condition could be either after a given number of repetitions, or if

\[
|x(n+1) - x(n)| \leq \varepsilon
\]

This is a fast and simple method.

nonsingular matrix Matrix in which no row or column is a linear combination of the other rows or columns, respectively. In other words, the rows or columns are linearly independent. This matrix has a unique inverse, and it has a nonzero determinant.
O

ODE

Ordinary Differential Equation.

optimal step

For a function that has many maxima, minima, or singularities, the standard method (using equidistant points) may yield incorrect results and in such cases the optimal step method is much better. This method starts off with taking equidistant points, but also checks the steepness of the curve between these points. In very steep portions (determined by the value of epsilon on the front panel) of the graph, a new point is generated in between. As a rule, the smaller the value of epsilon, the more the points that are generated, and the better the graph.

P

Pade approximation

Determines the coefficients of a rational polynomial to best suit a given set of first derivatives. Let $f$ be a given function with known values

$$f(0), f'(0), \ldots, f^{(n+m)}(0)$$

There exists a unique rational polynomial $(m \geq n)$

$$R(x) = \frac{a_0 + a_1 x + \ldots + a_m x^m}{1 + b_1 x + \ldots + b_n x^n}$$

with

$$R(0) = f(0), \ R'(0) = f'(0), \ldots, \ R^{(m+n)}(0) = f^{(m+n)}(0).$$

The rational polynomial can be determined by solving a special linear equation.

parametric plot

A plot of the variables (such as $x$ and $y$) which are given in terms of another variable $t$ (called a parameter), resulting in the parametric equations $x = f(t)$ and $y = g(t)$. Each value of $t$ determines a point $(x,y)$. As $t$ is varied, the point $(x,y) = (f(t), g(t))$ varies and traces a curve, that is plotted.
Glossary

parser
A VI used to scan a string to determine the function of each of the elements in the string.

partial derivative
A derivative taken with respect to one of two or more independent variables, with the others being treated as constants.

platform
Computer and operating system.

plot
A graphical representation of an array of data shown either on a graph or a chart.

Polak & Ribiere
See Conjugate Gradient Method.

power spectrum
The magnitude of the FFT.

fractional FFT

prime FFT
A special case of the FFT where the signal length is prime. The prime FFT VI is computationally more efficient than the classical FFT algorithms.

R

rational polynomial
A quotient of polynomials. The advantage that they have over polynomials for approximating functions is that they can be used to model functions with poles.

real Laplace transform
The Laplace transform for real s.

Ridders Method
Determines a zero of a 1D function in a given interval. The function has to be continuous and has to have different signs at the end points of the interval.

Let us be given the function \( f(x) \) with

\[
f(a) \cdot f(b) < 0.
\]

Ridders method determines

\[
c = \frac{a + b}{2}
\]

and calculates the new guess
\[
c_{\text{new}} = c + (c - a) \frac{\text{sign}(f(a) - f(b))f(c)}{\sqrt{f(c)^2 - f(a)f(b)}}
\]

The triplets start, \(c_{\text{new}}\), end are the base for the new iteration, depending on whether

\[f(\text{start}) \ast f(c_{\text{new}}) < 0\]

or

\[f(c_{\text{new}}) \ast f(\text{end}) < 0.\]

The algorithm stops, if

\[|a - b| < \text{accuracy}.
\]

Ridders’ method is fast and reliable.

**roots**

Zeros of a function, i.e. values of the variable(s) for which the function is equal to zero.

**Runge Kutta method**

A numerical method for solving ordinary differential equations with start conditions. The Runge Kutta method works with a fixed step rate but with a higher degree of accuracy than the common Euler method.

**S**

**scalar**

Number capable of being represented by a point on a scale. A single value as opposed to an array. Scalar Booleans and clusters are explicitly singular instances of their respective data types.

**simplex**

A simple geometrical body. A simplex in 2D is a triangle, a simplex in 3D is a tetrahedron and so on. Used in finding the minimum of a function. See the Downhill Simplex nD VI for more information.
Glossary

sine integral
The sine integral is defined by

\[ \text{si}(x) = \int_0^x \frac{\sin s}{s} \, ds \]

We have that \( \text{si}(-x) = -\text{si}(x) \) where \( x \) is any real number.

singular matrix
A square matrix in which a row or column is a linear combination of the other rows or columns, respectively. i.e. the rows or columns are linearly dependent. This matrix does not have an inverse, and its determinant is equal to zero.

sparse signals
Signals with a large number of zero values.

spectrogram
A particular representation of a signal that describes the distribution of the energy of the time waveform in the joint time-frequency domain.

spike function
The spike function is defined by

\[ \text{spike}(x) = \begin{cases} 1 & \text{if } 0 \leq x < 1 \\ 0 & \text{else} \end{cases} \]

square function
The square function is defined as

\[ \text{square}(x) = \begin{cases} 1 & \text{if } 2n \leq x < 2n + 1 \quad n = \ldots, -1, 0, 1, \ldots \\ 0 & \text{if } 2n + 1 \leq x < 2n + 2 \quad n = \ldots, -1, 0, 1, \ldots \end{cases} \]

where \( x \) is any real number.

step function
The step function is defined by

\[ \text{step}(x) = \begin{cases} 0 & \text{if } x < 0 \\ 1 & \text{else} \end{cases} \]

where \( x \) is any real number.

STFT
Short-Time Fourier Transform.

string
Representation of a value as text.
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Glossary

**X**

X0  A vector of start (initial) conditions.

**Z**

zeros  See roots.