# chemcode<sup>®</sup> chemical pocket calculator

mobile and fast calculations in chemistry and life sciences

# Manual



diginorm elektronik

and

Peter Barthel & Wilhelm Schmidthals GmbH



### **Important Notice**

- 1) The calculator switches off automatically two minutes after the last input. The last input of a
- compound formula or a

sequence of nucleotides

is memorized automatically and displayed again after switching on the calculator and entering into a menu.

 If troubles should occur during operation (e.g. caused by a strong electrical field): press the key "on" (reset).

Afterwards the calculator can be operated as usual. The last compound formula / sequence of nucleotides put in will appear again in the display after entering a menu.

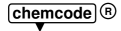
See chapter 9.1 explaining the automatic memory function.

# chemcode® chemical pocket calculator

mobile and fast calculations in chemistry and life sciences

Manual

# Please keep this manual always carefully together with your chemcode®!



chemcode<sup>®</sup> − chemical pocket calculator Mobile and fast calculations in chemistry and life sciences Manual

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### Introduction

Thank you very much for your confidence in our product, the chemical pocket calculator chemcode  $^{\tiny{\textcircled{\tiny 0}}}\!!$ 

We are sure of having delivered a valuable tool of great use. Stoichiometric calculations of your daily lab routine will be decisively simplified with this innovative product.

Everybody having to deal frequently with stoichiometric standard calculations will soon appreciate the chemcode® as an indispensable companion for educational and professional purposes.

Please make yourself acquainted with the functions of the calculator by studying this manual before use. You will quickly achieve routine in operating the chemcode<sup>®</sup>. Operation is easy and the display is self-explaining to a high degree. Should there be, however, any troubles in understanding a function you surely will find the answer to your problem in this manual.

You will find further informations about the project chemcode  $^{\tiny{\textcircled{\scriptsize 0}}}$  in the internet adressing our website:

### www.chemcode.com

There are also forms for giving us feedback, making proposals for the optimization of the chemcode® and for asking questions. Thus we will be able to make the chemcode® a still more attractive tool in the future.

We are looking forward to your feedback!

In representation of your chemcode®-team

Florian Ens - Dr. Robert Stark

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### **GETTING STARTED – KEYBOARD STRUCTURE**

#### 1.1 **LANGUAGE**

All terms and abbreviations of the display, the case and the keyboard follow the usage of English language.

Explanation of abbreviations: See chapter 10.

### TURN ON / OFF 1.2



← On: Press key "on"

M1

Off: "2nd" and "clr" (second function of "clr") →





### **SECOND FUNCTION OF KEYS** 1.3



A print above a key marks the second function of this key (except for prints on green lines). They are activated by pressing the key "2nd". Afterwards the symbol "2nd" appears in the top line of the display:

1st line

Molar Mass

2nd line →

M = 0 g/mol

Second functions are disactivated by pressing the key "2nd" again (or by pressing any other key without second function) and the symbol "2nd" in the top line disappears.

After the execution of a key's second function (print above a key) the primary key functions (symbolized by the print on the key) are automatically reactivated and the symbol "2nd" in the top line vanishes.

Please note: Nickel (Ni 28) is the second function of Selenium (Se 34), "Pd 46" of "Te  $^{52}$ ", and "Pt  $^{78}$ " of "Po  $^{84}$ ". The three elements, that belong to transition group VIII in the periods 4, 5, 6 and 7 are enclosed by a white frame. Nevertheless every element symbol on the case is a second function of the key below of this symbol.

### Example: Input of Fe<sub>2</sub>O<sub>3</sub>



Press key "Cm": The calculator changes to menu M1:

Line 2

M1 Molar Mass
M = 0 q/mol

Note: This example deals with the case, that the compound formula put in at last has been deleted by pressing the key "clr" (see ch. 3.2.1). If this has not been the case, the last compound formula put in would be visible in line 2. See p. 22, ch. 5.2 step 2a and 2b.



Press key "2nd" - the symbol "2nd" appears in the top line:

Top line

Line 1 →

M1 Molar Mass

Line 2

M = 0 g/mol



Press key "Co  $^{27}$ ": Its second function "Fe  $^{26}$ " has been activated before by pressing the key "2nd": Display:

• Top line: symbol "2nd" disappears.

• line 1: **Fe** 

• line 2: molar mass of Fe: 55.847 g/mol

Top line →
Line 1 →
Line 2 →

M1 Fe M = 55.847 g/mol

-

• Press key "2" for the subscript of "Fe" in Fe<sub>2</sub>O<sub>3</sub>



• Press key "O 8"



• Press key "3" (subscript of "O" in Fe<sub>2</sub>O<sub>3</sub>)

 $\bullet$  Press key " = ". The result is shown in the display:

Top line →
Line 1 →
Line 2 →

M1 Fe<sub>2</sub>O<sub>3</sub> M = 159.692 g/mol

Further details: See ch. 5.2.

### 1.4 PERIODIC SYSTEM KEYBOARD (WITH KEYS FOR NUCLEOTIDES OF DNA / RNA)

VIII b

All letters and numbers on the green lines of the case serve for orientation within the periodic system and do not mark second functions of keys (e.g. VIII b = transition group 8; further explanation see ch. 11.2).



Na 11 ← • Elements of the main groups: ...... black letters



• Lanthanoides and actinoides: ...... blue letters



Bases of DNA- / RNA-nucleotides: blue letters

These are second functions of the keys from "B 5" up to "O 8", that are automatically active in the menus GeD und GeR. In these menus the key "2nd" must not be pressed!

### MENU CONTROL 2

### **CALCULATOR MODE** 2.1

After switching on (key "on") the device is always in the calculator mode. Zero signals in line 2:1

Line 1 → Line 2 →



As long as the device is in the caculator mode the letters "Calculator" appear in line 1.

In the calculator mode usual calculation operations can be performed by using the numerical and operation keys like with any other pocket calculator (see chapter 8).

By pressing the key " $\mathbf{Cm}$ " the device changes to menu  $\mathbf{M1}$ .

In this manual signalling of symbols in the display is graphically symbolized by underlining these symbols.

### 2.2 KEY "Cm" (CHANGE MENU)



Pressing this key causes the device to change to the next menu (or from the calculator mode to menu **M1**). Each menu is symbolized by an abbrev. on the left side of line 1. This abbrev. lasts as long as the menu is not left:

Examples: • M = molar mass

- Sol = solution
- Explanation of further abbreviations: See chapter 10.1

Order of menus: calculator mode, M1, So1, C1, T, D, F, P1, Lib, GeD, GeR, calculator mode, M1, So1, ...

Each menu (except menu Lib - Library) offers:

- a single chemical calculation or
- a group of submenus each submenu for another calculation.

Each submenu is marked by a number following directly after the menu's abbreviation. After change into a menu consisting of a group of submenus, the submenu with number "1" appears at first.

The menus GeD (for DNA) and GeR (for RNA), however, are different. Though they have no number, they are the first submenus of two whole groups of submenus. In both groups only the following submenus (from the second on) are marked with numbers: Ge1, Ge2, etc. 1

### 2.3 KEY "Sm" (SUBMENU)



Pressing this key causes the device to change to the next submenu. After the last submenu the first one appears again.

Example: C1, C2, C3, then again C1, etc.

Graphic illustration of menu order: see chapter 4 (page 20).

The submenus Ge1 – Ge6 of GeD and GeR are in principle identical. There are, however, some necessary differences: E.g. the base Uracil is used instead of Thymine during calculations.

### KEY COMBINATION "2nd" AND "C m" 2.4





Pressing these two keys causes the device to change to the preceding menu. Thus the menus can be entered the other way round :

Menu order: calculator mode, GeR, GeD, Lib, P1, F, D, T, C1, Sol, M1, calculator mode, GeR, GeD, ...

### **NAVIGATING THROUGH MENUS AND SUBMENUS** 2.5



Within a menu or submenu the key " = " must be pressed in order to reach the next step in a menu. This could be:

- A new input / the affirmation of a displayed value
- Display of the result





Exception: Menus Lib and Ge6: Navigation by "scroll keys".

### **DISPLAY: STRUCTURE, INPUT, DELETING, SCROLLING**

### 3.1 STRUCTURE AND DATA INPUT

### Line 1

Top line Line 1 Line 2





• abbrev. (left) and name of the menu

a) Left Side: The menu's abbreviation, e.g. "M1" or "So1".

b) Right Side: Alternatively:

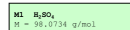
- 1) The  $\boldsymbol{menu's}$   $\boldsymbol{name} : e.g.$  "Solution" (when starting the menu);
- 2) A compound formula or a nucleotide sequence;
- 3) A formula or an explanation: e.g. " $n/V[mol/1] \rightarrow m[g]$ ".

- Calculator mode (see this ch., p. 12 under c).
- Menu **Lib**: The element's name appears (see p. 57).
- Menu Ge6: To the right there are the base tripletts by which a special amino acid is coded (see p. 69).

### c) Input into the right side of line 1 at menu start

• Compound formula - menus M1-M4, So1, C1-C3, P1-P2





A compound formula can also appear when memorized before (automatic memory or using the using the key "read memory").

Input: Change between element keys and numerical keys (for subscripts). Line 2: molar mass of the last element put in or of the whole compound after pressing the key " = ".

Usually the input of data is done when starting Menu M1. The compound formula put in will be automatically taken over by the other stoichiometric menus mentioned above. There it serves as the base for further calculations. This is not the case in the menus  $\mathbf{T}, \mathbf{D}, \mathbf{F}$ , where other data than compound formulas serve as calculation base.

Compound formulas can be replaced in every menu mentioned above (not in P3-P4) by writing the new formula over the old one.

• Nucleotide sequence - menus GeD and GeR





Starting these menus a nucleotide sequence can also be displayed by a memory function. See ch. 9, p. 78.

Input: Press the keys below the symbols for the bases (A, T / U, G, C). (In line 2 appears the corresponding amino acid sequence.)

The input of nucleotide sequences (or calling them up from memory) can only be performed in **GeD** und **GeR** (see ch. 8.2). Data delivered from memory functions can be replaced by new inputs!

The input of a nucleotide sequence will be taken over into the menus **Ge1** up to **Ge5**. There they serve as the base for further calculations. (**Ge6** is a lexical function: base tripletts that code a special amino acid.)

### d) Calculator mode (constants, atomic mass unit):

- In line 1 the letters: "Calculator" appear.
- This is also the case with the functions "C" ("2nd" and "5") and "u" ("2nd" and "6"). Called up values can be used in the calculator mode.
- By pressing any operation key, it is possible to get into the calculator mode. Then the letters "Calculator" appear.

### 3.1.2 Line 2

 a) Left Side: Before the sign " = " you find a formula sign for the value displayed or required.

```
Line 1 \rightarrow Sol n/V[mol/1] \rightarrow m[g] Exceptions: Menus GeD, GeR, P3, P4 Lib, Ge6.
```

**Examples** (a signalling zero is marked by underlining):

- "M = 98.0734 g/mol" (molar mass = 98.0734 g/mol)
- "[x] = 0 mol/l" (value x of moles per I = ... mol / I)
- "V =  $\underline{0}$   $\underline{1}$ " (Volume = ... liter) / "OD =  $\underline{0}$ " (optical density = ...)

Explanation of further abbreviations: see ch. 11.1.

P1 - P2: There is an element symbol on the display's left side without " = ".

### b) Right side of line 2

b1) Corresponding values appear in line 2 after an input in line 1 or if data are delivered by a memory function.

(see page 11, ch. 3.1.1 o):

```
Line 1 → M1 H<sub>2</sub>SO<sub>4</sub>
Line 2 → M = 98.0734 g/mol
```

GeR AUGUGUUUGCCGUGUGAU
STACysLeuProCysAsp

Molar masses after input of compound formulas in the menus M1-M4, Sol, C1-C3, P1-P4.

Amino acid sequences after input of nucleotide sequences in the menus **GeD** and **GeR**.

b2) An input is required (at the beginning or in the course of a menu)

• Zero is signalling:



Zero has to be written over by a number. In Menu **M1** zero signals in line 2 at the menu start (see ch. 3.1.1). The input, however, is displayed in line 1. In line 2 corresponding values appear [see example under b1) on this page or chapter 5.2 with more details].

### • A proposed value is signalling:

Line 1 →

```
 \begin{array}{ccc} & & & \\ \text{C3} & & \text{K[\$]}, \rho \rightarrow & \text{x[mol/1]} \\ \rho & & & \text{global} & \text{global} \\ & & \text{e} & \text{signalling is symbolized by underlining} \\ \end{array}
```

Example: In menu C3 (conversion: mass percentage into mol / I) the density of some acids / bases for commonly available concentrations is proposed (memorized values see page 37). The signalling value can be replaced or affirmed by pressing the key " = ".

Example in the display above: NH $_3$  aq [aqueous solution of ammonia] with a mass percentage of 25: density  $\rho=910$  g / l.)

### c) A result is indicated

After pressing the key " = " results are indicated on the right side of line 2. This value never signals!

Line 1 →

```
P1 H<sub>2</sub>SO<sub>4</sub>
S 32.6898 %
```

In line 2 of this example the percentage of the element Sulfur (S) in the compound  $H_2SO_4$  is indicated.

### d) Calculator mode (line 1: "Calculator")

Inputs and the display of results are performed in line 2 (see ch. 2.1). When you call up a constant in function "C" (press "2nd" and "5") or the atomic mass unit in function "u" (press "2nd" and "6") the value and its unit appear in line 2 (see ch. 6.1 and 6.2).

Calculator  $N_A = 6.02252e+23 \text{ mol}^{-1}$ 

Abbreviation before " = ": Symbol of a constant (here N<sub>A</sub>) or "u" for the atomic mass unit.

### e) Exactness of input / indication of results in line 2

There are six digits disposable for the input / display of numbers in line 2 (up to seven digits for values x, for which is valid: -1 < x < 1).

For numbers with absolute values (magnitudes) > 999999 or < 0.00001 the input or the indication of results is performed by expressing these numbers as powers of 10 (from  $10^{37}$  up to  $10^{437}$ ). The input of such values is done with the function "exp" (see ch. 8.2).

### 3.1.3 Top Line

# Top line → ← RM 2nd □ → GeD TGTTTGCCGTGTGATTGG

The left side of the top line contains the following symbols:

Arrow to the left (alone or with another arrow on the right side):
If this sign appears, it is possible to scroll the display to the left: The data input reaches beyond the left side of the display (see ch. 3.3).

RM Symbol "Read Memory": It appears, if the keys

- $\bullet\,$  "RM" (Read Memory; calling up data from a memory place) oder
- "2nd" and "RM"(second function of "M" [memory] is activated) are pressed (memory functions: see ch. 9).

2nd Symbol "Second Function": It appears, if the key "2nd" has been pressed and the second functions are activated (see ch. 1.3).

The right side of the top line contains the following symbols:

Battery symbol: not active in this version (see. ch. 10.1).

Arrow to the right (alone or with another arrow on the left side):

If this sign appears, it is possible to scroll the display to the right: The data input reaches beyond the right side of the display (see ch. 3.3).

### 3.2 DELETING

3.2.1 Key "clr" (clear)



- a) Stoichiometric Functions
- a1) Correction of Compound Formulas

In the menus M1, So1, C1-C3, P1-P2 a compound formula can be corrected during the data input.

I.e. before pressing the key " = ". Afterwards the element symbols and subscripts put in are treated as a compound formula and get processed).



Pressing the key "clr" deletes the last part of a compound formula during the input:

- the complete subscript (stoichiometric number) or
- all letters of an element symbol.

Remarks: Pressing an element key in the calculator mode or in the menus M2-M4 causes the device to chance automatically to menu M1. The input of compound formulas is not possible in the menus: T, D und F.

### a2)

### Deleting whole compound formulas and molar masses

There are two ways of deleting the complete and processed input of a compound formula (i.e. after pressing the key " = "):



- Press key "clr" (only possible in menu M1).
  - The molar mass (indicated in line 2) is deleted together with the compound formula (indicated in line 1).
- Write a new formula over the old one and press key " = ".

Afterwards this new compound formula will be the base for further calculations in all stoichiometric menus.

Deleting a whole compound formula with the key "clr" is only possible in menu M1, but not in the menus So1, C1–C3, P1–P2.

Instead of changing to menu M1 from these menus it is advisable to directly replace the compound fomula.

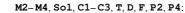
Deleting a whole input is also possible concerning:

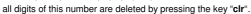
- Compound formulas called up from a memory space (see. ch. 9).
- The input of molar masses in line 2 without the input of a compound formula in line 1 (compound formula unknown; see ch. 5.2, 3c).
   This kind of input is only possible in menu M1.

### a3)

### Deleting the input of numbers

After the input of a number in the menus:





In the menu group P this function is only activated in menu P2 and P4 for the input of a substance's amount in "g". Other values are indicated automatically.



### b) Menus for Molecular Biology

### b1) Menus GeD / GeR (input of oligonucleotides)



Pressing this key deletes

- during the input of a sequence: the last nucleotide put in (line 1).
   After deleting the third nucleotide of a codon the three-letter-code of the corresponding amino acid (line 2) is deleted too.
- the whole oligonucleotide after input and processing, i.e. after pressing the key " = " (see ch. 5.18 and 5.19).
- the whole oligonucleotide called up by a memory function (see ch. 9).

The input of oligonucleotides in GeD / GeR (or oligonucleotides called up by a memory function) are taken over automatically into the menus Ge1-Ge5. In these menus it is not possible to delete them by pressing the key "clr". For this you must change to menu GeD / GeR.

### b2) Submenus Ge2-Ge5 (of GeD and GeR):



After the input of a number pressing this key deletes

• all digits of this number.

Submenus G1 and G6: The key "clr" is not active / needed.

### c) Calculator mode



After the input of a number pressing this key deletes

• all digits of this number.

### d) Other remarks concerning the key "clr"

Deleting a whole compound formula or oligonucleotide with the key "clr" removes these inputs also from the automatic memory function. After switching off the device respectively pressing the key "on" during use these inputs are lost (see. Kap. 9.1).

In some menus (e.g.  ${\tt Lib}$ ) or functions (e.g. Call-up of constants, see ch. 6.1) this key is not active.

### 3.2.2 Key "on" (master -clear-key)



Pressing the key "on" in any menu causes the device to set back to the calculator mode (see ch. 2.1). In line 2 zero is signalling.



The last input made in a menu (i.e. compound formulas in stoichiometric menus and oligonucleotides in menus for molecular biology) are always memorized automatically. They are displayed again when entering a menu.

This is also the case, if the device has been turned off in the meanwhile or if the key "on" has been pressed during use.

### 3.3 SCROLLING THE DISPLAY

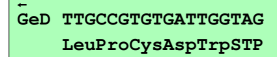
Inputs made in a menu, i.e.

- Compound formulas in the stoichiometric functions,
- Oligonucleotides menus for molecular biology,

may also reach beyond the left or right frame of the display.

Arrows at the left or the right side of the display's top line indicate, that this is the case:

← Arrow to the left: The input reaches beyond the display's left frame.





By pressing this key ("Scroll left") the legible part of the whole input is moved for one unit of the input (one nucleotide, symbol of an element or digit of a subscript [stoichiometric number]) to the left.

(The underlined part is also visible in the next illustration).

Example: Starting with the last display picture (page before) you get the following display after having six times pressed the key "Scroll left" (The display moves for six nucleotides to the left):

Arrow to the right: The input reaches beyond the display's right frame:

# GeD ATGTGTTTGCCGTGTGAT STACysLeuProCysAsp

The underlined part of the nucleotide sequence markes the part, that was also visible in the last display.

By pressing this key ("Scroll right") the legible part of the whole input is moved for one unit of the input (one nucleotide, symbol of an element or digit of a subscript [stoichiometric number]) to the right.

> Example: Starting with the last display picture (page before) you get the following display after having three times pressed the key "Scroll right" (The display moves for three nucleotides to the left):

Arrows at the left and right frame of the top line:

The input reaches on both sides beyond the frame of the display:

# GeD TGTTTGCCGTGTGATTGG CysLeuProCysAspTrp

The underlined part of the nucleotide sequence markes the part, that was also visi-

Analogous to the explanations above scrolling to the left or to the right is possible now.

### 4 SHORT INSTRUCTIONS FOR MENU CONTROL

**Preliminaries:** It is advisable to study ch. 1–3 before using this device for the first time. Afterwards these short instructions can serve as recollecting aid for the main points of operation (see backcover of the manual). The single menus are self-explanatory to a high degree. Details and variations of menu control, especially within the menus and further informations (e.g. formulas used) are delivered in chapter 5.

- a) On Press "on": The device is in the calculator mode (Calculator).
- b) Press "Cm" or an element key (e.g. Na 11): Change to menu M1.
- c) Press "Cm" for entering the following menus / menu groups.
- d) Sm "Sm": If there is a number behind a menu's abbreviation (or in GeD / GeR) submenus of this menu group are entered with the key "Sm":

(→)		Compounds / Analysis					0	ligo	os
Calculator →	→ <u>sol</u> →	$\rightarrow$ <u>C1</u> $\rightarrow$ T $\rightarrow$ D $\rightarrow$ F $\rightarrow$ <u>P1</u>			→ Lib →	GeD	÷	GeR	
Û	Ψ.		4		Ψ.	1	4		4
Press one time	M2		C2	Explanation	P2	Press-	Ge1		Ge1
Û	4		4	of the menus	4	element	4		
	<u>M3</u>		<u>C3</u>	abbreviations:	P3	key and	Ge2		Ge2
(Sm) ( <b>V</b> )	4			contents	4	scroll	4		4
	<u>M4</u>			(pp. 4 and 5)	P4		Ge3		Ge3
Input of compound formulas (possible in underlined menus): Change between						4			
element keys and numerical keys (for subscripts [stoichiometric numbers]).									
<ul> <li>Input of nucleotide s</li> </ul>	eque	ences (oligo	s) in	GeD / GeR: Press th	ne ke	ys with the	4		
second functions A, T	/ <b>U</b> ,	G, C, which	are a	automatically active i	n the	se menus.	Ge6		Ge6

- e) (E) Key " = ": Moving forward within a menu (new input, indicating a result).
- clr "clr": 1) Correcting inputs (elements, subscripts, nucleotides, numbers).
   2) Deleting whole compound formulas/oligonucleotides, that have been put in completely (or called up by a memory function) and are processed.
  - Scrolling: 1) If inputs reach beyond the display's frame (an arrow appears on the left or/and the right side of the display's top line), it is possible to scroll left or right by pressing these keys. 2) Navigating through Lib / Ge6.

### **MENU DESCRIPTIONS** – DETAILED INSTRUCTIONS

### 5.1 PRELIMINARIES

5

 a) The order of descriptions corresponds to the order of menus pressing the keys "Cm" and "Sm" (see graphic on p. 20 a. ch. 2.2 a. ch. 2.3).

If a menu consists out of a group of several submenus, all submenus are dealed with in the order, in which these submenus are entered by pressing the key "Sm".

All steps within a calculation are described in tables. Alternatives for the input of data are also listed.

- b) At the end of a menu's / submenu's description there are additional remarks and explanations for understanding how the menu is operating:
  - Values and formulas, that are used by the device for operations;
  - Possibilities of further processing with displayed results;
  - · Other comments.

### c) Symbolizing signalling values (underlining):

- If inputs are required in the display: Zero signals.
- If values are proposed (e.g. in menu C3 for densities): The value signals. In graphics signalling numbers are symbolized by underlining!
- d) Automatic memory function: The last inputs, i.e.
  - compound formulas in stoichiometric menus (input possible in M1, So1, C1-C3, P1-P2) and
  - nucleotide sequences in the menus GeD und GeR,

are automatically memorized. They are displayed when a menu is entered again and serve as base for further calculations.

This is not the case, if the last compound formula/nucleotide sequence put in has been deleted with the key "clr" before

- turning off the calculator or
- pressing the key "on" during use (master-clear-key).

# 5.2 M1 MOLAR MASS OF A SUBSTANCE

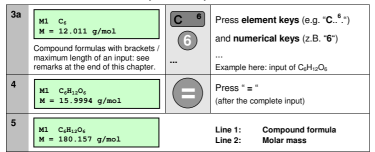
Display	Keys	Actions / Comments
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### Enter Menu M1

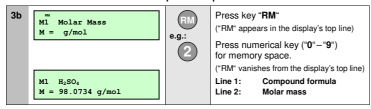
1a	Any display	Cm	Press (repeatedly) "Cm" (Change to the next menu) until menu M1 appears Continue with 2a or 2b
Alte	rnatives:		
1b	Any display	2nd Cm	Press "2nd" and "Cm" until menu M1 appears. The menus are entered in the opposite order. Continue with 2a or 2b
1c	Calculator mode  Calculator  0	e.g.:	Press an element key, e.g. "C 6" Continue with 3a

Disp	Display 2a or 2b appears					
2a	M1 Malau Mana	Zero is signalling.				
	M1 Molar Mass $M = 0 g/mol$	The last compound formula has been deleted with "cIr"; see. ch. 9.1.				
		Continue with 3a, 3b or 3c				
2b	M1 NH <sub>3</sub> M = 17.0304 g/mol	The last compound formula input appears (automatic memory call-up; see. ch. 8.1) – example here: <b>NH</b> <sub>3</sub> .				
If you want to continue with the automatically called-up compound for the transport of transpo						
	<ul> <li>If you want to continue with another compound formula:</li> <li>Replace it by writing over the old formula.</li> <li>Continue with step 3a</li> </ul>					

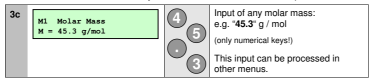
### Alternative of continuation 1: Input of a compound formula



# Alternative of continuation 2: Call-up of a compound formula



### Alternative of continuation 3: Input of a molar mass without a compound formula



### Remarks

### Step 3c

This possibility of input serves for calculations with substances, whose molar
mass is known, but not the parts respectively the percentage of the parts.

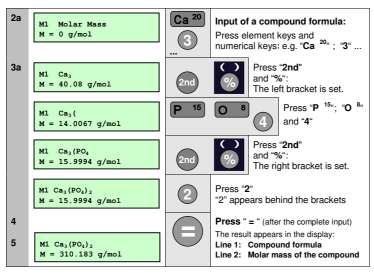
### Step 3a

- After each pressing of an element key appears the molar mass of the represented element in line 2 of the display. This value does not change after the input of a subscript (stoichiometric number) greater than "1". No provisional results are indicated in line 2. Only after pressing the key " = " (4) the molar mass of the whole compound is indicated (5).
- The molar masses of elements displayed are average values of the naturally occuring elements (natural isotope mixtures). Molar masses of isotopes cannot be called up.

### **Bracket Function**

The brackets (second function of the key "%") can be used for the input of compound formulas with brackets (only for one level of brackets, input of interlocked brackets is not possible).

Example:  $Ca_3(PO_4)_2$  (Calciumphosphate):



The bracket function is not disposable in the calculator mode. It can only be used for the input of compound formulas in the stoichiometric menus.

### Maximum Length for the Input of Compounds

The input of compounds may constist of a maximum of 20 elements and their subscripts (stoichiometric numbers). If the compound formula reaches beyond the length of the display, it is possible to scroll (scroll function: see page 18 ch. 3.3).

### Continuation with Results in Other Menus / in the Calculator Mode

- The compound formula (5 / 3b) respectively the input of a molar mass (3c) are automatically taken over into other stoichiometric menus (M2-M4, So1, C1-C3, P1-P4). There these data are used as the base for further calculations.
- In these menus the data taken over can be replaced by writing over with a new compound formula (input as described in 3a 5).
   Inputs according to step 3c (molar mass without compound formula) cannot be replaced in other stoichiometric menus. Only the input of a compound formula is possible there. For the input of a molar mass according to step 3c you must again enter menu M1. Then the new value can be taken over again into other menus.
- It is possible to perform further calculations with a result (5) in the calculator mode.
   After pressing an operation key in the right part of the keyboard (e.g. pressing the key " x ") the device changes automatically into the calculator mode.

### Symbolizing of Signalling Numbers (Underlining):

A zero is signalling, if the input of numbers is required. In the illustrations signalling of a number is marked by underlining: " $\underline{0}$ ".

### **Automatic Memory Function**

The last input of a compound formula is automatically memorized. It is again disposable when entering a stoichiometric menu and serves as base for further calculations. This is not the case, if the last input of a compound formula has been deleted with the key "clr" before turning off the device or pressing the key "on" (master-clear-key) during use.

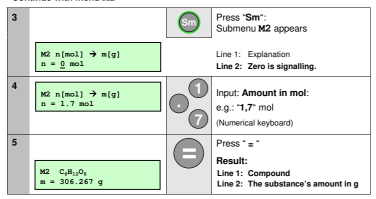
# 5.3 M2 CONVERTING A SUBSTANCE'S AMOUNT: mol into g

	Display	Keys	Actions / Comments
1	Any display	Cm	Press (repeatedly) "Cm" (Change to the next menu) until menu M1 appears

Continue with menu M1: Take over or put in a compound formula (molar mass)

2	M1 $C_6H_{12}O_6$ M = 180.157 g/mol	Example here: Display of the last input of a compound formula (see p. 22, menu M1, 2b)						
Input	Alternatively a zero might signal in line 2 (see p. 22, menu M1, 2a). Input respectively take-over or change of compound formulas / molar masses: see p. 22 f. menü M1, 3a – 3c.							

### Continue with menu M2



- 1) It is possible to perform further calculations with a result (5) in the calculator mode.
- 2) The result can be memorized if needed (see ch. 8.2).

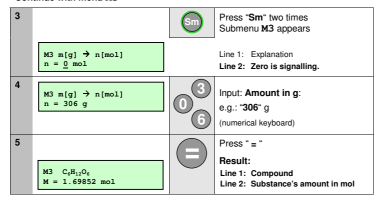
### 5.4 M3 CONVERTING A SUBSTANCE'S AMOUNT: g into mol

	Display	Keys	Actions / Comments
1	Any display	Cm	Press (repeatedly) "Cm" (change to the next menu) until menu M1 appears

### Continuation with menu M1: Take-over or input of a compound formula (molar mass)

2	M1 $C_6H_{12}O_6$ M = 180.157 g/mol	Example here: Display of the last input of a compound formula (see p. 22, menu M1, 2b)			
Input	Alternatively a zero might signal in line 2 (see p. 22, menu <b>M1</b> , <b>2a</b> ). Input respectively take-over or change of compound formulas / molar masses: see p. 22 f. menü <b>M1</b> , <b>3a</b> – <b>3c</b> .				

### Continue with menu M3



- 1) It is possible to perform further calculations with a result (5) in the calculator mode.
- 2) The result can be memorized if needed (see ch. 8.2).

# 5.5 M4 IDEAL GAS VOLUME OF A SUBSTANCE

	Display	Keys	Actions / Comments
1	Any display	Cm	Press (repeatedly) "Cm" (Change to the next menu) until menu M1 appears

Continue with menu M1: Take-over or input of a compound formula (molar mass)

2	M1 $C_6H_{12}O_6$ M = 180.157 g/mol	Example here: Display of the last input of a compound formula (see p. 22, menu <b>M1</b> , <b>2b</b> )		
Input	Alternatively a zero might signal in line 2 (see p. 22, menu M1, 2a). Input respectively take-over or change of compound formulas / molar masses: see p. 22 f. menu M1, 3a – 3c.			

### Input of the substance's amount in submenu M2 or M3

3		Sm	Press "Sm" once or twice Change to submenü M2 or M3
4 – 5	Now you have to put in the subs (in grammes) as described in ch		ount in menu <b>M2</b> (in moles) or <b>M3</b> (step <b>4</b> – <b>5</b> ). See p. 26 or 27.
		T	
6	Result:	Sm	Press "Sm" (1 x or 2 x) Change to submenu M4
	M4 n[mol] → V[1] V = 44.8 1		Line 1: Explanation Line 2: Ideal gas volume
	e.g. input of "C <sub>6</sub> H <sub>12</sub> O <sub>6</sub> " in <b>M1</b> and of "360 314" g in <b>M3</b>		of the substance according to the input in step 4 - 5

- 1) It is possible to perform further calculations with a result (6) in the calculator mode.
- 2) The result can be memorized if needed (see ch. 8.2).
- 3) Formula:  $V_m = n \mod \bullet V_{mol} \text{standard conditions presupposed};$  Ideal gas volume of the substance X (in: I) = amount of X (in: mol)  $\bullet$  mole volume ( $\approx 22,4 \text{ I/mol}$ )

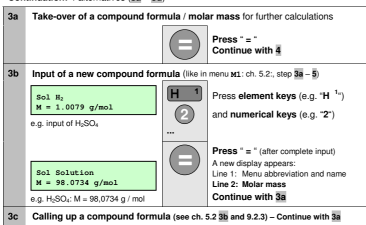
### 5.6 Sol SOLUTION: Mass of a solute in g

	Display	Keys	Actions / Comments
1	Any display	Cm	Press (repeatedly) "Cm" (change to the next menu) until menu SOL appears

### Continue with menu So1 (The display indicates 2a or 2b)

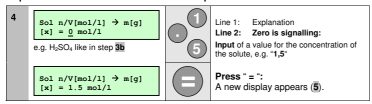
2a	Sol Solution M = 0 g/mol	Line 1: Line 2:	Abbreviation and name of the menu  0 g / mol; the last input of a compound formula / molar mass has been deleted with "clr" – Continue with step 3b
2b	Sol Solution M = 180.157 g/mol	Line 1: Line 2:	Abbreviation and name of the menu  Molar mass of the last input (substance)
	e.g. molar mass of C <sub>6</sub> H <sub>12</sub> O <sub>6</sub>	-	Continue with 3a

### Continuation: 4 alternatives (3a - 3d)

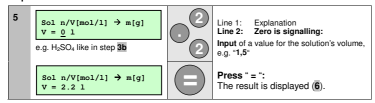


Input of molar mass without compound formula: Only possible in menu M1 (see ch. 5.2, 3c). Enter menu So1 again with key "Cm" — continue with step 3a.

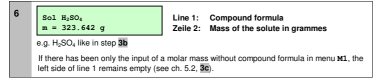
### Input of the solute's concentration in moles per volume



### Input of the solution's volume



### Result: The solute's mass



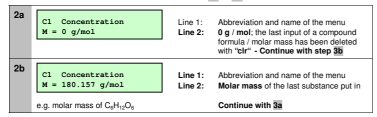
- 1) It is possible to perform further calculations with a result (6) in the calculator mode.
- 2) The result can be memorized if needed (see ch. 8.2).
- 3) Formula: m = M<sub>X</sub> c<sub>X</sub> × V needed mass (in: g) of a substance X (solute) = molar mass of X (in: g / mol) concentration of X (in: mol / I) × volume of solution (in: I)

# 5.7 C1 CONCENTRATION – CONVERSION:

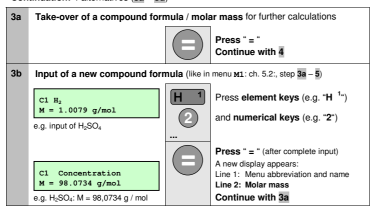
Mass concentration (g / I) into moles per volume (mol / I)

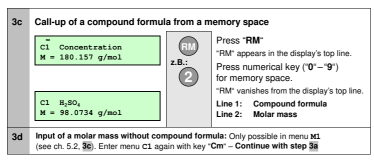
	Display	Keys	Actions / Comments
1	Any display	Cm	Press (repeatedly) "Cm" (change to the next menu) until menu C1 appears.

### Continue with menu C1 (the display indicates 2a or 2b)

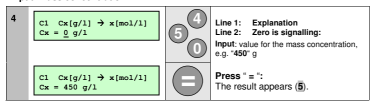


### Continuation: 4 alternatives (3a - 3d)

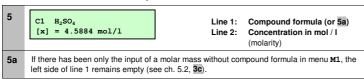




### Input: Mass concentration



### Result: Concentration in moles per volume



### Remarks

- 1) It is possible to perform further calculations with a result  $(\mathbf{5})$  in the calculator mode.
- 3) The result can be memorized if needed (see ch. 8.2).
- 4) Formula:  $[x] = c_X / M_X$

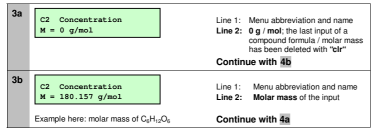
Concentration in moles per volume (molarity) of a substance X (in: mol / l) = mass concentration of X (in: g / l) / molar mass of X (in: g / mol)

## 5.8 C2 CONCENTRATION – CONVERSION:

Moles per volume (mol / I) into mass concentration (g / I)

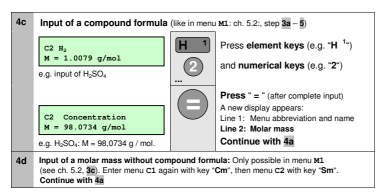
	Display	Keys	Actions / Comments						
Ente	Enter menu C2								
1	Any display	Cm	Press (repeatedly) "Cm" (change to the next menu) until menu C1 appears.						
	C1 Concentration M = 0 g/mol		Line 1: Menu abbreviation and name Line 2: Here "0 g / mol" (see 3a) A value for a molar mass could also be indicated (see 3b).						
2		Sm	Press "Sm" (1 ×): Submenu C2 appears						

## Continue with menu C2 (The display indicates 3a oder 3b)

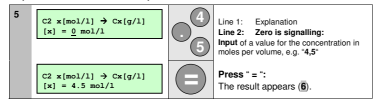


## Continuation: 4 alternatives (4a - 4d)

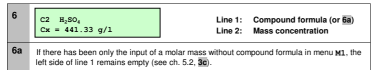
4a	Take-over of compound formula / molar mass for further calculations						
	Press " = " Continue with 5						
4b	Calling up a compound formula (see ch. 5.2 3b and 9.2.3) - Continue with 4a						



## Input: Concentration in moles per volume



## Result: Mass concentration (example here: H<sub>2</sub>SO<sub>4</sub>)

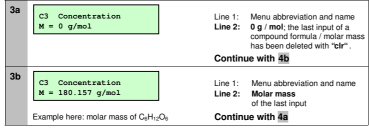


### Remarks

- 1) It is possible to perform further calculations with a result (6) in the calculator mode.
- 2) The result can be memorized if needed (see ch. 8.2).
- 3) Formula:  $c_X = M_X \bullet [x]$  mass concentration of substance X (in:  $g \mid I$ ) = molar mass of X (in:  $g \mid mol$ ) concentration in moles per volume (molarity) of X (in: mol  $\mid I$ )

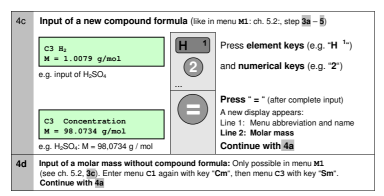
# 5.9 CONCENTRATION – CONVERSION: mass percentage into moles per volume (mol / I)

	Display	Keys	Actions / Comments					
Ente	er menu C3							
1	Any display	Cm	Press (repeatedly) "Cm" (change to the next menu) until menu C1 appears.					
	C1 Concentration M = 0 g/mol		Line 1: Menu abbreviation and name Line 2: Here: "0 g / mol" (see 3a); a value for a molar mass could also be indicated (see 3b).					
2		Sm	Press " <b>Sm</b> " (2 ×): Submenu <b>C3</b> appears					
Con	Continue with menu C3 (the display indicates 3a or 3b)							

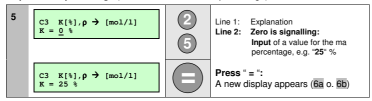


## Continuation: 4 Alternatives (4a - 4d)

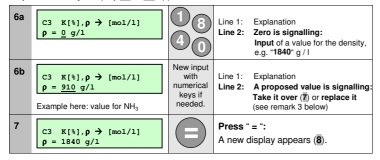
4a	Take-over of compound formula / molar mass for further calculations						
	Press " = " Continue with 5						
4b	Call up of a compound formula (ch. 5.2 3b and 9.2.3) – Continue with 4a						



## Input: Mass percentage (see remarks for volume percentage)



## Input of density: Display 6a or 6b appears



## Result: Concentration in moles per volume

8 C3 H<sub>2</sub>SO<sub>4</sub> Line 1: Compound formula (or 8a) Line 2: Concentration in mol/I

8a If there has been only the input of a molar mass without compound formula in menu M1, the left side of line 1 remains empty (see ch. 5.2, 3c).

## Remarks

- 1) It is possible to perform further calculations with a result (8) in the calculator mode.
- 2) The result can be memorized if needed (see ch. 8.2).
- 3) Densities are proposed for the following acids, bases and solutions (6b), that are listed in commonly available concentrations:

Name	Compound formula	concentration in mol / I (rounded)	mass percentage	Density g / cm <sup>3</sup> 20°/4°
Sulphuric acid	H <sub>2</sub> SO <sub>4</sub>	36	95-97	1,84
Formic acid / H-COOH	H <sub>2</sub> CO <sub>2</sub>	26	98 - 100	1,22
Hydrobromic acid	HBr	7	40	1,38
Acetic acid / CH <sub>3</sub> -COOH	H <sub>4</sub> C <sub>2</sub> O <sub>2</sub>	18	99 - 100	1,06
Hydriodic acid	HI	7,5	57	1,70
Phosporic acid (concentrated)	H <sub>3</sub> PO <sub>4</sub>	15	85	1,71
Nitric acid (concentrated)	HNO <sub>3</sub>	14	65	1,40
Hydrochloric acid (concentrated)	HCI	12	36	1,18
Ammonia (aqueous)	NH <sub>3</sub>	13,5	25	0,91
Potassium hydroxide solution	кон	7	30	1,30
Sodium hydroxide solution	NaOH	11	33	1,36

## 5) Formula: $[x] = K \cdot \rho / M_X$

Concentration in moles per volume (molarity) (in: mol / I) of a substance X = mass percentage (here defined as "K") • density of X (in: g / I) / molar mass of X (in: g / I)

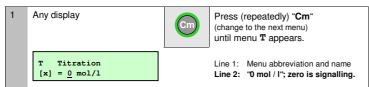
## 6) Approximate calculation of volume percentage

This is possible with the formula (4) above, if the density of a substance is known for a mass percentage of 100.

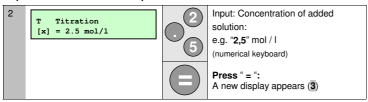
## 5.10 T TITRATION

Display	Keys	Actions / Comments
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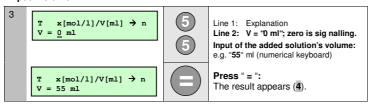
## Enter Menu T



## Input: Concentration in moles per volume



## Input: Volume



## Result: Amount of the added solution in mmol



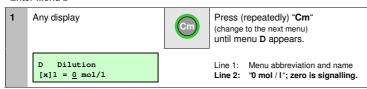
## Remarks

- 1) It is possible to perform further calculations with a result (4) in the calculator mode.
- 2) The result can be memorized if needed (see ch. 8.2).
- Formula: n = [x] V
   amount of the (added) substance X (in mmol) = concentration in moles per volume of X (in mol / I) Volumen (in: ml)
- 4) Conversion of the substance's amount into grammes :
  - Convert result (4) into moles (• 1000) and
  - Multiplicate this result with the molar mass of the substance.

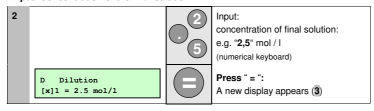
## 5.11 D DILUTION

Display	Kevs	Actions / Comments
Display	itcys	Actions / Comments

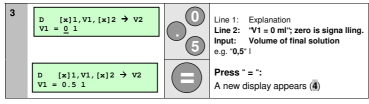
## Enter Menu D



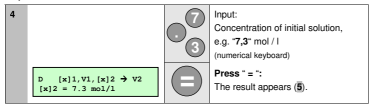
## Input: Concentration of the final solution



## Input: Volume of the final solution



## Input: Concentration of the initial solution



### Result: Initial volume in I

D [x]1,V1,[x]2  $\rightarrow$  V2 V2 = 0.171233 1 Line 1: Explanation
Line 2: Volume of initial solution Example here: "0,171233 I"

- $\bullet~$  You have to pipet 171 ml of an initial solution with a concentration of 7,3 mol / l.
- Fill this amount up with the solvens until a volume of 0,5 l is reached.
  Now you have a final solution with a concentration of 2,5 mol / l.

## Remarks

- 1) It is possible to perform further calculations with a result (5) in the calculator mode.
- 2) The result can be memorized if needed (see ch. 8.2).
- 3) Formula:  $V_2 = [x]_1 \cdot V_1 / [x]_2$

 $V_2$  = volume to be pipetted (initial volume) of a substance X (in: I)

 $[X]_1$  = desired concentration of X in the final solution (in: g / mol)

 $V_1$  = desired volume of the final solution in which X is the solute (in: I)

 $[X]_2$  = concentration of the initial solution with X as solute (g / mol)

## 5.12 F FORMULA – CALCULATING A COMPOUND FORMULA

Display	Keys	Actions / Comments
---------	------	--------------------

## Prepare the necessary values

- The following data have to be provided in order to be able to calculate the compound formula of a substance in this menu:
  - a) Mass percentage of the elements;
  - b) Molar mass of the substance and derived by it:
    - b1) The ideal gas volume of the substance at a specific ...
    - b2) ... mass of the compound (any mass could be chosen).

Provided that the substance's molar mass is known, it is possilbe to calculate the ideal gas volume corresponding to any mass in  ${\bf M4}$ .

### Procedure:

- Input of the molar mass in M1: Entering menu M1 by pressing the key "Cm" (repeatedly) until menu M1 appears. A compound formula possibly indicated automatically by the memory function has to be eliminated by pressing the button "clr". Now put in the molar mass with the numerical keys (see ch.5.2, step 3c, page. 22).
- Change to subenu M3 (press key "Sm" twice): put in any mass (note down this mass); press key " = " (see ch. 5.4 on page 25).
- Change to submenu M4 (press key "Sm" once): the ideal gas volume corresponding with the mass put in is indicated (see ch. 5.5 on S. 26).

Our examples: a) C: 52 %; H: 13 %; O: 35 %

b1) 4,86 l b2) 10 g

## Enter menu F

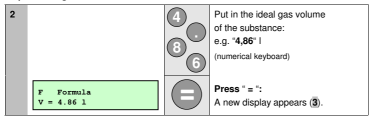
Any display

Press (repeatedly) "Cm"
(Change to the next menu)
until menu F appears.

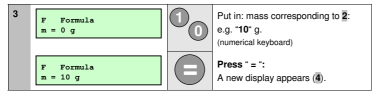
F Formula
V = 0 1

Line 1: Menu abbreviation and name
Line 2: "V = 0 I"; zero is signalling

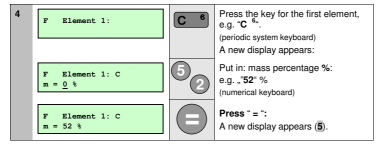
## Input: Ideal gas volume of the substance



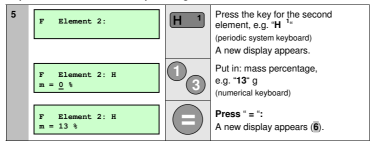
Input: Mass corresponding to the ideal gas volume



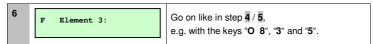
## Input: First element and its percentage



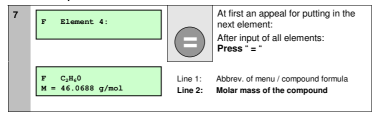
## Input: Second element and its percentage



## Input: Third element and others



## **Result: Compound formula**



### Remarks

- 1) The result can be memorized if needed (see ch. 8.2).
- 2) The order of the elements in the compound formula displayed depends upon the order of the elements during the input. The calculator does not arrange the elements according to the rules usually applied to formulas in chemistry. These rules should already be obeyed during the input.

## 3) Formula:

Starting with the relative atom masses of the substance's elements it is possible to calculate the molar mass of an unit X with entire stoichiometric numbers (Indices):

Element:	m(%) <sub>element</sub>	1	M <sub>element</sub>	=	relative number of atoms	
Our examp	le:					Index
С	m(%) <sub>C</sub> <b>52</b> %	1	<i>M</i> <sub>C</sub> 12	=	relative number of atoms: C 4,33	4,33 / 2,18 1,98 ≈ <b>2</b>
н	<i>т(%)</i> <sub>Н</sub> <b>13 %</b>	/	<i>M</i> <sub>H</sub> <b>1</b>	=	relative number of atoms: H	13 / 2,18 5,96 ≈ <b>6</b>
0	m(%)₀ <b>35</b> %	1	<i>М</i> о 16	=	relative number of atoms: O 2,18	2,18 / 2,18 <b>1</b>

The lowest relative number of atoms 2,18 corresponds with the value 1 for an index. The indices of the other elements can now be calculated by dividing their relative atom numbers with 2,18. The results are rounded off.

Thus the molar mass of an unit X can be calculated:

$$(2 \cdot 12 g/mol) + (6 \cdot 1 g/mol) + (1 \cdot 16 g/mol) = 46 g/mol$$

The ratio of the stoichiometric number corresponds already to the ratio of the formula searched for. You will, however, get the absolute values of the true indices only by multiplication with a factor n.

n = molar mass of compound / molar mass of the unit X

Our example:

46 g / mol / 46 g / mol = 1

In this case the stoichiometric numbers of the unit X and those of the compound formula searched for are already the same (multiplication with factor n=1).

Another example:

Molar mass of the unit X:  $CH_2O$  = 30 g / mol. Molar mass of the compound = 180 g / mol

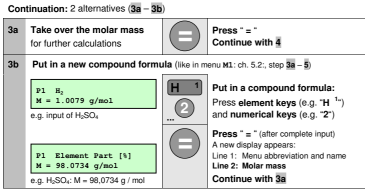
180 g / mol / 30 g / mol = 6

According to this the compound formula is:  $C_{1 \cdot 6} H_{2 \cdot 6} O_{1 \cdot 6} = C_6 H_{12} O_6$ 

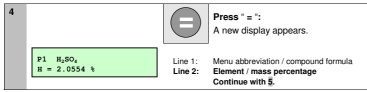
### P1 ELEMENT PARTITION 1: 5.13

Parts of a compound - each element in %

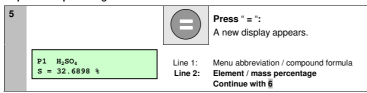
	Display	Keys	Actions / Comments				
Ente	er Menu P1						
1	Any display	Cm	Press (repeatedly) "Cm" (change to the next menu) until menu P1 appears.				
Continue with P1 (display 2a or 2b appears)							
2a	P1 Element Part [%] M = 0 g/mol	Line 1: Line 2:	Menu abbreviation and name 0 g / mol; the last input of a compound formula / molar mass has been deleted with "clr" – Continue with 3b				
2b	P1 Element Part [%] M = 180.157 g/mol e.g. molar mass of CoHtoOc	Line 1: Line 2:	Menu abbreviation and name Molar mass of the last input (substance) Continue with 3a or 3b				



## Input: Mass percentage of the first element



## Input: Mass percentage of the second element



## If necessary repeat step 5 (until all elements of the compound have been passed)



## Display: Molar mass like in 2b or 3b

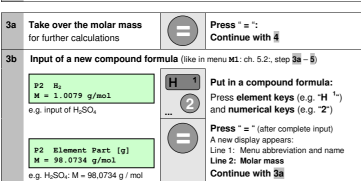


Formula:	<i>M</i> <sub>element</sub>	•	index	/	$M_{\rm compound}$	=	mass percentage
Our example:	Мн	•	index	/	MH <sub>2</sub> SO <sub>4</sub>	=	mass percent. of <b>H</b>
H in: H <sub>2</sub> SO <sub>4</sub>	1,008 g / mol	•	2	/	98,07 g / mol	=	0,02055 = 2,055 %

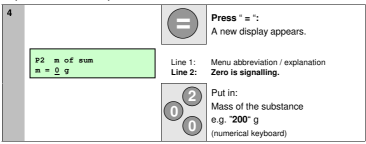
## 5.14 P2 ELEMENT PARTITION 2:

Parts of a compound – each element in g

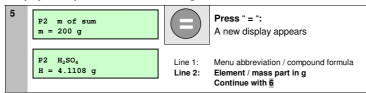
	Display	Keys	Actions / Comments					
Ente	er menu P2							
1	Any display	Cm	Press"Cm" (repeatedly) until menu P1 appears.					
	P1 Element Part [%] M = 0 g/mol	Sm	Press "Sm": Menu P2 appears (2a or 2b).					
Con	Continue with menu P2 (display 2a or 2b appears)							
2a	P2 Element Part [g] M = 0 g/mol	Line 2:	Menu abbreviation and name 0 g / mol; compound formula / molar mass deleted with "cir" – Continue with 3b					
2b	P2 Element Part [g] M = 180.157 g/mol	Line 1: Line 2:	Menu abbreviation and name  Molar mass of the lest input (substance)					
	e.g. molar mass of C <sub>6</sub> H <sub>12</sub> O <sub>6</sub>		of the last input (substance)  Continue with 3a or 3b					
3a	Take over the molar mass		Press " = ":					



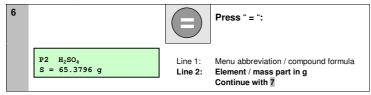
## Input: Mass of the compound



## Display: Mass part of the first element in g



## Display: Mass part of the second element in g



If necessary repeat step 6 (until all elements of the compound have been passed)



Display: Molar Mass like in 2b or 3b



## Formula 1

Mass parts of the elements expressed in parts of 1:

Formula:	<i>M</i> <sub>element</sub>	•	index	/	$M_{\rm compound}$	=	mass part
Our example:	Мн	•	index	/	<i>M</i> H₂SO <sub>4</sub>	=	mass part of <b>H</b>
H in H <sub>2</sub> SO <sub>4</sub>	1,008 g / mol	•	2	/	98,07 g / mol	=	0,02055

Taken for granted a certain mass of the compound in g (e.g. 200g) the mass part of an element in g can be calculated as follows:

Formula:	mass part in part of 1	•	mass in g (compound)	=	mass part in g
Our example:	part of <b>H</b>		H <sub>2</sub> SO <sub>4</sub>		part of <b>H</b> in g
H in H <sub>2</sub> SO <sub>4</sub>	0,02055	•	200 g	=	4,11 g

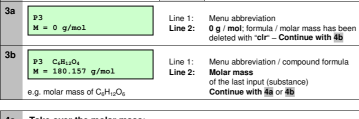
Thus the part of H in 200g of  $H_2SO_4$  is: 4,11 g (see.  $\overline{\textbf{5}}$ ).

<sup>&</sup>lt;sup>1</sup> In the tables: M = molar mass

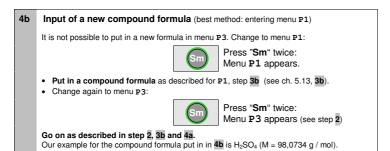
## 5.15 P3 FORMULA PARTITION 3:

Parts of a compound – several elements in %

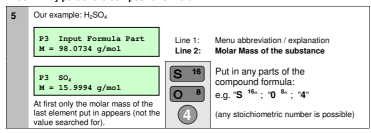
	Display	Keys	Actions / Comments
Ente	er menu P3		
1	Any display	Cm	Press"Cm" (repeatedly) until menu P1 appears.
	P1 Formula Part [%] M = 0 g/mol	Sm	Press "Sm" twice: Menu P3 appears (2).
Con	tinue with menu P3		
2	P3 Formula Part [%]		Line 1: Menu abbreviation and name Line 2: Nothing
			Press " = " Continue with 3a or 3b



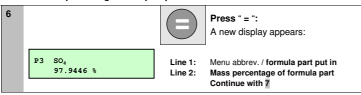




## Put in: Any part of the compound formula



## Result: Mass percentage of the part put in



## Repetition of steps 3b until 6 (if necessary with new values)



## Remarks

Step 5) It is possible to put in entire parts of the stoichiometric numbers.

After having pressed an element key in line 2 appears only the molar mass of this element. This value does not change after the input of a stoichiometric number. The input, however, is processed correctly.

## Formula

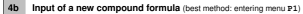
M <sub>element1</sub>	•	index	+	M <sub>element 2</sub>	•	index	/	$M_{ m compound}$	=	mass percentage
Our example:										
Ms	•	index	+	Mo		index	/	Мн <sub>2</sub> so <sub>4</sub>	=	mass percent. SO <sub>4</sub>
<b>32,06</b> g / mol	•	1	+	<b>15,9994</b> g / mol	•	4	/	<b>98,07334</b> g / mol	=	0,97446 = <b>97,446</b> %
			<b>,05</b> / m				1	<b>98,07334</b> g / mol	=	0,97446 = <b>97,446</b> %

Thus the mass percentage of SO<sub>4</sub> in H<sub>2</sub>SO<sub>4</sub> is 97,446 % (see 6).

## 5.16 P4 FORMULA PARTITION 4:

Parts of a compound – several elements in g

	Display	Keys	Actions / Comments
Ente	er menu P1		
1	Any display	Cm	Press"Cm" (repeatedly) until menu P1 appears.
	P1 Element Part [%] M = 0 g/mol	Sm	Press "Sm" three times: Menu P4 appears (2).
Con	tinue with menu P4		
2	P4 Formula Part [g]		Line 1: Menu abbreviation and name Line 2: Nothing
			Press " = " Continue with 3a or 3b
•			
3a	P4 M = 0 g/mol	Line 1: Line 2:	Menu abbreviation 0 g / mol; formula / molar mass has been deleted with "clr" – continue with $4b$ .
3b	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Line 1: Line 2:	Menu abbreviation / compound formula Molar mass of the last input (substance) Continue with 4a or 4b.
4a	Take over the molar mass:		
			Press " = " Continue with 5



It is not possible to put in a new formula in menu P4. Change to menu P1:



Press "Sm" once: Menu P1 appears.

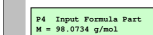
- Put in a compound formula as described for P1, step 3b (see ch. 5.13, 3b).
- Change again to menu P3:



Press "Sm" three times: Menu P1 appears.

Go on as described in step 2, 3b and 4a. Our example for the compound formula put in in 4b is  $\rm H_2SO_4$  (M = 98,0734 g / mol).

## Input: Any part of the compound formula



Our example: H<sub>2</sub>SO<sub>4</sub>:

Line 1: Line 2:

Menu abbreviation / explanation Molar Mass of the substance

P4 SO<sub>4</sub> M = 15.9994 g/mol At first only the molar mass of the last element put in appears.



Put in any parts of the compound formula: e.g. "S 16"; "0 8"; "4"

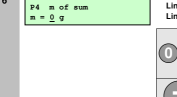
(any stoichiometric number is possible)



Press " = ":

A new display appears (6)

## Input: Mass of the compound



Line 1: Menu abbreviation / explanation Line 2: Zero is signalling.

Put in:

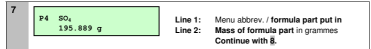


Mass of the compound e.g. "**200**" g (numerical keyboard)



The result appears (7).

## Result: Mass of part put in



## Repetition of step 3b to 6 (if necessary with new values)



## Remarks

Step 5) It is possible to put in entire parts of the stoichiometric numbers.

After having pressed an element key in line 2 appears only the molar mass of this element. This value does not change after the input of a stoichiometric number. The input, however, is processed correctly.

### Formula:

Calculation of the relative mass of the compound's part (our example:  $SO_4$  in  $H_2SO_4$ ):



Taken for granted a certain mass of the compound in g (e.g. 200 g) the mass of a part of this compound in g can be calculated as follows:

Formula:	relative mass of a part	•	mass in g (compound)	=	mass of part in g
Our example: SO <sub>4</sub> in H <sub>2</sub> SO <sub>4</sub>	rel. mass of SO <sub>4</sub> 0,97446	•	H₂SO₄ 200 g	=	mass of <b>SO<sub>4</sub></b> in g <b>195,89 g</b>

The part of  $SO_4$  in 200 g of  $H_2SO_4$  is: 195,89 g (see  $\overline{7}$ ).

### 5.17 Lib LIBRARY: DATA OF THE ELEMENTS

	Display	Keys	Actions / Comments				
Ente	ering menu Lib						
1	Any display	Cm	Press (repeatedly) "Cm" (change to the next menu) until menu Lib appears.				
	Lib Library	Line 1: Line 2:	<b>Menu abbreviation / menu name</b> Nothing				
2		H 1	Press an <b>element key</b> (e.g. "H 1"): A new display appears (1 <sup>st</sup> entry).				
	Lib Hydrogen year of discovery 1766	Line 1: Line 2:	Menu / name of the element Year of discovery: 1766				
Scro	Scrolling in the menu "Library"						
3	"Scroll left" (last entry)		Press Scroll-key: The last or the next entry is				

Line 1:

Line 2:

indicated in the display.

Menu / name of the element

Electronegativity after Pauling

The following entries are made for each element:

"Scroll right" (next entry)

Lib Hydrogen el.neg.(Paul.) 2.2

year of discovery

electronegativity after Pauling (in: electron volt) (in: gramm / cubic centimeter; gases in: g / dm³) boiling point (in: Kelvin) melting point (in: Kelvin) el.neg. [Paul.] (eV)

density (g / cm³) boiling pt (K) melting pt (K)

f.ion.pot. (eV) first ionization potential (in: electron volt)

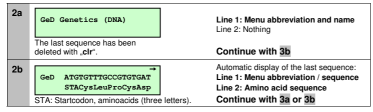
therm.conduct. thermic conductivity

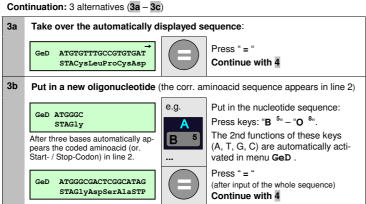
The order of this list corresponds with the order when calling up data with the key "Scroll right" (). Conversion of Kelvin into other temperature units: see ch. 7.

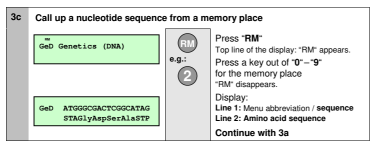
## GeD GENETICS (DNA) – Input of a Nucleotide Sequence Number of different nucleotides / percentage of GC

	Display	Keys	Actions / Comments
Ente	er menu GeD		
1	Any display	Cm	Press (repeatedly) "Cm" (change to the next menu) until menu GeD appears.

## Continue with menu GeD (display 2a or 2b appears)







## Display: Number of nucleotides of each type / percentage of GC

GeD ATGGGCGACTCGGCATAG

4A3T7G4C 61.11% GC

Line 1: Menu abbreviation and name

Line 2: Number of each nucleotide type

(symbolized by A, T, U, G, C),

percentage of GC

### Remarks

## Step 2b and 3b

## Scroll function

The input of a nucleotide sequence can reach beyond the display. This is indicated by arrows in the top line of the display at the left or right border. By using the scroll keys ( ) it is possible to move the displayed sequence. See ch. 3.3 with a detailled description.

## Display of start and stop codons and amino acids in line 2

Start- und stop codons are symbolized by capitals (STA, STP) whereas aminoacids that do not serve as start- or stop codon are symbolized by the three letter code with only the first letter beeing a capital (e.g. Cys, Leu, ...).

## Step 3b

## Maximal length of input possible:

The maximum length for the input of a oligonucleotide is 80 nucleotides.

## Further processing of sequences in menus Ge1 - Ge6

- The input of a nucleotide sequence in GeD oder GeR is taken over into menus Ge1 – Ge6 and remains the base for further calculations there.
- A data input in GeD or GeR is processed similar in Ge1 Ge6.
   Difference: Concerning key "C 6" (2nd function "T / U") thymine ("T") is activated in GeD, Uracil ("U") in GeR.



- The input of a sequence in **GeD** is also displayed in **GeR** and viceversa. Concerning display and calculations Thymine or Uracil replace each other.
- Only in GeD or GeR the input of a nucleotide sequence can be deleted by pressing the key "clr", not in the menus Ge1 – Ge6.

## 5.19 GeR GENETICS (RNA) – Input of a Nucleotide Sequence: Number of different nucleotides / percentage of GC %

Steps 1 - 4 follow exactly the schedule described in ch. 5.18 for menu **GeD**. All remarks concerning **GeD** (pages 59-60) are also valid for menu **GeR**.

T/II

The only differences are: • Instead of GeD the abbreviation GeR is used.



 Instead of Thymine ("T") in GeR Uracil ("U") is activated ("T / U" is 2nd function of key "C 6").

1	Any	display	Cm	Press (repeatedly) "Cm" until menu GeR appears.			
2a	GeF	Genetics (RNA)		The last sequence has been deleted by pressing the key "clr".			
2b	The last input of a sequence is automatically displayed (automatic memory)						
3	a) b) c)	b) Put in a new sequence					

4 Display: Number of each nucleotide types / percentage of GC

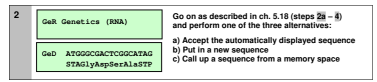
## 5.20 Ge1 MOLAR MASS (Nucleotide Strand, Coded Protein)

Display	Kevs	Actions / Comments
Display	itcys	Actions / Comments

## Enter menu GeD or GeR

1 Any display Press (repeatedly) "Cm" until menu GeD or GeR appears.

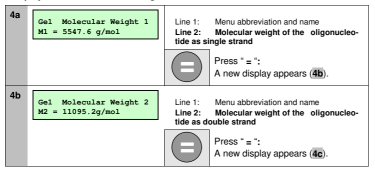
### Continue with menu GeD or GeR



## Enter Menü Ge1



## Display: Different molecular weights



4b Gel Molecular Weight 3 M3 = 461.475 g/mol

Line 1: Line 2: Menu abbreviation and name Molecular weight of the amino acidsequence coded by the oligonucleotide

## Repetition of step 4a - 4c

Press " = ":
A new display appears (4a)

### Enter Submenu Ge2

Press "Sm" once:
Menu Ge2 appears

### Formulas

**DNA-Oligonucleotide** (H at the 2'-end of the 2-Desoxy-D-Ribose):

Single strand:  $M_{\rm N} = 1 \cdot (a \cdot 312.2 + g \cdot 328.2 + c \cdot 288.2 + t \cdot 303.2 - 61) g/mol$ Double strand:  $M_{\rm N} = 2 \cdot (a \cdot 312.2 + g \cdot 328.2 + c \cdot 288.2 + t \cdot 303.2 - 61) g/mol$ 

RNA-Oligonucleotide (OH-group at the 2'-end of the D-Ribose; nucleotides with A, G and C have 16 g / mol more):

Single strand:  $M_N = 1 \cdot (a \cdot 328.2 + g \cdot 344.2 + c \cdot 304.2 + u \cdot 305.2 - 61) g/mol$ Double strand:  $M_N = 2 \cdot (a \cdot 328.2 + g \cdot 344.2 + c \cdot 304.2 + u \cdot 305.2 - 61) g/mol$ 

- a, g, c, t and u represent the nucleotides with Adenin, Guanin, Cytosin, Thymin, Uracil.
- The values for the bases are rounded to one significant number after the decimal point.
- The subtrahend "- 61" at the end of the formula is the rounded result of this calculation:
- MPO4 + MOH + MOH  $\approx -$  95 g/mol + 17 g/mol + 17 g/mol  $\approx -$  61 g/mol (no phosphate [PO4] at the 5'-end of a single strand, instead of this an OH-group [oH]; furthermore: OH-group at the 3'-end [OH], where no phosphate is bonded as ester)

### Amino acid sequence (coded protein):

M<sub>protein</sub> = M<sub>first amino acid</sub> - 18 g/mol + M<sub>second amino acid</sub> - 18 g/mol + ... + M<sub>last amino acid</sub>

- The amino acid is displayed in line 2 of menu GeD or GeR.
- The molar masses of the amino acids can be called up in menu Ge6.
- The calculation of the sequences' molar weight is based upon the values for amino acids as displayed in  ${\tt Ge6}$ . 18 g/mol (=  ${\tt MH}_2{\tt O}$ ) are subtracted for the lost water when two amino acids bond.

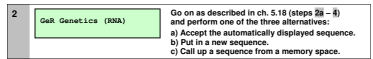
## 5.21 Ge2 CALCULATION: Optical density into n (nmol)

Display	Keys	Actions / Comments
Display	itcys	Actions / Comments

## Enter menu GeD or GeR

1 Any display Press (repeatedly) "Cm" until menu GeD or GeR appears.

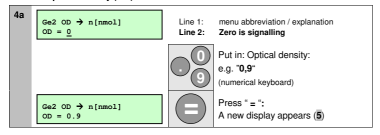
### Continue with menu GeD or GeR



## Enter menu Ge2



## Put in: Optical density (OD)



## Result: Amount of substance in nmol

5	Ge2 ATGGGCGACTCGGCATAG n = 0.162232 nmol	Line 1: Line 2:	Menu abbreviation / sequence Amount of substance in nmol	
---	--	--------------------	---	--

Formula: See end of ch. 5.22

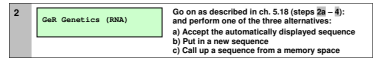
## 5.22 Ge3 CALCULATION: n (nmol) into optical density

Dienlay	Kevs	Actions / Comments
Display	Keys	Actions / Comments

## Enter menu GeD or GeR



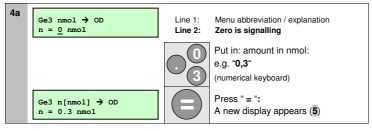
## Continue with menu GeD or GeR



## Enter menu Ge3



## Put in: the substance's amount in nmol



## Result: Optical density (OD)

5	Ge2 ATGGGCGACTCGGCATAG OD = 1,66428	Line 1: Line 2:	Menu abbreviation / sequence Optical density
		-	

Formula (for ch. 5.21 and ch. 5.22)

Calculation in ch. 5.21:

A substance's amount in nmol corresponding to a certain optical density (OD)

 $n = OD \bullet 1000 / \{M_N\} \text{ (nmol)}$ 

amount of substance (in: nmol) =

optical density (no dimension) • 1000 / value of molar mass of the nucleotide sequence

Calculation in ch. 5.22:

Optical density (OD) corresponding to a certain amount of substance in nmol (ch. 5.22):

 $OD = \{n\} \bullet \{M_{N}\} / 1000$ 

Optical density (no dimension) =

value of amount of substance • value of molar mass of the nucleotide sequence / 1000

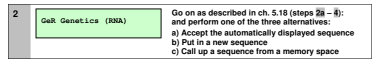
## 5.23 Ge4 CALCULATION: nmol into ng

Display	Kevs	Actions / Comments
Display	itcys	Actions / Comments

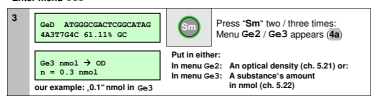
## Enter menu GeD or GeR

1 Any display Press (repeatedly) "Cm" until menu GeD or GeR appears.

### Continue with menu GeD or GeR



## Enter menu Ge3



## Enter menu Ge4 / result: mass of substance in ng



## Formula:

 $m_{\rm N} = n \cdot M_{\rm N}$ 

mass of the oligonucleotide (in: ng) = amount of substance (in: nmol)  $\bullet$  molar mass (in: ng / nmol)

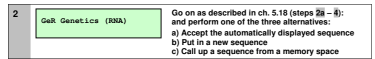
## Ge5 MELTING TEMPERATURE OF THE OLIGONUCLEOTIDE

Display Keys Actions / Comments	Display	Keys	Actions / Comments
---------------------------------	---------	------	--------------------

## Enter menu GeD or GeR

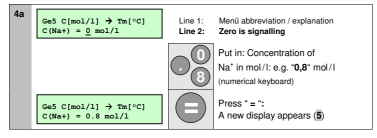
Any display Press (repeatedly) "Cm" until menu GeD or GeR appears.

### Continue with menu GeD or GeR





## Put in: Concentration of Na<sup>+</sup>



## Result: Melting temperature in °C

5	Ge5 ATGGGCGACTCGGCATAG Tm = 75.0736 °C	Line 1: Line 2:	Menu abbreviation / sequence Melting temperature in °C

### Remarks

## Conversion of the temperature into another unit

The temperature displayed in °C can be converted into Kevin oder °Fahrenheit (see ch. 7):





Press the keys " ${f 2nd}$ " and " ${f 4}$ " (function " ${f tu}$ " – temperature unit): Always after pressing these keys the temperature is displayed in another unit.

## Formula:

 $T_m = 81,5$ °C + 16,6 log [c(Na<sup>+</sup>)] + 0,41 (% G + C) - 500 / n

 $T_{\mathsf{m}}$ melting temperature

81,5°C Constante in °C

c(Na<sup>+</sup>) Concentration of Na<sup>+</sup> (input in Ge5 )

0,41 Constant

 $(\% \ G + C) \ \ percentage \ of Guanin and Cytosin$ 

The percentage of GC results from the input in  ${\tt GeD}$  /  ${\tt GeR}$  and is displayed there.

This value is taken over automatically into menu Ge5.

500 / **n n** represents the number of nucleotides in the oligo.

> The number of each type of nucleotide (different bases A, T / U, G and C) is displayed in menu GeD / GeR. These numbers are automatically summed up and taken over into

# 5.25 Ge6 CODONS (BASE TRIPLETTS) OF AMINO ACIDS AND MOLAR MASS OF AMINOACIDS

	Display	Keys	Actions / Comments				
Ent	Enter menu GeD or GeR						
1	Any display	Cm	Press (repeatedly) "Cm" until menu GeD or GeR appears.				
2	GeD Genetics (DNA)	is only a le	as to be performed in GeD / GeR, since Ge6 xical function complementary to the menus molecular biology.				
Ent	er menu Ge6						
3	GeD Genetics (DNA)	Sm	Press "Sm" six times: Menu Ge 6 appears (4).				
Mer	nu Ge6: First display						
4	Ge6 GCA GCC GCG GCU Alanine 89.093	Line 1: Line 2:	Menu abbreviation / base tripletts Coded amino acid / molar mass				
Scr	oll: Moving in Ge6						
3	"Scroll left" (last data) →  "Scroll right" (next data) →		Press a scroll-key once: The last or the next entry is displayed.				
	Ge6 AGA AGG CGA CGC Arginine 174.2	Line 1: Line 2:	Menu abbreviation / base tripletts Coded amino acid / molar mass				
	Ge6 CGG CGU Arginine 174.2	If the display is not big enough to show all base tri- pletts that code the same amino acid at once, the tripletts left over are shown in the next display					

**Remark:** See also the remarks at the end of ch. 5.20 (molar mass of amino acid sequences).

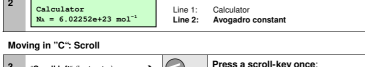
#### LEXICAL FUNCTIONS 6

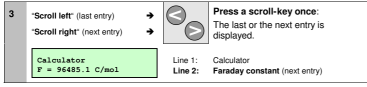
#### **CONSTANTS - FUNCTION "C"** 6.1

This function serves to call up some important constants, that can be also used for further calculations:

	Display	Keys	Actions / Comments					
Ente	Enter function "C"							
1	Any display	C 5	Press "2nd" and "5" (2" function of key "5": "C") Function "C" ist activated. A new display appears (2):					

#### Display of the first entry

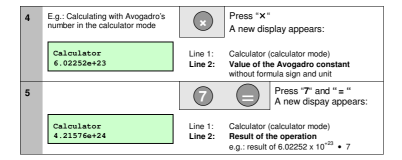




# Processing of data in the calculator mode

Press an operational key (e.g. "+" or "X"):

- The calculator leaves function "C"
  The value is displayed without formula sign and unit in line 2 (calculator mode). Example: see next page



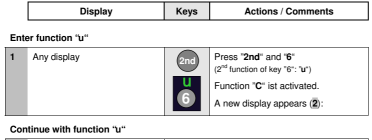
The following constants can be called up in this function:

N <sub>A</sub> =	6.02252 • 10 <sup>23</sup> mol <sup>-1</sup>	Avogadro constant (in: mol <sup>-1</sup> )
<b>F</b> =	96485,1 C / mol	Faraday constant (in: Coulomb / mol)
R =	8,31451 J / (K × mol)	(universal / molar) gas constant (in: Joule / Kelvin • mol)
h =	6,62608 • 10 <sup>-34</sup> Js	Planck constant (in: Joule • second)
k =	1,38066 • 10 <sup>-23</sup> J/K	Boltzmann constant (in: Joule / Kelvin)
<b>€</b> 0 =	8,854 • 10 <sup>-12</sup> F/m	Electrical field constant (in: Faraday-constant / meter)
m <sub>e</sub> =	9,10994 • 10 <sup>-28</sup> g	mass of an electron (in: gramm)
<i>m</i> <sub>n</sub> =	1,67493 • 10 <sup>-24</sup> g	mass of a neutron (in: gramm)
<i>m</i> <sub>p</sub> =	1,67262 • 10 <sup>-24</sup> g	masse of a Protons (in: gramm)
e =	1,60218 • 10 <sup>-19</sup> C	charge of an electron / elementary charge (in: Coulomb)
<i>V</i> <sub>m, n</sub> =	22,4136 I / mol	mole volume (Liter / mol)

**Remark:** Multiplications and other operations with two constants of "C" or of a constant of "C" with the atomic mass unit "u" (2<sup>nd</sup> function of key "6") are only possible after memorizing the first value in one of 10 memory places (see ch. 9.2.2 p. 79).

#### 6.2 ATOMIC MASS UNIT "u"

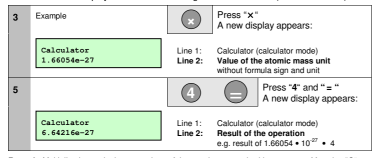
This function serves to call up the atomic mass unit "u", that can be also used for further calculations:



2	Calculator u = 1.66054e-27 kg	Line 1: Line 2:	Calculator Atomic mass unit u

#### Processing of data in the calculator mode

- Press an operational key (e.g. "+" or "X"): The calculator leaves function "C"
- The value is displayed without formula sign and unit in line 2 (calculator mode).



**Remark:** Multiplication and other operations of the atomic mass unit with constants of function "C" can only be performed after memorizing the first value in one of 10 memory places (see ch. 9.2.2 p. 79).

#### 6.3 LIBRARY - (Menu Lib)

In the menu **Lib** data of the chemical elements can be called up. Detailled description: ch. 5.17, p. 56.

# 6.4 AMINO ACIDS (Menu Ge 6)

In the menu **Ge6** base tripletts, that code an amino acid, and the molar mass of amino-acids can be called up. Detailled description: ch. 5.25, p. 68.

# 7 CONVERTING TEMPERATURES

Function "**u** – temperature unit" (2<sup>nd</sup> function of key "**4**"):

Conversion of temperatures in °Celsius, °Fahrenheit and Kelvin into one another.

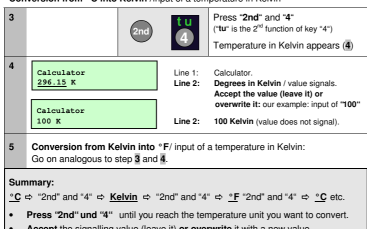
You can enter this function: • from the calculator mode (with "0" or any number put in)

• from a result in menu Ge5 (ch. 5.24)

Calculator mode: Enter 'tu" / put in: temperature in °C

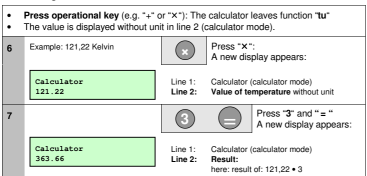
	Display	Keys	Actions / Comments
1	Calculator mode:  Calculator 0  Error appears (input is nonsense):  Calculator Error	2nd t u 4	Press "2nd" and "4"  ("tu" is the 2 <sup>nd</sup> function of key "4")  a) A new display appears (2):  b) "Error" appears:  The last value is nonsense, e.g.: "-300" (°C)  Press "2nd" and "4" again A new display appears (2):
2	Calculator 0 °C  Display after overwriting:  Calculator 55 °C	Line 1: Line 2:	Calculator  Degrees in °C/ signalling number (e.g. "0")  Accept the number (leave it) or write over: e.g. with "55"  55 °C (the new value does not signal).

#### Conversion from °C into Kelvin /input of a temperature in Kelvin



- Accept the signalling value (leave it) or overwrite it with a new value.
- Press "2nd" und "4" until the temperature is converted into the wished unit.

#### Processing of data in the calculator mode



#### Entering function "tu" from menu Ge5

# Display Keys Actions / Comments Display of melting temp. in °C in menu Ge5: / conversion into Kelvin Result from Ge5 (vgl. Kap. 5.23) Ge5 ATGGCGACTCGGCATAG Tm = 75.0736 °C Tm = 75.0736 °C Actions / Comments Press "2nd" and "4" ("tu" is the 2<sup>nd</sup> function of key "4") Line 2: Melting temperature in °C

Ge5 ATGGGCGACTCGGCATAG

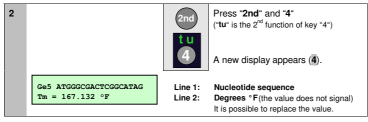
Line 1: Nucleotide sequence

Ge5 ATGGGCGACTCGGCATAG

Tm = 348.224 K

Line 1: Nucleotide sequence
Temp. in Kelvin (value does not signal)
It is possible to replace the value.

#### Conversion of melting temperature: Kelvin into ${^\circ}\text{F}$



#### Conversion of melting temperature: Kelvin into °F

3 Proceed analogous to step 2.

#### Order of keys:

 ${}^{\circ}\underline{\mathbf{C}} \Leftrightarrow {}^{\circ}2\text{nd}{}^{"}$  and  ${}^{\circ}4{}^{"} \Leftrightarrow \underline{\mathbf{Kelvin}} \Leftrightarrow {}^{\circ}2\text{nd}{}^{"}$  and  ${}^{\circ}4{}^{"} \Leftrightarrow {}^{\circ}\underline{\mathbf{F}} = {}^{\circ}2\text{nd}{}^{"}$  and  ${}^{\circ}4{}^{"} \Leftrightarrow {}^{\circ}\underline{\mathbf{C}} = {}^{\circ}\mathbf{C}$  etc. Like described on the previous page after step  $\overline{\mathbf{5}}$ .

# 8 CALCULATOR MODE

# 8.1 NUMERICAL KEYS

To the right of the periodic system keyboard are placed the numerical keys from "0" to "9" and a key for the input of decimal numbers ("  $\cdot$  .").

The input is performed analogous to common pocket calculators.

#### 8.2 OPERATION KEYS

To the right of the periodic system keyboard are the following operational keys:

Key	Function	Remarks	
<b>(+)</b>	Addition	If pressed within a menu: The device changes into the calculator mode ("Calculator").	
	Subtraction	If pressed within a menu: The device changes into the calculator mode ("Calculator").	
×	Multiplikation	If pressed within a menu: The device changes into the calculator mode ("Calculator").	
$\odot$	Division	If pressed within a menu: The device changes into the calculator mode ("Calculator").	
<u> </u>	Change of sign		
exp	Exponent of 10 (see ch. 3.1.2, e)	2 <sup>nd</sup> function of key " + "; press at first key "2nd"     A number, that has been put in before, is multiplicated with a power of 10 – possible from 10 <sup>37</sup> up to 10 <sup>+37</sup> .	
	Result		

The operational keys are used analogous to common pocket calculators.

The function for setting brackets ( $2^{nd}$  function of '%') is only active in the stoichiometric menus and can only be used for the input of compound formulas (see ch. 5.2 p. 24)!

# 8.3 FUNCTIONAL KEYS

To the right of the periodic system keyboard are the following functional keys:

Key	Function	Remarks	
on	Reset key	Deletes the whole input displayed.	
clr	Clear error / clear	<ul> <li>Deletes only the last input (for corrections of the entry).</li> <li>Deletes an automatically displayed entry completely.</li> </ul>	
2nd	Activates / deactivates 2 <sup>nd</sup> functions		
%	Percent		
1/x	Reciprocal value	<ul> <li>2<sup>nd</sup> function of key " 0"; press at first key "2nd".</li> <li>If pressed within a menu: The device changes into the calculator mode ("Calculator").</li> </ul>	
X <sup>2</sup>	2 <sup>nd</sup> power (square)	<ul> <li>2<sup>nd</sup> function of key " X "; press at first key "2nd"</li> <li>If pressed within a menu: The device changes into the calculator mode ("Calculator").</li> </ul>	
•	Square root	<ul> <li>2<sup>nd</sup> function of key " . " ; press at first key "2nd"</li> <li>If pressed within a menu: The device changes into the calculator mode ("Calculator").</li> </ul>	
y*	x <sup>th</sup> power of y	<ul> <li>2<sup>nd</sup> function of key "±"; press at first key "2nd"</li> <li>If pressed within a menu: The device changes into the calculator mode ("Calculator").</li> </ul>	
lg ÷	Common (base-10) logarithm	<ul> <li>2<sup>nd</sup> function of key " + "; press at first key "2nd"</li> <li>If pressed within a menu: The device changes into the calculator mode ("Calculator").</li> </ul>	
in _	Natural logarithm	2 <sup>nd</sup> function of key " - " ; press at first key "2nd"     If pressed within a menu: The device changes into the calculator mode ("Calculator").	

The functional keys are used analogous to common pocket calculators.

#### 9 MEMORY FUNCTIONS

#### 9.1 AUTOMATIC MEMORY

The following data put in by the user or called up from a memory place (vgl. Kap. 9.2.3 and 9.2.4) are automatically memorized:

- Compound formulas in den stoichiometric menus (Input possible in M1, C1 C3, P1 P4) and
- Nucleotide sequences in the menus GeD and GeR.

These data are automatically displayed, if one of the menus is entered again. This is not the case, if the last compound formula / nucleotide sequence has been deleted with key "clr"

- before the device is switched off (by pressing "2nd" and "off" respectively automatically after 2 minutes without any operation) or
- before pressing the key "on" during operation time (master-clear-key).

#### 9.2 AUTOMATIC MEMORY

#### 9.2.1 General Remarks

In every of the three functional areas of the device, i.e.

- · calculator mode,
- stoichiometric functions,
- · biochemical functions,

up to 10 memory places, that are independent from the automatic memory, can be used (alltogether 30 memory places). These memory places serve to memorize:

- Numbers in the calculator mode
- Compound formulas in the stoichiometric functions
- Nucleotid sequences in the biochemical functions

For memorizing data within one of these areas the user must assign the data to a memory place by pressing a numerical key:

Therefore the numerical keys "1" - "9" inclusive "0" can be used.

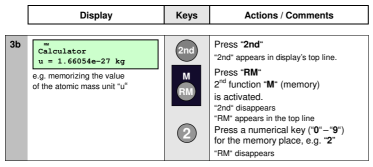
#### 9.2.2 Calculator Mode

#### Memorizing a number

In the calculator mode it is possible to memorize:

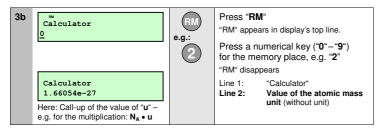
- Any numbers in the calculator mode.
- The values of constants called up in function "C" (without formula sign and unit) and the value of the atomic mass unit "u" (without unit).

Operations with two constants of "C" or of a constant of "C" with the atomic mass unit "u"  $(2^{nd}$  function of key "6") are only possible after memorizing the first value in one of the 10 memory places (see ch. 6.1 and 6.2).



The value of the atomic mass unit can now be called from memory place "2".

#### Calling up a memorized number



#### 9.2.3 Stoichiometric Menus

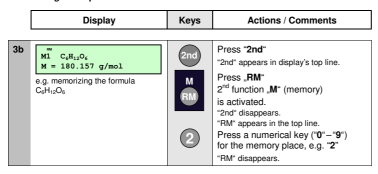
#### Preliminaries

A memorized compound formulas can be called up in all stoichiometric menus, in which it is possible to put in such a formula (see the schedule in ch. 4, respectively the backside of this manual).

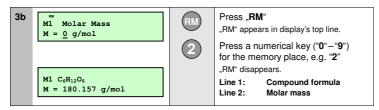
#### Memorizing a compound formula is only possible in menu M1.

It is not possible to memorize a molar mass without the input of a compound formula (see ch. 5.2, step 3c and remarks on p. 25).

#### Memorizing a compound formula in menu M1



#### Calling up a memorized compound formula



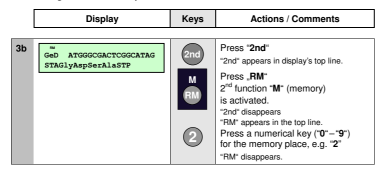
#### 9.2.4 Molecular Biology (PCR)

#### Preliminaries

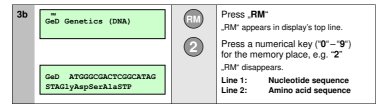
# Memorizing or calling up a nucleotide sequence from a memory place is only possible in the menus GeD / GeR!

A nucleotide sequence put in or called up from a memory space in these menus remains the base for all calculations in the menus **Ge1** – **Ge5**. Changing a nucleotide sequence as calculation base is only possible in **GeD** / **GeR**.

#### Memorizing a nucleotide sequence in menu GeD / GeR



#### Calling up a memorized nucleotide sequence



#### 10 BATTERY CHANGE – WARRANTY

#### 10.1 BATTERY CHANGE

#### You will get new batteries at your chemcode®-dealer!

#### Technical Data of the batteries

Type: CR 2025 Voltage: 2 x 3 V

Operation time: ca. 100 hours of operation

#### Keep the new batteries ready (2 pieces)

Change batteries, when the colour of displayed letters becomes too weak. After removing the two old batteries you have two minutes time during which all data memorized on one of the 30 memory places (see ch. 9.2) are kept.

Please keep the two new batteries ready before taking the old ones out!.

#### Open the battery case and remove the old batteries

Turn out the screw with a single slot on the left side of the battery case on the calculator's backside (a). Remove the cover of the battery case, the gum behind the batteries and afterwards the batteries.

#### Put in the new batteries

Take the batteries with the inscription on the upper side (plus-pole) push them forward and to the left and right side of the battery case (circles mark the position).

Before closing the battery case again it is important to put the black gum behind the batteries again (b)! This gum is necessary for optimal performance of the power source.

#### Close the batterie case again

Push the notch at the inner edge of the cover unter the case of the device. Then press the outer edge of the cover down until it fits exactly. Take the screw and fasten it again.

Attention! Users may only open the cover (srew with a single slot). The screws with crossed slots of the calculator's case may not be be opened. (see ch. 10.2)!







 b) Battery case opened: Plus-pole of batteries with inscription orientated towards the user; the gum behind batteries is marked grey.



c) Battery case opened: The gum behind batteries has been removed.



 Push the notch at the inner edge of the cover in the slot of the rim surrounding the opening of the battery case



e) Press down the outer edge of the cover until it fits exactly and fasten it with the screw again.

#### 10.2 WARRANTY

Each chemcode® has been checked thoroughly after production.

Should your item nonetheless have any defect not discovered by us, we will replace your item as quick as possible. Please turn to your chemcode<sup>®</sup>-dealer with the defect device.

In order to prove your claim please pay attention to the following points:

- a) Keep your receipt and bring it to the dealer together with the chemcode<sup>®</sup>.
- b) The identidy-code on a sticker below the batteries in the battery case should not have been removed.

There is no warranty for calculators, whose case has been opened by the user. Only the battery case may be opened (all screws with crossed slots must remain untouched). There is no warranty for devices, that have been opened or show other damages on the surface.

Warranty is orientated towards the legal statuses valid in Germany  $\!\!/$  the European Union.

# 11 EXPLANATION OF SYMBOLS AND ABBREVIATIONS

# 11.1 SYMBOLS AND ABBREVIATIONS IN THE DISPLAY

Abbrev. / Symbol	Explanation	See chapter					
Calcula	Calculator: Calculator mode 2.1/8						
C1	Menu C1 (Concentration 1)	5.7					
C2	Menu C2 (Concentration 2)	5.8					
C3	Menu C3 (Concentration 3)	5.9					
Сж	Mass concentration of a substance x						
D	Menu D (Dilution)	5.11					
Error	An input is false or has not been done						
F	Menu F (Formula)	5.12					
GeD	Menu GeD (Genetics DNA)	5.18					
GeR	Menu GeR (Genetics RNA)	5.19					
g	Grammes						
K	Concentration in mass percentage						
Lib	Menu Lib (Library)	5.17					
1	Liter						
M	Molar mass						
M1	Menu M1 (Molar mass)	5.2					
M2	Menu M2 (amount of a substance: mol in g)	5.3					
м3	Menu M3 (amount of a substance: g in mol)	5.4					
M4	Menu M4 (ideal gas volume of a substance)	5.5					
m	Mass						

Abbrev. / Symbol		Explanation	See chapter
mol	Unit "Mol"	- 1 Mol = 6,22 x 10 <sup>23</sup> particles	
n	Amount of	substance	
P1	Menu P1	(Element Partition 1): Part of single elements in %	5.13
P2	Menu P2	(Element Partition 2): Part of single elements in g	5.14
P3	Menu P3	(Formula Partition 3): Part of several elements in %	5.15
P4	Menu P4	(Formula Partition 4): Part of several elements in in g	5.16
Sol		(Solution) a substance in g for preparing a solution	5.6
T	Menu T (T	itration)	5.10
v	Volume		
V1	Final volur	me (Volumen 1)	5.11
V2	Initial volu	me (Volumen 2)	5.11
[x]	Concentra	Concentration in mol / I	
[x]1	Final cond	entration in mol / I	5.11
[x]2	Initial valu	e of concentration in mol / I	5.11
ρ	Density of	a substance	5.9
$\boldsymbol{\epsilon}_{0}$	Electrical	field constant	6.1

An underlined number in the display pictures symbolizes a signalling number.

# 11.2 ABBREVIATIONS ON KEYS AN THE CASE

# 11.2.1 Abbreviations on keys

Abbrev. / Symbol	Explanation	See chapter
2nd	The 2 <sup>nd</sup> function of a key is activated / disactivated	1.3
clr	Clear / error correction	3.2.1
Cm	Change menu (next menu group is entered)	2.2
on	Switch on / reset key	1.2 / 3.2.2
RM	Read Memory	9
Sm	Submenu (next submenu in a menu group is entered)	2.3

# 11.2.2 Abbreviations on the geen lines of the case

Abbrev. / Symbol	Explanation	See chapter	
1, 2,	Periods $1-7$ , elements of main groups	(black)	1.4
I a, II a,	Main groups	(black	1.4
1, 2,	Periods 1 – 7, transitory elements	(white)	1.4
l b, ll b,	Transition groups	(white)	1.4
В	Bases of the nucleotides	(blue)	1.4
L	Lanthanoids	(blue)	1.4
A	Actinoids	(blue)	1.4

# 11.2.3 Abbreviations on the case (second function of keys)

Abbrev. / Symbol	Explanation		See chapter
Α	Adenine	(blue)	5.18
T/U	Thymine / Uracil	(blue)	5.18
G	Guanine	(blue)	6.1
С	Cytosine	(blue)	5.18
М	Memory	(white)	9
off	Switch off	(white)	1.2
tu	Temperature unit	(green)	5.18
С	Constants	(green)	7

6.2

(green)

# 11.3 ABBREVIATIONS IN THE MANUAL'S TEXT

Atomic mass unit "u"

a. and

u

abbrev. abbreviation ch. chapter

corr. corresponding

e.g. exempli gratia (for example)

i.e. id est
p./pp. page/pages
resp. respective(ly)
stoichiom. stoichiometric
temp. temperature

#### 12 FEEDBACK

# Let us know about your opinion!

Having talked with many experts we know, that a product like the chemcode® meets the needs and interests of many persons.

Again and again we have used the informations collected in these conversations in order to improve the conception of the chemcode $^{\text{®}}$ .

E.g. all functions concerning biochemistry have been the result of the exchange with experts, to whom the integration of these features was of great importance.

We are convinced of the need for and the benefits of a product like the chemcode<sup>®</sup>. Nevertheless we are interested in getting feedback from our customers and want to know about their experiences in working with the chemcode<sup>®</sup> during daily routine.

Please use our website "website" in order to send us an e-mail:

#### www.chemcode.com

Or send us a message by fax:

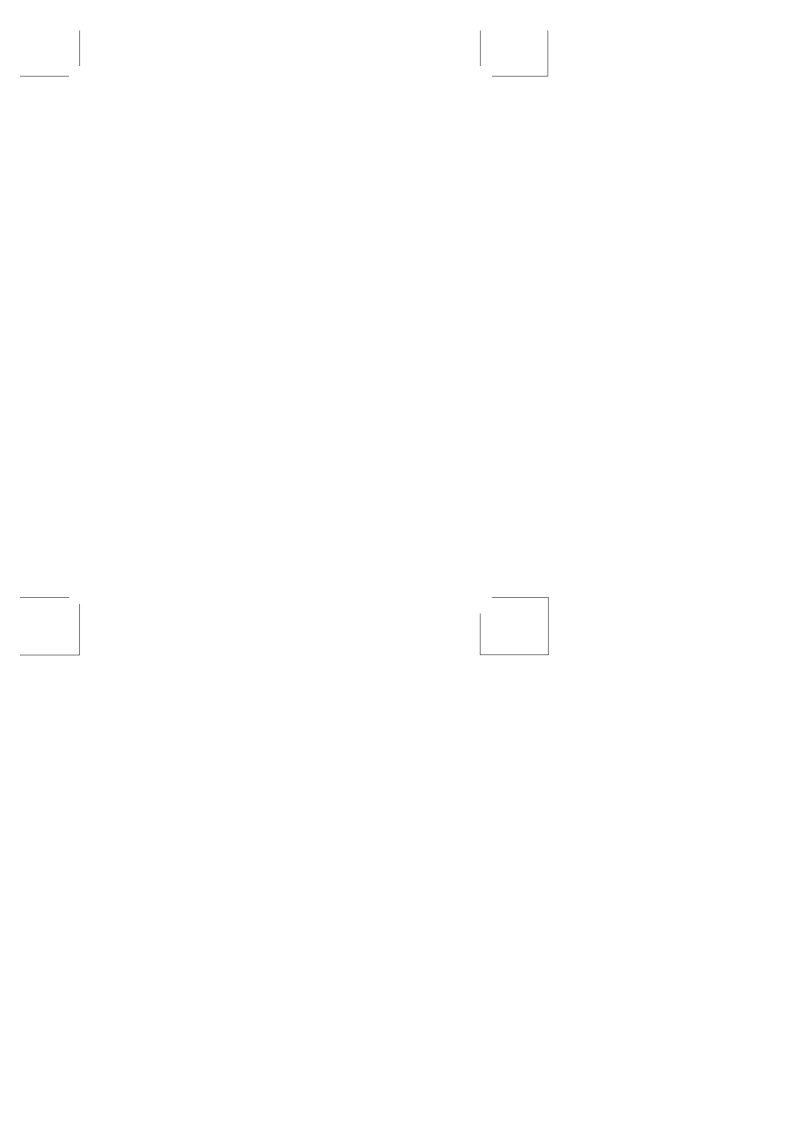
0049 / (0)89 / 3 50 75 03

You will find some questions we are especially interested in on the next pages

We are looking forward to your feedback.

Your chemcode®-team

1)	Did the chemcode <sup>®</sup> prove its worth during your daily routine in the laboratory and elswhere?							
2)	Do you have a need for the integration of further functions?  Stoichiometry:							
•	Molecular Biology:							
•	Others:							
Ple	Please contact us! We will always look very carefully at your concerns!							



# **SHORT INSTRUCTIONS**

It is advisable to study ch. 1-3 before using this device for the first time.

The single menus are self-explanatory to a high degree.

Details: see ch. 5.





Press "on": The device is in the calculator mode (Calculator).



Press "Cm" or an element key (e.g. Na 11): Change to menu M1.



Press "Cm" for entering the following menus/menu groups.



"Sm": If there is a number behind a menu's abbreviation (or in GeD / GeR) submenus of this menu group are entered with the key "Sm":

(→)	Compounds / Analysis				Data	Oligos			
Calculator →	<u>M1</u>	→ <u>sol</u> →	<u>C1</u>	$\rightarrow$ T $\rightarrow$ D $\rightarrow$ F $\rightarrow$	<u>P1</u>	→ Lib →	GeD	÷	GeR
Û	+		Ψ.		4	+	•		+
Press one time	<u>M2</u>		<u>C2</u>	Explanation	<u>P2</u>	Press-	Ge1		Ge1
Û	4		4	of the menus'	4	element	4		4
	<u>M3</u>		<u>C3</u>	see	P3	key and	Ge2		Ge2
( <b>Sm</b> ) ( <b>V</b> )	Ψ.			contents	Ψ.	scroll	•		<b>4</b>
	<u>M4</u>			(pp. 4 - 5)	<u>P4</u>		Ge3		Ge3
Input of compound formulas (possible in <u>underlined menus</u> ): Change betweer element keys and numeric keys (for subscripts [stoichiometric numbers]).									<b>+</b>
<ul> <li>Input of nucleotide sequences (oligos) in GeD / GeR: Press the keys with the second functions A, T / U, G, C, which are automatically active in these menus.</li> </ul>									







"clr": 1) Correcting inputs (elements, subscripts, nucleotides, numbers). 2) Deleting whole compound formulas/oligonucleotides, that have been put in completely (or called up by a memory function) and are processed.



Scrolling: 1) If inputs reach beyond the display's frame (an arrow appears on the left or/and the right side of the display's top line), it is possible to scroll left or right by pressing these keys. 2) Navigating through  ${ t Lib}$  /  ${ t Ge6}$ .