Programming in Scilab

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Abstract

In this document, we present programming in Scilab. In the first part, we present the management of the memory of Scilab. In the second part, we present various data types and analyze programming methods associated with these data structures. In the third part, we present features to design flexible and robust functions. In the last part, we present methods which allow to achieve good performances. We emphasize the use of vectorized functions, which allow to get the best performances, based on calls to highly optimized numerical libraries. Many examples are provided, which allow to see the effectiveness of the methods that we present.

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1 Introduction

This document is an open-source project. The LAT_EX sources are available on the Scilab Forge:

http://forge.scilab.org/index.php/p/docprogscilab

The LATEX sources are provided under the terms of the Creative Commons Attribution-ShareAlike 3.0 Unported License:

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The Scilab scripts are provided on the Forge, inside the project, under the scripts sub-directory. The scripts are available under the CeCiLL licence:

http://www.cecill.info/licences/Licence_CeCILL_V2-en.txt

2 Variable and memory management

In this section, we describe several Scilab features which allow to manage the variables and the memory. Indeed, we sometimes face large computations, so that, in order to get the most out of Scilab, we must increase the memory available for the variables.

We begin by presenting the stack which is used by Scilab to manage its memory. We present how to use the **stacksize** function in order to configure the size of the stack. Then we analyze the maximum available memory for Scilab, depending on the limitations of the operating system. We briefly present the **who** function, as a tool to inquire about the variables currently defined. Then we emphasize the portability variables and functions, so that we can design scripts which work equally well on various operating systems. We present the **clear** function, which allows to delete variables when there is no memory left. Finally, we present two functions often used when we want to dynamically change the behavior of an algorithm depending on the type of a variable, that is, we present the **type** and **typeof** functions.

The informations presented in this section will be interesting for those users who want to have a more in-depth understanding of the internals of Scilab. By the way, explicitly managing the memory is a crucial feature which allows to perform the most memory demanding computations. The commands which allow to manage variables and the memory are presented in figure 1.

In the first section, we analyze the management of the stack and present the **stacksize** function. Then we analyze the maximum available memory depending on the operating system. In the third section, we present the **who** function, which displays the list of current variables. We emphasize in the fourth section the use of portability variables and functions, which allows to design scripts which work equally well on most operating systems. Then we present the **clear** function, which allows to destroy an existing variable. In the final section, we present the **type** and **typeof** functions, which allows to dynamically compute the type of a variable.

| clear | kills variables |
|-------------|-----------------------------------|
| clearglobal | kills global variables |
| global | define global variable |
| isglobal | check if a variable is global |
| stacksize | set scilab stack size |
| gstacksize | set/get scilab global stack size |
| who | listing of variables |
| who_user | listing of user's variables |
| whos | listing of variables in long form |

Figure 1: Functions to manage variables.





Figure 2: The stack of Scilab.

2.1 The stack

In Scilab v5 (and previous versions), the memory is managed with a *stack*. At startup, Scilab allocates a fixed amount of memory to store the variables of the session. Some variables are already predefined at startup, which consumes a little amount of memory, but most of the memory is free and left for the user. Whenever the user defines a variable, the associated memory is consumed inside the stack, and the corresponding amount of memory is removed from the part of the stack which is free. This situation is presented in the figure 2. When there is no free memory left in the stack, the user cannot create a new variable anymore. At this point, the user must either destroy an existing variable, or increase the size of the stack.

There is always some confusion about bit and bytes, their symbols and their units. A bit (binary digit) is a variable which can only be equal to 0 or 1. A byte (denoted by B) is made of 8 bits. There are two different types of unit symbols for multiples of bytes. In the decimal unit system, one kilobyte is made of 1000 bytes, so that the symbols used are KB (10^3 bytes), MB (10^6 bytes), GB (10^9 bytes) and more (such as TB for terabytes and PB for petabytes). In the binary unit system, one kilobyte is made of 1024 bytes, so that the symbols are including a lower case letter "i" in their units: KiB, MiB, etc... In this document, we use only the decimal unit system.

The stacksize function allows to inquire about the current state of the stack. In the following session, executed after Scilab's startup, we call the stacksize in order to retrieve the current properties of the stack.

```
-->stacksize()
ans =
5000000. 33360.
```

The first number, 5 000 000, is the total number of 64 bits words which can be stored in the stack. The second number, 33 360, is the number of 64 bits words which are already used. That means that only 5 000 000 - 33 360 = 4 966 640 words of 64 bits are free for the user.

The number 5 000 000 is equal to the number of 64 bits double precision floating point numbers (i.e. "doubles") which could be stored, if the stack contained only that type of data. The total stack size 5 000 000 corresponds to 40 MB, because 5 000 000 * 8 = 40 000 000. This memory can be entirely filled with a dense square 2236-by-2236 matrix of doubles, because $\sqrt{5000000} \approx 2236$.

In fact, the stack is used to store both real values, integers, strings and more complex data structures as well. When a 8 bits integer is stored, this corresponds to 1/8th of the memory required to store a 64 bits word (because $8^*8 = 64$). In that situation, only 1/8th of the storage required to store a 64 bits word is used to store the integer. In general, the second integer is stored in the 2/8th of the same word, so that no memory is lost.

The default setting is probably sufficient in most cases, but might be a limitation for some applications.

In the following Scilab session, we show that creating a random 2300×2300 dense matrix generates an error, while creating a 2200×2200 matrix is possible.

```
-->A=rand(2300,2300)
!--error 17
rand: stack size exceeded
(Use stacksize function to increase it).
-->clear A
-->A=rand(2200,2200);
```

In the case where we need to store larger datasets, we need to increase the size of the stack. The **stacksize("max")** statement allows to configure the size of the stack so that it allocates the maximum possible amount of memory on the system. The following script gives an example of this function, as executed on a Gnu/Linux laptop with 1 GB memory. The **format** function is used so that all the digits are displayed.

```
-->format(25)
-->stacksize("max")
-->stacksize()
ans =
28176384. 35077.
```

We can see that, this time, the total memory available in the stack corresponds to 28 176 384 units of 64 bits words, which corresponds to 225 MB (because 28 176 384 * 8 = 225 411 072). The maximum dense matrix which can be stored is now 5308-by-5308 because $\sqrt{28176384} \approx 5308$.

In the following session, we increase the size of the stack to the maximum and create a 3000-by-3000 dense, square, matrix of doubles.

```
-->stacksize("max")
-->A=rand(3000,3000);
```

Let us consider a Windows XP 32 bits machine with 4 GB memory. On this machine, we have installed Scilab 5.2.2. In the following session, we define a 12 000-by-12 000 dense square matrix of doubles. This corresponds to approximately 1.2 GB of memory.

```
-->stacksize("max")

-->format(25)

-->stacksize()

ans =

152611536. 36820.

-->sqrt(152611536)

ans =

12353.604170443539260305

-->A=zeros(12000,12000);
```

We have given the size of dense matrices of doubles, in order to get a rough idea of what these numbers correspond to in practice. Of course, the user might still be able to manage much larger matrices, for example if they are sparse matrices. But, in any case, the total used memory can exceed the size of the stack.

Scilab version 5 (and before) can address $2^{31} \approx 2.1 \times 10^9$ bytes, i.e. 2.1 GB of memory. This corresponds to $2^{31}/8 = 268435456$ doubles, which could be filled by a 16 384-by-16 384 dense, square, matrix of doubles. This limitation is caused by the internal design of Scilab, whatever the operating system. Moreover, constraints imposed by the operating system may further limit that memory. These topics are detailed in the next section, which present the internal limitations of Scilab and the limitations caused by the various operating systems that Scilab can run on.

2.2 More on memory management

In this section, we present more details about the memory available in Scilab from a user's point of view. We separate the memory limitations caused by the design of Scilab on one hand, from the limitations caused by the design of the operating system on the other hand.

In the first section, we analyze the internal design of Scilab and the way the memory is managed by 32 bits signed integers. Then we present the limitation of the memory allocation on various 32 and 64 bits operating systems. In the last section, we present the algorithm used by the stacksize function.

These sections are rather technical and may be skipped by most users. But users who experience memory issues or who wants to know what are the exact design issues with Scilab v5 may be interested in knowing the exact reasons of these limitations.

2.2.1 Memory limitations from Scilab

Scilab version 5 (and before) addresses its internal memory (i.e. the stack) with 32 bits signed integers, whatever the operating system it runs on. This explains why the maximum amount of memory that Scilab can use is 2.1 GB. In this section, we

| Addr#0 | Addr#1 |
|--------|-------------------------|
| Addr#2 | Addr#3 |
| | |
| | |
| | |
| | |
| | Addr#2 ³¹ -1 |

Figure 3: Detail of the management of the stack. – Addresses are managed explicitly by the core of Scilab with 32 bits signed integers.

present how this is implemented in the most internal parts of Scilab which manage the memory.

In Scilab, a *gateway* is a C or Fortran function which provides a particular function to the user. More precisely, it connects the interpreter to a particular set of library functions, by reading the input arguments provided by the user and by writing the output arguments required by the user.

For example, let us consider the following Scilab script.

$$x = 3$$

 $y = sin(x)$

Here, the variables \mathbf{x} and \mathbf{y} are matrices of doubles. In the gateway of the sin function, we check the number of input arguments and their type. Once the variable \mathbf{x} is validated, we create the output variable \mathbf{y} in the stack. Finally, we call the sin function provided by the mathematical library and put the result into \mathbf{y} .

The gateway explicitly accesses to the addresses in the stack which contain the data for the x and y variables. For that purpose, the header of a Fortran gateway may contain a statement such as:

integer il

where il is a variable which stores the location in the stack which stores the variable to be used. Inside the stack, each address corresponds to one byte and is managed explicitly by the source code of each gateway. The integers represents various locations in the stack, i.e. various addresses of bytes. The figure 3 present the way that Scilab's v5 core manages the bytes in the stack. If the integer variable il is associated with a particular address in the stack, the expression il+1 identifies the next address in the stack.

Each variable is stored as a couple associating the header and the data of the variable. This is presented in the figure 4, which presents the detail of the header of a variable. The following Fortran source code is typical of the first lines in a legacy gateway in Scilab. The variable il contains the address of the beginning of the current variable, say x for example. In the actual source code, we have already checked that the current variable is a matrix of doubles, that is, we have checked



Figure 4: Detail of the management of the stack. – Each variable is associated with a header which allows to access to the data.

the type of the variable. Then, the variable **m** is set to the number of rows of the matrix of doubles and the variable **n** is set to the number of columns. Finally, the variable **it** is set to 0 if the matrix is real and to 1 if the matrix is complex (i.e. contains both a real and an imaginary part).

```
m=istk(il+1)
n=istk(il+2)
it=istk(il+3)
```

As we can see, we simply use expressions such as il+1 or il+2 to move from one address to the other, that is, from one byte to the next. Because of the integer arithmetic used in the gateways on integers such as il, we must focus on the range of values that can achieve this particular data type.

The Fortran integer data type is a signed 32 bits integer. A 32 bits signed integers ranges from $-2^{31} = 2147483648$ to $2^{31} - 1 = 2147483647$. In the core of Scilab, we do not use negative integer values to identifies the addresses inside the stack. This directly implies that no more that 2 147483647 bytes, i.e. 2.1 GB can be addressed by Scilab.

Because there are so many gateways in Scilab, it is not straightforward to move the memory management from a stack to a more dynamic memory allocation. This is the project of the version 6 of Scilab, which is currently in development and will appear in the coming years.

The limitations associated with various operating systems are analyzed in the next sections.

2.2.2 Memory limitations of 32 and 64 bit systems

In this section, we analyze the memory limitations of Scilab v5, depending on the version of the operating system where Scilab runs.

On 32 bit operating systems, the memory is addressed by the operating system with 32 bits unsigned integers. Hence, on a 32 bit system, we can address 4.2 GB, that is, $2^{32} = 4$ 294 967 296 bytes. Depending on the particular operating system

(Windows, Linux) and on the particular version of this operating system (Windows XP, Windows Vista, the version of the Linux kernel, etc...), this limit may or may not be achievable.

On 64 bits systems, the memory is addressed by the operating system with 64 bits unsigned integers. Therefore, it would seem that the maximum available memory in Scilab would be $2^{64} \approx 1.8 \times 10^{10}$ GB. This is larger than any available physical memory on the market (at the time this document is written).

But, be the operating system a 32 or a 64 bits, be it a Windows or a Linux, the stack of Scilab is still managed internally with 32 bits signed integers. Hence, no more than 2.1 GB of memory is usable for Scilab variables which are stored inside the stack.

In practice, we may experience that some particular linear algebra or graphical feature works on 64 bits systems while it does not work on a 32 bits system. This may be caused, by a temporary use of the operating system memory, as opposed to the stack of Scilab. For example, the developer may have used the malloc/free functions instead of using a part of the stack. It may also happen that the memory is not allocated by the Scilab library, but, at a lower level, by a sub-library used by Scilab. In some cases, that is sufficient to make a script pass on a 64 bits system, while the same script can fail on a 32 bits system.

2.2.3 The algorithm in stacksize

In this section, we present the algorithm used by the **stacksize** function to allocate the memory.

When the stacksize("max") statement is called, we first compute the size of the current stack. Then, we compute the size of the largest free memory region. If the current size of the stack is equal to the largest free memory region, we immediately return. If not, we set the stack size to the minimum, which allocates the minimum amount of memory which allows to save the currently used memory. The used variables are then copied into the new memory space and the old stack is de-allocated. Then we set the stack size to the maximum.

We have seen that Scilab can address $2^{31} \approx 2.1 \times 10^9$ bytes, i.e. 2.1 GB of memory. In practice, it might be difficult or even impossible to allocate such a large amount of memory. This is partly caused by limitations of the various operating systems on which Scilab runs, which is analyzed in the next section.

2.3 The list of variables and the function who

The following script shows the behavior of the **who** function, which shows the current list of variables, as well as the state of the stack.

-->who

Your variables are:

| WSCI | home | scinoteslib | modules_managerlib |
|------------------------|-----------------------|-------------|--------------------|
| atomsguilib | atomslib | matiolib | parameterslib |
| simulated_annealinglib | genetic_algorithmslib | umfpacklib | fft |
| scicos_autolib | scicos_utilslib | xcoslib | spreadsheetlib |
| demo_toolslib | development_toolslib | soundlib | texmacslib |

| SCI | the installation directory of the current Scilab installation |
|---------|---|
| SCIHOME | the directory containing user's startup files |
| MSDOS | true if the current operating system is Windows |
| TMPDIR | the temporary directory for the current Scilab's session |

Figure 5: Portability variables.

| compatibility_functilib | maple2scilablib | m2scilib | tclscilib |
|-------------------------|----------------------|------------------|----------------------------|
| stringlib | timelib | windows_toolslib | statisticslib w |
| %z | signal_processinglib | sparselib | special_functionslib |
| optimsimplexlib | overloadinglib | polynomialslib | %s |
| interpolationlib | optimizationlib | neldermeadlib | optimbaselib |
| iolib | output_streamlib | jvmlib | linear_algebralib |
| guilib | uitreelib | dynamic_linklib | integerlib |
| datatipslib | graphic_exportlib | cacsdlib | data_structureslib |
| elementary_functionslib | functionslib | fileiolib | graphicslib |
| PWD | corelib | helptoolslib | differential_equationlib |
| %F | MSDOS | %pvm | %tk |
| SCI | %inf | %nan | %т |
| %fftw | %gui | TMPDIR | SCIHOME |
| %eps | %f | %t | \$ |
| %pi | %e | %i | %io |
| | | 5000000. | using 7936 elements out of |
| | | f 9231. | and 80 variables out of |
| | | | Your global variables are: |
| %exportFileName | %driverName | demolist | %modalWarning |
| | | 999. | using 601 elements out of |
| | | f 767. | and 4 variables out of |

All the variables which names are ending with "lib" (as optimizationlib for example) are associated with Scilab's internal function libraries. Some variables starting with the "%" character, i.e. %i, %e and %pi, are associated with predefined Scilab variables because they are mathematical constants. Other variables which name start with a "%" character are associated with the precision of floating point numbers and the IEEE standard. These variables are %eps, %nan and %inf. Upcase variables SCI, SCIHOME, MSDOS and TMPDIR allow to create portable scripts, i.e. scripts which can be executed independently of the installation directory or the operating system. These variables are described in the next section.

2.4 Portability variables and functions

There are some predefined variables and functions which allow to design portable scripts, that is, scripts which work equally well on Windows, Linux or Mac. These variables are presented in the figures 5 and 6. These variables and functions are mainly used when creating external modules, but may be of practical value in a large set of situations.

In the following session, we check the values of some pre-defined variables on a Linux machine.

-->SCI SCI =

| [OS,Version]=getos() | the name of the current operating system |
|---------------------------------|--|
| <pre>f = fullfile(p1,p2,)</pre> | builds a file path from parts |
| s = filesep() | the Operating-System specific file separator |

Figure 6: Portability functions.

```
/home/myname/Programs/Scilab-5.1/scilab-5.1/share/scilab
-->SCIHOME
SCIHOME =
/home/myname/.Scilab/scilab-5.1
-->MSDOS
MSDOS =
F
-->TMPDIR
TMPDIR =
/tmp/SD_8658_
```

The TMPDIR variable, which contains the name of the temporary directory, is associated with the current Scilab session: each Scilab session has a unique temporary directory.

Scilab's temporary directory is created by Scilab at startup (and is not destroyed when Scilab quits). In practice, we may use the **TMPDIR** variable in test scripts where we have to create temporary files. This way, the file system is not polluted with temporary files and there is a little less chance to overwrite important files.

We often use the $\tt SCIHOME$ variable to locate the $\tt.startup$ file on the current machine.

In order to illustrate the use of the SCIHOME variable, we consider the problem of finding the .startup file of Scilab.

For example, we sometimes load manually some specific external modules at the startup of Scilab. To do so, we insert **exec** statements in the .startup file, which is automatically loaded at the Startup of Scilab. In order to open the .startup file, we can use the following statement.

```
editor(fullfile(SCIHOME,".scilab"))
```

Indeed, the fullfile function creates the absolute directory name leading from the SCIHOME directory to the .startup file. Moreover, the fullfile function automatically uses the directory separator corresponding to the current operating system: / under Linux and \ under Windows. The following session shows the effect of the fullfile function on a Windows operating system.

```
-->fullfile(SCIHOME,".scilab")
ans =
C:\DOCUME~1\Root\APPLIC~1\Scilab\scilab-5.3.0-beta-2\.scilab
```

The function **filesep** returns a string representing the directory separator on the current operating system. In the following session, we call the **filesep** function under Windows.

```
-->filesep()
ans =
\
```

Under Linux systems, the filesep function returns "/".

There are two features which allow to get the current operating system in Scilab: the getos function and the MSDOS variable. Both features allow to create scripts which manage particular settings depending on the operating system. The getos function allows to manage uniformly all operating systems, which leads to an improved portability. This is why we choose to present this function first.

The getos function returns a string containing the name of the current operating system and, optionally, its version. In the following session, we call the getos function on a Windows XP machine.

```
-->[OS,Version]=getos()
Version =
XP
OS =
Windows
```

The getos function may be typically used in select statements such as the following.

```
OS=getos()
select OS
case "Windows" then
   disp("Scilab on Windows")
case "Linux" then
   disp("Scilab on Linux")
case "Darwin" then
   disp("Scilab on MacOs")
else
   error("Scilab on Unknown platform")
end
```

A typical use of the MSDOS variable is presented in the following script.

```
if ( MSDOS ) then
   // Windows statements
else
   // Linux statements
end
```

In practice, consider the situation where we want to call an external program with the maximum possible portability. Assume that, under Windows, this program is provided by a ".bat" script while, under Linux, this program is provided by a ".sh" script. In this situation, we might write a script using the MSDOS variable and the unix function, which executes an external program. Despite its name, the unix function works equally under Linux and Windows.

```
if ( MSDOS ) then
   unix("myprogram.bat")
else
   unix("myprogram.sh")
end
```

The previous example shows that it is possible to write a portable Scilab program which works in the same way under various operating systems. The situation might be more complicated, because it often happens that the path leading to the program is also depending on the operating system. Another situation is when we want to compile a source code with Scilab, using, for example, the ilib_for_link or the tbx_build_src functions. In this case, we might want to pass to the compiler some particular option which depends on the operating system. In all these situations, the MSDOS variable allows to make one single source code which remains portable across various systems.

The SCI variable allows to compute paths relative to the current Scilab installation. The following session shows a sample use of this variable. First, we get the path of a macro provided by Scilab. Then, we combine the SCI variable with a relative path to the file and pass this to the 1s function. Finally, we concatenate the SCI variable with a string containing the relative path to the script and pass it to the editor function. In both cases, the commands do not depend on the absolute path to the file, which make them more portable.

```
-->get_function_path("numdiff")
ans =
/home/myname/Programs/Scilab-5.1/scilab-5.1/...
share/scilab/modules/optimization/macros/numdiff.sci
-->filename=fullfile(SCI,"modules","optimization",..
-->"macros","numdiff.sci");
-->ls(filename)
ans =
/home/myname/Programs/Scilab-5.1/scilab-5.1/...
share/scilab/modules/optimization/macros/numdiff.sci
-->editor(filename)
```

In common situations, it often seems to be more simple to write a script only for a particular operating system, because this is the one that we use at the time that we write it. In this case, we tend to use tools which are not available on other operating systems. For example, we may rely on the particular location of a file, which can be found only on Linux. Or we may use some OS-specific shell or file management instructions. In the end, a useful script has a high probability of being used on an operating system which was not the primary target of the original author. In this situation, a lot of time is wasted in order to update the script so that it can work on the new platform.

Since Scilab is itself a portable language, it is most of the time a good idea to think of the script as being portable right from the start of the development. If there is a really difficult and unavoidable issue, it is of course reasonable to reduce the portability and to simplify the task so that the development is shortened. In practice, this happens not so often as it appears, so that the usual rule should be to write as portable a code as possible.

2.5 Destroying variables : clear

A variable can be created at will, when it is needed. It can also be destroyed explicitly with the **clear** function, when it is not needed anymore. This might be useful when a large matrix is currently stored in memory and if the memory is becoming a problem.

In the following session, we define a large random matrix A. When we try to define a second matrix B, we see that there is no memory left. Therefore, we use the clear function to destroy the matrix A. Then we are able to create the matrix B.

type Returns a string
typeof Returns a floating point integer

Figure 7: Functions to compute the type of a variable.

```
-->A = rand(2000,2000);

-->B = rand(2000,2000);

!--error 17

rand: stack size exceeded

(Use stacksize function to increase it).

-->clear A

-->B = rand(2000,2000);
```

The clear function should be used only when necessary, that is, when the computation cannot be executed because of memory issues. This warning is particularly true for the developers who are used to compiled languages, where the memory is managed explicitly. In the Scilab language, the memory is managed by Scilab and, in general, there is no reason to manage it ourselves.

An associated topic is the management of variables in functions. When the body of a function has been executed by the interpreter, all the variables which have been used in the body ,and which are not output arguments, are deleted automatically. Therefore, there is no need for explicit calls to the **clear** function in this case.

2.6 The type and type of functions

Scilab can create various types of variables, such as matrices, polynomials, booleans, integers and other types of data structures. The type and typeof functions allow to inquire about the particular type of a given variable. The type function returns a floating point integer while the typeof function returns a string. These functions are presented in the figure 7.

The figure 8 presents the various output values of the type and typeof functions.

In the following session, we create a 2×2 matrix of doubles and use the type and typeof to get the type of this matrix.

```
-->A=eye(2,2)
A =
1. 0.
0. 1.
-->type(A)
ans =
1.
-->typeof(A)
ans =
constant
```

These two functions are useful when processing the input arguments of a function. This topic will be reviewed later in this document, in the section 4.2.5, when we consider functions with variable type input arguments.

When the type of the variable is a tlist or a mlist, the value returned by the typeof function is the first string in the first entry of the list. This topic is reviewed

| type | typeof | Detail |
|------|---------------------------|--|
| 1 | "constant" | real or complex constant matrix |
| 2 | "polynomial" | polynomial matrix |
| 4 | "boolean" | boolean matrix |
| 5 | "sparse" | sparse matrix |
| 6 | "boolean sparse" | sparse boolean matrix |
| 7 | "Matlab sparse" | Matlab sparse matrix |
| 8 | "int8", "int16", | matrix of integers stored |
| | "int32","uint8", | on $1 2$ or 4 bytes |
| | "uint16" or "uint32" | |
| 9 | "handle" | matrix of graphic handles |
| 10 | "string" | matrix of character strings |
| 11 | "function" | un-compiled function (Scilab code) |
| 13 | "function" | compiled function (Scilab code) |
| 14 | "library | function library |
| 15 | "list" | list |
| 16 | "rational", "state-space" | typed list (tlist) |
| | or the type | |
| 17 | "hypermat", "st", "ce" | matrix oriented typed list (mlist) |
| | or the type | |
| 128 | "pointer" | sparse matrix LU decomposition |
| 129 | "size implicit" | size implicit polynomial used for indexing |
| 130 | "fptr" | Scilab intrinsic (C or Fortran code) |

Figure 8: The returned values of the type and typeof function.

in the section 3.6, where we present typed lists.

The data types cell and struct are special forms of mlists, so that they are associated with a type equal to 17 and with a typeof equal to "ce" and "st".

2.7 Notes and references

It is likely that, in future versions of Scilab, the memory of Scilab will not be managed with a stack. Indeed, this features is a legacy of the ancestor of Scilab, that is Matlab. We emphasize that Matlab does not use a stack since a long time, approximately since the 1980s, at the time where the source code of Matlab was re-designed[41]. On the other hand, Scilab kept this rather old way of managing its memory.

Some additional details about the management of memory in Matlab are given in [34]. In the technical note [35], the authors present methods to avoid memory errors in Matlab. In the technical note [36], the authors present the maximum matrix size available in Matlab on various platforms. In the technical note [37], the authors presents the benefits of using 64-bit Matlab over 32-bit Matlab.

2.8 Exercises

Exercise 2.1 (*Maximum stack size*) Check the maximum size of the stack on your current machine.

Exercise 2.2 (who_user) Start Scilab, execute the following script and see the result on your machine.

```
who_user()
A=ones(100,100);
who_user()
```

Exercise 2.3 (whos) Start Scilab, execute the following script and see the result on your machine.

whos()

3 Special data types

In this section, we analyze Scilab data types which are the most commonly used in practice. We review strings, integers, polynomials, hypermatrices, lists and tlists. We present some of the most useful features of the overloading system, which allows to define new behaviors for typed lists. In the last section, we briefly review the cell, the struct and the mlist, and compare them with other data structures.

3.1 Strings

Although Scilab is not primarily designed as a tool to manage strings, it provides a consistent and powerful set of functions to manage this data type. A list of commands which are associated with Scilab strings is presented in figure 9.

In order to create a matrix of strings, we can use the " " "character and the usual syntax for matrices. In the following session, we create a 2×3 matrix of strings.

```
-->x = ["11111" "22" "333"; "4444" "5" "666"]
x =
!11111 22 333 !
! ! !
!4444 5 666 !
```

In order to compute the size of the matrix, we can use the **size** function, as for usual matrices.

```
-->size(x)
ans =
2. 3.
```

The length function, on the other hand, returns the number of characters in each of the entries of the matrix.

```
-->length(x)
ans =
5. 2. 3.
4. 1. 3.
```

| string | conversion to string |
|------------------------------|--|
| sci2exp | converts an expression to a string |
| ascii | string ascii conversions |
| blanks | create string of blank characters |
| $\operatorname{convstr}$ | case conversion |
| emptystr | zero length string |
| grep | find matches of a string in a vector of strings |
| justify | justify character array |
| length | length of object |
| part | extraction of strings |
| regexp | find a substring that matches the regular expression string |
| strcat | concatenate character strings |
| strchr | find the first occurrence of a character in a string |
| strcmp | compare character strings |
| $\operatorname{strcmpi}$ | compare character strings (case independent) |
| $\operatorname{strcspn}$ | get span until character in string |
| $\operatorname{strindex}$ | search position of a character string in an other string |
| $\operatorname{stripblanks}$ | strips leading and trailing blanks (and tabs) of strings |
| $\operatorname{strncmp}$ | copy characters from strings |
| strrchr | find the last occurrence of a character in a string |
| strrev | returns string reversed |
| $\operatorname{strsplit}$ | split a string into a vector of strings |
| strspn | get span of character set in string |
| strstr | locate substring |
| $\operatorname{strsubst}$ | substitute a character string by another in a character string |
| strtod | convert string to double |
| strtok | split string into tokens |
| tokenpos | returns the tokens positions in a character string |
| tokens | returns the tokens of a character string |
| str2code | return Scilab integer codes associated with a character string |
| code2str | returns character string associated with Scilab integer codes |

Figure 9: Scilab string functions.

Perhaps the most common string function that we use is the **string** function. This function allows to convert its input argument into a string. In the following session, we define a row vector and use the **string** function to convert it into a string. Then we use the **typeof** function and check that the **str** variable is indeed a string. Finally, we use the **size** function and check that the **str** variable is a 1×5 matrix of strings.

```
-->x = [1 2 3 4 5];

-->str = string(x)

str =

!1 2 3 4 5 !

-->typeof(str)

ans =

string

-->size(str)

ans =

1. 5.
```

The string function can take any type of input argument.

We will see later in this document that a tlist can be used to define a new data type. In this case, we can define a function which makes so that the string function can work on this new data type in a user-defined way. This topic is reviewed in the section 3.8, where we present a method which allows to define the behavior of the string function when its input argument is a typed list.

The strcat function concatenates its first input argument with the separator defined in the second input argument. In the following session, we use the strcat function to produce a string representing the sum of the integers from 1 to 5.

```
-->strcat(["1" "2" "3" "4" "5"],"+")
ans =
1+2+3+4+5
```

We may combine the string and the strcat functions to produce strings which can be easily copied and pasted into scripts or reports. In the following session, we define the row vector \mathbf{x} which contains floating point integers. Then we use the strcat function with the blank space separator to produce a clean string of integers.

```
-->x = [1 2 3 4 5]

x =

1. 2. 3. 4. 5.

-->strcat(string(x)," ")

ans =

1 2 3 4 5
```

The previous string can be directly copied and pasted into a source code or a report. Let us consider the problem of designing a function which prints data in the console. The previous combination of function is an efficient way of producing compact messages. In the following session, we use the mprintf function to display the content of the x variable. We use the %s format, which corresponds to strings. In order to produce the string, we combine the strcat and string functions.

```
-->mprintf("x=[%s]\n",strcat(string(x)," "))
x=[1 2 3 4 5]
```

| isalphanum | check if characters are alphanumerics |
|------------|--|
| isascii | check if characters are 7-bit US-ASCII |
| isdigit | check if characters are digits between 0 and 9 |
| isletter | check if characters are alphabetics letters |
| isnum | check if characters are numbers |

Figure 10: Functions related to particular class of strings.

The sci2exp function converts an expression into a string. It can be used with the same purpose as the previous method based on strcat, but the formatting is less flexible. In the following session, we use the sci2exp function to convert a row matrix of integers into a 1×1 matrix of strings.

```
-->x = [1 2 3 4 5];

-->str = sci2exp(x)

str =

[1,2,3,4,5]

-->size(str)

ans =

1. 1.
```

Comparison operators, such as "i" or "i" for example, are not defined when strings are used, i.e., the statement "a" < "b" produces an error. Instead, the strcmp function can be used for that purpose. It returns 1 if the first argument is lexicographically less than the second, it returns 0 if the two strings are equal, or it returns -1 if the second argument is lexicographically less than the first. The behavior of the strcmp function is presented in the following session.

```
-->strcmp("a","b")

ans =

- 1.

-->strcmp("a","a")

ans =

0.

-->strcmp("b","a")

ans =

1.
```

The functions presented in the table 10 allow to distinguish between various classes of strings such as ASCII characters, digits, letters and numbers.

For example, the isdigit function returns a matrix of booleans, where each entry i is true if the character at index i in the string is a digit. In the following session, we use the isdigit function and check that "0" is a digit, while "d" is not.

```
-->isdigit("0")

ans =

T

-->isdigit("12")

ans =

T T

-->isdigit("d3s4")

ans =

F T F T
```

| polynomial | a polynomial, defined by its coefficients |
|------------|---|
| rational | a ratio of two polynomials |

Figure 11: Data types related to polynomials.

A powerful regular expression engine is available from the **regexp** function. This feature has been included in Scilab version 5. It is based on the PCRE library [22], which aims at being PERL-compatible. The pattern must be given as a string, with surrounding slashes of the form "/x/", where x is the regular expression.

We present a sample use of the **regexp** function in the following session. The **i** at the end of the regular expression indicates that we do not want to take the case into account. The first **a** letter forces the expression to match only the strings which begins with this letter.

```
-->regexp("AXYZC","/a.*?c/i")
ans =
1.
```

Regular expressions are an extremely powerful tool for manipulating text and data. Obviously, this document cannot even scratch the surface of this topic. For more informations about the **regexp** function, the user may read the help page provided by Scilab. For a deeper introduction, the reader may be interested in Friedl's [20]. As expressed by J. Friedl, "regular expressions allow you to code complex and subtle text processing that you never imagined could be automated".

The exercise 3.1 presents a practical example of use of the regexp function.

3.2 Polynomials

Scilab allows to manage univariate polynomials. The implementation is based on a vector containing the coefficients of the polynomial. At the user's level, we can manage a matrix of polynomials. Basic operations like addition, subtraction, multiplication and division are available for polynomials. We can, of course, compute the value of a polynomial p(x) for a particular input x. Moreover, Scilab can perform higher level operations, such as computing the roots, factoring or computing the greatest common divisor or the least common multiple of two polynomials. When we divide one polynomial by another polynomial, we obtain a new data structure representing the rational function.

In this section, we make a brief review of these topics. The polynomial and rational data types are presented in the figure 11. Some of the most common functions related to polynomials are presented in the figure 12. A complete list of functions related to polynomials is presented in figure 13.

The poly function allows to define polynomials. There are two ways to define them: by its coefficients, or by its roots. In the following session, we create the polynomial p(x) = (x-1)(x-2) with the poly function. The roots of this polynomial are obviously x = 1 and x = 2 and this is why the first input argument of the poly function is the matrix [1 2]. The second argument is the symbolic string used to display the polynomial.

| poly | defines a polynomial |
|---------|------------------------------------|
| horner | computes the value of a polynomial |
| coeff | coeffcients of a polynomial |
| degree | degree of a polynomial |
| roots | roots of a polynomial |
| factors | real factorization of polynomials |
| gcd | greatest common divisor |
| lcm | least common multiple |

Figure 12: Some basic functions related to polynomials.

| bezout | clean | cmndred | coeff | coffg | colcompr | degree |
|--------|----------|---------|---------|-----------|----------|---------|
| denom | derivat | determ | detr | diophant | factors | gcd |
| hermit | horner | hrmt | htrianr | invr | lcm | lcmdiag |
| ldiv | numer | pdiv | pol2des | pol2str | polfact | residu |
| roots | rowcompr | sfact | simp | simp_mode | sylm | systmat |

Figure 13: Functions related to polynomials.

```
-->p=poly([1 2],"x")
p = 2
2 - 3x + x
```

In the following session, we call the typeof function and check that the variable p is a polynomial.

```
-->typeof(p)
ans =
polynomial
```

We can also define a polynomial based on its coefficients, in increasing order. In the following session, we define the q(x) = 1 + 2x polynomial. We pass the third argument "coeff" to the poly function, so that it knows that the [1 2] matrix represents the coefficients of the polynomial.

```
-->q=poly([1 2],"x","coeff")
q =
1 + 2x
```

Now that the polynomials p and q are defined, we can perform algebra with them. In the following session, we add, subtract, multiply and divide the polynomials p and q.

```
-->p+q
ans =
2
3 - x + x
-->p-q
ans =
2
1 - 5x + x
```

When we divide the polynomial p by the polynomial q, we produce the rational function r. This is illustrated in the following session.

```
-->typeof(r)
ans =
rational
```

In order to compute the value of a polynomial p(x) for a particular value of x, we can use the **horner** function. In the following session, we define the polynomial p(x) = (x - 1)(x - 2) and compute its value for the points x = 0, x = 1, x = 3 and x = 3, represented by the matrix [0 1 2 3].

```
-->p=poly([1 2],"x")

p = 2

2 - 3x + x

-->horner(p,[0 1 2 3])

ans =

2. 0. 0. 2.
```

The name of the **horner** function comes from the mathematician Horner, who designed the algorithm which is used in Scilab to compute the value of a polynomial. This algorithm reduces the number of multiplications and additions required for this evaluation (see [26], section 4.6.4, "Evaluation of Polynomials").

If the first argument of the poly function is a square matrix, it returns the characteristic polynomial associated with the matrix. That is, if A is a real $n \times n$ square matrix, the poly function can produce the polynomial det(A - xI) where I is the $n \times n$ identity matrix. This is presented in the following session.

```
-->A = [1 2;3 4]
A =
1. 2.
3. 4.
-->p = poly(A,"x")
p =
2
- 2 - 5x + x
```

We can easily check that the previous result is consistent with its mathematical definition. First, we can compute the roots of the polynomial **p** with the **roots** function, as in the previous session. On the other hand, we can compute the eigenvalues of the matrix **A** with the **spec** function.

```
- 0.3722813
5.3722813
-->spec(A)
ans =
- 0.3722813
5.3722813
```

There is another way to get the same polynomial \mathbf{p} . In the following session, we define the polynomial \mathbf{px} which represents the monomial x. Then we use the det function to compute the determinant of the matrix B = A - xI. We use the eye function to produce the 2×2 identity matrix.

```
->px = poly([0 1],"x","coeff")

px =

x

-->B = A-px*eye()

B =

1 - x 2

3 4 - x

-->det(B)

ans =

2

- 2 - 5x + x
```

If we compare this result to the previous one, we see that they are equal. We see that the det function is defined for the polynomial A-px*eye(). This is an example of the fact that many functions are defined for polynomial input arguments.

In the previous session, the matrix B was a 2×2 matrix of polynomials. Similarly, we can define a matrix of rational functions, as presented in the following session.

```
-->x=poly(0,"x");
-->A = [1 x; x 1+x^2];
-->B=[1/x 1/(1+x);1/(1+x) 1/x^2]
В
            =
!
    1
                  1
                         i
                ____
  ____
                        1
!
Т
                1 + x
    х
Т
I.
      1
                1
                         I
               _ _ _
                         I
                 2
                         I
Т
Т
    1 + x
                x
```

This is a very useful feature of Scilab for systems theory. The link between polynomials, rational functions data types on one hand, and control theory on the other hand, is analyzed briefly in the section 3.14.

3.3 Hypermatrices

A matrix is a data structure which can be accessed with two integer indices (e.g. i and j). Hypermatrices are a generalized type of matrices, which can be addressed with more than two indices. This feature is familiar to Fortran developers, where the multi-dimensional array is one of the basic data structures. Several functions related to hypermatrices are presented in the figure 14.

| hypermat | Creates an hypermatrix |
|----------|---|
| zeros | Creates an matrix or hypermatrix of zeros |
| ones | Creates an matrix or hypermatrix of ones |
| matrix | Create a matrix with new shape |
| squeeze | Remove dimensions with unit size |

Figure 14: Functions related to hypermatrices.

In most situations, we can manage an hypermatrix as a regular matrix. In the following session, we create the $4 \times 3 \times 2$ hypermatrix of doubles A with the ones function. Then, we use the **size** function to compute the size of this hypermatrix.

```
-->A=ones(4,3,2)
 A =
(:,:,1)
                    1.
            1.
    1.
    1.
            1.
                    1.
            1.
    1.
                    1.
    1.
            1.
                    1.
(:,:,2)
    1.
            1.
                    1.
    1.
            1.
                    1.
    1.
            1.
                    1.
    1.
            1.
                    1.
-->size(A)
 ans
      =
                    2.
    4.
            З.
```

In the following session, we create a 4-by-2-by-3 hypermatrix. The first argument of the hypermat function is a matrix containing the number of dimensions of the hypermatrix.

```
-->A=hypermat([4,3,2])
 A =
(:,:,1)
    Ο.
            0.
                   0.
    Ο.
            Ο.
                   0.
    0.
            0.
                   0.
    0.
            Ο.
                   0.
(:,:,2)
            Ο.
                   0.
    0.
    Ο.
            Ο.
                   0.
    0.
            0.
                   0.
    0.
            Ο.
                   0.
```

To insert and to extract a value from an hypermatrix, we can use the same syntax as for a matrix. In the following session, we set and get the value of the (3,1,2) entry.

```
-->A(3,1,2)=7
A =
(:,:,1)
```

Ο. 0. 0. Ο. 0. 0. 0. 0. 0. 0. 0. 0. (:,:,2)0. 0. 0. Ο. 0. 0. 7. 0. 0. Ο. Ο. 0. -->A(3,1,2)ans = 7.

The colon ":" operator can be used for hypermatrix, as in the following session.

```
-->A(2,:,2)
ans =
0. 0. 0.
```

Most operations which can be done with matrices can also be done with hypermatrices. In the following session, we define the hypermatrix B and add it to A.

```
-->B=2 * ones(4,3,2);
-->A + B
 ans =
(:,:,1)
    3.
            З.
                    З.
    З.
            З.
                    З.
    З.
            З.
                    З.
    З.
            З.
                    З.
(:,:,2)
            З.
                    З.
    З.
    З.
            З.
                    З.
    З.
            З.
                    З.
    3.
            З.
                    З.
```

The hypermat function can be used when we want to create an hypermatrix from a vector. In the following session, we define an hypermatrix with size $2 \times 3 \times 2$, where the values are taken from the set $\{1, 2, ..., 12\}$.

```
-->hypermat([2 3 2],1:12)
 ans =
(:,:,1)
    1.
           З.
                  5. !
1
           4.
                  6. !
!
    2.
(:,:,2)
1
    7.
           9.
                   11. !
    8.
           10.
                   12. !
```

We notice the particular order of the values in the produced hypermatrix. This order correspond to the rule that the leftmost indices vary first. This is just an extension of the fact that, for 2 dimensional matrices, the values are stored column-by-column.

An hypermatrix can also contain strings, as shown in the following session.

```
-->A=hypermat([3,1,2],["a","b","c","d","e","f"])
A =
(:,:,1)
```

!a 1 ! i ! b L 1 T ! c (:,:,2)!d Ţ 1 ļ ! e Ţ ! 1 !f Ţ

An important property of hypermatrices is that all entries must have the same type. For example, if we try to insert a double into the hypermatrix of strings created previously, we get an error message such as the following.

3.4 Type and dimension of an extracted hypermatrix

In this section, we present the extraction of hypermatrices for which one dimension has a unit size. We present the effect that this may have on performances and describe the squeeze function.

We may extract slices of hypermatrices, producing various shapes of output variables. For example, let us consider a 3 dimensional hypermatrix, with size 2-by-4-by-3. It is sometimes useful to use the colon : operator to extract full rows or columns of matrices or hypermatrices. For example, the statement A(1,:,:) extracts the values associated with the first index equal to 1, and produces an hypermatrix with size 1-by-4-by-3. Similarly, the statement A(:,1,:) extracts the values associated with the second index equal to 1, and produces an hypermatrix with size 2-by-1-by-3. On the other hand, the statement A(:,:,1) extracts the values associated with the third index equal to 1, but produces a matrix with size 4-by-3. Hence, the statement A(:,:,1) does not produce an hypermatrix, but produces a matrix, that is, the type of the variable changes. Moreover, we see that the last dimension, which should be equal to one, is just removed from the output variable. Hence, both the type and the shape of an extracted hypermatrix can be expected to change, depending on the indices that we extract.

This fact is explored in the following session.

```
-->A=hypermat([2,4,3],[1:24])
 A =
(:,:,1)
                          7.
    1.
           З.
                   5.
    2.
           4.
                   6.
                          8.
(:,:,2)
                     13.
    9.
            11.
                             15.
```

| 10. | 12. | 14. | 16. |
|---------|-----|-----|-----|
| (:,:,3) | | | |
| 17. | 19. | 21. | 23. |
| 18. | 20. | 22. | 24. |

We first experiment the extraction statement B=A(1,:,:).

```
-->B = A(1, :, :)
 в =
(:,:,1)
            З.
                   5.
                          7.
    1.
(:,:,2)
    9.
            11.
                    13.
                             15.
(:,:,3)
    17.
             19.
                     21.
                              23.
-->typeof(B)
 ans =
 hypermat
-->size(B)
 ans
            4.
                   З.
    1.
```

We may experiment the statement B=A(:,1,:) and we would get a similar behavior. Instead, we experiment the statement B=A(:,:,1). We see that, in this case, the extracted variable is a matrix, instead of an hypermatrix.

```
-->B = A(:,:,1)
 в =
    1.
           З.
                  5.
                          7.
    2.
           4.
                  6.
                          8.
-->typeof(B)
 ans
      =
 constant
-->size(B)
 ans =
    2.
           4.
```

The general rule is that, if the last dimension of an extracted hypermatrix is unity, it is removed. This rule can be applied, again, on the result of the extraction, eventually producing either an hypermatrix, or a regular 2D matrix. More precisely, when we extract an hypermatrix with shape n_1 -by-...-by- n_j -by- n_{j+1} -by-...-by- n_k , where $n_j \neq 1$ and $n_{j+1} = n_{j+2} = \dots = n_k = 1$, we get an hypermatrix with shape n_1 -by-...-by- n_j . Moreover, if the extracted hypermatrix is a regular, i.e. two dimensional, matrix, that is, if $j \leq 2$, therefore the extracted variable is a matrix instead of an hypermatrix. These two rules ensures that Scilab is compatible with Matlab on the extraction of hypermatrices.

This behavior may have a significant impact on performances of hypermatrices extraction. In the following script, we create a 2-by-4-by-3 hypermatrix of doubles and measure the performances of the extraction of either the second or the third index. We combine the extraction and the matrix statement, in order to produce a regular 2-dimensional matrix B in both cases. Since this extraction is extremely fast, we do this 100 000 times within a loop and measure the user time with the tic and toc functions. We can see that extracting the second dimension is much slower than extracting the third dimension.

```
-->A=hypermat([2,4,3],1:24);
-->B=matrix(A(:,2,:),2,3)
 В
   =
    3.
          11.
                  19.
    4.
          12.
                  20
-->tic();for i=1:100000;B=matrix(A(:,2,:),2,3);end;toc()
 ans =
    7.832
-->B=matrix(A(:,:,2),2,4)
B =
    9.
           11.
                   13.
                          15.
    10.
           12.
                   14.
                          16.
-->tic();for i=1:100000;B=matrix(A(:,:,2),2,4);end;toc()
 ans =
    0.88
```

The reason for this performance difference is that A(:,2,:) is a 2-by-1-by-3 hypermatrix while A(:,:,2) is a regular 2-by-4 matrix.

Indeed, the statement matrix(A(:,2,:),2,3) makes the matrix function to convert the 2-by-1-by-3 hypermatrix A(:,2,:) into a 2-by-3 matrix, which requires an extra step from the interpreter. On the other hand, the statement A(:,:,2) is already a 2-by-4 matrix. Hence, the statement matrix(A(:,:,2),2,4) does not require any processing from the matrix function, which is a no-op in this case.

We may want to just ignore intermediate sub-matrices with size 1 which are created by the hypermatrix extraction system. In this case, we can use the squeeze function, which removes dimensions with unit size.

```
-->A=hypermat([3,1,2],1:6)
A =
(:,:,1)
    1.
    2.
    3.
(:,:,2)
    4.
    5.
    6.
-->size(A)
 ans =
    З.
           1.
                  2.
-->B=squeeze(A)
 R
   =
           4.
    1.
    2.
           5.
    З.
           6.
-->size(B)
 ans
     =
    3.
           2.
```

3.5 The list data type

In this section, we describe the list data type, which is used to manage a collection of objects of different types. We often use lists when we want to gather in the same

| list | create a list |
|--------|---|
| null | delete the element of a list |
| lstcat | concatenate lists |
| size | for a list, the number of elements (for a list, same as length) |

Figure 15: Functions related to lists.

object a set of variables which cannot be stored into a single, more basic, data type. A list can contain any of the already discussed data types (including functions) as well as other lists. This allows to create nested lists, which can be used to create a tree of data structures. Lists are extremely useful to define structured data objects. Some functions related to lists are presented in the figure 15.

There are, in fact, various types of lists in Scilab: ordinary lists, typed lists and mlists. This section focuses on ordinary lists. Typed lists will be reviewed in the next section. The mlist data type is not presented in this document.

In the following session, we define a floating point integer, a string and a matrix. Then we use the list function to create the list mylist containing these three elements.

Once created, we can access to the i-th element of the list with the mylist(i) statement, as in the following session.

```
-->mylist(1)

ans =

12.

->mylist(2)

ans =

foo

-->mylist(3)

ans =

1. 2. 3. 4.
```

The number of elements in a list can be computed with the size function.

```
-->size(mylist)
ans =
3.
```

In the case where we want to get several elements in the same statement, we can use the colon ":" operator. In this situation, we must set as many output arguments as there are elements to retrieve. In the following session, we get the two elements at indices 2 and 3 and set the s and m variables.

-->[s,m] = mylist(2:3) m = 1. 2. 3. 4. s = foo

The element #3 in our list is a matrix. Suppose that we want to get the 4th value in this matrix. We can have access to it directly with the following syntax.

```
-->mylist(3)(4)
ans =
4.
```

This is much faster than storing the third element of the list into a temporary variable and to extract its 4th component. In the case where we want to set the value of this entry, we can use the same syntax and the regular equal "=" operator, as in the following session.

```
-->mylist(3)(4) = 12

mylist =

mylist(1)

12.

mylist(2)

foo

mylist(3)

1. 2. 3. 12.
```

Obviously, we could have done the same operation with several intermediate operations: extracting the third element of the list, updating the matrix and storing the matrix into the list again. But using the mylist(3)(4) = 12 statement is in fact both simpler and faster.

In order to browse the elements of a list, we can use a straightforward method or a method based on a particular feature of the **for** statement. First, the straightforward method is to count the number of elements in the list with the **size** function. Then we access to the elements, one by one, as in the following session.

```
for i = 1:size(mylist)
  e = mylist(i);
  mprintf("Element #%d: type=%s.\n",i,typeof(e))
end
```

The previous script produces the following output.

```
Element #1: type=constant.
Element #2: type=string.
Element #3: type=constant.
```

There is a simpler way, which uses directly the list as the argument of the for statement.

```
-->for e = mylist
--> mprintf("Type=%s.\n",typeof(e))
-->end
Type=constant.
Type=string.
Type=constant.
```

| tlist | create a typed list |
|---------------|--|
| typeof | get the type of the given typed list |
| fieldnames | returns all the fields of a typed list |
| definedfields | returns all the fields which are defined |
| setfield | set a field of a typed list |
| getfield | get a field of a typed list |

Figure 16: Functions related to tlists.

We can fill a list dynamically by appending new elements at the end. In the following script, we define an empty list with the list() statement. Then we use the \$+1 operator to insert new elements at the end of the list. This produces exactly the same list as previously.

```
mylist = list();
mylist($+1) = 12;
mylist($+1) = "foo";
mylist($+1) = [1 2 3 4];
```

3.6 The tlist data type

Typed lists allow to define new data structures which can be customized depending on the particular problem to be solved. These new data structures behave like basic Scilab data types. In particular, any regular function such as size, disp or string can be overloaded so that it has a particular behavior when its input argument is the new tlist. This allows to extend the features provided by Scilab and to introduce new objects. Actually, typed list are also used internally by numerous Scilab functions because of their flexibility. Most of the time, we do not even know that we are using a list, which shows how much this data type is convenient.

In this section, we will create and use directly a tlist to get familiar with this data structure. In the next section, we will present a framework which allows to emulate object oriented programming with typed lists.

The figure 16 presents all the functions related to typed lists.

In order to create a typed list, we use the tlist function, which first argument is a matrix of strings. The first string is the *type* of the list. The remaining strings define the *fields* of the typed list. In the following session, we define a typed list which allows to store the informations of a person. The fields of a **person** are the first name, the name and the birth year.

At this point, the person **p** is created, but its fields are undefined. In order to set the fields of **p**, we use the dot ".", followed by the name of the field. In the following script, we define the three fields associated with an hypothetical Paul Smith, which birth year is 1997.

```
p.firstname = "Paul";
p.name = "Smith";
p.birthyear = 1997;
```

All the fields are now defined and we can use the variable name **p** in order to see the content of our typed list, as in the following session.

```
-->p
p =
    p(1)
!person firstname name birthyear !
    p(2)
Paul
    p(3)
Smith
    p(4)
    1997.
```

In order to get a field of the typed list, we can use the p(i) syntax, as in the following session.

```
-->p(2)
ans =
Paul
```

But it may be more convenient to get the value of the field firstname with the p.firstname syntax, as in the following session.

```
-->fn = p.firstname
fn =
Paul
```

We can also use the **getfield** function, which takes as input arguments the index of the field and the typed list.

```
->fn = getfield(2,p)
fn =
Paul
```

The same syntax can be used to set the value of a field. In the following session, we update the value of the first name from "Paul" to "John".

We can also use the **setfield** function to set the first name field from "John" to "Ringo".

```
-->setfield(2,"Ringo",p)
-->p
p =
p(1)
```

```
!person firstname name birthyear !
    p(2)
Ringo
    p(3)
Smith
    p(4)
    1997.
```

It might happen that we know the values of the fields at the time when we create the typed list. We can append these values to the input arguments of the tlist function, in a consistent order. In the following session, we define the person **p** and set the values of the fields at the creation of the typed list.

```
-->p = tlist( .. 
-->
     ["person","firstname","name","birthyear"], ...
-->
     "Paul", ..
     "Smith", ..
-->
-->
     1997)
 р
       p(1)
!person firstname
                      name
                            birthyear
                                         !
       p(2)
 Paul
       p(3)
 Smith
       p(4)
    1997.
```

An interesting feature of a typed list is that the typeof function returns the actual type of the list. In the following session, we check that the type function returns 16, which corresponds to a list. But the typeof function returns the string "person".

```
-->type(p)
ans =
16.
-->typeof(p)
ans =
person
```

This allows to dynamically change the behavior of functions for the typed lists with type "person". This feature is linked to the overloading of functions, a topic which will be reviewed in the section 3.8.

We will now consider functions which allow to dynamically retrieve informations about typed lists. In order to get the list of fields of a "person", we can use the p(1) syntax and get a 1×4 matrix of strings, as in the following session.

```
-->p(1)
ans =
!person firstname name birthyear !
```

The fact that the first string is the type might be useful or annoying, depending on the situation. If we only want to get the fields of the typed list (and not the type), we can use the fieldnames function.

```
-->fieldnames(p)
ans =
```

```
!firstname !
! !
!name !
!
!birthyear !
```

When we create a typed list, we may define its fields without setting their values. The actual value of a field might indeed be set, dynamically, later in the script. In this case, it might be useful to know if a field is already defined or not. The following session shows how the **definedfields** function returns the matrix of floating point integers representing the fields which are already defined. We begin by defining the person **p** without setting any value of any field. This is why the only defined field is the number 1. Then we set the "firstname" field, which corresponds to index 2.

```
-->p = tlist(["person","firstname","name","birthyear"])
p =
      p(1)
!person firstname
                           birthyear
                                       !
                     name
-->definedfields(p)
 ans
     =
    1.
-->p.firstname = "Paul";
-->definedfields(p)
 ans
     =
    1.
          2.
```

As we can see, the index 2 has been added to the matrix of defined fields returned by the definedfields function.

The functions that we have reviewed allows to program typed lists in a very dynamic way. We will now see how to use the **definedfields** functions to dynamically compute if a field, identified by its string, is defined or not. This will allow to get a little bit more practice with typed lists. Recall that we can create a typed list without actually defining the values of the fields. These fields can be defined afterward, so that, at a particular time, we do not know if all the fields are defined or not. Hence, we may need a function **isfielddef** which would behave as in the following session.

```
-->p = tlist(["person","firstname","name","birthyear"]);
-->isfielddef ( p , "name" )
ans =
F
-->p.name = "Smith";
-->isfielddef ( p , "name" )
ans =
T
```

The topic of the exercise 3.2 is to dynamically compute if the field associated with a given field, identified by its string, exists in a typed list.

3.7 Emulating OO Programming with typed lists

In this section, we review how typed lists can be used to emulate Object Oriented Programming (OOP). This discussion has been partly presented on the Scilab wiki [7].
We present a simple method to emulate OOP with current Scilab features. The suggested method is classical when we want to emulate OOP in a non-OOP language, for example in C or Fortran. In the first part, we analyze the limitations of functions, which use positional arguments. Then, we present a method to emulate OOP in Scilab using typed lists.

In the notes associated with this section, we present similar methods in other languages. We emphasize the fact that Object Oriented Programming has been used for decades in non-OOP languages, like C or Fortran for example by methods which are very similar to the one that we are going to present.

3.7.1 Limitations of positional arguments

Before going into the details, we first present the reasons why emulating OOP in Scilab is convenient and, sometimes, necessary. Indeed, the method we advocate may allow to simplify many functions, which are based on optional, positional, arguments. The fact that the execution, based on the location of the input and output arguments of a function is a severe limitation in some contexts, as we are going to see.

For example, the optim primitive is a built-in function which performs unconstrained and bound constrained numerical optimization. This function has 20 arguments, some of them being optional. The following is the header of the function, where square brackets [...] indicate optional parameters.

```
[f [,xopt [,gradopt [,work]]]]= ..
optim(costf [,<contr>],x0 [,algo] [,df0 [,mem]] [,work] ..
[,<stop>] [,<params>] [,imp=iflag])
```

This complicated calling sequence makes the practical use of the optim difficult (but doable), especially when we want to customize its arguments. For example, the carams> variable is a list of optional four arguments:

```
'ti', valti ,'td', valtd
```

Many parameters of the algorithm can be configured, but, surprisingly enough, many more cannot be configured by the user of the optim function. For example, in the case of the Quasi-Newton algorithm without constraints, the Fortran routine allows to configure a length representing the estimate of the distance to the optimum. This parameter cannot be configured at the Scilab level and the default value 0.1 is used. The reason behind this choice is obvious: there are already too many parameters for the optim function and adding other optional parameters would lead to an unusable function.

Moreover, users and developers may want to add new features to the optim primitive, but that may lead to several difficulties.

• Extending the current optim gateway is very difficult, because of the complicated management of the 20 input optional arguments. Moreover, maintaining and developing the interface during the life of the Scilab project is difficult, because the order of the arguments matters. For example, we may realize that one argument may be unnecessary (because, for example, the same information may be computed from another variable). In this case, we cannot remove the argument from the calling sequence, because it would break the backward compatibility of all the scripts which are using the function.

- Extending the list of output arguments is difficult. For example, we may be interested by an integer representing the status of the optimization (e.g. convergence reached, maximum number of iterations reached, maximum number of function calls, etc...). We might also be interested in the number of iterations, the number of function calls, the final value of the approximation of the Hessian matrix, the final value of the gradient, and many other informations. The limited number of output arguments actually limits the number of informations that the user can extract out of a simulation. Moreover, if we want to get the output argument #6, for example, we must call the function with all the arguments from #1 to #6. This might generate a lot of unnecessary data.
- We might want to set the optional input argument #7, but not the optional argument #6, which is not possible with the current interface. This is because the argument processing system is based on the order of the arguments.

All in all, the fact that the input and the output arguments are used explicitly, and based on their order, is very inconvenient when the number of arguments is large.

If an Oriented Object Programming environment was available for Scilab, the management of the arguments would be solved with less difficulty.

3.7.2 A "person" class in Scilab

In this section, we give a concrete example of the method, based of the development of a "person" class.

The method that we present in this document to emulate Object Oriented Programming is classical in other languages. Indeed, it is common to extend a nonobject language into an OOP framework. A possible method is:

- we create an abstract data type (ADT) with the language basic data structures,
- we emulate methods with functions, where the first argument, named this, represents the current object.

We will use this method and emulate a sample "person" class as an example. The "person" class is made of the following functions.

- The person_new function, the "constructor", creates a new person.
- The person_free function, the "destructor", destroys an existing person.
- The person_configure and person_cget functions, allow to configure and quiery the fields of an existing person.
- The person_display function, a "method", displays the current object into the console.

In these functions, the current object will be stored in the variable this. To implement our class, we use a typed list.

The following function person_new, returns a new person this. This new person is defined by its name, first name, phone number and email. We choose to use the default empty string value for all the fields.

```
function this = person_new ()
this = tlist(["TPERSON","name","firstname","phone","email"])
this.name=""
this.firstname=""
this.phone=""
this.email=""
endfunction
```

The following person_free function destroys an existing person.

```
function this = person_free (this)
    // Nothing to be done.
endfunction
```

Since there is nothing to be done for now, the body of the function person_free is empty. Still, for consistency reasons, and because the actual body of the function may evolve later during the development of the component, we create this function anyway.

We emphasize that the variable **this** is both an input and an output argument of the **person_free**. Indeed, the current person is, in principle, modified by the action of the **person_free** function.

The function person_configure allows to configure a field of the current person. Each field is identified by a string, the "key", which corresponds to a value. Hence, this is simply a "key-value" mapping. The function sets the value corresponding to the given key, and returns the updated object this. For the "person" class, the keys are "-name", "-firstname", "-phone" and "-email".

```
function this = person_configure (this,key,value)
select key
case "-name" then
   this.name = value
case "-firstname" then
   this.firstname = value
case "-phone" then
   this.phone = value
case "-email" then
   this.email = value
else
   errmsg = sprintf("Unknown key %s",key)
   error(errmsg)
end
endfunction
```

We made the choice of prefixing each key by a minus "-" character. In calling sequences, this allows to easily distinguish a key from a value.

We emphasize that the variable **this** is both an input and an output argument of the **person_configure** function. This is similar to the **person_free** function that we have previously created. Similarly, the person_cget function allows to get a given field of the current person. The person_cget returns the value corresponding to the given key of the current object.

```
function value = person_cget (this,key)
select key
case "-name" then
value = this.name
case "-firstname" then
value = this.firstname
case "-phone" then
value = this.phone
case "-email" then
value = this.email
else
errmsg = sprintf("Unknown key %s",key)
error(errmsg)
end
endfunction
```

More precisely, the person_cget function allows to get the value of a configurable key. The "c" in "cget" refers to the first letter of "configure". If, instead, we want to create a function which returns the value of a non-configurable key, we may name it person_get.

Now that our class is setup, we can create a new "method". The following person_display function displays the current object this in the console.

```
function person_display (this)
mprintf("Person\n")
mprintf("Name: %s\n", this.name)
mprintf("First name: %s\n", this.firstname)
mprintf("Phone: %s\n", this.phone)
mprintf("E-mail: %s\n", this.email)
endfunction
```

We now present a simple use of the class "person" that we have just developed. In the following script, we create a new person by calling the **person_new** function. Then we call the **person_configure** function several times in order to configure the various fields of the person.

```
p1 = person_new();
p1 = person_configure(p1,"-name","Backus");
p1 = person_configure(p1,"-firstname","John");
p1 = person_configure(p1,"-phone","01.23.45.67.89");
p1 = person_configure(p1,"-email","john.backus@company.com");
```

In the following session, we call the **person_display** function and prints the current person.

```
-->person_display(p1)
Person
Name: Backus
First name: John
Phone: 01.23.45.67.89
E-mail: john.backus@company.com
```

We can also quiery the name of the current person, by calling the person_get function.

```
-->name = person_cget(p1,"-name")
name =
Backus
```

Finally, we destroy the current person.

```
p1 = person_free(p1);
```

3.7.3 Extending the class

In this section, we discuss a way to extend a class based on the emulation method that we have presented. We aim at being able to manage more complex components, with more fields, more methods or more classes.

First, we emphasize that the management of the options of the class is secure. Indeed, the system that we have just designed simply maps a key to a value. Since the list of keys is defined once for all, the user cannot configure or get the value of a key which does not exist. If we try to, the person_configure or person_cget functions generate an error.

It is straightforward to add a new key into the class. We first have to update the **person_new** function, adding the new field to the typed list. We can decide which default value to use for the new field. Notice that existing scripts using the "person" class will work without any modification, using the default value of the new field. If required, a script may be updated in order to configure the new key to a non-default value.

We may decide to distinguish between public fields, which can be configured by the user of the class, and private fields, which cannot. In order to add a new private field into the "person" class, say "bankaccount", we modify the person_new function and add the corresponding string into the typed list. Since we have not made it available neither in the person_configure, nor in the person_cget functions, the user of the class cannot access to it.

We may be a little bit more flexible, allowing the user of the class to get the value of the field, without the possibility of changing the value. In this case, we should create a separated **person_get** function (notice the lack of the "c" letter). This allows to separate configurable options and non-configurable options.

We can create even more complex data structures, by nesting the classes. This is easy, since a typed list can contain a typed list, which can contain a typed list, etc... at any level of nesting that is required. Hence, we can emulate a limited inheritance, an idea which is one of the main topics in OOP.

For example, assume that we want to create a "company" class. That "company" is specified by its name, its address, its purpose and the list of all the "persons" working in it. The following company_new function suggests a possible implementation.

```
function this = company_new ()
this = tlist(["TCOMPANY","name","address","purpose",..
"employees"])
this.name=""
this.address=""
this.purpose=""
this.employees = list()
endfunction
```

The following company_addperson function allows to add a new person to the list of employees.

```
function this = company_addperson ( this , person )
  this.employees($+1) = person
endfunction
```

In fact, all the methods which are commonly used in OOP can be applied using this emulation scheme. This allows to create stand-alone components, providing a clear public interface and avoiding the need for complex functions using a large number of positional arguments.

3.8 Overloading typed lists

In this section, we will see how to overload the **string** function so that we can convert a typed list into a string. We will also review how to overload the **disp** function, i.e. the printing system, which allows to customize the printing of typed lists.

The following %TPERSON_string function returns a matrix of strings containing a description of the current person. It first creates an empty matrix. Then it adds the strings one after the other, by using the output of the sprintf function, which allows to format the strings.

The %TPERSON_string function allows to overload the string function for any object with type TPERSON.

The following %TPERSON_p function prints the current person. It first calls the string function in order to compute a matrix of strings describing the person. Then it makes a loop over the rows of the matrix and display them one after the other.

```
function %TPERSON_p ( this )
  str = string(this)
  nbrows = size(str,"r")
  for i = 1 : nbrows
    mprintf("%s\n",str(i))
  end
endfunction
```

The %TPERSON_p function allows to overload the printing of the objects with type TPERSON (the letter "p" stands for "print").

In the following session, we call the string function with the person p1.

```
-->p1 = person_new();
-->p1 = person_configure(p1,"-name","Backus");
-->p1 = person_configure(p1,"-firstname","John");
-->p1 = person_configure(p1,"-phone","01.23.45.67.89");
-->p1 = person_configure(p1,"-email",..
-->"john.backus@company.com");
-->string(p1)
ans =
!Person:
                                 I
                                 I
1
Т
!Name: Backus
!First name: John
!Phone: 01.23.45.67.89
Ţ.
!E-mail: john.backus@company.com
                                 I
```

As we can see in the previous session, the **string** function has automatically called the **%TPERSON_string** function that we have previously defined.

In the following session, we simply type the name of the variable p1 (and immediately press the enter key in the console), as with any other variable. This displays the content of our p1 variable.

In the previous session, the printing system has automatically called the "TPERSON_p function that we have previously defined. Notice that the same output would have been produced if we had used the disp(p1) statement.

Finally, we destroy the current person.

p1 = person_free(p1);

There are many other functions which can be defined for typed lists. For example, we may overload the + operator so that we can add two persons. This is described in more detail in the help page devoted to overloading:

help overloading

3.9 The mlist data type

In this section, we present the **mlist** data type, which is a matrix-oriented type of list. In the first part of this section, we present the main motivation for the **mlist** data type, by comparison with the **tlist**. Then we present a sample example of an **mlist**, where we define the extraction function.

The main difference between a tlist and an mlist is with respect to the extraction and insertion functions. Indeed, for a tlist M, the index-based extraction, i.e. the statement x=M(2), for example, is defined by default. It can be overloaded by the user, but this is not mandatory, as we are going to see.

In the following script, we define a tlist with type "V". This typed list has two fields, "name" and "age".

```
M=tlist(["V","name","age"],["a","b";"c" "d"],[1 2; 3 4]);
```

As expected from a tlist, the statement M(2) allows to extract the second field, i.e. the "name", of the variable.

```
-->M(2)
ans =
!a b !
! !
```

This is the same for the insertion (i.e. M(2)=x) into a list (or tlist).

On the other hand, for an mlist, the extraction and insertion functions *must* be defined. If not, an error is generated. In the following script, we define a mlist with type "V". As previously, this matrix list has two fields, "name" and "age".

```
M=mlist(["V","name","age"],["a","b";"c" "d"],[1 2; 3 4]);
```

In the following session, we see that we cannot directly extract the second field of this list.

```
-->M(2)
!--error 144
Undefined operation for the given operands.
check or define function %1_e for overloading.
```

The error message tells us that we are not able to extract the second entry of the variable M and that some function, that we are going to see later in this section, must be defined.

Let us reconsider the previous example and assume that M is a tlist. The specific issue with M is that the second entry M(2) may not correspond to what we need. Indeed, the variable M(2) is the matrix of strings ["a", "b"; "c" "d"]. Similarily, the variable M(1) is the matrix of strings ["V", "name", "age"]. But there might be situations where we would like to express that the "second entry" represents a particular entry which does not correspond to the particular meaning which is associated to a tlist. For example, we may want to concatenate the name with the value to form the string "Name: b, age: 2", which would be the second entry of our matrix M. Hence, it seems that we need to redefine the extraction (or insertion) of values for a tlist. But this is not possible, because the overloading system only allows to define functions which *do not exist*: in this case, the extraction function already exists, so that we cannot define it. This is where the mlist is useful: we can define the extraction and insertion functions for a mlist and customize its behavior. In fact, we are even forced to do so: as we have seen before, it is mandatory.

In order to define the extraction function for the mlist with type "V", we must define the function $%V_e$, where the "e" letter stands for "extraction". This is done in the following script. The header of the extraction function must be

[x1,..,xm]=%<type_of_a>_e_(i1,...,in,a) where x1,...,xm are the extracted values, i1,...,in are the indices of the values to be extracted and a is the current variable. Hence, the statement M=varargin(\$) allows to set into M the current variable to extract the values from. Then, the indices can be retrieved with the syntax varargin(1:\$-1), which creates a list of variables containing the actual indices. Finally, we create a matrix of strings, by concatenating the name with the string representing the age.

```
function r=%V_e(varargin)
M = varargin($)
r= "Name: " + M.name(varargin(1:$-1)) + ..
", Age: " + string(M.age(varargin(1:$-1)))
endfunction
```

In the following script, we extract the second entry of M.

```
-->M(2)
ans =
Name: c, Age: 3
```

We can also use a syntax which is similar to matrices, as in the following script.

```
-->M(2:3)
 ans
!Name: c, Age: 3
                  !
                  1
!Name: b, Age: 2
                 _ !
-->M(2,:)
ans
!Name: c, Age: 3 Name: d, Age: 4 !
-->M(:,1)
 ans =
!Name: a, Age: 1
                  !
1
                   I
!Name: c, Age: 3 !
```

3.10 The struct data type

In this section, we briefly present the **struct**, which is a data structure with unordered items of heterogeneous type. We compare its features against the **list**.

Inside a struct, all the items can be extracted by their name. In the following session, we define the variable d as a struct, with three fields "day", "month" and "year".

```
-->d=struct("day",25,"month" ,"DEC","year",2006)
d =
day: 25
month: "DEC"
year: 2006
```

Notice that the fields of the **struct** do not have the same types: the first field is a string, while the second field is a double.

In order to extract a value from a **struct**, we simply use the name of the variable, followed by a dot "." and the name of the field.

```
-->d.month
ans =
DEC
```

In order to insert a value into a struct, we simply use the equal "=" operator.

```
-->d.month='AUG'
d =
day: 25
month: "AUG"
year: 2006
```

A struct can contain another struct, which can lead to nested data structures. For example, in the following session, we create a week-end from two consecutive days.

```
-->d1=struct("day",01,"month" ,"JAN","year",2011);
-->d2=struct("day",02,"month" ,"JAN","year",2011);
-->weekend = struct("Sat",d1,"Sun",d2)
weekend =
    Sat: [1x1 struct]
    Sun: [1x1 struct]
-->weekend.Sat
ans =
    day: 01
    month: "JAN"
    year: 2011
```

From the flexibility point of view, a tlist is more powerful than a struct. This is because we can overload the functions so that their behaviour can be customized for a tlist. Indeed, customizing the behaviour of a struct is not possible. This is why most users tend to favor the tlist data type.

In fact, the main advantage of the **struct** is to be compatible with Matlab and Octave.

3.11 The array of structs

An array of structs is an array where each index is associated to a struct. There are several ways to create an array of structs. In the first part of this section, we concatenate several structures to create an array. In the second part, we initialize a whole array of structures in one call, then fill the fields one after the other.

If several structures have the same fields, and if all these fields have the same size, then they can be concatenated into one array of structures. In the following session, we create four separate structs.

```
s1=struct("firstname","John","birthyear",1940);
s2=struct("firstname","Paul","birthyear",1942);
s3=struct("firstname","George","birthyear",1943);
s4=struct("firstname","Ringo","birthyear",1940);
```

Notice that the fields do not have the same type: the first field is a string, while the second field is a double. Then we concatenate them into one array of structs, using the [] syntax, which is similar to matrices.

-->s=[s1,s2,s3,s4]

```
s =
1x4 struct array with fields:
    firstname
    name
```

We can extract the third structure out of the array, by using the s(3) syntax, which is similar to matrices.

```
-->s(3)
ans =
firstname: "George"
name: "Harrisson"
```

We can access to the "firstname" field of all the structures, as in the following script. This produces a list of strings.

```
-->s.firstname

ans =

ans(1)

John

ans(2)

Paul

ans(3)

George

ans(4)

Ringo

-->typeof(s.firstname)

ans =

list
```

For performance reasons, we do not advise to let structures grow dynamically. This is because this forces the interpreter to dynamically allocate more and more memory, and can lead to slow scripts. Instead, whenever possible, we should predefine an array of structures, and then fill the existing entries.

In the following script, we define a 4-by-1 array of **structs**, where each structure contains an empty firstname and an empty name.

```
t(1:4)=struct("firstname",[],"birthyear",[])
```

Then we can fill each structure, as in the following script.

```
t(1).firstname="John";
t(1).birthyear=1940;
t(2).firstname="Paul";
t(2).birthyear=1942;
t(3).firstname="George";
t(3).birthyear=1943;
t(4).firstname="Ringo";
t(4).birthyear=1940;
```

We can check that this produces exactly the same array of structs as before.

```
-->t
t =
4x1 struct array with fields:
   firstname
   birthyear
-->t(1)
ans =
```

| cell | Create a cell array with empty cells. |
|----------|---|
| cell2mat | Convert a cell array into a matrix. |
| iscell | Test if variable a is a cell array. |
| makecell | Create a cell array and initiate its cells. |

Figure 17: Functions related to cells.

```
firstname: "John"
birthyear: 1940
```

In the previous scripts, we have only seen arrays with 1 index. It must be clear that an array of **structs** is actually a 2-index array, i.e. a matrix. By the way, we are going to review another method to initialize an array of **structs**.

In the following script, we initialize a 2-by-2 structure array.

```
-->u(2,2).firstname=[]
u =
2x2 struct array with fields:
   firstname
-->u(2,2).birthyear=[]
u =
2x2 struct array with fields:
   firstname
   birthyear
```

Then we can fill the entries with a syntax which is similar to matrices.

```
u(1,1).firstname="John";
u(1,1).birthyear=1940;
u(2,1).firstname="Paul";
u(2,1).birthyear=1942;
u(1,2).firstname="George";
u(1,2).birthyear=1943;
u(2,2).firstname="Ringo";
u(2,2).birthyear=1940;
```

Once done, we can extract the structure associated with the indices (2,1), for example.

```
-->u(2,1)
ans =
firstname: "Paul"
birthyear: 1942
```

3.12 The cell data type

In this section, we briefly present the cell, which is an heterogeneous array of variables. Then we compare its features with the features against the hypermatrix and the list. The figure 17 presents several functions related to cells.

The cell function allows to create an array. This cell can contain any other type of variables, including doubles, integers, strings, etc... In the following session, we create a 2-by-3 cell.

-->c = cell(2,3) c = !{} {} {} ! ! ! !{} {} !

The size of a cell can be computed with the **size** function.

-->size(c) ans = 2. 3.

In order to insert a value into the cell, we cannot use the same syntax as for matrices.

Instead, we can use the entries field of the (2,1) entry, as in the following session.

```
-->c(2,1).entries=12
c =
!{} {} {} !
! !
!12 {} {} !
```

In the following session, we insert a string into the (1,3) entry.

```
-->c(1,3).entries="5"
c =
!{} {} "5" !
! ! !
!12 {} {} !
```

In order to delete the (1,3) entry, we can use the empty matrix [].

```
-->c(1,3).entries=[]
c =
!{} {} {} !
! !
!12 {} {} !
```

We can extract a sub-part of the cell, using a syntax similar to matrices. Notice that the result is a cell.

```
-->x=c(1:2,1)
x =
!{} !
!12 !
-->typeof(x)
ans =
ce
```

On the other hand, when we extract one particular entry with the entries field, we get the same data type as this particular entry. In the following session, we extract the (2,1) entry into x, and check that the variable x is a double.

```
-->x = c(2,1).entries
x =
12.
-->typeof(x)
ans =
constant
```

If we extract several entries at once with the **entries** field, we get a list. In the following session, we extract all the entries in the first column.

We can create multi-indexed arrays by using cells. For example, the statement c=cell(2,3,4) creates a 2-by-3-by-4 array. This feature can be compared to hypermatrices, with the additionnal advantage that the entries of a cell can have different types, while all the entries of an hypermatrix must have the same type.

The behaviour of a **cell** can also be compared to a **list**. But the entries of a **cell** can be accessed with a syntax similar to matrices, which can be convenient in some situations.

One of the features of the cell is that it is partially compatible with Matlab and Octave. But the cell in Scilab does not completely behave the same way, so that this compatibility is only partial. For example, the (2,1) entry of a cell can be extracted with the c $\{2,1\}$ syntax (notice the braces " $\{\}$ "): this syntax is not available in Scilab.

The syntax to extract values out of a cell might be particularly useful in some situations. This feature is shared with the tlist and the mlist, where the extraction can be overloaded.

3.13 Comparison of data types

In this section, we compare the various data types that we have presented. The figure 18 presents an overview of these data types.

The figure 19 presents the various data types that we have met so far and suggests other data types for replacement.

Replacing one data type by another might be interesting, for example in situations where performance matters. In this situation, experimenting various implementations and measuring the performance of each one may allow to improve the performance of a given algorithm.

Some data types are heterogeneous, i.e. these data types can contain variables which can have different types. This is the situation of the **struct**, the **mlist** and the **cell**. Other data types can only contain one particular type of variables: this is the situation of the matrix and the hypermatrix. If all the variables that the data

| Data type | Purpose | Advantage | Drawback |
|-------------|------------------------|----------------------|-----------------|
| matrix | 2-indices matrix | Efficient, Simple | Homogeneous |
| hypermatrix | Multi-index matrix | | Homogeneous |
| list | Ordered set of items | | No overloading. |
| tlist | Typed list | Can be overloaded. | |
| mlist | Matrix-oriented list | Extraction/Insertion | |
| | | can be user-defined. | |
| struct | Unordered set of items | Compatible | |
| | | with Matlab. | No overloading. |
| cell | | Partly compatible | |
| | | with Matlab. | No overloading. |

Figure 18: Scilab data structures.

| Data | Implemen- | Hetero- | Nesting |
|-------------|-----------|---------|---------|
| Type | tation | geneous | |
| matrix | built-in | No | No |
| hypermatrix | mlist | No | No |
| list | built-in | Yes | Yes |
| tlist | built-in | Yes | Yes |
| mlist | built-in | Yes | Yes |
| struct | mlist | Yes | Yes |
| cell | mlist | Yes | Yes |

Figure 19: Comparison of data types. The "Implementation" column refers to the implementation of these data types in Scilab v5.

type must manage have the same type, there is no doubt that the matrix or the hypermatrix should be chosen.

Nesting can be useful when we want to create trees of data structures. In this case, neither the matrix, nor the hypermatrix can be used. This is actually one of the main practical uses of the list. The "Nesting" column of the figure 19 presents this feature for all the data structures.

One of the reasons which may explain performance differences is the implementation of the various data types. This is presented in the "Implementation" column of the figure 19, where we present the implementation of these data types in Scilab v5.

For example, we have presented the hypermatrices in the section 3.3. In the section 3.4, we have already analyzed the performance impact that extraction of hypermatrices may have. In Scilab v5, hypermatrices are implemented with mlists, supported by both compiled source code and Scilab macros. For example, the extraction of hypermatrices of doubles is based on a compiled source code, while the extraction of hypermatrices of strings is based on a macro: this may also lead to performance differences.

3.14 Notes and references

In the first part of this section, we present some notes and references about the emulation of OOP in Scilab. In the second part, we describe the link between the polynomial data type and the control features in Scilab.

The abstract data structure to emulate OOP depends on the language.

- In C the abstract data structure of choice is the struct. This method of extending C to provide an OOP framework is often used [47, 45] when using C++ is not wanted or not possible. This method is already used inside the source code of Scilab, in the graphics module, for the configuration of graphic properties. This is known as "handles" : see the ObjectStructure.h header file [43].
- In Fortran 77, the "common" statement is sufficient (but it is rarely used to emulate OOP).
- In Fortran 90, the "derived type" was designed for that purpose (but is rarely used to truly emulate OOP). The Fortran 2003 standard is a real OOP Fortran based on derived types.
- In Tcl, the "array" data structure is an appropriate ADT (this is used by the STOOOP package [19] for example). But this may also be done with the "variable" statement, combined with namespaces (this is done in the SNIT package [17] for example).

The "new" constructor is emulated by returning an instance of the ADT. The "new" method may require to allocate memory. The "free" destructor takes "this" as first argument and frees the memory which have have been allocated previously. This approach is possible in Fortran, C, and other compiled languages, if the first argument "this" can be modified by the method. In C, this is done by passing a pointer to the ADT. In Fortran 90, this is done by adding "intent(inout)" to the declaration (or nothing at all).

We now discuss the link between the polynomial data type and control. When the former Matlab open-source project was considered by the researchers at IRIA (French Institute for Research in Computer Science and Control) in the early 1980s, they wanted to create a Computer Aided Control System Design (C.A.C.S.D.) software. At this time, the main developers were Franois Delebecque and Serge Steer. In the context of control theory, we analyze the behavior of dynamical systems. In the case of a single-input-single-output (SISO) control system, we may use the Laplace transform on the variables, which leads to the transfer function. Hence, the rational data type has been introduced to support the analysis of the transfer functions produced for linear systems. There are many other functions related to CACSD in Scilab, and presenting them is far beyond the scope of this document.

3.15 Exercises

Exercise 3.1 (*Searching for files*) Scripting languages are often used to automate tasks, such as finding or renaming files on a hard drive. In this situation, the **regexp** function can be used to

locate the files which match a given pattern. For example, we can use it to search for Scilab scripts in a directory. This is easy, since Scilab scripts have the ".sci" (for a file defining a function) or the ".sce" (for a file executing Scilab statements) extensions.

Design a function **searchSciFilesInDir** which searches for these files in a given directory. We suggest to provide the following header.

filematrix = searchSciFilesInDir(directory,funname)

If such a file is found, we add it to the filematrix matrix of strings, which is returned as an output argument. Moreover, we call the function funname back. This allows for us to process the file as we need. Hence, we can use this feature to display the file name, move the file into another directory, delete it, etc...

Design a function mydisplay with the following header:

mydisplay(filename)

and use this function in combination with the **searchSciFilesInDir** in order to display the files in a directory.

Exercise 3.2 (*Querying typed lists*) The functions that we have reviewed allows to program typed lists in a very dynamic way. We will now see how to use the **definedfields** functions to dynamically compute if a field, identified by its string, is defined or not. This will allow to get a little bit more practice with typed lists. Recall that we can create a typed list without actually defining the values of the fields. These fields can be defined afterward, so that, at a particular time, we do not know if all the fields are defined or not. Hence, we may need a function isfielddef which would behave as in the following session.

```
-->p = tlist(["person","firstname","name","birthyear"]);
-->isfielddef ( p , "name" )
ans =
F
-->p.name = "Smith";
-->isfielddef ( p , "name" )
ans =
T
```

Write the implementation of the isfielddef function. To do so, you may combine the find and definedfields functions.

4 Management of functions

In this section, we review the management of functions and present features to design flexible functions. We present methods to inquire on functions and to separate the macros from the primitives. We notice that functions are not reserved in Scilab and warn about issues of the **funcprot** function. We present the use of callbacks, which allow to let the user of a function customize a part of an algorithm. We analyze methods to design functions with a variable number of input or output arguments, based on the **argn**, **varargin** and **varargout** statements. We show common ways to provide default values to input arguments. We present how to use the empty matrix [] to overcome the problem caused by positional input arguments. We also consider the definition of functions where the input arguments can have various types. Then we present features which allow to design robust functions. We present practical examples based on the **error**, **warning** and **gettext** functions. We present the **parameters** module, which allows to solve the problem of designing a function with a large number of parameters. The scope of variable through the call stack

| type | typeof | Description |
|------|------------|------------------|
| 11 | "function" | Uncompiled macro |
| 13 | "function" | Compiled macro |
| 130 | "fptr" | Primitive |

Figure 20: Various types of functions.

is analyzed in depth. The issues caused by poor uses of this feature are analyzed based on typical use cases. We present issues with callbacks and analyze methods to solve them. We present a method based on lists to provide additional arguments to callbacks. Finally, we present meta-programming tools based on the execstr and deff functions.

4.1 Advanced function management

In this section, we present advanced function management. In the first section, we present the differences between macros and primitives. Then we emphasize that functions are not reserved, which implies that it is possible to redefine a function which already exists. We show that this is caused by the fact that functions are variables. We present the **funcprot** function and show how to use callbacks.

4.1.1 How to inquire about functions

In this section, we present the difference between macros and primitive. We analyze the various values returned by the type and typeof functions for functions input arguments. We introduce the deff function, which allows to dynamically define a new function, based on strings. Finally, we present the get_function_path function, which returns the path to the file defining a macro.

There are two main types of functions:

- macros, which are written in the Scilab language,
- primitives, which are written in in a compiled language, like C, C++ or Fortran for example.

Most of the time, there is not direct way to distinguish between these two types of functions, and this is a good feature: when we use a function, we do not care about the language in which the function is written since only the result matters. This is why we usually use the name *function*, without further details. But sometimes, this is important, for example when we want to analyze or debug a particular function: in this case, it is necessary to know where the function comes from.

There are several features which allow to inquire about a particular function. In this section, we focus on the type, typeof, deff and get_function_path functions.

The type of a macro is either 11 or 13 and that the type of a primitive is 130. These types are summarized in the figure 20.

In the following session, we define a function with the function statement. Then we compute its type with the type and typeof functions.

```
-->function y = myfunction ( x )
--> y = 2 * x
-->endfunction
-->type(myfunction)
ans =
    13.
-->typeof(myfunction)
ans =
function
```

The eye function is a primitive which returns identity matrices. In the following session, we exercise the type and typeof functions with the eye input argument.

```
-->type(eye)
ans =
130.
-->typeof(eye)
ans =
fptr
```

The deff function allows to dynamically define a function based on strings representing its definition. This allows to dynamically define a new function, for example in order to encapsulate a function into another. The deff function accepts an optional input argument which allows to change how the function is processed by the interpreter. Such a function can be compiled, compiled and profiled or not compiled. The compilation process allows to generate a faster intermediate bytecode, which can be directly used by the interpreter without additional treatment. But it also makes the debugging impossible, and this is why this feature can be disabled. We will review the deff in more detail in the section 4.7 of this document.

By default, the deff function create compiled macros. In the following session, we define an uncompiled macro with the deff function. The first argument of the deff function is the header of the function. In our case, the myplus function takes the two input arguments y and z and returns the output argument x. The second argument of the deff function is the body of the function. Here, we simply set x to the sum of y and z. Finally, the third argument of the deff function is a string defining the type of function which is to be created. We can choose between "c", for a compiled macro, "p", for a compiled and profiled macro, and "n", for an uncompiled macro.

```
-->deff("x=myplus(y,z)","x=y+z","n")
-->type(myplus)
ans =
    11.
-->typeof(myplus)
ans =
    function
```

The previous session allows to check that the type of an uncompiled function is 11.

When a function is provided in a *library*, the get_function_path function allows to get the path to the function. For example, the *optimization* module provided with Scilab contains both primitives (for example the function optim) and macros (for example, the function derivative). Optimization macros are collected in the library *optimizationlib*, which is analyzed in the following session.

| >optimiz | zationlib | | | | |
|-----------|------------|-------------|-----------|-------------|-----------|
| optimizat | ionlib = | | | | |
| Functions | files loca | ation : SCI | \modules\ | optimizatio | $n \in .$ |
| aplat | bvodeS | datafit | derivativ | <i>r</i> e | fit_dat |
| karmarkar | | leastsq | linpro | list2vec | |
| lmisolver | | lmitool | mps2linpı | 0 | NDcost |
| numdiff | pack | pencost | qpsolve | unpack | vec2list |

The previous session does not tell us if any of the functions listed within the optimizationlib variable is a macro or a primitive. For that purpose, we can call the typeof function, as in the following session, which shows that the derivative function is a macro.

```
-->typeof(derivative)
ans =
function
```

In the following session, we compute the path leading to the derivative function by using the get_function_path function. Then we combine the get_function_path and the editor functions to edit this macro.

```
-->get_function_path("derivative")
ans =
D:/Programs/SCILAB~1.1-B\modules\optimization\
    macros\derivative.sci
-->editor(get_function_path("derivative"))
```

Notice that the get_function_path function does not work when the input argument is a function provided in a compiled language, e.g. the optim function.

```
-->typeof(optim)
ans =
fptr
-->get_function_path("optim")
WARNING: "optim" is not a library function
ans =
[]
```

4.1.2 Functions are not reserved

It is possible to redefine a function which already exists. Often, this is a mistake which makes Scilab generate an error message. In the following session, we define the **rand** function as a regular function and check that we can call it as any other user-defined function.

```
-->function y = rand(t)
--> y = t + 1
-->endfunction
Warning : redefining function: rand.
  Use funcprot(0) to avoid this message
-->y = rand(1)
  y =
    2.
```

The warning about the fact that we are redefining the function **rand** should make us feel uncomfortable with our script. It tells us that the **rand** function already exists in Scilab, which can be easily verified with the help rand statement. Indeed, the built-in rand function allows to generate random numbers and we certainly do not want to redefine this function.

In the present case, the error is obvious, but practical situations might be much more complicated. For example, we can use a complicated nested tree of functions, where one very low level function raises this warning. Examining the faulty function and running it interactively allows most of the time to discover the issue. Moreover, since there are many existing functions in Scilab, it is likely that creating a new program initially produces name conflicts. In all cases, we should fix this bug without re-defining any existing function.

We will review this topic more deeply in the next section, where we present an example where temporarily disabling the function protection is really necessary. That is, we will present the **funcprot** function which is mentioned in the warning message of the previous session.

4.1.3 Functions are variables

In this section, we show that function are variables and introduce the funcprot function.

A powerful feature of the language is that functions are variables. This implies that we can store a function into a variable and use that variable as a function. In compiled languages, this feature is often known as "function pointers".

In the following session, we define a function f. Then we set the content of the variable fp to the function f. Finally, we can use the function fp as a regular function.

```
-->function y = f ( t )

--> y = t + 1

-->endfunction

-->fp = f

fp =

[y]=fp(t)

-->fp ( 1 )

ans =

2.
```

This feature allows to use a very common programming tool, known as "callbacks". A callback is a function pointer which can be configured by the user of a component. Once configured, the component can call back the user-defined function, so that the algorithm can have, at least partly, a customized behavior.

Since functions are variables, we can set a variable containing a function several times. In fact, in order to protect users against unwanted redefinition of functions, a warning message can appear, as shown in the following example.

We begin by defining two functions f1 and f2.

```
function y = f1 ( x )
  y = x^2
endfunction
function y = f2 ( x )
  y = x^4
endfunction
```

| prot=funcprot() | Get the current function protection mode |
|-----------------|--|
| funcprot(prot) | Set the function protection mode |
| prot==0 | no message when a function is redefined |
| prot==1 | warning when a function is redefined (default) |
| prot==2 | error when a function is redefined |

Figure 21: The funcprot function.

Then, we can set the variable **f** two times successively, as in the following session.

```
-->f = f1
f =
[y]=f(x)
-->f = f2
Warning : redefining function: f.
Use funcprot(0) to avoid this message
f =
[y]=f(x)
```

We have already seen this warning message in the section 4.1.2, in the case where we tried to redefine a built-in function. But in the present situation, there is no reason to prevent us from setting the variable **f** to a new value. Unfortunately, there is no possibility for Scilab to distinguish between these two situations. Fortunately, there is a simple way of disabling the warning message temporarily. The **funcprot** function, presented in figure 21, allows to configure the protection mode of functions.

In general, it is not recommended to configure the function protection mode *permanently*. For example, we should never write the funcprot(0) statement in the .scilab startup file. This is because this would prevent us from getting warning messages and, therefore, may let us use bugged scripts without knowing it.

But, in order to avoid the unnecessary message of the previous session, we can *temporarily* disable the protection. In the following session, we make a backup of the function protection mode, set it temporarily to zero and then restore the mode to the backup value.

```
-->oldfuncprot = funcprot()
oldfuncprot =
    1.
-->funcprot(0)
-->f = f2
f =
[y]=f(x)
-->funcprot(oldfuncprot)
```

4.1.4 Callbacks

In this section, we present a method to manage callbacks, that is, we consider the case where an input argument of a function is itself a function. As an example, we consider the computation of numerical derivatives by finite differences.

In the following session, we use the (built-in) derivative function in order to compute the derivative of the function myf. We first define the myf function which

squares its input argument x. Then we pass the function myf to the derivative function as a regular input argument in order to compute the numerical derivative at the point x=2.

```
-->function y = myf ( x )
--> y = x^2
-->endfunction
-->y = derivative ( myf , 2 )
y =
4.
```

In order to understand the behavior of callbacks, we can implement our own simplified numerical derivative function as in the following session. Our implementation of the numerical derivative is based on an order 1 Taylor expansion of the function in the neighborhood of the point \mathbf{x} . It takes as input arguments the function \mathbf{f} , the point \mathbf{x} where the numerical derivative is to be computed and the step \mathbf{h} which must be used.

```
function y = myderivative ( f , x , h )
y = (f(x+h) - f(x))/h
endfunction
```

We emphasize that this example is not to be used in practice, since the builtin derivative function is much more powerful than our simplified myderivative function.

Notice that, in the body of the myderivative function, the input argument f is used as a regular function.

In the following session, we use the myf variable as an input argument to the myderivative function.

```
-->y = myderivative ( myf , 2 , 1.e-8 )
y =
4.
```

We made the assumption that the function f has the header y=f(x). We may wonder what happens if this is not true.

In the following script, we define the myf2 function, which takes both x and a as input arguments.

```
function y = myf2 ( x , a )

y = a * x^2

endfunction
```

The following session shows what happens if we try to compute the numerical derivative of the myf2 function.

We may still want to compute the numerical derivative of our function, even if it has two arguments. We will see in the section 4.5.1 how the scope of variables can be used to let the myf2 function know about the value of the a argument. The

| argn | Returns the current number of input and output arguments. |
|-----------|---|
| varargin | A list storing the input arguments. |
| varargout | A list storing the output arguments. |

Figure 22: Functions related to variable number of input and output arguments.

programming method which we will review is not considered as a "clean" software practice. This is why we will present in the section 4.6.4 a method to manage callbacks with extra arguments.

4.2 Designing flexible functions

In this section, we present the design of functions with a variable number of input and output arguments. This section reviews the **argn** function and the **varargin** and **varargout** variables. As we are going to see, providing a variable number of input arguments allows to simplify the use of a function, by providing default values for the arguments which are not set by the user.

The functions related to this topic are presented in the figure 22.

In the first part, we analyze a simple example which allows to see the **argn** function in action. In the second part, we consider the implementation of a function which computes the numerical derivatives of a given function. Then we describe how to solve the problem generated with ordered input arguments, by making a particular use of the empty matrix syntax. In the final section, we present a function which behavior depends on the type of its input argument.

4.2.1 Overview of argn

In this section, we make an overview of the argn function and the varargin and varargout variables. We also present a simple function which allows to understand how these function perform.

We begin by analyzing the principle of the management of a function with a variable number of input and output arguments. The typical header of such a function is the following.

```
function varargout = myargndemo ( varargin )
  [lhs,rhs]=argn()
```

The varargout and varargin arguments are lists which represent the output and input arguments. In the body of the function definition, we call the argn function, which produces the two following outputs:

- lhs, the number of output arguments,
- rhs, the number of input arguments.

Both the varargin and varargout variables are defined when the function is called. The number of elements in these two lists is the following.

- On input, the number of elements in varargin is rhs.
- On input, the number of elements in varargout is zero.
- On output, the number of elements in varargout must be lhs.

The actual number of elements in **varargin** depends on the arguments provided by the user. On the contrary, the **varargout** list is always empty on input and the task of defining its content is left to the function.

Let us now consider the following myargndemo function, which will make this topic more practical. The myargndemo function displays the number of output arguments lhs, the number of input arguments rhs and the number of elements in the varargin list. Then, we set the output arguments in varargout to 1.

```
function varargout = myargndemo ( varargin )
  [lhs,rhs]=argn()
  mprintf("lhs=%d, rhs=%d, length(varargin)=%d\n",..
    lhs,rhs,length(varargin))
  for i = 1 : lhs
    varargout(i) = 1
  end
endfunction
```

In our function, we simply copy the input arguments into the output arguments. The following session shows how the function performs when it is called.

```
-->myargndemo();
lhs=1, rhs=0, length(varargin)=0
-->myargndemo(1);
lhs=1, rhs=1, length(varargin)=1
-->myargndemo(1,2);
lhs=1, rhs=2, length(varargin)=2
-->myargndemo(1,2,3);
lhs=1, rhs=3, length(varargin)=3
-->y1 = myargndemo(1);
lhs=1, rhs=1, length(varargin)=1
-->[y1,y2] = myargndemo(1);
lhs=2, rhs=1, length(varargin)=1
-->[y1,y2,y3] = myargndemo(1);
lhs=3, rhs=1, length(varargin)=1
```

The previous session shows that the number of elements in the list varargin is rhs.

We notice that the minimum number of output arguments is 1, so that we always have $lhs \ge 1$. This is a property of the interpreter of Scilab, which forces a function to always return at least one output argument.

We emphasize that a function can be defined with both the varargin and varargout variables, or only with the varargin variable or even only with the varargout variable. This decision is left to the designer of the function.

The myargndemo function can be called with any number of input arguments. In practice, not all calling sequences will be authorized, so that we will have to insert error statements which will limit the use of the function. This topic will be reviewed in the next section.

4.2.2 A practical issue

We now turn to a more practical example, which involves the computation of numerical derivatives. This example is quite complex, but represents a situation where both the number of input and output arguments is variable.

The derivative function in Scilab allows to compute the first (Jacobian) and second (Hessian) derivatives of a multivariate function. In order to make the discussion of this topic more vivid, we consider the problem of implementing a simplified version of this function. The implementation of the derivative function is based on finite differences and uses finite differences formulas of various orders (see [46] for more details on this subject).

We assume that the function is smooth and that the relative error in the function evaluation is equal to the machine precision $\epsilon \approx 10^{-16}$. The total error which is associated with any finite difference formula is the sum of the truncation error (because of the use of a limited number of terms in the Taylor expansion) and a rounding error (because of the limited precision of floating point computations in the function evaluation). Therefore, the optimal step which minimizes the total error can be explicitly computed depending on the machine precision.

The following function myderivative1 computes the first and second derivatives of an univariate function. Its input arguments are the function to differentiate f, the point x where the derivative is to be evaluated, the step h and the order order of the formula. The function provides an order one forward formula and an order 2 centered formula. The output arguments are the value of the first derivative fp and second derivative fpp. In the comments of the function, we have written the optimal steps h, as a reminder.

We now analyze how the **myderivative1** function behaves in practice. We are going to discover that this function has several drawbacks and lacks of flexibility and performance.

We consider the computation of the numerical derivative of the cosine function at the point x = 0. We define the function myfun, which computes the cosine of its input argument x.

```
function y = myfun ( x )
y = cos(x)
endfunction
```

In the following session, we use the myderivative1 function in order to compute the first derivative of the cosine function.

```
-->format("e",25)
-->x0 = %pi/6;
```

```
-->fp = myderivative1 ( myfun , x0 , 1 , %eps^(1/2) )
fp =
- 5.000000074505805969D-01
-->fp = myderivative1 ( myfun , x0 , 2 , %eps^(1/3) )
fp =
- 5.00000000026094682D-01
```

The exact value of the first derivative is $-\sin(\pi/6) = -1/2$. We see that the centered formula, associated with order=2, is, as expected, more accurate than the order=1 formula.

In the following session, we call the myderivative1 function and compute both the first and the second derivatives of the cosine function.

The exact value of the second derivative fpp is $-\cos(\pi/6) = -\sqrt{3}/2 \approx -8.6602 \cdot 10^{-1}$. Again, we see that the second order formula is more accurate than the first order formula.

We have checked that our implementation is correct. We are now going to analyze the design of the function and introduce the need for the variable number of arguments.

The myderivative1 function has three drawbacks.

- It may make unnecessary computations, which is a performance issue.
- It may produce unaccurate results, which is an accuracy issue.
- It does not allow to use default values for order and h, which is a flexibility issue.

The performance issue is caused by the fact that the two output arguments fp and fpp are computed, even if the user of the function does not actually required the second, optional, output argument fpp. We notice that the computation of fpp requires some additional function evaluations with respect to the computation of fp. This implies that, if the output argument fp only is required, the output argument fpp will still be computed, which is useless. More precisely, if the calling sequence:

fp = myderivative1 (f , x , order , h)

is used, even if fpp does not appear as an output argument, internally, the fpp variable will still be computed.

The accuracy issue is cause by the fact that the optimal step can be used either for the first derivative computation, or for the second derivative computation, but not both. Indeed, the optimal step is different for the first derivative and for the second derivative. More precisely, if the calling sequence:

 $[fp, fpp] = myderivative1 (myfun , x0 , 1 , %eps^(1/2))$

is used, then the step $eps^{(1/2)}$ is optimal for the first derivative fp. But, if the calling sequence:

```
[fp, fpp] = myderivative1 ( myfun , x0 , 1 , %eps^(1/3) )
```

is used, then the step $\ensuremath{\sc keps^{(1/3)}}$ is optimal for the second derivative fpp. In all cases, we must choose and cannot have a good accuracy for both the first and the second derivative.

The flexibility issue is caused by the fact that the user must specify both the order and the h input arguments. The problem is that the user may not know what value to use for these parameters. This is particularly obvious for the h parameter, which is associated with floating point issues which might be completely unknown to the user. Hence, it would be convenient if we could use default values of these input arguments. For example, we may want to use the default order=2, since it provides a more accurate derivative with the same number of function calls. Given the order, the optimal step h could be computed with either of the optimal formulas.

Now that we know what problems are associated with our myderivative1 function, we are going to analyze an implementation based on a variable number of input and output arguments.

4.2.3 Using variable arguments in practice

The goal of this section is to provide an implementation of the function which allows to use the following calling sequences.

```
fp = myderivative2 ( myfun , x0 )
fp = myderivative2 ( myfun , x0 , order )
fp = myderivative2 ( myfun , x0 , order , h )
[fp,fpp] = myderivative2 ( ... )
```

The main advantage of this implementation are

- to be able to provide default values for order and h,
- to compute **fpp** only if required by the user,
- to use the optimal step h for both the first and second derivatives, if it is not provided by the user.

The following function myderivative2 implements a finite difference algorithm, with optional order order and step h.

```
function varargout = myderivative2 ( varargin )
1
2
    [lhs,rhs]=argn()
3
    if ( rhs < 2 | rhs > 4 ) then
      error ( msprintf(..
4
         "%s: Expected from %d to %d input arguments, "+..
5
6
         "but %d are provided",..
         "myderivative2",2,4,rhs))
7
8
    end
```

```
9
     if ( lhs > 2 ) then
10
        error ( msprintf(..
          "%s: Expected from %d to %d output arguments, "+..
11
12
          "but %d are provided",..
13
          "myderivative2",0,2,lhs))
14
     end
15
     f = varargin(1)
16
     x = varargin(2)
17
     if ( rhs \geq 3 ) then
18
        order = varargin(3)
19
     else
20
        order = 2
21
     end
     if ( rhs \ge 4 ) then
22
23
       h = varargin(4)
24
       hflag = %t
25
     else
26
        hflag = \%f
27
     end
28
     if ( order == 1 ) then
29
        if ( ~hflag ) then
30
          h = %eps^{(1/2)}
31
        end
32
        fp = (f(x+h) - f(x))/h
33
     else
34
        if ( ~hflag ) then
35
          h = %eps^{(1/3)}
36
        {\tt end}
        fp = (f(x+h) - f(x-h))/(2*h)
37
38
     end
39
     varargout(1) = fp
      if ( lhs \geq 2 ) then
40
        if ( order == 1 ) then
41
42
          if ( ~hflag ) then
43
            h = %eps^{(1/3)}
44
          end
45
          fpp = (f(x+2*h) - 2*f(x+h) + f(x))/(h^2)
46
        else
          if ( ~hflag ) then
47
            h = %eps^{(1/4)}
48
49
          end
50
          fpp = (f(x+h) - 2*f(x) + f(x-h))/(h^2)
51
        end
52
        varargout(2) = fpp
53
      end
54
   endfunction
```

As the body of this function is quite complicated, we are now going to analyze its most important parts.

The line #1 defines the function as taking the varargin variable as input argument and the varargout variable as output argument.

The line #2 makes use of the argn function, which returns the actual number of input and output arguments. For example, when the calling sequence fp = myderivative2 (myfun , x0) is used, we have rhs=2 and lhs=1. The following script presents various possible calling sequences.

The lines #2 to #12 are designed to check that the number of input and output arguments is correct. For example, the following session shows the error which is generated in the case where the user wrongly calls the function with 1 input argument.

The lines #13 and #14 shows how to directly set the values of the input arguments f and x, which always are forced to be present in the calling sequence. The lines #15 to #19 allows to set the value of the **order** parameter. When the number of input arguments **rhs** is greater than 3, then we deduce that the value of the **order** variable is given by the user, and we use directly that value. In the opposite case, we set the default value of this parameter, that is we set **order=2**.

The same processing is done in the lines #20 to #25 in order to set the value of h. Moreover, we set the value of the hflag boolean variable. This variable is set to %t if the user has provided h, and to %f if not.

The lines #26 to #36 allow to compute the first derivative, while the second derivatives are computed in the lines #38 to #51. In each case, if the user has not provided h, that is, if hflag is false, then the optimal step is used.

The number of output arguments is necessary greater or equal to one. Hence, there is no need to test the value of lhs before setting varagout(1) at the line #37.

One important point is that the second derivative **fpp** is computed only if it required by the user. This is ensured by the line #38, which checks for the number of output arguments: the second derivative is computed only of the second output argument **fpp** was actually written in the calling sequence.

The following session shows a simple use of the myderivative2 function, where we use the default value of both order and h.

```
-->format("e",25)
-->x0 = %pi/6;
-->fp = myderivative2 ( myfun , x0 )
fp =
- 5.00000000026094682D-01
```

We notice that, in this case, the second derivative has not been computed, which may save a significant amount of time.

In the following session, we set the value of **order** and use the default value for h.

```
-->fp = myderivative2 ( myfun , x0 , 1 ) fp =
```

- 5.00000074505805969D-01

We can also call this function with two output arguments, as in the following session.

```
-->[fp,fpp] = myderivative2 ( myfun , x0 , order )
fpp =
- 8.660254031419754028D-01
fp =
- 5.00000000026094682D-01
```

We notice that, in this case, the optimal step **h** has been used for both the first and the second derivatives.

We have seen in this section how to manage a variable number of input and output arguments. This method allows to design flexible and efficient functions.

But, in some situations, this is not sufficient and still suffers from limitations. For example, the method that we have presented is limited by the order of the arguments, that is, their position in the calling sequence. Indeed, the function myderivative2 cannot be called by using the default value for the order argument and setting a customized value for the h argument. This limitation is caused by the order of the arguments: the h input argument comes after the order argument. In practice, it would be convenient to be able to use the default value of an argument #i, and, still, set the value of the argument #i+1 (or any other argument on the right of the calling sequence). The next section allows to solve this issue.

4.2.4 Default values of optional arguments

In this section, we describe how to manage optional arguments with default values. We show how to solve the problem generated with ordered input arguments, by making a particular use of the empty matrix [] syntax.

Indeed, we saw in the previous paragraph that a basic management of optional arguments forbid us to set an input argument #i+1 and use the default value of the input argument #i. In this section, we present a method where the empty matrix is used to represent a default parameter.

Let us consider the function myfun, which takes x, p and q as input arguments and returns the output argument y.

```
function y = myfun ( x , p , q )
y = q*x^p
endfunction
```

If we both set x, p and q as input arguments, the function performs perfectly, as in the following session.

```
-->myfun(3,2,1)
ans =
9.
```

For now, our function myfun is not very flexible, so that if we pass only one or two arguments, the function generates an error.

```
-->myfun(3)
!--error 4
Undefined variable: q
```

```
at line 2 of function myfun called by :
myfun(3)
-->myfun(3,2)
!--error 4
Undefined variable: q
at line 2 of function myfun called by :
myfun(3,2)
```

It would be convenient if, for example, the parameter p had the default value 2 and the parameter q had the default value 1. In that case, if neither p nor q are provided, the statement foo(3) would return 9.

We can use the **varargin** variable in order to have a variable number of input arguments. But, used directly, this does not allow to set the third argument and use the default value of the second argument. This issue is caused by the ordering of the input arguments. In order to solve this problem, we are going to make a particular use of the empty matrix [].

The advantage of the method is to be able to use the default value of the argument p, while setting the value of the argument q. In order to inform the function that a particular argument must be set to its default value, we set it to the empty matrix. This situation is presented in the following session, where we set the argument p to the empty matrix.

```
-->myfun2(2,[],3) // same as myfun2(2,2,3)
ans =
12.
```

In order to define the function, we use the following method. If the input argument is not provided, or if it is provided, but is equal to the empty matrix, then we use the default value. If the argument is provided and is different from the empty matrix, then we use it directly. This method is presented in the myfun2 function.

```
function y = myfun2 (varargin)
  [lhs,rhs]=argn()
  if ( rhs < 1 | rhs > 3 ) then
    msg=gettext("%s: Wrong number of input arguments: "+..
        "%d to %d expected.\n")
    error(msprintf(msg,"myfun2",1,3))
  end
  x = varargin(1)
  pdefault = 2
  if (rhs \ge 2) then
    if (varargin(2) <> [] ) then
      p = varargin(2)
    else
      p = pdefault
    end
  else
   p = pdefault
  end
  qdefault = 1
  if ( rhs >= 3 ) then
    if (varargin(3) <> [] ) then
      q = varargin(3)
    else
      q = qdefault
```

```
end
else
q = qdefault
end
y = q * x^p
endfunction
```

From the algorithm point of view, we could have chosen another reference value, different from the empty matrix. For example, we could have considered the empty string, or any other particular value. The reason why the empty matrix is preferred in practice is because the comparison against the empty matrix is faster than the comparison with any other particular value. The actual content of the matrix does not even matter, since only the size of the matrices are compared. Hence the performance overhead caused by the management of the default values is as low as possible.

Of course, we can still use the optional arguments as usual, which is presented in the following session.

```
-->myfun2(2)

ans =

4.

-->myfun2(2,3)

ans =

8.

-->myfun2(2,3,2)

ans =

16.
```

In practice, this method is both flexible and consistent with the basic management of input arguments, which remains mainly based on their order.

This method is still limited if the number of input arguments is large. In this case, there are other solutions which allow to avoid to use ordered arguments. In the section 4.4, we analyze the **parameters** module, which allows to configure an unordered set of input arguments separately from the actual use of the parameters. Another solution is to emulate Object Oriented Programming, as presented in the section 3.7.

4.2.5 Functions with variable type input arguments

In this section, we present a function which behavior depends on the type of its input argument.

The following myprint function provides a particular display for a matrix of doubles, and another display for a boolean matrix. The body of the function is based on a if statement, which switches to different parts of the source code depending on the type of the X variable.

```
function myprint ( X )
if ( type(X) == 1 ) then
  disp("Double matrix")
elseif ( type(X) == 4 ) then
  disp("Boolean matrix")
else
  error ( "Unexpected type." )
```

```
end
endfunction
```

In the following session, we call the myprint function with two different types of matrices.

```
->myprint ( [1 2] )
Double matrix
-->myprint ( [%T %T] )
Boolean matrix
```

We can make the previous function more useful, by defining separated formatting rules for each type of data. In the following function, we group of three values is separated by a large blank space, which visually creates groups of values.

```
function myprint ( X )
  if (type(X) == 1) then
    // Real matrix
    for i = 1 : size(X, "r")
      for j = 1 : size(X,"c")
        mprintf("%-5d ",X(i,j))
        if (j == 3) then
          mprintf("
                        ")
        end
      end
      mprintf("\n")
    end
  elseif ( type(X) == 4 ) then
    // Boolean matrix
    for i = 1 : size(X, "r")
      for j = 1 : size(X, "c")
        mprintf("%s ",string(X(i,j)))
        if ( j == 3 ) then
          mprintf(" ")
        end
      end
      mprintf("\n")
    end
  else
    error ( "Unexpected type for input argument var." )
  end
endfunction
```

The following session gives a sample use of the previous function. First, we display a matrix of doubles.

```
-->X = [
-->1 2 3 4 5 6
-->7 8 9 10 11 12
-->13 14 15 16 17 18
-->];
-->myprint ( X )
      2
            3
                        4
                              5
                                     6
1
      8
             9
7
                        10
                              11
                                     12
13
      14
             15
                        16
                              17
                                     18
```

Then we display a boolean matrix.

| error | Sends an error message and stops the computation. |
|---------|---|
| warning | Sends a warning message. |
| gettext | Get text translated into the current locale language. |

Figure 23: Functions related to error messages.

```
-->X = [
-->%T %T %F %F %F %F %F
-->%T %F %F %T %T %T
-->%F %T %F %F %T %F
-->];
-->myprint ( X )
T T F F F F F
T F F T T T
F T F F T F
```

Many built-in function are designed on this principle. For example, the **roots** function returns the roots of a polynomial. Its input argument can be either a polynomial or a matrix of doubles representing its coefficients. In practice, being able to manage different types of variables in the same function provides an additional level of flexibility which is much harder to get in a compiled language such as C or Fortran.

4.3 Robust functions

In this section, we present some rules which can be applied to all functions which are designed to be robust against wrong uses. In the next section, we present the warning and error functions which are the basis of the design of robust functions. Then we present a general framework for the checks used inside robust functions. We finally present a function which computes the Pascal matrix and show how to apply these rules in practice.

4.3.1 The warning and error functions

It often happens that the input arguments of a function can have only a limited number of possible values. For example, we might require that a given input double is positive, or that a given input floating point integer can have only three possible values. In this case, we can use the **error** or **warning** functions, which are presented in the figure 23. The **gettext** function is related to localization and is described later in this section.

We now give an example which shows how these functions can be used to protect the user of a function against wrong uses. In the following script, we define the mynorm function, which is a simplified version of the built-in norm function. Our mynorm function allows to compute the 1, 2 or infinity norm of a vector. In other cases, our function is undefined and this is why we generate an error.

```
function y = mynorm ( A , n )
if ( n == 1 ) then
y = sum(abs(A))
```

```
elseif ( n == 2 ) then
  y=sum(A.^2)^(1/2);
elseif ( n == "inf" ) then
  y = max(abs(A))
else
  msg = msprintf("%s: Invalid value %d for n.","mynorm",n)
  error ( msg )
end
endfunction
```

In the following session, we test the output of the function mynorm when the input argument n is equal to 1, 2, "inf" and the unexpected value 12.

```
-->mynorm([1 2 3 4],1)

ans =

10.

-->mynorm([1 2 3 4],2)

ans =

5.4772256

-->mynorm([1 2 3 4],"inf")

ans =

4.

-->mynorm([1 2 3 4],12)

!--error 10000

mynorm: Invalid value 12 for n.

at line 10 of function mynorm called by :

mynorm([1 2 3 4],12)
```

The input argument of the **error** function is a string. We could have used a more simple message, such as "Invalid value n.", for example. Our message is a little bit more complicated for the following reason. The goal is to give to the user a useful feedback when the error is generated at the bottom of a possibly deep chain of calls. In order to do so, we give as much information as possible about the origin of the error.

First, it is convenient for the user to be informed about what exactly is the value which is rejected by the algorithm. Therefore, we include the actual value of the input argument n into the message. Furthermore, we make so that the first string of the error message is the name of the function. This allows to get immediately the name of the function which is generating the error. The msprintf function is used in that case to format the string and to produce the msg variable which is passed to the error function.

We can make so that the error message can be translated into other languages if necessary. Indeed, Scilab is *localized* so that most messages appear in the language of the user. Hence, we get English messages in the United States and Great Britain, French messages in France, etc... In order to do so, we can use the **gettext** function which returns a translated string, based on a localization data base. This is done for example in the following script.

```
localstr = gettext ( "%s: Invalid value %d for n." )
msg = msprintf ( localstr , "mynorm" , n )
error ( msg )
```

Notice that the string which is passed to the gettext function is not the output but the input of the msprintf function. This is because the localization system
provides a map from the string "%s: Invalid value %d for n." into localized strings such as the French message "%s: Valeur %d invalide pour n.", for example. These localized messages are stored in ".pot" data files. A lot more information about localization is provided at [28].

There are cases when we do want to generate a message, but we do not want to interrupt the computation. For example, we may want to inform our user that the **mynorm** function is rather poor compared to the built-in **norm** function. In this case, we do not want to interrupt the algorithm, since the computation is correct. We can use the **warning** function, as in the following script.

```
function y = mynorm2 ( A , n )
msg = msprintf("%s: Please use norm instead.","mynorm2")
warning(msg)
if ( n == 1 ) then
    y = sum(abs(A))
elseif ( n == 2 ) then
    y=sum(A.^2)^(1/2)
elseif ( n == "inf" ) then
    y = max(abs(A))
else
    msg = msprintf("%s: Invalid value %d for n.","mynorm2",n)
    error ( msg )
end
endfunction
```

The following session shows the result of the function.

```
-->mynorm2([1 2 3 4],2)
Warning : mynorm2: Please use norm instead.
ans =
5.4772256
```

4.3.2 A framework for checks of input arguments

A robust function should protect the user against wrong uses of the function. For example, if the function takes a matrix of doubles as an input argument, we may get error messages which are far from being clear. We may even get no error message at all, for example if the function silently fails. In this section, we present a general framework for the checkings in a robust function.

The following pascalup_notrobust is function which returns the upper triangular Pascal matrix P, depending on the size n of the matrix.

```
function P = pascalup_notrobust ( n )
P = eye(n,n)
P(1,:) = ones(1,n)
for i = 2:(n-1)
    P(2:i,i+1) = P(1:(i-1),i)+P(2:i,i)
end
endfunction
```

In the following session, we compute the 5×5 upper triangular Pascal matrix.

```
-->pascalup_notrobust ( 5 )
ans =
1. 1. 1. 1. 1. 1.
```

| 0. | 1. | 2. | 3. | 4. |
|----|----|----|----|----|
| 0. | 0. | 1. | 3. | 6. |
| 0. | 0. | 0. | 1. | 4. |
| Ο. | 0. | 0. | 0. | 1. |

The pascalup_notrobust function does not make any checks on the input argument n. In the case where the input argument is negative or is not a floating point integer, the function does not generate any error.

```
-->pascalup_notrobust(-1)

ans =

[]

-->pascalup_notrobust(1.5)

ans =

1.
```

This last behavior should not be considered as "normal", as it silently fails. Therefore, the user may provide wrong input arguments to the function, without even noticing that something wrong happened.

This is why we often check the input arguments of functions so that the error message generated to the user is as clear as possible. In general, we should consider the following checks:

- number of input/output arguments,
- type of input arguments,
- size of input arguments,
- content of input arguments,

which may be shortened as "number/type/size/content". These rules should be included as a part of our standard way of writing public functions.

4.3.3 An example of robust function

In this section, we present an example of a robust function which computes the Pascal matrix. We present examples where the checkings that we use are useful to detect wrong uses of the function.

The following pascalup is an improved version, with all necessary checks.

```
error ( msprintf(lstr,"pascalup",1,"constant",typeof(n)))
  end
 11
 // Check size of arguments
 if ( size(n, "*") \iff 1 ) then
    lstr=gettext("%s: Wrong size for input argument #%d: "+..
        "%d entries expected, but %d provided.")
    error ( msprintf(lstr,"pascalup",1,1,size(n,"*")))
 end
 11
 // Check content of arguments
 if (imag(n) <>0) then
   lstr = gettext("%s: Wrong content for input argument #%d: "+..
        "complex numbers are forbidden.")
    error ( msprintf(lstr,"pascalup",1))
 end
 if (n < 0) then
    lstr=gettext("%s: Wrong content for input argument #%d: "+..
        "positive entries only are expected.")
    error ( msprintf(lstr,"pascalup",1))
 end
 if ( floor(n) <> n ) then
    lstr=gettext("%s: Wrong content of input argument #%d: "+..
        "argument is expected to be a flint.")
    error ( msprintf(lstr,"specfun_pascal",1))
 end
 11
 P = eye(n,n)
 P(1,:) = ones(1,n)
 for i = 2:(n-1)
   P(2:i,i+1) = P(1:(i-1),i)+P(2:i,i)
 end
endfunction
```

In the following session, we run the **pascalup** function and produce various error messages.

```
-->pascalup ( )
!--error 10000
pascalup: Wrong number of input arguments:
    1 to 1 expected, but 0 provided.
              8 of function pascalup called by :
at line
pascalup ( )
-->pascalup ( -1 )
!--error 10000
pascalup: Wrong content for input argument #1:
    positive entries only are expected.
at line
            33 of function pascalup called by :
pascalup ( -1 )
-->pascalup ( 1.5 )
!--error 10000
specfun_pascal: Wrong content of input argument #1:
    argument is expected to be a flint.
at line
             37 of function pascalup called by :
pascalup ( 1.5 )
```

The rules that we have presented are used in most macros of Scilab. Numerous

| add_param | Add a parameter to a list of parameters |
|--------------|---|
| get_param | Get the value of a parameter in a parameter list |
| init_param | Initialize the structure which will handles the parameters list |
| is_param | Check if a parameter is present in a parameter list |
| list_param | List all the parameters name in a list of parameters |
| remove_param | Remove a parameter and its value from a list of parameters |
| set_param | Set the value of a parameter in a parameter list |

Figure 24: Functions from the parameters module.

experiments have proved that this method provides an improved robustness, so that users are less likely to use the functions with wrong input arguments.

4.4 Using parameters

The goal of the **parameters** module is to be able to design a function which has a possibly large amount of optional parameters. Using this module allows to avoid to design a function with a large number of input arguments, which may lead to confusion. This module has been introduced in Scilab v5.0, after the work by Yann Collette to integrate optimization algorithms such as Genetic Algorithms and Simulated Annealing. The functions of the **parameters** module are presented in the figure 24.

In the first section, we make an overview of the module and describe its direct use. In the second section, we present a practical example, based on a sorting algorithm. The last section focuses on the safe use of the module, protecting the user against common mistakes.

4.4.1 Overview of the module

In this section, we present the **parameters** module and give an example of its use, in the context of a merge-sort algorithm. More precisely, we present the **init_param**, **add_param** and **get_param** functions.

The module is based on a mapping from keys to values.

- Each key corresponds to a field of the list of parameters and is stored as a string. The list of available keys is defined by the developer of the function. There is no limit in the number of keys.
- The type of each value depends on the particular requirements of the algorithm. Virtually any type of value can be stored in the data structure, including (but not limited to) matrices of doubles, strings, polynomials, etc...

Hence, this data structure is extremely flexible, as we are going to see.

In order to understand the **parameters** module, we can separate the situation between two points of views:

• the user's point of view,

• the developer's point of view.

Therefore, in the following discussion, we will present first what happens when we want to use the **parameters** module in order to call a function. Then, we will discuss what the developer has to do in order to manage the optional parameters.

In order to make the discussion as applied as possible, we will take the example of a sorting algorithm and will examine it throughout this section. We consider a mergesort function, which defines a sorting algorithm based on a combination of recursive sorting and merging. The mergesort function is associated with the following calling sequences, where x is a matrix of doubles to be sorted, params is a list of parameters and y is a matrix of sorted doubles.

```
y = mergesort ( x )
y = mergesort ( x , params )
```

The actual implementation of the **mergesort** function will be defined later in this section. The **params** variable is a list of parameters which defines two parameters:

- direction, a boolean which is true for increasing order and false for decreasing order (default direction = %t),
- compfun, a function which defines a comparison function (default compfun = compfun_default).

The compfun_default function is the default comparison function and will be presented later in this section.

We first consider the user's point of view and present a simple use of the functions from the **parameters** module. We want to sort a matrix of doubles into increasing order. In the following script, we call the **init_param** function, which creates the empty list of parameters **params**. Then, we add the key "direction" to the list of parameters. Finally, we call the **mergesort** function with the variable **params** as the second input argument.

```
params = init_param();
params = add_param(params,"direction",%f);
x = [4 5 1 6 2 3]';
y = mergesort ( x , params );
```

We now consider the developer's point of view and analyze the statements which may be used in the body of the mergesort function. In the following session, we call the get_param function in order to get the value corresponding to the key "direction". We pass the %t value to the get_param function, which is the default value to be used in the case where this key is not defined.

```
-->direction = get_param(params,"direction",%t)
direction =
F
```

Since the "direction" key is defined and is false, we simply get back our value. We may use an extended calling sequence of the get_param function, with the output argument err. The err variable is true if an error has occurred during the processing of the argument.

```
-->[direction,err] = get_param(params,"direction",%t)
err =
   F
direction =
   F
```

In our case, no error is produced and this is why the variable err is false.

We can analyze other combinations of events. For example, let us consider the situation where the user defines only an empty list of parameters, as in the following script.

```
params = init_param();
```

In this case, in the body of the **mergesort** function, a straightforward call to the **get_param** function generates a warning.

```
-->direction = get_param(params,"direction",%t)
WARNING: get_param: parameter direction not defined
direction =
T
```

The warning indicates that the "direction" key is not defined, so that the default value %t is returned. If we use the additional output argument err, there is no warning anymore, but the err variable becomes true.

```
-->[direction,err] = get_param(params,"direction",%t)
err =
T
direction =
T
```

As a last example, let us consider the case where the user defines the **params** variable as an empty matrix.

```
params = [];
```

As we are going to see later in this section, this case may happen when we want to use the default values. The following session shows that an error is generated when we call the get_param function.

```
-->direction = get_param(params,"direction",%t)
!--error 10000
get_param: Wrong type for input argument #1: plist expected.
at line 40 of function get_param called by :
direction = get_param(params,"direction",%t)
```

Indeed, the variable params is expected to have the plist data type. This is because the init_param function creates variables with type plist, which stands for "parameters list". In fact, the actual implementation of the init_param function creates a typed list with type plist. In order to avoid to generate the previous error, we can use the err output argument, as in the following session.

```
-->[direction,err] = get_param(params,"direction",%t)
err =
T
direction =
T
```

We see that the **err** variable is true, indicating that an error has been detected during the processing of the variable **params**. As before, in this case, the **get_param** function returns the default value.

4.4.2 A practical case

In this section, we consider the actual implementation of the **mergesort** function presented in the previous section. The sorting algorithm that we are going to present is based on an algorithm designed in Scilab by Bruno Pinçon [44]. The modifications included in the implementation presented in this section include the management of the "direction" and "compfun" optional arguments. Interested readers will find many details about sorting algorithms in [27].

The following mergesort function provides a sorting algorithm, based on a merge-sort method. Its first argument, x, is the matrix of doubles to be sorted. The second argument, params, is optional and represents the list of parameters. We make a first call to the get_param function in order to get the value of the direction parameter, with %t as the default value. Then we get the value of the compfun parameter, with compfun_default as the default value. The compfun_default function is defined later in this section. Finally, we call the mergesortre function, which actually implements the recursive merge-sort algorithm.

```
function y = mergesort ( varargin )
  [lhs,rhs]=argn()
  if ( and( rhs <> [1 2] ) ) then
    errmsg = sprintf(..
      "%s: Unexpected number of arguments : " + ...
      "%d provided while %d to %d are expected.",..
      "mergesort", rhs, 1, 2);
    error(errmsg)
  end
  x = varargin(1)
  if ( rhs < 2 ) then
   params = []
  else
    params = varargin(2)
  end
  [direction,err] = get_param(params,"direction",%t)
  [compfun,err] = get_param(params,"compfun",compfun_default)
  y = mergesortre ( x , direction , compfun )
endfunction
```

The following mergesortre function implements the recursive merge-sort algorithm. The algorithm divides the matrix x into two sub-parts x(1:m) and x(m+1:n), where m is an integer in the middle of 1 and n. Once sorted by a recursive call, we merge the two ordered sub-parts back into x.

```
function x = mergesortre ( x , direction , compfun )
n = length(x)
indices = 1 : n
if ( n > 1 ) then
m = floor(n/2)
p = n-m
x1 = mergesortre ( x(1:m) , direction , compfun )
```

```
x2 = mergesortre ( x(m+1:n) , direction , compfun )
x = merge ( x1 , x2 , direction , compfun )
end
endfunction
```

We notice that the mergesort function does not call itself recursively: the mergesort function calls mergesortre, which performs the algorithm recursively. The important fact is that the mergesortre function has fixed input arguments. This allows to reduce the overhead caused by the processing of the optional arguments.

The following merge function merges its two sorted input arguments x1 and x2 into its sorted output argument x. The comparison operation is performed by the compfun function, which returns -1, 0 or 1, depending on the order of its two input arguments. In the case where the direction argument is false, we invert the sign of the order variable, so that the order is "decreasing" instead of the default "increasing" order.

```
function x = merge ( x1 , x2 , direction , compfun )
   n1 = length(x1)
   n2 = length(x2)
   n = n1 + n2
   x = []
    i = 1
    i1 = 1
    i2 = 1
    for i = 1:n
      order = compfun (x1(i1), x2(i2))
      if ( ~direction ) then
        order = -order
      end
      if ( order <=0 ) then
        x(i) = x1(i1)
        i1 = i1+1
        if (i1 > m) then
          x(i+1:n) = x2(i2:p)
          break
        end
      else
        x(i) = x2(i2)
        i2 = i2+1
        if (i2 > p) then
          x(i+1:n) = x1(i1:m)
          break
        end
      end
    end
endfunction
```

The following compfun_default function is the default comparison function. It returns order=-1 if x<y, order=0 if x==y and order=1 if x>y.

```
function order = compfun_default ( x , y )
if ( x < y ) then
order = -1
elseif ( x==y ) then</pre>
```

```
order = 0
else
order = 1
end
endfunction
```

We now analyze the behavior of the mergesort function by calling it with particular inputs. In the following session, we sort the matrix \mathbf{x} containing floating point integers into increasing order.

```
-->mergesort ( x )'
ans =
1. 2. 3. 4. 5. 6.
```

In the following session, we configure the direction parameter to %f, so that the order of the sorted matrix is decreasing.

```
-->params = init_param();
-->params = add_param(params,"direction",%f);
-->mergesort ( x , params )'
ans =
6. 5. 4. 3. 2. 1.
```

We now want to customize the comparison function, so that we can change the order of the sorted matrix. In the following session, we define the comparison function mycompfun which allows to separate even and odd floating point integers.

```
function order = mycompfun (x, y)
  if (modulo(x,2) == 0 \& modulo(y,2) == 1) then
    // even < odd
    order = -1
 elseif ( modulo(x,2) == 1 \& modulo(y,2) == 0 ) then
    // odd > even
    order = 1
  else
    // 1<3 or 2<4
    if (x < y) then
      order = -1
    elseif ( x==y ) then
      order = 0
    else
      order = 1
    end
  end
endfunction
```

We can configure the **params** variable so that the sort function uses our customized comparison function **mycompfun** instead of the default comparison function.

```
-->params = init_param();

-->params = add_param(params,"compfun",mycompfun);

-->mergesort ( [4 5 1 6 2 3]', params )'

ans =

2. 4. 6. 1. 3. 5.
```

We did not configure the **direction** parameter, so that the default "increasing" order was used. As we can see, in the output argument, the even numbers are before the odd numbers, as expected.

4.4.3 Issues with the parameters module

There is an issue with the **parameters** module, which may make the user consider default values instead of the expected ones. The **parameters** module does not protect the user against unexpected fields, which may happen if we make a mistake in the name of the key when we configure a parameter. In this section, we show how the problem may appear from a user's point of view. From the developer's point of view, we present the **check_param** function which allows to protect the user from using a key which does not exist.

In the following session, we call the mergesort function with the wrong key "compfun" instead of "compfun".

```
-->params = init_param();
-->params = add_param(params,"compffun",mycompfun);
-->mergesort ( [4 5 1 6 2 3]', params )'
ans =
1. 2. 3. 4. 5. 6.
```

We see that our wrong "compffun" key has been ignored, so that the default comparison function is used instead: the matrix has simply been sorted in to increasing order. The reason is that the **parameters** module does not check that the "compffun" key does not exist.

This is why we suggest to use the following check_param function, which takes as intput arguments the list of parameters params and a matrix of strings allkeys. The allkeys variable stores the list of all keys which are available in the list of parameters. The algorithm checks that each key in params is present in allkeys. The output arguments are a boolean noerr and a string msg. If there is no error, the noerr variable is true and the string msg is an empty matrix. If one key of params is not found, we set noerr to false and compute an error message. This error message is used to actually generate an error only if the output argument msg was not used in the calling sequence.

```
function [noerr,msg] = check_param(params,allkeys)
 if ( typeof(params) <> "plist" ) then
    lclmsg = gettext("%s: Wrong type for "+..
        "input argument #%d: %s expected.\n")
    error(sprintf(lclmsg, "check_param", 1, "plist"))
 end
 if ( typeof(allkeys) <> "string" ) then
    lclmsg = gettext("%s: Wrong type for "+..
        "input argument #%d: %s expected.\n")
    error(sprintf(lclmsg, "check_param", 2, "string"))
 end
 currkeys = getfield(1,params)
 nkeys = size(currkeys,"*")
 noerr = %t
 msg = []
 // The key #1 is "plist".
 for i = 2 : nkeys
   k = find(allkeys==currkeys(i))
    if (k == []) then
      noerr = %f
      lclmsg = gettext("%s: Unexpected key ""%s"" "+..
```

```
"in parameter list.")
msg = msprintf(lclmsg,"check_param",currkeys(i))
if ( lhs < 2 ) then
error ( msg )
end
break
end
end
end
end
end</pre>
```

The following version of the mergesort function uses the check_param function in order to check that the list of available keys is the matrix ["direction" "compfun"].

```
function y = mergesort ( varargin )
  [lhs,rhs]=argn()
  if ( and( rhs <> [1 2] ) ) then
    errmsg = sprintf(..
      "%s: Unexpected number of arguments : "+..
      "%d provided while %d to %d are expected.",..
      "mergesort", rhs, 1, 2);
    error(errmsg)
  end
  x = varargin(1)
  if ( rhs < 2 ) then
    params = []
  else
    params = varargin(2)
  end
  check_param(params,["direction" "compfun"])
  [direction,err] = get_param(params,"direction",%t)
  [compfun,err] = get_param(params,"compfun",compfun_default)
  y = mergesortre ( x , direction , compfun )
endfunction
```

In the following session, we use, as previously, the wrong key "compffun" instead of "compfun".

```
-->params = init_param();
-->params = add_param(params,"compffun",mycompfun);
-->mergesort ( [4 5 1 6 2 3]', params )
!--error 10000
check_param: Unexpected key "compffun" in parameter list.
at line 75 of function check_param called by :
at line 16 of function mergesort called by :
mergesort ( [4 5 1 6 2 3]', params )
```

With our fixed function, an error message is generated, which allows to see our mistake.

4.5 The scope of variables in the call stack

In this section, we analyze the scope of variables and how this can interact with the behavior of functions. As this feature may lead to unexpected bugs, we warn against the use of this feature in functions which may be designed in a better way. Then we present two different cases where the scope of variables is poorly used and how this can lead to difficult bugs.

4.5.1 Overview of the scope of variables

In this section, we present the scope of variables and how this can interact with the behavior of functions.

Assume that a variable, for example **a**, is defined in a script and assume that the same variable is used in a function called directly or indirectly. What happens in this case depends on the first statement reading or writing the variable **a** in the body of the function.

- If the variable **a** is first read, then the value of the variable at the higher level is used.
- If the variable **a** is first written, then the local variable is changed (but the higher level variable a is not changed).

This general framework is now presented in several examples.

In the following script, we present a case where the variable is read first. The following function f calls the function g and evaluates an expression depending on the input argument x and the variable a.

```
function y = f ( x )
y = g ( x )
endfunction
function y = g ( x )
y = a(1) + a(2)*x + a(3)*x^2
endfunction
```

Notice that the variable **a** is *not* an input argument of the function **g**. Notice also that, in this particular case, the variable **a** is only read, but not written.

In the following session, we define the variables \mathbf{a} and \mathbf{x} . Then we call the function \mathbf{f} with the input argument \mathbf{x} . When we define the value of \mathbf{a} , we are at the calling level #0 in the call stack, while in the body of the function \mathbf{g} , we are at the calling level #2 in the call stack.

```
-->a = [1 2 3];
-->x = 2;
-->y = f (x)
y =
17.
-->a
a =
1. 2. 3.
```

We see that the function f was evaluated correctly, using, in the body of the function g, at the calling level #2, the value of a defined at the calling level #0. Indeed, when the interpreter comes to the body of g, the variable a is used, but is not defined. This is because it is neither an input argument, nor locally defined in the body of g. Therefore, the interpreter searches for a at higher levels in the call stack. In the body of f, at the level #1, there is no variable a. Hence, the interpreter searches at a higher level. At the level #0, the interpreter finds the variable a, with the content



Figure 25: Scope of variables. – When a variable is used, but not defined at a lower level, the interpreter searches for the same variable at a higher level in the call stack. The first variable which is found is used.

[1 2 3]. This variable is used to evaluate the expression $y = a(1) + a(2)*x + a(3)*x^2$, which finally allows to return the variable y with the correct content. This process is presented in the figure 25.

In practice, we should change the design of the function g in order to make the use of the variable a more clear. The following script defines a modified version of the function g, where the input argument a makes clear the fact that we use the variable a.

```
function y = gfixed ( x , a )
y = a(1) + a(2)*x + a(3)*x^2
endfunction
```

The function **f** should be updated accordingly, in order to provide to the function **gfixed** the argument that it needs. This is presented in the following script.

```
function y = ffixed ( x , a )
y = gfixed ( x , a )
endfunction
```

The client source code should be also updated, as in the following session.

```
-->a = [1 2 3];
-->x = 2;
-->y = ffixed ( x , a )
y =
17.
```

The functions **ffixed** and **gfixed** are, from a software engineering point of view, much clearer that their respective previous versions.

We now present a case where the variable a is written first. The following function f2 calls the function g2 and evaluates an expression depending on the input argument x and the variable a.

```
function y = f2 ( x )
y = g2 ( x )
endfunction
function y = g2 ( x )
a = [4 5 6]
y = a(1) + a(2)*x + a(3)*x^2
endfunction
```

Notice that the variable **a** is written before it is used.

The following session uses the same script as before.

```
-->a = [1 2 3];
-->x = 2;
-->y = f2 ( x )
y =
38.
-->a
a =
1. 2. 3.
```

We notice that the value returned by the function f2 is 38, instead of the previous value 17. This confirms that the local variable content $a = [4 \ 5 \ 6]$, defined at level #2 was used, and not the variable $a = [1 \ 2 \ 3]$, defined at level #0. Moreover, we notice that the variable a defined at level #0 has not changed after the call to f2: its content remains $a = [1 \ 2 \ 3]$.

The call stack may be very deep, but the same behavior will always occur: whatever the level in the call stack, if a variable is first read and was defined at a higher level, then the value will be used directly. This is a feature which is seen by some users as convenient. We emphasize that it might lead to bugs which may be invisible and hard to detect. Indeed, the developer of the function g may have inadvertently misused the variable a instead of another variable. This leads to issues which are presented in the next section.

4.5.2 Poor function: an ambiguous case

In the two next sections, we present cases where wrong uses of the scope of variables lead to bugs which are difficult to analyze. In the first case, we present two nested function which produce the wrong result because the function at level #2 in the call stack contains a bug and uses the variable defined at level #1. In the second case, a function using a callback silently fails, because it uses the same variable name as the user.

An ambiguous case is presented in the following perverse example. We first define the function f3, which takes x and a as input arguments. This function defines the variable b, then calls the function g3. The function g3 takes x and a as input arguments, but, surprisingly, does not use the value of a. Instead, the expression involves the variable b.

```
function y = f3 ( x , a )
b = [4 5 6]
```

y = g3 (x, a) endfunction function y = g3 (x, a) y = b(1) + b(2)*x + b(3)*x^2 endfunction

In the following session, we define the value of a and x and call the function f3.

```
-->a = [1 2 3];

-->x = 2;

-->y = f3 ( x , a )

y =

38.

-->expected = a(1) + a(2)*x + a(3)*x^2

expected =

17.
```

The value returned by f3 corresponds to the parameters associated with the variable b=[4 5 6]. Indeed, the variable b used in g3 has been defined in f3, at a higher level in the call stack.

It is obvious that, in this simple case, the function g3 is badly designed and may contain a bug. Indeed, we see that the input argument a is never used, while b is used, but is not an input argument. The whole problem is that, depending on the context, this may or may not be a bug: we do not know.

4.5.3 Poor function: a silently failing case

In this section, we present a typical failure caused by a wrong use of the scope of variables.

The following function myalgorithm, provided by the developer, uses an order 1 numerical derivative formula, based on a forward finite difference. The function takes the current point x, the function f and the step h, and returns the approximate derivative y.

```
// From the developer
function y = myalgorithm ( x , f , h )
y = (f(x+h) - f(x))/h
endfunction
```

The following function myfunction, written by the user, evaluates a degree 2 polynomial by a straightforward formula. Notice that the evaluation of the output argument y makes use of the point x, which is an input argument, and the parameter a, which is not an input argument.

```
// From the user
function y = myfunction ( x )
y = a(1) + a(2)*x + a(3)*x^2
endfunction
```

In the following session, we set a, x, h and call myalgorithm in order to compute a numerical derivative. We compare with the exact derivative and get a good agreement.

 $-->a = [1 \ 2 \ 3];$ -->x = 2;

```
-->h = sqrt(%eps);
-->y = myalgorithm ( x , myfunction , h )
y =
    14.
-->expected = a(2) + 2 * a(3)*x
expected =
    14.
```

The scope of the variable a allows to evaluate the function myfunction. Indeed, the variable a is defined by the user at the level #0 in the call stack. When the body of myfunction is reached, the interpreter is at the level #2, where the variable a is used, but not set. Therefore, the interpreter searches for a variable a at a higher level in the call stack. There is no such variable at the level #1, which is why the variable a defined at the level #0 is used, finally producing the expected result.

In fact, this way of using the scope of variables is a dangerous programming practice. Indeed, the user make assumptions about the internal design of the myalgorithm function, and these assumptions may be false, leading to wrong results.

We change slightly the function provided by the developer, and renamed the step as **a**, instead of the previous **h**.

```
// From the developer
function y = myalgorithm2 ( x , f , a )
y = (f(x+a) - f(x))/a
endfunction
```

In the following session, we execute the same script as previously.

```
-->a = [1 2 3];
-->x = 2;
-->h = sqrt(%eps);
-->y = myalgorithm2 ( x , myfunction , h )
!--error 21
Invalid index.
at line 2 of function myfunction called by :
at line 2 of function myalgorithm2 called by :
y = myalgorithm2 ( x , myfunction , h )
```

The "Invalid index" error is the consequence of the fact that the variable **a** has been overwritten in the body of myalgorithm2, at the level #1 in the call stack. As previously, when the body of the function myfunction is reached, at the level #2 in the call stack, the interpreter searches for a variable **a** at a higher level. At the level #1, in myalgorithm2, the interpreter finds the variable **a**, which contains the step of the finite difference formula. This variable **a** is a matrix of doubles with only 1 entry. When the interpreter tries to evaluate the statement **a**(1) + **a**(2)***x** + **a**(3)***x**², this fails since neither **a**(2) nor **a**(3) exist, that is, the integers 2 and 3 are invalid indices of the variable **a**.

The previous error was, in fact, a nice one. Indeed, it warns us that something wrong happened, so that we can change our script. In the next case, the script will pass, but with a wrong result and this is a much more dangerous situation.

We change again the body of the algorithm provided by the developer. This time, we set the variable **a** to an arbitrary matrix of doubles, as in the following myalgorithm3 function.

```
// From the developer
function y = myalgorithm3 ( x , f , h )
    a = [4 5 6]
    y = (f(x+h) - f(x))/h
endfunction
```

The body is somewhat strange, since we do not make use of the variable **a**. But the function is still valid. Actually, this represent more complicated cases, where the implementation of the myalgorithm3 is using a full, possibly large, set of variables. In this case, the probability of using the same variable name as the user is much higher.

In the following session, we call the myalgorithm3 function and compare with the expected result.

```
-->a = [1 2 3];
-->x = 2;
-->h = sqrt(%eps);
-->y = myalgorithm3 ( x , myfunction , h )
y =
29.
-->expected = a(2) + 2 * a(3)*x
expected =
14.
```

We see that the script did not produce an error. We also see that the expected result is completely wrong. As previously, the variable **a** has been re-defined at the level #1, in the body of the myalgorithm3. Therefore, when the expression $a(1) + a(2)*x + a(3)*x^2$ is evaluated, the value [4 5 6] is used, instead of the matrix that the user provided at a higher level.

In our particular case, it is easy to debug the problem, because we have the exact formula for the derivative. In practice, we probably do not have the exact formula (which is why we use a numerical derivative...), so that it would be much more difficult to detect and, if detected, solve the issue.

Using a variable defined at a higher level in the call stack may be considered as a bad programming practice. In the case where this was not wanted by the developer of the function, this may lead to bugs which are difficult to detect and may stay unnoticed for a long time before getting fixed.

Used in this way, the scope of variables may be considered as a bad programming practice. Still, it may happen that a function needs more than one input argument to be evaluated. This particular issue is addressed in the section 4.6.4, where we present a method to provide additional input arguments to a callback.

4.6 Issues with callbacks

In this section, we analyze particular issues with callbacks. We first consider interactions between the names of the functions of the user and the developer. We present two different types of issues in specific examples. Then we present methods which allows to partially solve this issue. In the final section, we present a method which allows to manage callbacks with additional arguments. The method that we advocate is based on lists, which provide a good flexibility, since a list can contain any other data type. We emphasize that the two first issues that we are going to present in the sections 4.6.1 and 4.6.2 are not bugs of the interpreter. As we are going to see, these issues are generated by the scope of variables which has been presented in the previous section. It is likely that these issues will remain in future versions of the interpreter. Therefore, it is worth to analyze them in some detail, so that we can understand and solve this particular problem efficiently.

4.6.1 Infinite recursion

In this section, we present an issue which generates an infinite recursion. This issue occurs when there is an interaction between the name of the function chosen by the developer and the user of an algorithm. This is why our analysis must separate the developer's point of view on one side, and the user's point of view on the other side.

The following myalgorithm function, provided by a developer, takes the variables x and f as input arguments and returns y. The variable f is assumed to be a function and the expression y=f(x) is evaluated in a straightforward way.

```
// At the developer level
function y = myalgorithm ( x , f )
y = f(x)
endfunction
```

The following myfunction function, provided by a user, takes x as an input argument and returns y. In order to compute y, the user applies a second function f that he has written for that purpose.

```
// At the user level
function y = myfunction ( x )
y = f ( x )
endfunction
function y = f ( x )
y = x(1)^2 + x(2)^2
endfunction
```

In the following session, the user sets the variable x and calls the myalgorithm function with the input arguments x and myfunction.

```
-->x = [1 2];
-->y = myalgorithm ( x , myfunction )
!--error 26
Too complex recursion! (recursion tables are full)
at line 2 of function myfunction called by :
at line 2 of function myfunction called by :
at line 2 of function myfunction called by :
[...]
```

This recursion, which is in theory infinite, is, in fact, finite, because Scilab authorizes only a limited number of recursive calls.

The cause of this failure is the conflict of the names of the developer's and user's function which, in both cases, uses the variable name f. From the user's point of view, the function myfunction simply calls f. But the interpreter does not see the script like this. From the interpreter's point of view, the symbols myfunction and f are variables which are stored in the interpreter's internal data structures. The

variable myfunction has a particular data type: it is a function. The same is true for the user's f. Both these variables are defined at the level #0 in the call stack.

Hence, before the call to myalgorithm, at the level #0, the function f is defined by the user, with the body " $y = x(1)^2 + x(2)^2$ ". When myalgorithm is called, we enter in the level #1 and the variable **f** is considered as an input argument: its content is f=myfunction. Hence, the function f, which was previously defined by the user, has been overwritten and has lost its original value. The interpreter evaluates the expression y = f(x), which calls back myfunction. In the body of myfunction, at the level #2, the interpreter finds the expression y = f(x). Then the interpreter searches for a variable **f** at the current level in the call stack. There is no such variable f. Hence, the interpreter searches for a variable f at a higher level in the call stack. At the level #1, the variable **f** is defined, with content f=myfunction. This value is then used at the level #2. The interpreter evaluates the expression y = myfunction (x) and enters at level #3. Then the interpreter searches for a variable f at the current level. There is no such variable at this level. As previously, the interpreter searches for a variable f at a higher level in the call stack. Again, the interpreter calls f, which is, in fact the function myfunction itself. This process repeats over and over again, until the interpreter reaches the maximum authorized size of the call stack and generates an error. The following pseudo-session details the sequence of events.

```
-->y = myalgorithm ( x , myfunction )
-1->f = myfunction
-1->y = f(x)
-1->y = myfunction(x)
-2->y = f ( x )
-2->y = myfunction ( x )
etc...
```

We emphasize that the problem is a consequence of the scope of variables in the Scilab language. It is a straightforward implication of the following features of the language.

- A function is stored as any other variable (but with a particular data type).
- If a variable is unknown at level #k, it is searched at higher levels in the call stack until it is either found or an error is generated (the exact error is "Undefined variable").

Notice that there is no mean for the user to know that the developer has used the variable name **f**. The converse is also true for the developer, who cannot predict that the user will use the variable name **f**. Notice that there is little way for the user to change this situation. The **myalgorithm** function may be provided by an external module. In this case, changing the name of the variable used in the developer's function may be impossible for the user. As we are going to see later in this section, the user can still find a solution, based on a small source code reorganization.

4.6.2 Invalid index

In this section, we present a situation where the interpreter generates a "Invalid index" error, because of a conflict between the user and the developer's function names.

In the following myalgorithm function, provided by the developer, we set the local variable f to 1. Then we call back the userf function with the input argument x, and return the output argument y.

```
// At the developer level
function y = myalgorithm ( x , userf )
  f = 1
  y = userf(x)
endfunction
```

Notice that the variable **f** is set, but not used. The function is still valid and this does not change our analysis.

The following myfunction function, provided by the user, calls the function f with the input argument x and returns the output argument y.

```
// At the user level
function y = myfunction ( x )
y = f ( x )
endfunction
function y = f ( x )
y = x(1)^2 + x(2)^2
endfunction
```

In the following session, we set the variable x and call the myalgorithm function. This generates a "Invalid index" error in the body of myfunction.

```
-->x = [1 2];
-->y = myalgorithm ( x , myfunction )
!--error 21
Invalid index.
at line 2 of function myfunction called by :
at line 4 of function myalgorithm called by :
y = myalgorithm ( x , myfunction )
```

The explanation is the following. At the level #0 in the call stack, the function f is defined as the user expects. In the body of myalgorithm, we are at level #1 in the call stack. The variable f is updated with the f=1 statement: f is now a matrix of doubles. Then, the interpreter evaluates the expression y = userf(x). Since userf is myfunction, the interpreter calls the function myfunction and enters in the level #2 in the call stack. At this level, the interpreter evaluates the expression y = f (x). The interpreter then searches for the variable f at the level #2: there is no such variable at this level. The interpreter finds the variable f, which is a matrix of doubles. In this context, the expression y = f (x) has no sense: the variable x is interpreted as the index of the matrix of doubles f. Since $x=[1 \ 2]$, the interpreter looks for the entries at indices 1 and 2 in f. But f contains a 1-by-1 matrix of double, that is 1. Hence, there is only one entry in f, and this is why the error "Invalid index" error is generated.

As in the previous section, the user cannot predict the variable names chosen by the developer and the converse is true for the developer.

In the next section, we suggest methods to solve these callback problems.

4.6.3 Solutions

In this section, we present solutions for the callback issues that we have presented earlier. Solving the problem can be done by two different ways.

- Change the developer's function, so that there is less chance to get a conflict with user's variable names.
- Change the user's function, so that he can use the developer's function as is.

Obviously, we may also do both. In general, it is the task of the developer to provide functions which are well enough designed so that the user does not have to be troubled by strange bugs. This is why we present this update first. The update of the user's function is presented in the second part of this section.

The following modified function presents the method that the developer can use to write the myalgorithm function. We can see that the second argument __myalgorithm_f__ has a long and complicated name. This reduces the chances of getting a conflict between the developer's and the user's variable names.

```
function y = myalgorithm ( x , __myalgorithm_f__ )
y = __myalgorithm_f__(x)
endfunction
```

This is a workaround: there is still the possibility that the user get an error. Indeed, if the user chose the variable name __myalgorithm_f__, there is still a conflict between the developer's and the user's variable names. But this is much less likely to happen.

The following modified function presents the method that the user can use to write the myfunction function. Instead of defining the function f outside of the body of myfunction, we define it inside. In the new version, the variable f is now defined locally, inside myfunction. This limits the visibility of the local function f, which can only be accessed by myfunction (and lower levels), and does not appear at the global scope anymore. Hence, when we call f, the interpreter does not search at higher levels in the call stack: the variable f is defined at the current level.

```
function y = myfunction ( x )
function y = f ( x )
y = x(1)^2 + x(2)^2
endfunction
y = f ( x )
endfunction
```

In the following session, we call the myalgorithm function with the updated function myfunction.

```
-->x = [1 2];
-->y = myalgorithm ( x , myfunction )
y =
5.
```

As we can see, the issue is now solved and we get the expected result.

4.6.4 Callbacks with extra arguments

In this section, we present a method which allows to solve issues associated with callbacks with extra arguments. The method that we advocate is based on lists, which provide a good flexibility, since it can contain any other data type. When we consider an algorithm which takes a callback as input argument, the header of the function being called back is generally definitely fixed, by a choice of the developer of the algorithm. But the user of the algorithm may want to provide a function which does not exactly matches the required header. For example, the algorithm may expect a function with the header y=f(x). But the user has a function which requires the additional parameter a. In order to avoid confusions created by the use of the scope of variables, the user may chose to update the header of the function so that the variable a is now an input argument. This leads to the header y=g(x,a). In this case, we assume that the additional argument a is constant, which means that the algorithm will not modify its content. Simply providing the function g may not work, because the header does not match. In order to illustrate this situation, we analyze a case where we want to compute a numerical derivative.

The following myderivative1 function uses a forward finite difference formula to evaluate a numerical derivative.

```
function fp = myderivative1 ( __myderivative_f__ , x , h )
fp = (__myderivative_f__(x+h) - __myderivative_f__(x))/h
endfunction
```

We consider the function myfun, which computes the cosine of a degree 2 polynomial.

```
function y = myfun (x)

y = cos(1+2*x+3*x^2)

endfunction
```

In the following session, we compare the numerical derivative with the exact derivative.

As the two values consistently match, we are now confident in our implementation.

But it would be clearer if the parameters of the polynomial were stored in a matrix of doubles. This leads to the following function myfun2, which takes the point x and the parameter a as input arguments.

function y = myfun2 (x , a) y = $cos(a(1)+a(2)*x+a(3)*x^2)$ endfunction

In order to manage this situation, we modify the implementation of the finite difference algorithm and create the myderivative2 function, which will be detailed later in this section. In the following session, we call myderivative2 and provide the list list(myfun2,a) as the first argument. The myderivative2 function assume that the first element of the list is the name of the function and that the remaining elements in the list are the additional arguments of the function to be evaluated. Here, the only additional argument is a.

```
-->x = %pi/6;
-->a = [1 2 3];
-->h = %eps^(1/2);
-->fp = myderivative2 ( list(myfun2,a) , x , h )
fp =
    - 1.380975857377052307D+00
-->expected = -sin(a(1)+a(2)*x+a(3)*x^2) * (a(2)+2*a(3)*x)
expected =
    - 1.380976033975957140D+00
```

The following myderivative2 function provides a flexible numerical derivative implementation, which allows for additional arguments in the header of the function to be differenciated.

```
function fp = myderivative2 ( __myderivative_f__ , x , h )
1
\mathbf{2}
     tyfun = typeof(__myderivative_f__)
3
     if ( and(tyfun<>["list" "function" "fptr"]) ) then
4
       error ( msprintf("%s: Unknown function type: %s",...
          "myderivative2",tyfun))
5
6
     end
     if ( tyfun == "list" ) then
7
8
       nitems = length(__myderivative_f__)
       if ( nitems < 2 ) then
9
         error ( msprintf("%s: Too few elements in list: %d",...
10
            "myderivative2", nitems))
11
12
       end
13
       __myderivative_f__fun__ = __myderivative_f__(1)
14
       tyfun = typeof(__myderivative_f__fun__)
       if ( and(tyfun<>["function" "fptr"]) ) then
15
         error ( msprintf("%s: Unknown function type: %s",...
16
            "myderivative2",tyfun))
17
18
       end
       fxph=__myderivative_f__fun__(x+h, __myderivative_f__(2:$))
19
       fx = __myderivative_f__fun__ ( x , __myderivative_f__(2:$))
20
21
     else
22
       fxph = \__myderivative_f\__ (x + h)
23
       fx = __myderivative_f__ ( x )
24
     end
25
     fp = (fxph - fx)/h
26
   endfunction
```

At line #2, we compute the type of the input argument __myderivative_f_... If this argument is neither list, nor a function (i.e. a macro), nor a function pointer (i.e. a primitive), then we generate an error. Then the code considers two cases. If the argument is a list, then we check in the lines #8 to #12 the number of items in the list. Then, at line #13, we store in a separate variable the first element, which is assumed to be the function. In the lines #14 to #18, we check the type of this variable and generate an error if the variable is not a function. Then, at lines #19 and #20, we evaluate the two function values fxph and fx. Notice that we use the expression __myderivative_f_(2:\$) in order to provide the additional arguments to the called function. Because of the special way that the elements are extracted from a list, this creates the number of input arguments which is required by header of the function which is being called.

In practice, the method that we have presented is extremely flexible. More that

| deff | on-line definition of function |
|---------|--------------------------------|
| execstr | execute statements in string |
| evstr | evaluation of string |

Figure 26: Meta programming functions.

one additional argument can be provided by the function. Actually, the number of additional arguments may be reduced by using a list gathering all the necessary parameters. Indeed, lists can be nested, which allows to gather all the required parameters into one single additional variable.

4.7 Meta programming: execstr and deff

In this section, we present functions which allow to execute statements which are defined as strings. As these strings can be dynamically generated, we can make programs which have a flexibility level which cannot be achieved by other means. In the first part, we present the **execstr** function and give a sample use of this extremely powerful function. In the second part, we present the **deff** function, which allows to dynamically create functions based on two strings containing the header and the body of the function. In the final part, we present a practical use of the **execstr** function, where we make the glue between two modules which cannot be modified by the user and which do not match exactly.

The figure 26 present the functions which allow to dynamically execute statements based on strings.

4.7.1 Basic use for execstr

In this section, we present the execstr function.

The execstr function takes as its first input argument a string which contains a valid Scilab statement. Then, the function executes the statement as if it was a part of the original script, at the same level in the call stack.

In the following session, we dynamically generate a string, containing a statement which displays the string "foo". We first define the variable s, which contains the string "foo". Then we set the variable instr, by concatenating the variable s with the strings disp(" and "). This is performed with the + operator, which allows to concatenate its operands. The double quotes " are doubled, which makes so that the expression "" is interpreted as a " character inside the target string. Indeed, if this quote does not appear twice, it is interpreted as the end of the string: doubling the quote allows to "escape" the quote character. Finally, we run the execstr function, which displays "foo".

```
-->s = "foo"
s =
foo
-->instr = "disp(""" + s + """)"
instr =
disp("foo")
-->execstr(instr)
```

foo

The argument of execstr can contain any valid Scilab statement. For example, we can dynamically set a new variable, with a dynamically created content. In the following session, we create a string instr containing the statement z=1+2. Then we evaluate this expression and check that the variable z has, indeed, been set to 3.

```
-->x = 1

x =

1.

-->y = 2

y =

2.

-->instr = "z="+string(x)+"+"+string(y)

instr =

z=1+2

-->execstr(instr)

-->z

z =

3.
```

As another example of the use of execstr, we can read a collection of data files. Assume that the file datafile1.txt contains the lines

1 2 3 4 5 6

and the file datafile2.txt contains the lines

7 8 9 10 11 12

The following script allows to read these two files in sequence. The **read** function takes a string representing a file containing a matrix of doubles, and the number of rows and columns in the matrix. The script sets the variable **data**, which contains a list of matrices containing the data that have been read in the two files.

```
data = list();
for k = 1 : 2
  instr = "t = read(""datafile" + string(k) + ".txt"",2,3)"
  execstr(instr)
  data($+1)=t
end
```

The following session shows the dynamic of the previous script.

```
instr
       =
t = read("datafile1.txt",2,3)
data =
      data(1)
   1.
       2.
               З.
   4.
         5.
               6.
instr
       =
t = read("datafile2.txt",2,3)
data =
      data(1)
```

| 1. | 2. | З. |
|-----|---------|-----|
| 4. | 5. | 6. |
| | data(2) | |
| 7. | 8. | 9. |
| 10. | 11. | 12. |

4.7.2 Basic use for deff

The deff function allows to dynamically define a function, from two strings containing the header and the body of the function.

In the following session, we define the function myfun, which takes the matrix of doubles x as input argument and returns the output argument y. First, we define the header of the function as the string "y=myfun(x)" and set the header variable. Then we define the body of the function and, finally, we call the deff function, which creates the required function.

```
-->header = "y=myfun(x)"
header =
y=myfun(x)
-->body = [
-->"y(1) = 2*x(1)+x(2)-x(3)^2"
-->"y(2) = 2*x(2) + x(3)"
-->]
body =
!y(1) = 2*x(1)+x(2)-x(3)^2 !
!
!y(2) = 2*x(2) + x(3) !
-->deff(header,body)
```

The following session shows that the new function **myfun** can be used as any other function.

```
-->x = [1 2 3]
x =
1. 2. 3.
-->y = myfun(x)
y =
- 5.
7.
```

The only difference is the type of a function created by deff, as presented in the section 4.1.1. In the following session, we show that a function created by deff has type 13, which corresponds to a compiled macro.

```
-->type(myfun)
ans =
13.
-->typeof(myfun)
ans =
function
```

If we add the "n" argument to the call of deff, this tells the interpreter to create an uncompiled macro instead, which produces a function with type 11.

```
-->deff(header,body,"n")
Warning : redefining function: myfun.
```

```
Use funcprot(0) to avoid this message
-->type(myfun)
ans =
    11.
-->typeof(myfun)
ans =
function
```

We emphasize that the execstr function could be used instead of deff to create a new function. For example, the following script shows a way of defining a new function with the execstr, by separately defining the header, body and footer of the function. These strings are then concatenated into a single instruction, which is finally executed to define the new function.

```
header = "function y = myfun(x)"
body = [
"y(1) = 2*x(1)+x(2)-x(3)^2"
"y(2) = 2*x(2) + x(3)"
]
footer = "endfunction"
instr = [
header
body
footer
]
execstr(instr)
```

4.7.3 A practical optimization example

In this section, we present a practical use-case for the methods that we have presented in previous sections.

In this example, we use the **optim** function, which is a numerical optimization solver. We present a situation where we need a function to define an adapter (i.e. a "glue") function which connects **optim** to a module providing a collection of optimization problems. In the first method, we define an adapter function and pass a callback with extra arguments to the **optim** function. In the second method, we use the **execstr** function to create a function which body is dynamically generated.

The example that we have chosen might seem complicated. In practice, though, it represents a practical software engineering problem where only non-trivial solutions can be used.

Assume that we are interested in the evaluation of the performance of the nonlinear optimization function optim provided by Scilab. In this case, we can use the Atoms module "uncprb", which provides a collection of 23 unconstrained optimization test problems known as the More, Garbow and Hillstrom collection. In order to install this module, we use the statement:

```
atomsInstall("uncprb")
```

and restart Scilab.

We emphasize that we consider here the point of view of a user which cannot modify any of these packages, that is, cannot modify neither the optim function, nor the uncprb module. As we will soon see, there is not an exact match between the header of the objective function which is required by optim and the header of the test functions which are provided by the uncprb module. More precisely, the number and type of input and output arguments which are required by optim do not match the number and type of input and output arguments which are provided by uncprb. This is not caused by a poor design of both tools: it is in fact impossible to provide a "universal" header, matching all the possible needs. Therefore, we have to create an intermediate function, which makes the "glue" between these two components.

The uncprb module provides the function value, the gradient, the function vector and the Jacobian for all the test problems, and provides the Hessian matrix for 18 problems. More precisely, the module provides the following functions, where nprob is the problem number, from 1 to 23.

```
[n,m,x0]=uncprb_getinitf(nprob)
f=uncprb_getobjfcn(n,m,x,nprob)
g=uncprb_getgrdfcn(n,m,x,nprob)
```

The uncprb_getinitf function returns the size n of the problem, the number m of functions and the initial guess x0. Indeed, the objective function is the sum of the squares of m functions. The uncprb_getobjfcn function returns the value of the objective function while the uncprb_getgrdfcn function returns the gradient.

The optim function is a nonlinear unconstrained optimization solver, providing several algorithms for this class of problems. It can manage unconstrained or bound constrained problems. The simplest calling sequence of the optim function is

```
[fopt,xopt]=optim(costf,x0)
```

where costf is the cost (i.e. objective) function to minimize, x0 is the starting point (i.e. initial guess), fopt is the minimum function value and xopt is the point which achieves this value. The cost function costf must have the header

```
[f,g,ind]=costf(x,ind)
```

where \mathbf{x} is the current point, ind is a floating point integer representing what is to be computed, \mathbf{f} is the function value and \mathbf{g} is the gradient. On output, ind is a flag sent by the function to the optimization solver, for example to interrupt the algorithm.

We must first get the problem parameters, so that we can setup the glue function. In the following script, we get the parameters of the first optimization problem.

```
nprob = 1;
[n,m,x0] = uncprb_getinitf(nprob);
```

The question is: how to glue these two components, so that optim can use the uncprb module? Obviously, we cannot pass neither the uncprb_getobjfcn nor the uncprb_getgrdfcn functions as input arguments to optim, since the objective function must compute both the function value and the gradient. The solution to this problem is to define an intermediate function, which has the header required by optim and which calls the uncprb_getobjfcn and uncprb_getgrdfcn functions. The only problem is the management of the nprob, n and m, which depends on the problem to solve. There may be two different ways to do this.

• We can define a function with the extra arguments nprob, n and m and use a special feature of optim to manage them.

• We can define a function which is dynamically defined with execstr so that, in the end, the variables nprob, n and m are constants.

In the following, we explore both approaches.

The first method is based on the management of extra arguments of a callback which has been presented in the section 4.6.4. We define the objective function costfun, which has the expected calling sequence, with the extra arguments nprob, n and m.

```
function [f,g,ind]=costfun(x,ind,nprob,n,m)
    [n,m,x0] = uncprb_getinitf(nprob)
    // Make a column vector
    x = x(:)
    f = uncprb_getobjfcn(n,m,x,nprob)
    g = uncprb_getgrdfcn(n,m,x,nprob)
endfunction
```

We can then use a special feature of the optim function, which manages the case where the objective function needs extra input arguments. The argument costf can also be the list (myfun,a1,a2,...). In this case, myfun, the first element in the list, must be a function and must have the header:

[f,g,ind]=myfun(x,ind,a1,a2,...)

We use this feature in the following script.

```
nprob = 1;
[n,m,x0] = uncprb_getinitf(nprob);
myobjfun = list(costfun,nprob,n,m);
[fopt,xopt]=optim(myobjfun,x0)
```

The previous script produces the following output.

```
-->[fopt,xopt]=optim(myobjfun,x0)
xopt =
1.
1.
fopt =
0.
```

We know that the global minimum of this problem is $\mathbf{x}^* = (1, 1)$, so that the previous result is correct.

The second method is based on the dynamic creation of the objective function with the **execstr** function.

In the following script, we define the objfun function, which will be passed as an input argument to the optim function. The body of objfun is dynamically defined by using the variables nprob, m and n which have been previously defined. The header, body and footer of the function are then concatenated into the instr matrix of strings.

```
header = "function [fout,gout,ind]=objfun(x,ind)";
body = [
"fout=uncprb_getobjfcn("+string(n)+","+string(m)+..
",x,"+string(nprob)+")"
"gout=uncprb_getgrdfcn("+string(n)+","+string(m)+..
",x,"+string(nprob)+")"
```

```
];
footer = "endfunction"
instr = [
header
body
footer
]
```

The previous source code is rather abstract. In fact, the generated function is simple, as shown by the following session.

```
-->instr
instr =
!function [fout,gout,ind]=objfun(x,ind) !
!fout=uncprb_getobjfcn(2,2,x,1) !
!gout=uncprb_getgrdfcn(2,2,x,1) !
!endfunction !
```

As we can see, the input argument \mathbf{x} of the objfun function is the only one remaining: the other parameters have been replaced by their actual value for this particular problem. Hence, the objfun function is cleanly defined, and does not use its environment to get the value of nprob, for example. We emphasize this particular point, which shows that using the scope of variables through the call stack, as presented in the section 4.5.1, can be avoided by using the execstr function.

In order to define the objective function, we finally use execstr.

```
execstr(instr)
```

In the following session, we call the **optim** function and compute the solution of the first test problem.

```
-->[fopt,xopt]=optim(objfun,x0)
xopt =
1.
1.
fopt =
0.
```

4.8 Notes and references

The parameters module was designed by Yann Collette as a tool to provide the same features as the optimset/optimget functions in Matlab. He noticed that the optimset/optimget functions could not be customized from the outside: we have to modify the functions in order to add a new option. This is why the parameters module was created in a way which allows to allow the user to manage as many options as required.

The scope of variables has been presented in the section 4.5.1. This topic is also presented in [50].

The issues with callbacks have been presented in section 4.6. This topic has been analyzed in the bug reports #7102, #7103 and #7104 [12, 11, 9].

In [49], Enrico Segre describe several features related to functions.

In the section 4.3, we have presented a template for the design of robust functions. Using this template allows to be compatible with the code convention presented by Pierre Maréchal in [33].

| tic, toc | measure user time |
|----------|---------------------|
| timer | measure system time |

Figure 27: Performance functions.

In the section 4.3, we have presented rules to write robust functions. Writing such a robust function requires a large number of checks and may lead to code duplication. The "apifun" module [8] is an experimental attempt to provide an API to check for input arguments with more simplicity.

5 Performances

In this section, we present a set of Scilab programming skills which allow to produce scripts which are *fast*. These methods are known as *vectorization* and is at the core of most efficient Scilab functions.

We show how to use the tic and toc functions to measure the performance of an algorithm. In the second section, we analyze a naive algorithm and check that vectorization can dramatically improve the performance, typically by a factor from 10 to 100, but sometimes more. Then we analyze compiling methods and present the link between the design of the interpreter and the performances. We present the profiling features of Scilab and give an example of the vectorization of a Gaussian elimination algorithm. We present vectorization principles and compare the performances of loops against the performances of vectorized statements. In the next section, we present various optimization methods, based on practical examples. We present the linear algebra libraries used in Scilab. We present BLAS, LAPACK, AT-LAS and the Intel MKL numerical libraries which provide optimized linear algebra features and can make a sensitive difference with respect to performance of matrix operations. In the last section, we present various performance measures of Scilab, based on the count of floating point operations.

The figure 27 presents some Scilab functions which are used in the context of analysing the performance of a Scilab script.

5.1 Measuring the performance

In this section, we describe the functions which allow to measure the time required by a computation. Indeed, before optimizing any program, we should precisely measure its current performances.

In the first section, we present the tic, toc and timer functions. Then we compare these functions and emphasize their difference on multi-core machines. We present the profiling features of Scilab which allow to analyse the parts of an algorithm which cost the most. Finally, we present the benchfun function which allows to produce easy and reliable performance analyses.

5.1.1 Basic uses

In order to measure the time required by a computation, we can use the tic and toc functions which measure the *user* time in seconds. The user time is the wall clock time, that is, the time that was required by the computer from the start of the task to its end. This includes the computation itself, of course, but also all other operations of the system, like refreshing the screen, updating the file system, letting other process do a part of their work, etc... In the following session, we use the tic and toc functions to measure the time to compute the eigenvalues of a random matrix, based on the spec function.

```
-->tic(); lambda = spec(rand(200,200)); t = toc()
t =
0.1
```

The call to the **tic** function starts the counter and the call to the **toc** function stops it, returning the number of elapsed seconds. In this case, we typed the statements on the same line, using the ; separator: indeed, if we typed the instructions interactively on several lines in the console, the measured time would mainly be the time required by the typing of the statements on the keyboard.

The previous time measure is not very reliable in the sense that if we perform the same statements several times, we do not get the same time. The following is an example of this behavior.

```
-->tic(); lambda = spec(rand(200,200)); t = toc()
t =
0.18
-->tic(); lambda = spec(rand(200,200)); t = toc()
t =
0.19
```

A possible solution for this problem is to perform the same computation several times, say 10 times for example, and to average the results. In the following script, we perform the same computation 10 times, and then display the minimum, the maximum and the average time.

```
for i = 1 : 10
   tic();
   lambda = spec(rand(200,200));
   t(i) = toc();
end
[min(t) mean(t) max(t)]
```

The previous script produces the following output.

```
-->[min(t) mean(t) max(t)]
ans =
0.141 0.2214 0.33
```

If another program on the computer uses the CPU at the same time that Scilab performs its computation, the user time may increase. Therefore, the tic and toc functions are not used in situations where only the CPU time matters. In this case, we may use instead the timer function, which measures the system time. This corresponds to the time required by the CPU to perform the specific computation required by Scilab, and not by other processes.

The following script presents an example of the timer function.

```
for i = 1 : 10
   timer();
   lambda = spec(rand(200,200));
   t(i) = timer();
end
[min(t) mean(t) max(t)]
```

The previous script produces the following output.

```
-->[min(t) mean(t) max(t)]
ans =
0.100144 0.1161670 0.1602304
```

5.1.2 User vs CPU time

In this section, we analyze the difference between the user and CPU time on single and multi-core systems.

Let us consider the following script, which performs the product between two square matrices. We measure the user time with the tic and toc functions and the CPU time with the timer function.

```
stacksize("max")
n = 1000;
A = rand(n,n);
B = rand(n,n);
timer();
tic();
C = A * B;
tUser = toc();
tCpu = timer();
disp([tUser tCpu tCpu/tUser])
```

The following session presents the result when we run this script on a Linux machine with one single core at 2 GHz.

```
-->disp([tUser tCpu tCpu/tUser])
1.469 1.348084 0.9176882
```

As we can see, the two times are rather close and show that 91% of the wall clock time was spent by the CPU on this particular computation.

Now, consider the following session, performed on a Windows machine with 4 cores, where Scilab makes use of the multi-threaded Intel MKL.

-->disp([tUser tCpu tCpu/tUser]) 0.245 1.0296066 4.2024759

We see that there is a significant difference between the CPU and the wall clock time. The ratio is close to 4, which is the number of cores.

Indeed, on a multi-core machine, if Scilab makes use of a multi-threaded library, the timer function reports the sum of the times spent on each core. This is why, in most situations, we should use the tic and toc functions to measure the performances of an algorithm.

| add_profiling | Adds profiling instructions to a function |
|------------------|---|
| plotprofile | Extracts and displays execution profiles |
| profile | Extract execution profiles |
| remove_profiling | Removes profiling instructions |
| reset_profiling | Resets profiling counters |
| showprofile | Extracts and displays execution profiles |

Figure 28: Scilab commands to profile functions.

5.1.3 Profiling a function

In this section, we present the profiling features of Scilab. The profile of a function reveals the parts of the function which cost the most. This allows to let us focus on the parts of the source code which are the most interesting to improve.

We first consider a naive, slow, implementation of a Gaussian elimination algorithm and analyze its profile. We then use vectorization in order to speed up the computations and analyze its updated profile.

The functions in table 28 allow to manage the profiling of a function.

We consider in this section the resolution of systems of linear equations $A\mathbf{x} = \mathbf{b}$, where A is a $n \times n$ real matrix and **b** is a $n \times 1$ column vector. One of the most stable algorithm for this task is the Gaussian elimination with row pivoting. The Gaussian elimination with pivoting is used by Scilab under the backslash operator \backslash . In order to illustrate the profiling of a function, we are not going to use the backslash operator. Instead, we will develop our own Gaussian elimination algorithm (which is, obviously, not a good idea in practice). Let us recall that the first step of the algorithm requires to decompose A into the form PA = LU, where P is a permutation matrix, L is a lower triangular matrix and U is an upper triangular matrix. Once done, the algorithms proceeds by performing a forward and then a backward elimination.

The algorithm is both short and complex, and, in fact, rather fascinating. We do not aim to provide a thorough presentation of this algorithm in this document (see Golub and Van Loan's [21] for a more complete analysis). This algorithm is a good candidate for profiling, since it is more complex than an average function. Hence, it is not straightforward to obtain a good performance for it. This is the type of situations where profiling is the most useful. Indeed, the intuition is sometimes not sufficiently accurate to detect the "guilty lines", that is, the lines which are a bottleneck for the performance. All in all, measuring the performance is often better than guessing.

The following gausspivotalnaive function is a naive implementation of the Gaussian elimination with row pivoting. It takes the matrix A and right-hand side b as input arguments, and returns the solution x. It combines the PA = LU decomposition with the forward and backward substitutions in one single function. The terms of the L matrix are stored into the lower part of the matrix A, while the terms of the matrix U are stored in the upper part of the matrix A.

```
1 function x = gausspivotalnaive ( A , b )
2     n = size(A,"r")
```

```
3
     // Initialize permutation
     P=zeros(1,n-1)
4
5
     // Perform Gauss transforms
6
     for k=1:n-1
7
       // Search pivot
8
       mu = k
9
        abspivot = abs(A(mu,k))
10
       for i=k:n
11
          if (abs(A(i,k))>abspivot) then
12
            mu = i
13
            abspivot = abs(A(i,k))
14
          end
15
        end
16
        // Swap lines k and mu from columns k to n
17
        for j=k:n
18
          tmp = A(k,j)
          A(k,j) = A(mu,j)
19
20
          A(mu,j) = tmp
21
        end
22
       P(k) = mu
23
        // Perform transform for lines k+1 to n
24
        for i=k+1:n
25
          A(i,k) = A(i,k) / A(k,k)
26
        end
27
        for i = k+1:n
28
          for j = k+1:n
29
            A(i,j) = A(i,j) - A(i,k) * A(k,j)
30
          end
31
        end
32
     end
33
     // Perform forward substitution
34
     for k=1:n-1
        // Swap b(k) and b(P(k))
35
36
        tmp = b(k)
37
       mu = P(k)
       b(k) = b(mu)
38
39
       b(mu) = tmp
40
        // Substitution
       for i=k+1:n
41
42
          b(i) = b(i) - b(k) * A(i,k)
43
        end
44
     end
45
     // Perform backward substitution
     for k=n:-1:1
46
47
       t = 0.
48
        for j=k+1:n
          t = t + A(k, j) * b(j)
49
50
        end
51
       b(k) = (b(k) - t) / A(k,k)
52
     end
53
     x = b
54
   endfunction
```

Before actually measuring the performance, we must test for correctness, since there is no point at making faster a function which has bugs. That seems obvious, but it is often neglected in practice. In the following session, we generate a 10×10 matrix **A** with real entries randomly uniform in [0, 1). We selected this test case for its very small condition number. We select the expected solution **e** with unity entries and compute the right-hand side **b** from the equation **b=A*e**. Finally, we use our gausspivotalnaive function to compute the solution **x**.

```
n = 10;
A = grand(n,n,"def");
e = ones(n,1);
b = A * e;
x = gausspivotalnaive ( A , b );
```

Then, we check that the condition number of the matrix is not too large. In the following session, we compute the 2-norm condition number of the matrix A and check that it is roughly equal to 10^1 , which is rather small (only the order of magnitude matters). We also check that the relative error on x has the same magnitude as $\text{%eps} \approx 10^{-16}$, which is excellent.

```
-->log10(cond(A))
ans =
1.3898417
-->norm(e-x)/norm(e)
ans =
6.455D-16
```

A more complete testing would be required, but we now assume that we can trust our function.

In the following session, we call the add_profiling function in order to profile the gausspivotalnaive function.

```
-->add_profiling("gausspivotalnaive")
Warning : redefining function: gausspivotalnaive
Use funcprot(0) to avoid this message
```

The goal of the add_profiling function is to redefine the function to be profiled, so that Scilab can store the number of times that each line is executed. As indicated by the warning, this actually modifies the profiled function, making it slower. This does not matter, provided that we do not take too seriously the execution times in the profile plot: we will focus, instead, only on the number of times that a particular line of the source code has been executed.

In order to measure the performance of our function, we must execute it, as in the following session.

x = gausspivotalnaive (A , b);

During the run, the profiling system has stored profiling data that we can now analyze. In the following script, we call the **plotprofile** function in order to plot the profile of the function.

plotprofile(gausspivotalnaive)

The previous script produces the figure 29. The graphics is made of three histograms. Our function has 54 lines, which are represented by the X axis of each histogram. The Y axis depends on the histogram. On the first histogram, the Y axis represents the number of calls of each corresponding line in the profiled function. This is simply measured by updating a counter each time a line is executed. On the


Figure 29: Profile of the gausspivotalnaive function.

second histogram, the Y axis represents the complexity of each corresponding line in the profiled function. This is a measure of the interpreter effort for one execution of the corresponding line. The Y axis of the third histogram represents the CPU time, in seconds, required by the corresponding line. In practice, the CPU time measure is unreliable when it is not sufficiently large.

The plotprofile function also opens the editor and edits the profiled function. With the mouse, we can left-click on the colored vertical bars of the histogram. In the editor, this immediately moves the cursor to the corresponding line. We first click on the largest bar in the first histogram, which correspond to the lines which are executed more than 250 times. The editor then moves the cursor to the lines #23 and #24 in our function. This allows to immediately let us focus on the following source code lines.

```
for i = k+1:n
  for j = k+1:n
        A(i,j) = A(i,j) - A(i,k) * A(k,j)
```

In the remaining of this section, we are going to make a consistent use of the profiler. By analyzing the various parts of the function which cost the most, we are able to measure the part which have the most impact on the performance. This allows to apply vectorization principles only on the source code lines which matter and let us save a huge amount of time. Once done, we click on the "Exit" menu in the graphics window, which closes the profile plot.

We now analyze the previous nested loops on the variables i and j. We can make the loop on j faster, by using a vectorized statement. Using the colon : operator, we replace the variable j by the k+1:n statement, as in the following script.

We can push the principle further, by vectorizing also the loop on the *i* variable, again using the colon operator. This leads to the following source code.

A(k+1:n,k+1:n) = A(k+1:n,k+1:n) - A(k+1:n,k) * A(k,k+1:n)

The effect of the previous statement is to update the sub-matrix A(k+1:n,k+1:n)in one statement only. The key point is that the previous statement updates many entries of the matrix in just one single Scilab call. In the current situation, the number of updated entries is $(n-k)^2$, which may be large if n is large. All in all, we have replaced roughly n^2 calls to the interpreter processing 1-by-1 matrices, by one single call processing roughly a n-by-n matrix. We notice that the multiplication A(k+1:n,k)*A(k,k+1:n) is the multiplication of a column vector by a row vector, which produces a $(n-k) \times (n-k)$ matrix in just one statement.

Similarly, we can vectorize other statements in the algorithm.

The loop in the following source code is an obvious candidate for vectorization.

```
for i=k+1:n
    A(i,k) = A(i,k)/ A(k,k)
```

The previous algorithm is mathematically equivalent, but much slower than the following vectorized statement.

A(k+1:n,k) = A(k+1:n,k) / A(k,k)

The search for a pivot, in the following part of the algorithm, seems a less obvious candidate for vectorization.

```
mu = k
abspivot = abs(A(mu,k))
for i=k:n
    if (abs(A(i,k))>abspivot) then
        mu = i
        abspivot = abs(A(i,k))
    end
end
```

Essentially, this algorithm is a search for the largest magnitude of the entries in one sub-column of the matrix A. Hence, we can make use of the max function, which returns both the maximum entry of its argument and the index which achieves that maximum value. It remains to combine several vectorized functions to reach our goal, which is nontrivial.

By analyzing how the i index is updated during the loop, we see that the submatrix to be considered is A(k:n,k). The abs function is elementwise, that is, performs its computation element by element. Hence, the statement abs(A(k:n,k))computes the absolute values that we are interested in. We can finally call the max function to perform the search, as in the following script.

```
[abspivot,murel]=max(abs(A(k:n,k)))
mu = murel + k - 1
```

Notice that the max function only performs the search on a sub-part of the whole column of the matrix A. Hence, the murel variable holds the row achieving the maximum, relatively to the block A(k:n,k). This is why we use the statement mu = murel + k - 1, which allows to recover the global row index into mu.

The following loop allows to swap the rows k and mu of the matrix A.

```
for j=k:n
   tmp = A(k,j)
   A(k,j) = A(mu,j)
   A(mu,j) = tmp
end
```

It is now obvious for us to replace the previous script by the following vectorized script.

```
tmp = A(k,k:n)
A(k,k:n) = A(mu,k:n)
A(mu,k:n) = tmp
```

This is already a great improvement over the previous script. Notice that this would create the intermediate vector tmp, which is not necessary if we use the following statements.

```
A([k mu],k:n) = A([mu k],k:n)
```

Indeed, the matrix [mu k] is a valid matrix of indices and can be directly used to specify a sub-part of the matrix A.

All in all, the following gausspivotal function gathers all the previous improvements and uses only vectorized statements.

```
function x = gausspivotal (A, b)
 n = size(A, "r")
 // Initialize permutation
 P=zeros(1,n-1)
 // Perform Gauss transforms
 for k=1:n-1
    // Search pivot
    [abspivot,murel] = max(abs(A(k:n,k)))
    // Shift mu, because max returns with respect to (k:n,k)
   mu = murel + k - 1
    // Swap lines k and mu from columns k to n
   A([k mu],k:n) = A([mu k],k:n)
   P(k) = mu
    // Perform transform for lines k+1 to n
   A(k+1:n,k) = A(k+1:n,k) / A(k,k)
   A(k+1:n,k+1:n) = A(k+1:n,k+1:n) - A(k+1:n,k) * A(k,k+1:n)
 end
 // Perform forward substitution
 for k=1:n-1
    // Swap b(k) and b(P(k))
   mu = P(k)
   b([k mu]) = b([mu k])
    // Substitution
   b(k+1:n) = b(k+1:n) - b(k) * A(k+1:n,k)
```



Figure 30: Profile of the gausspivotal function.

```
end
// Perform backward substitution
for k=n:-1:1
    b(k) = (b(k) - A(k,k+1:n) * b(k+1:n)) / A(k,k)
end
x = b
endfunction
```

Again, we can check that our function works, as in the following session.

```
-->x = gausspivotal ( A , b );
-->norm(e-x)/norm(e)
ans =
6.339D-16
```

We can update the profile by launching the plotprofile function on the improved gausspivotal function. This produces the figure 30.

We see that the profile of the number of calls is flat, which means that there is no line which is called more often than the others. As this is an excellent result, we can stop there our vectorization process.

In the exercise 5.1, we compare the actual performances of these two functions. We show that, on a 100-by-100 matrix (which is modest), the average performance improvement can be greater than 50. In general, using vectorization allows to improve the performances by orders of magnitude.

5.1.4 The benchfun function

In this section, we present a function which may be used for the performance analysis of algorithms. We present the variability of the CPU time required by the computation of the upper triangular Pascal matrix, and present a more reliable way of measuring the performance of an algorithm.

We try to measure the performance of the following pascalup_col function, which computes the upper Pascal matrix.

```
function c = pascalup_col (n)
    c = eye(n,n)
    c(1,:) = ones(1,n)
    for i = 2:(n-1)
        c(2:i,i+1) = c(1:(i-1),i)+c(2:i,i)
    end
endfunction
```

The algorithm presented here is based on a column-by-column access to the matrix, which is especially fast because this is exactly the way that Scilab stores the entries of a matrix of doubles. This topic is presented in more depth in the section 5.3.6.

In the following session, we measure the time required to compute the 1000-by-1000 upper triangular Pascal matrix.

```
-->tic(); pascalup_col(1000); toc()

ans =

0.172011

-->tic(); pascalup_col(1000); toc()

ans =

0.15601
```

As we can see, there is some randomness in the measure of the elapsed time. This randomness may be explained by several facts, but, whatever the reasons, we need a more reliable way of measuring the time required by an algorithm. A common method is to run the algorithm several times, and to average the measure. This method is used in the following script, where we run the pascalup_col function 10 times. Then we display the minimum, mean and maximum times required during the experiment.

```
-->for k = 1 : 10
--> tic();
--> pascalup_col(1000);
--> t(k) = toc();
-->end
-->disp([min(t) mean(t) max(t)])
0.108006 0.1168072 0.16801
```

The previous method can be automated, which leads to the benchfun function, which is presented below. This function runs an algorithm for which we want to get reliable performance measures. The input arguments are the name of the function to be tested, the function fun to run, a list iargs containing the input arguments of fun, the number of output arguments nlhs and the number of iterations to perform kmax. The body of benchfun is based on the execution of the function fun, which is executed kmax times. At each iteration, the performance of fun is measured based on the tic() and toc() functions, which measure the elapsed (wall clock) time

required by the function. The benchfun function returns a kmax-by-1 matrix of double t which contains the various timings recorded during the execution of fun. It also returns the string msg which contains a user-friendly message describing the performance of the algorithm under test.

```
function [t,msg] = benchfun(name,fun,iargs,nlhs,kmax)
 // Compute the intstruction string to be launched
 ni = length ( iargs )
  instr = ""
 // Put the LHS arguments
 instr = instr + "["
 for i = 1 : nlhs
    if (i > 1) then
      instr = instr + ","
    end
    instr = instr + "x" + string(i)
 end
  instr = instr + "]"
 // Put the RHS arguments
  instr = instr + "=fun("
 for i = 1 : ni
    if ( i > 1 ) then
      instr = instr + ","
    end
    instr = instr + "iargs("+string(i)+")"
 end
 instr = instr + ")"
 // Loop over the tests
 for k = 1 : kmax
    tic()
    ierr = execstr ( instr , "errcatch" )
    t(k) = toc()
  end
 msgfmt = "%s: %d iterations, mean=%f, min=%f, max=%f\n"
 msg=msprintf(msgfmt,name,kmax,mean(t),min(t),max(t))
 mprintf("%s\n",msg)
endfunction
```

In the following session, we run the pascalup_col function 10 times in order to produce the 2000-by-2000 upper triangular Pascal matrix.

```
-->benchfun("pascalup_col",pascalup_col,list(2000),1,10);
pascalup_col: 10 iterations, mean=0.43162,
min=0.41059, max=0.47067
```

Although the **benchfun** function could be made more robust and more flexible, it is sufficient for many cases. We will use it often throughout this section.

5.2 Vectorization principles

In this section, we present the principles of vectorization. This programming method is the best way to achieve good performances in Scilab. The first section presents the principles of the interpreter as well as the link between a primitive and a gateway. We compare the performance of **for** loops with vectorized statements. We finally



Figure 31: Scilab is an interpreter.

present an example of performance analysis, where we transform a naive, slow, algorithm into a fast, vectorized, algorithm.

5.2.1 The interpreter

In this section, we briefly present compilation methods so that we can have clear ideas about the consequences of the interpreter for performances issues. We also present the link between the interpreter and the gateways, which are the functions which connect Scilab to the underlying libraries.

In order to understand what exactly happens when a statement as y=f(x) is executed, we must take a look at the structure of a compiler. Indeed, this will allow us to see the link between the statements at the interpreter level, and the execution of a possibly compiled source code at the library level.

A compiler[5] is a program that can read a program in one language and translate it into an equivalent program. If the target program is an executable written in machine language, it can be called by the user to process inputs and produce outputs. Typical compilers of this type are the Fortran and C/C++ compilers (among others).

Contrary to the previous scheme, Scilab is not a compiled language: it is an *interpreted* language. Instead of producing a machine-language program, Scilab both uses the source program and the input data and directly produces the output. The figure 31 presents an interpreter. Typical interpreters of this type are Scilab, Matlab and Python (among others).

The machine-language program produced by a compiler is generally faster that an interpreter at mapping inputs to outputs. An interpreter, however, can usually give better diagnostics than a compiler, because it executes the source program statement by statement. Moreover, Scilab uses highly optimized numerical libraries which, in some situations, may be faster than a naive implementation based on a compiled language. This topic will be reviewed in the section 5.4.1, where we present the numerical linear algebra libraries used in Scilab.

A compiler is typically structured as a sequence of operations. The first operation involves reading the characters of the source code and performing the lexical analysis. This step consists in grouping the characters into meaningful sequences called lexemes. The tokens are then passed to the syntax analyzer (or parser), which uses the tokens and creates a tree representation of the source, which is associated to the grammatical structure of the source. In the semantic analysis, the compiler uses the syntax tree and check the source for consistency with the language definition.



Figure 32: Scilab connects the interpreter to a collection of libraries, via gateways.

This step involves checking the types of the operands of each operator.

Several intermediate operations may then be involved, leading to the final call to a library routine. In Scilab, this final step is implemented in *gateways*. Indeed, each Scilab function or operator is uniquely associated with a computational routine implemented in a compiled language, generally in C or Fortran. The gateway reads the input arguments, check their types and consistency, performs the required computations, and then push back the output arguments into the interpreter. This structure is presented in figure 32.

5.2.2 Loops vs vectorization

In this section, we present a simple example of vectorization and analyze why for loops are good candidates for vectorization.

Let us consider the following matrix of doubles, made of n=5 entries.

x=[1 2 3 4 5];

Assume that we want to compute the sum of the 5 numbers stored in the variable x. We are going two compare two scripts which produce exactly the same correct result but which have a very different performance.

In the first script, we call the sum function and sets the variable y.

y = sum(x);

This requires only one call to the interpreter and involves only one call to one particular gateway. Inside the gateway, the numerical library performs a loop of size n=5 in a compiled and optimized source code.

In the second script, we use a for loop.

```
y = 0;
n = size(x,"*");
for i = 1 : n
  y = y + x(i);
end
```

The previous script involves the interpreter at least 2+n times, where n is the size of x. With n=5, this is 7 calls to the interpreter, but this grows linearly with the size of x. Inside the gateway, the numerical library performs one single sum each time it is called. When the number of loops is large, the extra cost of using a for statement instead of a vectorized statement is very expensive.

This is why *vectorized* statements are much faster than loops. In general, we should make a minimum number of calls to the interpreter and let it work on sufficiently large data sets.

We now emphasize the conclusion of the previous discussion.

- If Scilab has a built-in function, we should consider using it first before creating our own function. From a performance point of view, the built-in functions are, most of the time, well designed, that is, they make the maximum possible use of vectorized statements. Moreover, most of the time, they take into account for floating point number issues and accuracy problems as well, so that we can get both a fast computation and a good accuracy.
- If we must create our own algorithm, we should use vectorized statements as much as possible. Indeed, the speedup for vectorized statements is typically 10-100, but can be as large as 1000 sometimes. Several examples of performance improvement are presented in the remaining of this document.

But there is more, because Scilab may call optimized libraries, which make an efficient use of the processor when it must perform floating point computations. Scilab scripts can be as efficient as a compiled source code, but they can even be faster than a naive compiled source code. For example, the linear algebra libraries used in Scilab take into account the size of the cache of the processor and, moreover, can perform multi-threaded operations. This is central in Scilab, which is mainly a matrix language. The section 5.4 presents the optimized linear algebra libraries used by Scilab.

Not all programs can make profit from linear algebra operations and this is why we first focus on vectorization methods which can be applied in most situations. Indeed, our experience is that vectorization requires some practice. The next section presents a practical example of performance analysis based on vectorization.

5.2.3 An example of performance analysis

In this section, we compare the performances of a naive and a vectorized algorithm. We check that the speedup, in this particular case, can be as large as 500.

Assume that we are given a column vector and that we want to compute the mean of this vector. We have two possibilities to perform this task:

- use the mean function provided by Scilab,
- create our own algorithm.

The mean function can manage various shape of input matrices (row vectors, column vectors or general matrices) and various norms (1-norm, 2-norm, infinity-norm and Frobenius norm). This is why, in general, we should not redefine our own functions when Scilab already provides an implementation. Moreover, the performance of the mean function is excellent, as we are going to see. But, for the purpose of illustrating performance issues, we will develop our own algorithm.

The following mynaivemean function computes the mean of the input vector \mathbf{v} . It performs a loop on the data with a straightforward cumulated sum.

```
function y = mynaivemean ( v )
y = 0
n = length(v)
for i = 1:n
y = y + v(i)
```

```
end
y = y / n
endfunction
```

Before optimizing our algorithm, we check that it performs its task correctly: indeed, it would be a loss of time to make faster an algorithm which contains a bug... The algorithm seems to be correct, at least from the standpoint of the following session.

```
-->y = mynaivemean(1:9)
y =
5.
```

In order to test our algorithm, we generate large input vectors \mathbf{v} . For this purpose, we use the **rand** function, because we know that the performances of our algorithm do not depend on the actual values of the input vector. We emphasize this detail, because practical algorithms may exhibit faster or slower performances depending on their input parameters. This is not the case here, so that our analysis is simplified.

We generate vectors with size $n = 10^j$ with j = 1, 2, ..., 6. To generate the integers n, we use the logspace function, which returns values based on the base-10 logarithm function. The following script performs a loop from $n = 10^1$ up to $n = 10^6$ and measures the time required by our naive algorithm.

```
n_data = logspace(1,6,6);
for i = 1:6
  n = n_data(i);
  v = rand(n,1);
  tic();
  ymean = mynaivemean ( v );
  t = toc();
  mprintf("i=%d, n=%d, mynaivemean=%f, t=%f\n",i,n,ymean,t);
end
```

The previous script produces the following output.

```
i=1, n=10, mynaivemean=0.503694, t=0.000000
i=2, n=100, mynaivemean=0.476163, t=0.000000
i=3, n=1000, mynaivemean=0.505503, t=0.010014
i=4, n=10000, mynaivemean=0.500807, t=0.090130
i=5, n=100000, mynaivemean=0.499590, t=0.570821
i=6, n=1000000, mynaivemean=0.500216, t=5.257560
```

We see that the time grows very fast and is quite large for $n = 10^6$.

It is, in fact, possible to design a much faster algorithm. In the previous script, we replace the mynaivemean function by the built-in mean function. This produces the following output.

i=1, n=10, mean=0.493584, t=0.000000 i=2, n=100, mean=0.517783, t=0.000000 i=3, n=1000, mean=0.484507, t=0.000000 i=4, n=10000, mean=0.501106, t=0.0000000 i=5, n=100000, mean=0.502038, t=0.0000000 i=6, n=1000000, mean=0.499698, t=0.010014

As we see, the mean function is much faster than our mynaivemean function. In this case, we observe that, for n=1000000, the speed ratio is approximately 5.25/0.01 = 525.

In order to improve our naive algorithm, we can use the sum function, which returns the sum of the elements of its input argument. The following myfastmean function performs the computation based on the sum function.

```
function y = myfastmean ( v )
y = sum(v);
n = length(v);
y = y / n;
endfunction
```

The following session presents the performance of our improved function.

```
i=1, n=10, mean=0.419344, t=0.000000
i=2, n=100, mean=0.508345, t=0.000000
i=3, n=1000, mean=0.501551, t=0.000000
i=4, n=10000, mean=0.501601, t=0.000000
i=5, n=100000, mean=0.499188, t=0.000000
i=6, n=1000000, mean=0.499992, t=0.010014
```

The performance of our fast function is now approximately the same as the performance of the built-in mean function. In fact, the mean function is a macro, which performs mainly the same algorithm as our function, that is, it uses the sum function.

Why does the sum function is much faster than an algorithm based on a for loop? The reason is that the sum function is a *primitive*, based on a compiled source code. This can be easily checked by getting the type of this function, as in the following session.

```
-->typeof(sum)
ans =
fptr
```

Hence, the loop is performed by Scilab inside the sum function by a compiled source code. This is indeed much faster than a for loop performed by the interpreter and explains the performance difference between our two previous implementations.

5.3 Optimization tricks

In this section, we present common tricks to get faster algorithms and emphasize the use of vectorized statements. We present the danger of using matrices which size grow dynamically during the algorithm. We present general purpose functions, such as sorting and searching vectorized functions, which can be combined to create efficient algorithms. We also discuss the orientation of matrices and give an example of performance improvement based on this trick.

5.3.1 The danger of dymanic matrices

In this section, we analyze the performance issues which are caused by the use of matrices which size dynamically grow. We show a numerical experiment where predefining a matrix with a known size is much faster than letting it grow dynamically.

Let us consider a situation where we must compute a matrix but do not know the final size of this matrix in advance. This happens when, for example, the matrix is read from a data file or interactively input by the user of the program. In this case, we can use the fact that the size of a matrix can dynamically grow. But this has a cost, which can be sensitive in cases where the number of dynamic updates is large. This cost is associated with the internal behavior of the update of the size of the matrix. Indeed, increasing the size of the matrix implies that the interpreter has to update its internal data structures. Hence, the entire matrix content must be copied, which may require a significant amount of time. Furthermore, while the loop is proceeding, the matrix size is actually growing, which, in turn, imply slower and slower matrix copies.

In order to analyze this performance issue, we compare three functions with different implementations, but with the same output. The goal is to produce a matrix $A=[1,2,3,\ldots,n]$ where n is a large integer. For pure efficiency purposes, we could use the colon ":" operator and the statement A=(1:n). In order to illustrate the particular issue which is the topic of this section, we do not use the colon operator and, instead, we use the following implementations.

The following growingmat1 function takes n as an input argument and performs a dynamic computation of the matrix A. First, the algorithm initialize the matrix A to the empty matrix. Then, it makes use of the \$+1 syntax to dynamically increase the size of the matrix and store the new entry i.

```
function growingmat1 ( n )
  A = []
  for i = 1 : n
      A($+1) = i
   end
endfunction
```

We do not return the matrix A as an output argument, since this is not necessary for our demonstration purpose.

The following growingmat2 function also performs a dynamic computation of the matrix A, but uses a different syntax. This time, the matrix is dynamically updated with the [A,i] syntax, which creates a new row matrix with the entry i added at the end.

```
function growingmat2 ( n )
  A = []
  for i = 1 : n
      A = [A,i]
  end
endfunction
```

This creates a row matrix. A variant of this algorithm would be to create a column matrix with the [A;i] syntax. This variant would not fundamentally change the behavior of the algorithm.

The following growingmat3 function is completely different from the previous ones. Instead of initializing the matrix to the empty matrix, it pre-creates a column matrix of zeros. Then the entries are filled one after the other.

```
function growingmat3 ( n )
A = zeros(n,1)
for i = 1 : n
A(i) = i
end
endfunction
```

| 11 <i>(1)</i> | 14 <i>(4)</i> | 17 (7) | 20 (10) |
|---------------|---------------|---------------|-----------------|
| 12 (2) | 15 <i>(5)</i> | 18 <i>(8)</i> | 21 <i>(11</i>) |
| 13 <i>(3)</i> | 16 (6) | 19 <i>(9)</i> | 22 (12) |

Figure 33: Linear indexing of a 3-by-4 matrix. For example, the value "21" is at subscripts (i, j) = (2, 4) and has the linear index k = 11. The linear indices are behaving as if the entries were stored column-by-column.

In the following script, we compare the performances of the three previous functions with n=20000 and repeat the timing 10 times to get a more reliable estimate of the performance. We make use of the benchfun function, which has been presented in the section 5.1.4.

```
stacksize("max")
benchfun("growingmat1",growingmat1,list(20000),0,10);
benchfun("growingmat2",growingmat2,list(20000),0,10);
benchfun("growingmat3",growingmat3,list(20000),0,10);
```

In the following session, we get the timing of the three algorithms.

-->exec bench_dynamicmatrix.sce; growingmat1: 10 iterations, mean=1.568255 growingmat2: 10 iterations, mean=0.901296 growingmat3: 10 iterations, mean=0.023033

We see that pre-creating the matrix A with its known size n is much faster. The ratio from the slowest to the fastest is larger than 50.

5.3.2 Linear matrix indexing

In this section, we present how a regular matrix can be accessed with one single *linear* index k instead of a pair of subscripts (i,j). This feature works both for matrices and hypermatrices, although we focus in this section only on regular matrices.

If the matrix A has m rows, the linear index k corresponding to the subscripts (i, j) is k = i + (j-1)m. In this case, the statements A(i,j) and A(k) are equivalent. We can think of this feature as if the entries were stored column-by-column (see section 5.3.6 for more details on this topic).

In the following example, we create a 3-by-4 matrix of doubles. Then we set the entry at linear index k=11 to zero. This entry corresponds to the entry at subscripts (i, j) = (2, 4).

```
-->A = matrix(10+(1:12),3,4)
 Α
   =
           14.
                           20.
                   17.
    11.
    12.
            15.
                   18.
                           21.
    13.
            16.
                   19.
                           22.
-->A(11)=0
 Δ =
           14.
                  17.
    11.
                           20.
```

| 12. | 15. | 18. | 0. |
|-----|-----|-----|-----|
| 13. | 16. | 19. | 22. |

The previous example is presented in the figure 33.

The functions sub2ind and ind2sub convert from subscripts to linear indices and conversely. These functions are presented in the figure 34. We emphasize that these functions work both for matrices and hypermatrices, but we focus in the remaining of this section on regular matrices.

For example, we can convert a linear index into a pair of subscripts with the statement [i,j]=ind2sub(dims,k), where dims is a 1-by-2 matrix containing the dimensions of the matrix. The following session presents an example of the ind2sub function. We consider a 3-by-4 matrix (as in the figure 33), and check that the linear index k=11 corresponds to the subscripts i=2 and j=4.

```
-->[i,j]=ind2sub([3,4],11)
j =
4.
i =
2.
```

We now give a simple example of how the sub2ind function can be used in a practical case where it can be used in a vectorized computation. We consider a 3-by-4 matrix and want to set the first diagonal above the main diagonal to zero. In our particular case, this corresponds to the values 14, 18 and 22, which are located at the rows 1, 2 and 3 and the columns 2, 3 and 4. Of course, we could use a for loop, but this is not a vectorized computation. We cannot neither use directly the row and column subscripts, because this would set all the corresponding 3-by-3 sub-matrix to zero, as show in the following session.

```
-->A = matrix(10+(1:12),3,4)
 А
    =
     11.
              14.
                       17.
                                20.
     12.
              15.
                       18.
                                21.
     13.
              16.
                       19.
                                22.
-->A([1 \ 2 \ 3], [2 \ 3 \ 4]) = 0
 Α
              0.
                      0.
                              0.
     11.
                      0.
     12.
              Ο.
                              0.
     13.
              0.
                      0.
                              0.
```

Therefore, we use the sub2ind function to convert these subscripts into a linear index vector \mathbf{k} and then set these entries to zero.

-->A = matrix(10+(1:12),3,4)А 11. 14. 17. 20. 12. 15. 18. 21. 13. 16. 19. 22. -->k = sub2ind([3,4],[1 2 3],[2 3 4]) = k 4. 8. 12. -->A(k) = 0А = 11. 0. 17. 20. 12. 15. 0. 21.

| ind2sub | convert linear index k into subscripts i1,i2, |
|---------|---|
| sub2ind | convert subscripts $i1, i2, \ldots$ into linear index k |

Figure 34: Scilab function to convert from subscripts to linear indices.

13. 16. 19. 0.

The previous example can be used as a basis for algorithms which access to diagonals of matrices, such as in the **spdiags** function for example.

We will review an example of linear indexing in the section 5.3.5, where we will use the find function in order to access to particular entries of a matrix.

5.3.3 Boolean matrix access

In this section, we present how a regular matrix can be accessed with indices defined by a boolean matrix.

Operators which create boolean matrices, such as == or >, can be used to access to matrices with vectorized statements. Assume that A is a m-by-n matrix of doubles and that B is a m-by-n matrix of booleans. Therefore, the statement A(B) accesses to all entries of A where B is true. This is shown in the following example, where we compute all entries of A which are equal to 1, and set them to zero.

```
-->A = [1 \ 2 \ 1 \ 2; \ 2 \ 1 \ 2 \ 1; \ 1 \ 3 \ 1 \ 3]
 А
    =
             2.
                             2.
     1.
                     1.
     2.
             1.
                     2.
                             1.
             З.
                             З.
     1.
                     1.
-->B = (A==1)
 В
  TFTF
  FTFT
  TFTF
-->A(B)
          = 0
 Α
    =
     Ο.
             2.
                     0.
                             2.
     2.
             Ο.
                     2.
                             0.
     0.
             3.
                     0.
                             З.
```

The previous sequence of operations can be simplified into one single statement, with the statement A(A==1)=0. This is shown in the following example.

 $-->A = [1 \ 2 \ 1 \ 2; \ 2 \ 1 \ 2 \ 1; \ 1 \ 3 \ 1 \ 3]$ А = 1. 2. 1. 2. 2. 1. 2. 1. 1. З. 1. 3. -->A(A==1)=0А = 0. 2. 0. 2. 2. 0. 2. 0. 0. З. 0. З.

In the situation where the search for particular entries of A is too complex to be managed with this method, we can use the find function, as presented in the 5.3.5 section.

5.3.4 Repeating rows or columns of a vector

In this section, we present how to create copies of a row or column vector with vectorized statements.

An interesting feature of matrices is that the matrix extraction can be used to repeat the entries of one matrix. Indeed, if **A** is a matrix, we can repeatedly extract the first row, which creates copies of this row.

```
-->A = 1:3
 A =
     1.
            2.
                    3.
-->B = A([1 \ 1 \ 1],:)
 В
    =
             2.
                     З.
     1.
                    З.
     1.
             2.
                     З.
     1.
             2.
```

The interesting fact here is that the creation of B did not require a loop: this is vectorized.

This trick is used in the **repmat** function, which may be integrated into Scilab 5.4.

This topic is explored in the exercise 5.2.

5.3.5 Combining vectorized functions

In order to get good performances in Scilab, we can use vectorized statements. In this section, we present how to combine vectorized functions.

Vectorized scripts make a more consistent use of the optimized libraries provided by Scilab. The main rule is to avoid **for** loops. The second rule is to use matrix operations as often as possible, because they use the highly optimized linear algebra libraries which are provided with Scilab. But there are situations where it is not obvious to perform a vectorized computation. In these cases, we may *combine* vectorized functions: the purpose of this section is to present relatively advanced vectorized functions and to suggest how to combine them.

The table 35 present the most common vectorized functions that we can use to get good performances.

The algebra operators +,-,*,/ are presented here because they offer excellent performances in Scilab. In order to use these operators, we must generate matrices on which we can perform linear algebra. We may use the **zeros**, **ones** and other matrix-oriented functions, but a useful operator to generate matrices is the Kronecker product. In the following session, we multiply two 3-by-2 matrices with the Kronecker product, which creates a 4-by-9 matrix.

-->[1 2 3;4 5 6] .*. [7 8 9;1 2 3] ans = 9. 27. 7. 8. 14. 16. 18. 21. 24. 1. 2. З. 2. 4. 6. 3. 6. 9.

| +,-,*,/,\ | algebra operators |
|-----------|--|
| .*. | the Kronecker product |
| min,max | the minimum, the maximum |
| sum | the sum of the entries of a matrix |
| cumsum | the cumulated sum of the entries of a matrix |
| prod | the product of the entries of a matrix |
| cumprod | the cumulated product of the entries of a matrix |
| find | search for true indices in a boolean matrix |
| gsort | sorts the entries of a matrix |
| matrix | generate a matrix from the entries of a matrix |

Figure 35: Scilab function often used during a vectorization.

| 28. | 32. | 36. | 35. | 40. | 45. | 42. | 48. | 54 |
|-----|-----|-----|-----|-----|-----|-----|-----|----|
| 4. | 8. | 12. | 5. | 10. | 15. | 6. | 12. | 18 |

The Kronecker product can be, for example, used in combinatorial algorithms where we want to produce combinations of values from a given set. This topic is explored in the exercise 5.2.

We now review the functions find, gsort, min and max, which have special input and output arguments which make them very worthwhile when we are looking for fast scripts.

Using the find function is a common optimization method, which may replace search algorithms. If B is a m-by-n matrix of booleans, then the statement k=find(B) returns the linear indices k for which B is true.

If the search fails, that is, if no entry corresponds to the condition, then the find function returns the empty matrix. We may think that it is then necessary to explicitly manage this particular case. But the statement A([])=0 does nothing, since the empty matrix [] corresponds to an empty set of indices. Hence, on output of the find function, most of the time, there is no need to process the failure case with a separate if statement.

In the following example, we set to zero all the entries of the matrix **A** which are equal to 1.

```
-->A = [1 \ 2 \ 1 \ 2; \ 2 \ 1 \ 2 \ 1; \ 1 \ 3 \ 1 \ 3]
 А
   =
             2.
                            2.
     1.
                    1.
                    2.
     2.
             1.
                            1.
                    1.
                            З.
     1.
            З.
-->B = (A == 1)
 В
    =
  TFTF
  FTFT
  TFTF
-->k = find(B)
 k
    =
                            7.
     1.
             З.
                    5.
                                    9.
                                            11.
-->A(k) =
            0
 A =
     0.
            2.
                    0.
                            2.
```

2.0.2.0.0.3.0.3.

The previous session can be shortened by combining the functions into one single statement, as shown below.

```
-->A = [1 \ 2 \ 1 \ 2; \ 2 \ 1 \ 2 \ 1; \ 1 \ 3 \ 1 \ 3]
 Α
    =
               2.
                       1.
                                2.
     1.
     2.
               1.
                       2.
                                1.
     1.
               З.
                       1.
                                 З.
-->A(find(A==1)) = 0
 А
    =
               2.
     0.
                       Ο.
                                2.
     2.
               Ο.
                       2.
                                0.
     0.
              З.
                       Ο.
                                З.
```

Of course, as presented in the section 5.3.3, we could use the shorter statement A(A==1)=0.

Then we experiment what happens when the value which is searched for is not found.

```
-->A = [1 \ 2 \ 1 \ 2; \ 2 \ 1 \ 2 \ 1; \ 1 \ 3 \ 1 \ 3]
 A =
              2.
                               2.
     1.
                       1.
                       2.
     2.
              1.
                               1.
     1.
              З.
                       1.
                               З.
-->k = find(A==999)
 k
    =
       []
-->A(k)=0
 Α
   =
              2.
                               2.
     1.
                       1.
     2.
              1.
                       2.
                               1.
     1.
              З.
                       1.
                               3.
```

There are cases where the linear indices k must be processed before using them to set a matrix. Moreover, there are options of the **find** function which can be used to create more complex vectorized operations, as we are going to see.

If not all the entries matching a condition are to be used, but only a limited number of these, we may use the second input argument of the find function. For example, the statement k=find(A(:,2)>0,1) returns the first index of the second column of A which is positive.

The min and max functions can be used with two output arguments. Indeed, the statement [m,k]=min(A) returns the minimum value m and the index k in A which achieves the minimum value. In other words, the index k is such that A(k)==m. For example, the statement [m,k]=min(abs(A)) combines min and the elementwise abs functions. It returns the index k where the matrix A has the minimum absolute value.

The gsort function has also a second argument which can be useful to get good performances. Indeed, the statement [B,k]=gsort(A) returns the sorted matrix B and the rank k. In other words, the matrix B is such that B(:)==A(k). In practice, we may therefore apply the same re-ordering to a second matrix C, with the statement C=C(k).

5.3.6 Column-by-column access is faster

We may get faster performances by accessing to the elements of a matrix of doubles column-by-column, rather than row-by-row. In this section, we present an example of performance improvement by a change of orientation of the algorithm and present an explanation for this behavior.

The topic that we are going to analyze was discussed in the bug report #7670 [13]. The problem is to compute the Pascal matrix, which can be derived from the binomial formula. The number of distinct subsets with j elements which can be chosen from a set A with n elements is the binomial coefficient and is denoted by $\binom{n}{i}$. It is defined by

$$\binom{n}{j} = \frac{n.(n-1)\dots(n-j+1)}{1.2\dots j}.$$
 (1)

For integers n > 0 and 0 < j < n, the binomial coefficients satisfy

$$\binom{n}{j} = \binom{n-1}{j} + \binom{n-1}{j-1}.$$
(2)

The following two scripts allow to compute the Pascal matrix. In the first implementation, we compute the Pascal lower triangular matrix, and perform the algorithm row by row.

```
// Pascal lower triangular: Row by row version
function c = pascallow_row ( n )
    c = eye(n,n)
    c(:,1) = ones(n,1)
    for i = 2:(n-1)
        c(i+1,2:i) = c(i,1:(i-1))+c(i,2:i)
    end
endfunction
```

The previous implementation was suggested to me by Calixte Denizet in the discussion associated with a bug report, but a similar idea was presented by Samuel Gougeon in the same thread. In the second implementation, we compute the Pascal upper triangular matrix, and perform the algorithm column by column.

```
// Pascal upper triangular: Column by column version
function c = pascalup_col (n)
    c = eye(n,n)
    c(1,:) = ones(1,n)
    for i = 2:(n-1)
        c(2:i,i+1) = c(1:(i-1),i)+c(2:i,i)
    end
endfunction
```

The following session shows that both implementations are mathematically correct.

-->pascalup_col (5) ans = 1. 1. 1. 1. 1. 1.



Figure 36: Comparing algorithm by column and by row for the Pascal matrix.

| 0. | 1. | 2. | З. | 4. |
|--------|--------|-------|----|----|
| 0. | 0. | 1. | З. | 6. |
| 0. | Ο. | 0. | 1. | 4. |
| 0. | Ο. | 0. | 0. | 1. |
| >pasca | llow_r | ow (5 |) | |
| ans = | | | | |
| 1. | Ο. | 0. | 0. | 0. |
| 1. | 1. | 0. | 0. | 0. |
| 1. | 2. | 1. | 0. | 0. |
| 1. | З. | 3. | 1. | 0. |
| 1. | 4 | 6 | 4 | 1. |

But there is a significant performance difference between these two algorithms. The figure 36 presents the performances of these two algorithms for various matrix sizes. The figure presents the ratio between the by-column and the by-row implementation.

As we can see, the by-column implementation is, in general, 10% faster than the by-row implementation. Except in rare experiments, this fact is true for all matrix sizes.

The reason behind this is the internal data structure for matrices of doubles, which is Fortran-oriented. Therefore, the entries of a matrix are stored column-bycolumn, as presented in the figure 37. Hence, if we order the operations so that they are performed by-column, we have a fast access to the entries, because of the data locality. This allows to make the best possible use of the BLAS routine DCOPY which is used to create the intermediate vectors used in the algorithm. Indeed, the expression c(2:i,i) creates an intermediate column vector, while the expression c(i,2:i) creates an intermediate row vector. This requires to create temporary vectors, which must be filled with data copied from the matrix c. When the column



Figure 37: The storage of a m-by-n matrix of doubles in Scilab. Two consecutive entries in the same column of the matrix are stored consecutively in the memory. Two consecutive entries in the same row of the matrix are separated by m addresses in memory.

vector c(2:i,i) is created, all the entries are next to the other. Instead, the row vector c(i,2:i) requires to skip all the entries which are between two consecutive elements in the source matrix c. This is why extracting a row vector from a matrix requires more time than extracting a column vector made of consecutive entries: it requires less memory movements and makes a better use of the various levels of cache. Therefore, if a given algorithm can be changed so that it can access or update a matrix columnwise, this orientation should be preferred.

5.4 Optimized linear algebra libraries

In this section, we present the linear algebra libraries which are used in Scilab. In the next section, we present BLAS and LAPACK, which are the building block of linear algebra in Scilab. We also present the ATLAS and the Intel MKL numerical libraries. We present the method to install these optimized linear algebra libraries on the Windows and Linux operating systems. We present the effect of the change of library on the performance of the general dense matrix-matrix product.

5.4.1 BLAS, LAPACK, ATLAS and the MKL

In this section, we briefly present the BLAS, LAPACK, ATLAS and MKL libraries, in order to have a global overview on these libraries. These libraries are used by Scilab in order to provide maximum performances on each system.

The BLAS (Basic Linear Algebra Subprograms) library [2] is a collection of Fortran routines providing low level operations such as vector additions, dot products and matrix - matrix multiplications. The BLAS library is divided in three levels.

- The Level 1 BLAS perform scalar, vector and vector-vector operations.
- The Level 2 BLAS perform matrix-vector operations.
- The Level 3 BLAS perform matrix-matrix operations.



Figure 38: Memory levels in a computer system.

The LAPACK (Linear Algebra Package) library [3] is a collection of Fortran routines for solving high level linear algebra problems. This includes solving systems of simultaneous linear equations, least-square solutions of linear systems, eigenvalue problems and singular value problems.

The ATLAS (Automatically Tuned Linear Algebra Software) library [1] provides optimized linear algebra BLAS routines and a small subset of LAPACK optimized routines. The library focuses on applying empirical techniques in order to provide portable performance. The ATLAS library provides in many cases sensible performance improvements over a "basic" BLAS/LAPACK system.

The Intel Math Kernel Library [24] is a library of highly optimized, extensively threaded math routines for science, engineering, and financial applications that require maximum performance. Under Windows, Scilab currently uses only the linear algebra routines of the MKL, in replacement of BLAS.

On Windows, we should select the Intel MKL for installation (this is the default in Scilab v5). On Linux systems, optimized version of the ATLAS library should be used if performance matters. This topic is reviewed in the sections 5.4.3 for Windows and 5.4.4 for Linux.

The ATLAS and Intel MKL libraries use various low-level optimization methods in order to get floating point performances. This topic is reviewed in the next section.

5.4.2 Low-level optimization methods

In this section, we briefly present low-level optimization methods used in numerical linear algebra libraries such as ATLAS and the Intel MKL. We present the various levels of memory in a computer. We review the algorithms used in ATLAS to improve the use of the various levels of cache. We briefly present the instruction sets provided by processors and used by optimized linear algebra libraries.

The main reason for the performance improvement of vectorized functions is that Scilab is able to perform the computation on a *collection* of data, instead of processing each computation separately. Indeed, the performance of a computation is largely driven by the fact that the CPU can access to the data which is available in various types of memories. From the fastest to the slowest, the memories used in modern computers are the various levels of cache (built in the processor itself) and the Random Access Memory (RAM) (we do not consider here the hard drive as a memory unit, although it may be involved in the management of the virtual memory). The figure 38 presents the various levels of memories typically used by a computer.

Consider now the following Scilab script, where we perform a matrix-matrix multiplication.

A = rand(10,10); B = rand(10,10); C = A * B;

In the corresponding gateway, Scilab uses the BLAS routine *DGEMM* to perform the required computation. If the user has chosen the ATLAS or MKL libraries at installation time, the computation is performed in an optimized way.

As the source code of the Intel MKL is not open, we cannot know what are the exact methods used in this library. Instead, ATLAS is open-source and we can analyze it in detail.

The ATLAS approach [52] consists in isolating machine-specific features of the routines, all of which deal with performing an optimized on-chip, cache contained (i.e. in Level 1 cache) matrix multiply. The on-chip multiply is automatically created by a code generator which uses timings to determine the correct blocking and loop unrolling factors to perform an optimized on-chip multiply.

The matrix multiply is decomposed in two parts:

1. the higher level, general size, off-chip multiply, which is platform-independent,

2. the lower level, fixed size, on-chip multiply, which is machine-specific.

The higher level algorithm is based on a blocked matrix multiply. Assume that A is a m-by-k matrix and B is a k-by-n matrix. We assume that the integer NB, the size of the blocks, divides both m,n and k. The blocked matrix multiply algorithm could be written in the Scilab language as following.

In the ATLAS implementation, the statement C(I, J) = C(I, J) + A(I,K)*B(K, J) is performed by the on-chip multiply function.

The main parameter of the off-chip algorithm is NB, but other factors are taken into account. Indeed, ATLAS may use other implementations of this algorithm, based, for example, on other orderings of the three nested loops. In all these implementations, the k-loop is always the innermost loop. But the outer loop may be on m (the rows of A) or on n (over the columns of B). In order to find the optimal value of the parameters (including NB, the order of the loops, and other parameters), ATLAS performs an automated heuristic search, via timings. For each set of the explored parameters, ATLAS uses a code generator and measures the performances with these settings. In the end, the best implementation is kept, and the associated source code is generated, then compiled.

The on-chip multiply is sensitive to several factors including the cache reuse, the instruction cache overflow, the order of floating point instructions, the loop overhead, the exposure of possible parallelism and the cache misses. Moreover, the on-chip multiply may use special instructions available on the processor, such as Single Instruction Multiple Data (SIMD) instructions. These instructions allow to perform the same operation on multiple data simultaneously and exploit data parallelism. There are several SIMD instructions sets, including

- MMX, designed by Intel in 1996,
- 3DNow!, designed by AMD in 1998,
- Streaming SIMD Extension (SSE), designed by Intel in 1999,
- SSE2, designed by Intel in 2001,
- SSE3, designed by Intel in 2004,
- Advanced Vector Extensions (AVX), a future extension proposed by Intel in 2008.

For example, the MULPD instruction can be used to perform an SIMD multiply of two packed double-precision floating-point values. A similar instruction is ADDPD, which performs an SIMD addition of two packed double-precision floating point values.

In practice, generating the binary for the ATLAS library may require several hours of CPU time before the optimized settings are identified automatically by the ATLAS system.

These low-level methods are not the primary concern of Scilab users. Still, it allows to understand why specific libraries are shipped with Scilab and are different for a Pentium III, a Pentium 4 or a Dual or Quad Core. More importantly, it allows to know the level of optimization that we can get from floating point computations and especially linear algebra, which is the primary target of the Scilab language.

5.4.3 Installing optimized linear algebra libraries for Scilab on Windows

In this section, we present the method to install optimized linear algebra libraries for Scilab on Windows.

On Windows, we usually install Scilab after downloading it from http://www.scilab.org (but some users have Scilab pre-installed on their machines). When we download the Scilab installer for Windows and launch it, we can choose among three possibilities :

- Full Installation (Default), where all the modules are installed,
- Installation, where some modules are disabled,
- Custom Installation, where the user can select the modules to enable or disable.

By default, Scilab is installed with the Intel MKL, but this choice can be customized by the user in the installation dialog which is presented in the figure 39.

In the "CPU Optimization for Scilab" section of the installation dialog, we can choose between the following three options.

- Download Intel Math Kernel Library for Scilab. This installation is the default. This library contains optimized BLAS libraries and a subset of LAPACK, provided by Intel. This library is not the complete Intel MKL, but only the subset of functions which are used by Scilab for BLAS and LAPACK.
- Atlas library for Windows. This library has been compiled by the Scilab Consortium to provide optimum performance on a wide range of modern computers.
- BLAS, LAPACK Reference library for Windows. This library is the reference implementations provided on http://www.netlib.org/blas and http://www.netlib.org/lapack.

These options have a direct impact on two dynamic libraries which are stored in the **bin** directory of Scilab:

- bin\blasplus.dll : the dynamic library for BLAS,
- bin\lapack.dll : the dynamic library for LAPACK.

In practice, we can have the same version of Scilab installed with different linear algebra libraries. Indeed, it is straightforward to customize the installation directory, and then to customize the particular library that we want to use. Therefore, we can install Scilab in three different directories, each with one particular library, so that we can compare their behavior and performances. This method is used in the section 5.4.5, where we present the difference of performances between these three libraries on Windows on a classical benchmark.

5.4.4 Installing optimized linear algebra libraries for Scilab on Linux

In this section, we present the method to install optimized linear algebra libraries for Scilab on the Gnu/Linux operating system.

On Linux, it is more complicated to describe the installation process, because there are many different Linux distributions. Still, there are mainly two ways of getting Scilab for Linux:

- downloading Scilab from http://www.scilab.org,
- using the packaging system of the Linux distribution, e.g. Debian, Ubuntu, etc...

Whatever the source, Scilab comes with the Reference BLAS and LAPACK, compiled from http://www.netlib.org/blas and http://www.netlib.org/lapack.

To install an optimized ATLAS library, we should compile our own version of ATLAS. Because this is slightly more complicated and takes CPU time (typically one or two hours of compiling), this is reviewed at the end of this section.

| Which components should be i | nstalled? | Sci |
|--|--|------------------------|
| Select the components you wa install Sink Next when you are | et to install; clear the component to install; clear the component | nts you do not want to |
| Custom installation | | |
| CPU Optimization for Scilat | | 0,2 MB 🔺 |
| - O Download Intel Math K | ernel Library for Scilab | 0,1 MB |
| - O Atlas Library for Scilab | | 6,3 MB |
| 📙 🕒 🖸 Blas, Lapack Referenc | e libraries for Scilab | 0,1 MB |
| Signal processing Module | | 0,9 MB |
| Loterpolation Module | | 0,3 MB |
| Optimization Medule | | 1,8 MB |
| XCos 1.0 | | 42,9 MB |
| IMI ARnoldi PACKage Toolbox | <u> </u> | 0.3 MB 🖿 |
| | A MARCHAND A MUL | |

Figure 39: Choosing the linear algebra library under Windows (Scilab v5.2.2). We can choose between Reference BLAS, ATLAS and the Intel MKL.

A simpler solution is to use the binaries from the Debian distribution. The ATLAS binaries in Ubuntu are coming from the Debian distribution, so we can get ATLAS binaries on Ubuntu as well. Under Ubuntu, we can use, for example, the Synaptic package manager to install and remove binary packages. In this case, we can use the libatlas3gf-base "ATLAS generic shared" binary which is provided by Debian.

Using an optimized ATLAS binary on Linux requires some simple manual changes that we are going to describe. In the directory scilab/lib/thirdparty, we find the following files.

```
$ ls scilab-5.3.0/lib/thirdparty
-rw-r--r- 1 mb mb   6.3M 2010-09-23 14:45 liblapack.so.3gf.0
-rw-r--r- 1 mb mb   539K 2010-09-23 14:45 libblas.so.3gf.0
```

When we remove the libblas.so.3gf.0 file from the Scilab installation directory, then Scilab uses the BLAS library from the Linux system. Therefore, to make sure that Scilab uses the ATLAS library that we have installed on the system, we simply remove the libblas.so.3gf.0 file, as in the following session.

```
$ rm scilab-5.3.0/lib/thirdparty/libblas.so.3gf.0
```

In order to get good performances, we should not use the binary provided by Debian. Instead, we should compile our own ATLAS library on the particular machine we use. This is because the particular settings which allow ATLAS to improve its performances are detected at compile time. Hence, the ATLAS binary which is provided in Debian is reasonably optimized for the machine that the package maintainer has used to compile ATLAS. This works quite well and is easy to use, but does not provide the maximum performances that we can get on another machine.

Fortunately, the sources are available at http://math-atlas.sourceforge.net, so that anyone on a Linux is able to customize his own ATLAS library to get good linear algebra performances. Moreover, on Debian, the process is made more easy because this particular Linux distribution provides all the necessary tools. More details on this topic are given in the section 5.6.

5.4.5 An example of performance improvement

In this section, we present a simple experiment on Windows which shows the benefit of using an optimized linear algebra library in Scilab. As a benchmark, we consider here the multiplication of two square, dense, real matrices of doubles.

We define in the following script the mymatmul function, which takes the size of the matrices **n** as an input argument and performs the multiplication of two square matrices filled with unity entries.

```
function mymatmul ( n )
A = ones(n,n)
B = ones(n,n)
C = A * B
endfunction
```

The multiply operator is, in this case, connected to the BLAS function named DGEMM. This function is often used to measure the performance of optimized linear algebra libraries.

| Library | Ν | Mean (s) |
|-------------------|------|----------|
| REF-LAPACK | 100 | 0.003004 |
| REF-LAPACK | 200 | 0.033048 |
| REF-LAPACK | 400 | 0.279402 |
| REF-LAPACK | 1000 | 4.118923 |
| ATLAS | 100 | 0.001001 |
| ATLAS | 200 | 0.011016 |
| ATLAS | 400 | 0.065094 |
| ATLAS | 1000 | 0.910309 |
| Intel MKL | 100 | 0.001001 |
| Intel MKL | 200 | 0.010014 |
| Intel MKL | 400 | 0.063091 |
| Intel MKL | 1000 | 0.834200 |

Figure 40: The sensitivity of the performances of a matrix multiplication depending on the linear algebra library. The benchmark is a matrix-matrix multiplication. It is performed on Scilab v5.2.2 under Windows XP 32 bits. The CPU is a AMD Athlon 3200+ at 2 GHz and the system runs with 1 GB of memory.

The following script makes use of the **benchfun** function that we have presented in the section 5.1.4. We set the size of the stack to the maximum, then compute the multiplication of dense matrices of increasing size.

```
stacksize("max")
benchfun ( "matmul", mymatmul , list(100) , 0 , 10 );
benchfun ( "matmul" , mymatmul , list(200) , 0 , 10 );
benchfun ( "matmul" , mymatmul , list(400) , 0 , 10 );
benchfun ( "matmul" , mymatmul , list(1000) , 0 , 10 );
```

The version of Scilab used is Scilab v5.2.2 under Windows XP 32 bits. The CPU is a AMD Athlon 3200+ at 2 GHz and the system runs with 1 GB of memory. In the following session, we have executed the benchmark script with Scilab and the Atlas library.

```
-->exec bench_matmul.sce;
matmul: 10 iterations, mean=0.00300, min=0.00000, max=0.01001
matmul: 10 iterations, mean=0.03304, min=0.02002, max=0.08011
matmul: 10 iterations, mean=0.27940, min=0.26037, max=0.37053
matmul: 10 iterations, mean=4.11892, min=4.08587, max=4.19603
```

The figure 40 presents the results that we got from the three optimized libraries. The best performance is obtained from the Intel MKL, which is closely followed by the Atlas library. The performances of the Reference BLAS-LAPACK library are clearly lower.

By using optimized linear algebra libraries, users of Scilab can get fast linear algebra computations. On Windows, Scilab is shipped with the Intel MKL, which provides most of the time excellent performances. Notice that this library is commercial, and is provided free of charge for Scilab users, because the Scilab Consortium has a licence for the Intel MKL. On Linux, Scilab is shipped with the reference BLAS-LAPACK library. Optimized Atlas libraries are available from many Gnu/Linux distributions, so that it is easy to customize our Scilab installation to make use of Atlas. Moreover, on Gnu/Linux, we can compile our own Atlas library and create a customized library which is specifically tuned for our computer.

5.5 Measuring flops

A common practice in scientific computing is to count the number of floating point operations which can be performed during one second. This leads to the *flops* acronym, which stands for FLoating Point Operations by Second. In this section, we present two standard benchmarks in Scilab involving the computation of the product of two dense matrices and the computation of the solution of linear equations by LU decomposition. In both cases, we compare the performances of the Reference BLAS, ATLAS and the Intel MKL linear algebra libraries provided with Scilab on Windows.

5.5.1 Matrix-matrix product

In this section, we consider the performance of the matrix-matrix product on a Windows machine with several linear algebra libraries.

Let us consider the product of A and B, two dense n-by-n matrices of doubles using the * operator in Scilab. The number of floating point operations is straightforward to compute, based on the definition:

$$C_{ij} = \sum_{k=1}^{n} A_{ik} B_{kj},\tag{3}$$

for i, j = 1, 2, ..., n. To compute one entry C_{ij} of the matrix C, we have to perform n multiplications and n additions. All in all, there are 2n floating point operations. Since there are n^2 entries in the matrix C, the total number of floating point operations is $2n^3$.

In this case, Scilab makes use of the BLAS DGEMM routine, which is a part of the level 3 BLAS. The foundations of this subset of BLAS are described in [15], where the authors emphasize that a general guideline is that efficient implementations are likely to be achieved by reducing the ratio of memory traffic to arithmetic operations. This allows to make a full use of vector operations (if available) and allows to exploit parallelism (if available). In the case of DGEMM, the number of memory references is $3n^2$ and the number of flops is $2n^3$. The ratio is therefore 2n/3, which is one of the highest in the BLAS library. For example, the ratio for the dot product (level 1) is 1, while the ratio for the matrix-vector product (level 2) is 2. Hence, we expect to get a good performance for the DGEMM routine.

Moreover, in [52], the authors emphasize that many level 3 BLAS routines can be efficiently implemented given an efficient matrix-matrix product. This is why a good performance of DGEMM is a key point in the overall performance of BLAS. In general, the matrices A, B and C are too large to fit in the cache of the processor. This is why the authors of ATLAS [52] suggest to use block-partitioned algorithms. Indeed, it is possible to arrange the operations to make so that the most part of the operations are performed with data in cache, by dividing the matrix into blocks. In order to measure the performance of the matrix-matrix product in Scilab, we use the following script. We use the random number generator rand to produce matrices which entries are computed from the normal distribution function. We know that the random number generator behind rand is of poor statistical quality and that the grand function may be used as a replacement. But this has no effect on the performance of the matrix-matrix product, so that we keep it for simplicity. The 1.e6 factor in the computation of the mflops variable converts flops into mega-flops.

```
stacksize("max");
rand( "normal" );
n = 1000;
A = rand(n,n);
B = rand(n,n);
tic();
C = A * B;
t = toc();
mflops = 2*n^3/t/1.e6;
disp([n t mflops])
```

The benchmark is performed on Scilab v5.2.2 under Windows XP 32 bits. The CPU is a AMD Athlon 3200+ at 2 GHz and the system runs with 1 GB of memory. We compare here the performances of the three linear algebra libraries provided with Scilab, that is Reference BLAS, ATLAS and the Intel MKL. In order to keep the timings doable, we had to select separate matrix sizes for the various libraries. The results are presented in the figure 41.

In this experiment, the fastest libraries are the Intel MKL and ATLAS (more than 2000 Mflops), which are much faster than the Reference BLAS library (about 500 Mflops). The matrix sizes **n** that we used in the experiment are depending on the actual performance of the libraries, for a reason that we now analyze. We selected the size **n** for which the library produced timings between 0.1 second and 8 seconds. This has the main advantage of making the benchmark as fast as possible, while maintaining good reproductibility. Indeed, if **n** is too small, it may happen that the measured time **t** is zero. In this case, the computation of **mflops** is impossible, because of the division by zero. Conversely, the Reference BLAS library is so slow, and the timing grows so fast (as expected by the $2n^3$ formula), that running the algorithm for large matrix sizes leads to very large timings, making the benchmark process difficult in practice.

We now compare the actual performance of the libraries to the peak performance that we can expect for this particular CPU. The CPU is a AMD Athlon 3200+, code name Venice, for the Socket 939. The frequency of this CPU is 2 GHz. If the CPU is able to perform one floating point operation by cycle, the 2 GHz frequency implies that the peak performance should be greater than 2 Gflops, i.e. 2000 Mflops. The previous numerical experiment shows that ATLAS and the Intel MKL are able to reach this peak performance, and even a little bit more. This proves that the CPU was able to perform from 1 to 2 floating point operations by CPU cycle. This processor is a single-core CPU which supports MMX, 3DNow!, SSE, SSE2 and SSE3. The fact that the CPU supports these instruction sets may explain the fact that the actual performance was slightly above 2000 Mflops.



Figure 41: Performances of real, square, dense, matrix-matrix multiplication in Scilab with various libraries on Windows. Higher is better.

Let us consider the sensitivity of the performance on the size of the cache. It may happen that the linear algebra library exhibits an increased performance for matrices which can be entirely stored inside the cache. The goal of the ATLAS and Intel MKL libraries is to make so that this does not happen. Hence, the performances should be kept independent of the cache size. Let us check that this actually happens in this particular experiment. The L1 data cache is 64 KBytes while the L2 cache is 512 KBytes. As a double precision floating point number requires 8 bytes (i.e. 64 bits), the number of doubles which can be stored in the L1 cache is 64000/8 = 8000 doubles. This corresponds to a square 89-by-89 dense matrix of doubles. The L2 cache, on the other hand, can store a 252-by-252 dense, square, matrix of doubles. Since our matrices ranges from 500-by-500 to 2000-by-2000, we see that the total number of doubles cannot be (by far) entirely stored in the cache. Hence, we conclude that the algorithms used in the Intel MKL and ATLAS are not too sensitive to the size of the cache.

5.5.2 Backslash

In this section, we measure the performance of the backslash operator in Scilab on Linux. This benchmark is often called the "LINPACK benchmark", because this benchmark was created when the LINPACK project was in development. The LINPACK benchmark is still used nowadays, for example as a measure of the performance of High Performance Computers, such as the machines from the Top 500 [4], for example. This library is superseded by BLAS and LAPACK, which are used by Scilab.

Let us consider a square, dense, real, *n*-by-*n* matrix of doubles A and a *n*-by-1 real vector of doubles b. The backslash operator \setminus allows to compute the solution **x** of the linear equation A*x=b. Internally, if the matrix is reasonably conditionned, Scilab computes the LU decomposition of the matrix A, using row pivoting. Then, we perform the forward and backward substitutions using the LU decomposition. These operations are based on the LAPACK library, which, internally, makes use of BLAS. The precise names of the LAPACK routines used by the backslash operator are presented in the section 5.6.

Solving linear systems of equations is a typical use-case for linear algebra libraries. This is why this particular benchmark is often used to measure the performance of floating point computations on a particular machine [39, 16, 42]. The number of floating point operations for the whole process is $\frac{2}{3}n^3 + 2n^2$. Since this number grows as quickly as n^3 while the number of entries in the matrix is only n^2 , we can expect a good performance on this test.

The following script allows to measure the performance of the backslash operator in Scilab. We configure the random number generator from the **rand** function so that it generates matrices with entries computed from the normal distribution function. Then we measure the time with the **tic** and **toc** functions. Finally, we compute the number of floating point operations and divide by **1.e6** in order to get mega-flops.

```
n = 1000;
rand( "normal" );
A = rand(n,n);
b = rand(n,1);
tic();
x = A\b;
t = toc();
mflops = (2/3*n<sup>3</sup> + 2*n<sup>2</sup>)/t/1.e6;
disp([n t mflops])
```

We perform this test with scilab-5.3.0-beta-4 on Linux Ubuntu 32 bits. The CPU is an Intel Pentium M processor at 2GHz with 1 GB memory. The cache size is 2 MB. The CPU supports the MMX, SSE and SSE2 instruction sets. The figure 42 presents the results with increasing matrix sizes and compare the performances of the Reference BLAS and ATLAS. The version of ATLAS that we use is a precompiled binary which is provided by Ubuntu. For practical limitations reasons, these experiments are stopped when the time required to execute one backslash is larger than 8 seconds.

As we can see, the ATLAS library improves the performances of the backslash operator by a factor roughly equal to 2. Moreover, the Mflops regularly increases with the matrix size, while the performance of the Reference BLAS library seems to stay the same. The CPU frequency is 2 GHz, which implies that we can expect a peak performance greater than 2000 Mflops. The actual performance of ATLAS



Figure 42: Performances of real, square, dense, linear equations solution computation with backslash in Scilab with various libraries on Linux.

on these matrices is in the [1000,1400] range, which is somewhat disappointing. As this performance is increasing, this peak performance may be reached for larger matrices, but this has not been checked here, because the timings grows very fast, making the experiments longer to perform.

5.5.3 Multi-core computations

In this section, we show that Scilab can perform multi-core floating point computations if we use, for example, the Intel MKL linear algebra library which is provided with Scilab on Windows.

We used for this test a Scilab v5.3.0-beta-4 version on a Windows Vista Ultimate 32 bits operating system. The processor is an Intel Xeon E5410 with 4 cores at 2.33 GHz [25, 53]. The processor can manage 4 threads (i.e. one thread by core) and has 12 MB L2 cache. The operating system can use 4 GB of physical memory. The figure 43 presents the results of the matrix-matrix product benchmark that we have presented in the section 5.5.1.

Given that this processor has 4 cores at 2.33 GHz, we expect a performance greater than 9.33 Gflops. The actual performance of the Intel MKL is 26.494 Gflops, which shows that this library has made use of the 4 cores, performing 2.8 floating

| Library | Ν | User Time (s) | Mflops |
|-----------|------|-----------------|--------|
| Intel MKL | 4766 | 8.172 | 26494 |
| ATLAS | 1595 | 3.698 | 2194 |
| Ref. BLAS | 444 | 0.125 | 1400 |
| | | | |

Figure 43: The performance of the matrix-matrix product on a 4 cores system on Windows with various libraries.

point operations by cycle. On the other side, the ATLAS and Ref-BLAS libraries do not use the 4 cores of this machine and get much lower results.

5.6 Notes and references

In the section 5.2.3, we analyzed the performances of a summation algorithm. It could be pointed that the accuracy of a sum may depend on the size of the data and on the particular values in the dataset. This is analyzed, for example, by Wilkinson [54] and Higham [23]. More precisely, it can be shown that a dataset can be ill-conditionned with respect to the sum, for example if the values are of mixed signs (that is, some are positive while some others are negative), and of very different magnitude. In this case, we may observe that the relative error of the sum may be large.

In this document, we have focused mainly on performance and refer to the previous references for the accuracy issues. In practice, it is difficult to completely separate the two topics. Indeed, it is more easy to get better performances if we completely neglect the accuracy issues. This is why it is safe to consider optimizations only after we have a sufficiently large basis of tests, including both correctness and accuracy. This ensures that, during the "optimization" process, we do not introduce a bug or, worse, an unnecessary error in the result.

A topic which is closely associated with vectorization of Scilab algorithms is matrix indexing, that is, selecting a subset of elements within a matrix. The article by Eddins and Shure [18] focuses on this topic within the context of Matlab (\mathbb{R}) , where the authors presents indexing of vectors, indexing of matrices with two subscripts, linear indexing and logical indexing based on boolean expressions. Matlab is a registered trademark of The MathWorks.

In the section 5.2.1, we have presented several methods associated with compiling principles. The "Dragon Book" [5] is a classical reference on the subject of compilers. The book provides a thorough introduction to compiler design, including lexical analysis, regular expressions, finite state machines and parsing methods.

In the section 5.4.1, we briefly presented the ATLAS project. Whaley and Dongarra [51] describe an approach for the automatic generation and optimization of numerical software for processors with deep memory hierarchies and pipelined functional units.

In the section 5.1.3, we presented a Gaussian elimination algorithm and stated that this algorithm (but not the particular implementation that we presented) is used in Scilab, behind the backslash operator. The actual behavior of the backslash operator in Scilab is the following. First, we perform a call to the DGETRF LAPACK routine, which decomposes the matrix A into PA = LU, where P is a permutation matrix, L is a lower triangular matrix and U is an upper triangular matrix. The diagonal entries of L are unity. Then we call the DGECON LAPACK routine, which approximates the inverse of the 1-norm condition number of the matrix A. At this point, there are two cases. If the condition number of the matrix is not too large, then the system of linear equations is not ill-conditionned (this is not exact, because the condition is only approximated, but the algorithm makes the hypothesis that the estimation can be trusted). In this case, we call the DGETRS LAPACK routine, which solves the linear system of equations by using the PA = LU decomposition. More precisely, this step uses row permutations, and then the forward and backward substitutions in order to compute the solution \mathbf{x} depending on the right-hand side \mathbf{b} . In the other case, that is, if the matrix is ill-conditionned, we use a modified version of the DGELSY LAPACK routine, which solves the associated least-squares problem. This choice may reduce the accuracy for matrices of intermediate condition [10].

In [39], Cleve Moler analyze the performances of Matlab in various situations, including the computation of the solution of a linear system of equations with the backslash operator. To measure the performance of the computation, he counts the number of flops, by measuring the elapsed time in second with Matlab's tic and toc functions, and estimating the number of floating point operations for this particular computation. The author uses a random matrix with entries computed from the normal distribution function and uses the randn function in Matlab. By increasing the size of the matrix, he observes a peak in the performance, which corresponds to the size of the cache of the SPARC-10 computer he uses for this test. This machine has a one megabyte cache, which corresponds to a square matrix of order

More precisely, Moler uses the definition that associates one megabytes to 2^{20} bytes, but the result is roughly the same. The peak megaflop rate is obtained for a matrix that just fits in the cache. He states that this cache effect is a consequence of the LINPACK library. He emphasize that this is one of the primary motivations for the LAPACK project. This paper was written in 1994: Matlab uses the LAPACK library since 2000 [40]. In [16], Jack Dongarra collects various performances on this benchmark. Related resources are available on netlib [42].

General resources related to performance improvement for Matlab are available in [38].

In the section 5.4.3, we have discussed the installation of Scilab on Windows. This topic is discussed further in [14], where Allan Cornet, who is a developer from the Scilab Consortium, presents the full installation process of Scilab 5.2.0 on Windows.

In the section 5.4.4, we have presented the installation of a binary version of ATLAS for Scilab on Linux/Ubuntu. Since ATLAS is optimized at build time, it is better to compile ourselve ATLAS on the machine where we plan to use it. This guarantees that the binary that we use is really optimized with respect to the parameters of the current machine (for example, the size of the cache). In [48], Sylvestre Ledru, who is a developper from the Scilab Consortium and maintains the

Debian Science package, provides informations about the installation of ATLAS on Debian. This topic is presented in more depth in [30].

There are other references which may help Scilab users on Linux. In [31], Ledru describes the installation process for Scilab 5.1.1 on Linux. In [32], Ledru discusses the update of the management of BLAS and LAPACK packages under Debian. In [29], Ledru presents a method which allows to switch between different optimized linear algebra libraries.

In the section 5.1.4, we presented the **benchfun** function, which measures the performances of a given function. Alternatively, we can use the Scibench module [6], which is provided on ATOMS. The goal of the Scibench project is to provide tools to measure the performances of a function. Moreover, the module provides a collection of benchmark scripts to measure the performance of Scilab. Within the Scibench module, the scibench_benchfun function has the same goal as benchfun, which has been presented in the section 5.1.4.

5.7 Exercises

Exercise 5.1 (*Gauss pivoting***)** In the section 5.1.3, we have presented two functions gausspivotalnaive and gausspivotal which both implement the Gaussian elimination algorithm with row pivoting. Use the benchfun function and compare the actual performances of these two functions.

Exercise 5.2 (*Combinations*) Consider two 1-by-n matrices of doubles x and y. We would like to create a function generating at 2-by-m matrix c containing all the combinations of the vectors x and y, with $m = n^2$. That is, we would like to produce the couples [x(1);y(1)], [x(1);y(2)], [x(1);y(3)], ..., [x(n);y(n)]. For example, consider the following matrices x and y.

 $-->x = [10 \ 11 \ 12];$ $-->y = [13 \ 14 \ 15];$

We would like to generate the following matrix c:

-->c c = 10. 10. 10. 12. 12. 11. 11. 11. 12. 14. 15. 13. 13. 14. 15. 13. 14. 15.

- Create a naive implementation to produce the matrix c.
- Use the Kronecker product .*. and create a vectorized function to produce the matrix c.
- Use matrix extractions to create a vectorized function to produce the matrix c.

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7 Answers to exercises

7.1 Answers for section 2

Answer of Exercise 2.1 (*Maximum stack size*) The result of the algorithm which sets the size of the stack depends on the operating system. This is the result on a Windows machine :

```
-->format(25)
-->stacksize("max")
-->stacksize()
ans =
110054096. 34881.
```

and this is the result on a Linux machine

Answer of Exercise 2.2 (who_user) The following session is the result on a Linux machine.

```
-->who_user()
User variables are:
home
Using 7 elements out of 4990798
-->A=ones(100,100);
-->who_user()
User variables are:
A home
Using 10009 elements out of 4990796
-->home
home =
/home/mb
```

Actually, there may be other user-defined variable which may be displayed by the who_user function. For example, I have installed the ATOMS "uncprb" module, and here is what I get at startup.

```
-->who_user()
User variables are:
uncprblib
Using 217 elements ouf of 4990785
```

This is because the uncprblib variable contains the library associated with this particular module.

Answer of Exercise 2.3 (whos) The following session presents the output of the whos function, which displays the name, type, size and bytes required by all the variables in the current environment.

| >whos | | | |
|-------------|------------|--------|-------|
| Name | Туре | Size | Bytes |
| \$ | polynomial | 1 by 1 | 56 |
| %driverName | string* | 1 by 1 | 40 |
| %e | constant | 1 by 1 | 24 |
| %eps | constant | 1 by 1 | 24 |
| [] | | | |

| guilib | library | | 488 |
|------------------|---------|--------|------|
| helptoolslib | library | | 728 |
| home | string | 1 by 1 | 56 |
| integerlib | library | | 1416 |
| interpolationlib | library | | 336 |
| [] | | | |

7.2 Answers for section 3

Answer of Exercise 3.1 (Searching for files) The following function searchSciFilesInDir searches for these files in the given directory.

```
function filematrix=searchSciFilesInDir(directory,funname)
filematrix = []
lsmatrix = ls ( directory )';
for f = lsmatrix
    issci = regexp(f,"/(.*).sci/");
    if ( issci <> [] ) then
        scifile = fullfile ( directory , f )
        funname ( scifile )
        filematrix($+1) = scifile
    end
end
end
endfunction
```

In our example, we will simply display the file name. The following function mydisplay uses the mprintf function to display the file name into the console. In order to print a brief message, we skip the directory name from the displayed string. In order to separate the name of the directory from the name of the file, we use the fileparts function.

```
function mydisplay ( filename )
  [path,fname,extension]=fileparts(filename)
  mprintf ("%s%s\n",fname,extension)
endfunction
```

In the following session, we use the searchSciFilesInDir function to print the scilab scripts from the Scilab directory containing the macros associated with polynomials. The SCI variable contains the name of the directory containing the Scilab directory. In order to compute the absolute filename of the directory containing the macros, we use the fullfile function, which gathers its input arguments into a single string, separating directories with the separator corresponding to the current operating system (i.e. the slash "/" under Linux and the backslash "\" under Windows).

```
-->directory = fullfile(SCI,"modules","polynomials","macros")
directory =
/home/username/scilab-5.2.2/share/scilab/modules/
    polynomials/macros
-->filematrix = searchSciFiles ( directory , mydisplay );
invr.sci
polfact.sci
cmndred.sci
[...]
horner.sci
rowcompr.sci
htrianr.sci
```

Answer of Exercise 3.2 (*Querying typed lists*) We have already seen that the definedfields function returns the floating point integers associated with the locations of the fields in the data structure. This is not exactly what we need here, but it is easy to build our own function over this building block. The following isfielddef function takes a typed list tl and a string fieldname as input arguments and returns a boolean bool which is true if the field is defined and false if not. First, we compute ifield, which is the index of the field fieldname. To do this, we search the string fieldname in the complete list of fields defined in tl(1) and use find to find the required index. Then, we use the definedfields function to get the matrix of defined indices df. Then we search the index ifield in the matrix df. If the search succeeds, then the variable k is not empty, which means that the field is defined.

```
function bool = isfielddef ( tl , fieldname )
    ifield = find(tl(1)==fieldname)
    df = definedfields ( tl )
    k = find(df==ifield)
    bool = ( k <> [] )
endfunction
```

Although the program is rather abstract, it remains simple and efficient. Its main virtue is to show how typed lists can lead to very dynamic data structures. \Box

7.3 Answers for section 5

Answer of Exercise 5.1 (*Gauss pivoting*) We use the following script, which creates a 100by-100 matrix of doubles and use it to compare the performances of gausspivotalnaive and gausspivotal.

```
stacksize("max");
n = 100;
A = grand(n,n,"def");
e = ones(n,1);
b = A * e;
benchfun("naive",gausspivotalnaive,list(A,b),1,10);
benchfun("fast",gausspivotal,list(A,b),1,10);
```

The previous script produces the following output.

```
naive: 10 iterations, mean=1.541300, min=1.482000, max=2.03
fast: 10 iterations, mean=0.027000, min=0.010000, max=0.070000
```

In this case, for this matrix size, the average performance improvement is 1.5413/0.027=57.08. In fact, it could be larger, provided that we use sufficiently large matrices. For n=200, we have observed a performance ratio larger than 100.

Answer of Exercise 5.2 (*Combinations*) We consider two vectors x and y, with n entries. The goal of this exercise is to create a function able to generate a 2-by-m matrix c containing all the combinations of the vectors x and y, where $m = n^2$.

Our first idea is to use two nested loops: this idea is implemented in the following function. The first loop on ix enumerates the entries in x, while the second loop on iy enumerates the entries in y. The algorithm is initialized with k=1. Each time we create a new combination, we store it in the kth column of c, and then we increase k.

```
function c = combinatevectorsSlow ( x , y )
    cx=size(x,"c")
    cy=size(y,"c")
    c=zeros(2,cx*cy)
    k = 1
    for ix = 1 : cx
```

The previous function works, but is slow because it requires to perform n^2 loops. Therefore, the goal is to remove these two nested loops and to generate the combinations with vectorized statements.

Our second idea is to use the Kronecker product to generate all the combinations in just one statement. The following session shows the basic idea to generate all the combinations of the vectors [10 11 12] and [13 14 15].

```
-->[10 11 12] .*. [1 1 1]
 ans =
   10.
        10.
                 10.
                                             12.
                                                    12.
                                                           12.
                        11.
                               11.
                                      11.
-->[1 1 1] .*. [13 14 15]
 ans
   13.
          14.
                 15.
                        13.
                               14.
                                      15.
                                             13.
                                                    14.
                                                           15.
```

The following implementation is a direct generalization of the previous session.

```
function c = combinatevectorsKron ( x , y )
    cx=size(x,"c")
    cy=size(y,"c")
    c(1,:) = x .*. ones(1,cy)
    c(2,:) = ones(1,cx) .*. y
endfunction
```

Our third idea is to use the matrix extraction operations. Indeed, the first row in c is just the copy of the vector x, three times. This is why we create a matrix of repeated indices, based on matrix extraction. The repeated indices are also created with repeated indices with matrix extraction. These ideas are presented in the following session, which computes the first row in c. In this session, we first create the indices i1 from 1 to 3. Then we create the indices j1 by repeatedly extracting the row i1. Then we create k1 by using the matrix function, which converts the j1 matrix into a row vector of indices. Finally, we use k1 as indices of the columns in x.

```
-->x = [10 \ 11 \ 12];
-->i1 = 1:3
 i1 =
    1.
            2.
                   З.
-->j1 = i1([1;1;1],:)
 j1
                    З.
    1.
            2.
            2.
                   3.
    1.
    1.
            2.
                    З.
-->k1 = matrix(j1, 1, 9)
 j1
          1.
                1.
                      2.
                            2.
                                   2.
                                         З.
                                               З.
                                                     З.
   1.
-->x(:,k1)
 ans
                  10.
                                                 12.
                                                        12.
   10.
           10.
                          11.
                                  11.
                                         11.
                                                               12.
```

We can use the same principles to generate the second row in c. But the entries are not just repeated, they are also alternated. In order to do this, we use the same idea as previously, but transpose the indices before we convert the matrix of indices into a vector of indices.

```
-->y = [13 14 15];
-->i2 = 1:3
```

```
i2
    =
            2.
    1.
                   З.
-->j2 = i2([1;1;1],:)
 j2
     =
            2.
                    З.
    1.
    1.
            2.
                   3.
            2.
                   З.
    1.
-->k2 = matrix(j2',1,9)
 k2
     =
   1.
         2.
                З.
                      1.
                            2.
                                  З.
                                         1.
                                               2.
                                                     З.
-->y(:,k2)
 ans
      =
           14.
                  15.
                          13.
                                 14.
                                         15.
                                                13.
                                                        14.
                                                               15.
   13.
```

In the previous session, it is important to notice the single quote operator in the expression matrix(j2',1,9). The following script is a direct generalization of the previous ideas.

```
function c = combinatevectorsExtr ( x , y )
   // Returns all combinations of the two row vectors x and y
   cx=size(x,"c")
   cy=size(y,"c")
   11
   i1 = 1:cx
   j1 = i1(ones(cy, 1), :)
  k1 = matrix(j1, 1, cx * cy)
   11
   i2 = 1:cy
   j2 = i2(ones(cx, 1), :)
  k2 = matrix(j2', 1, cx*cy)
   //
   c(1,:)=x(:,k1)
   c(2,:)=y(:,k2)
endfunction
```

In the following session, we use the **benchfun** function to compare the performances of the previous functions.

```
n = 300;
x=(1:n);
y=(1:n);
benchfun("Slow",combinatevectorsSlow,list(x,y),1,10);
benchfun("Kron.",combinatevectorsKron,list(x,y),1,10);
benchfun("Extr.",combinatevectorsExtr,list(x,y),1,10);
```

The previous script typically produces the following output.

```
Slow: 10 iterations, mean=0.24010, min=0.23700, max=0.25000
Kron.: 10 iterations, mean=0.01440, min=0.01300, max=0.02200
Extr.: 10 iterations, mean=0.00960, min=0.00700, max=0.01500
```

We see that the implementation based on the Kronecker product is more than 10 times faster on the average. We notice that there is no significant difference between the combinatevectorsKron and combinatevectorsExtr functions, although combinatevectorsExtr seems a little faster. This may be explained by the fact that combinatevectorsKron performs many floating point multiplications, which are unnecessary since one of the operands is unity.

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