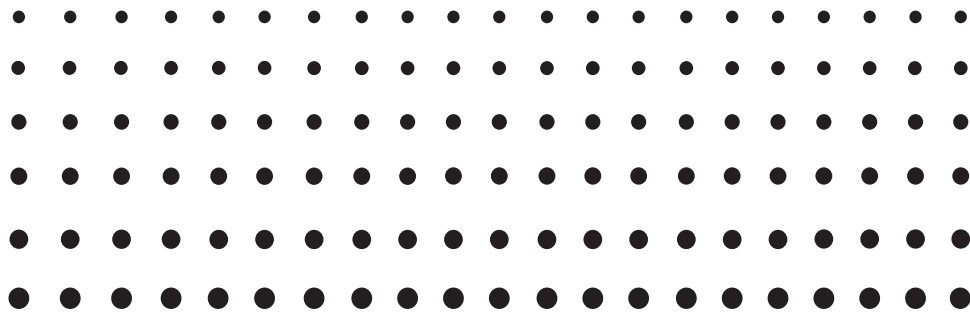
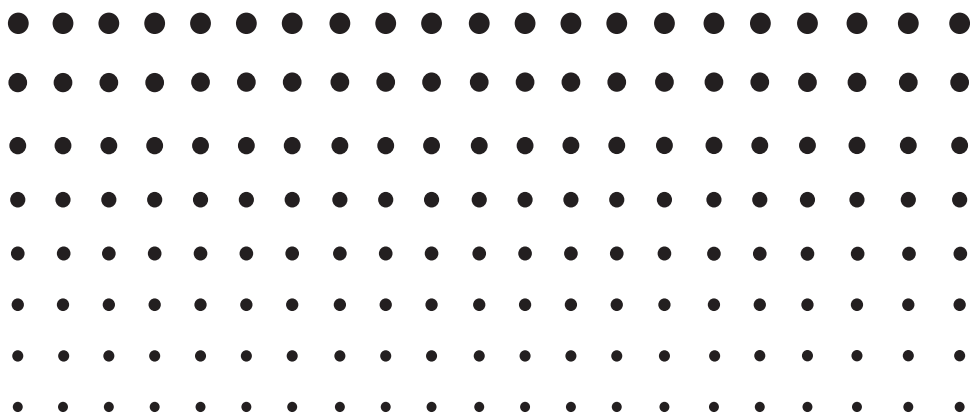


For fx-9860G Series/
GRAPH 95/75+/75/85 Series/
fx-9860GIII/fx-9750GIII/GRAPH35+ E II



Physium
Application
User's Guide



CASIO Worldwide Education Website

<https://edu.casio.com>

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- 6 Precautions**

1 Physium Overview

- The Physium application provides you with the following capabilities.

Periodic Table of Elements

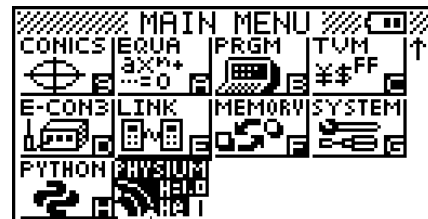
- The application can be used to display periodic table of elements.
- Table shows element atomic number, atomic symbol, atomic weight, etc.
- Elements can be searched for based on element name, atomic symbol, atomic number or atomic weight.

Fundamental Physical Constants

- The application can be used to display fundamental physical constants, which are grouped for easy referencing.
- Physical constants can be edited and saved as required.
- Physical constants can be stored in Alpha memory and used in the **RUN • MAT** mode.

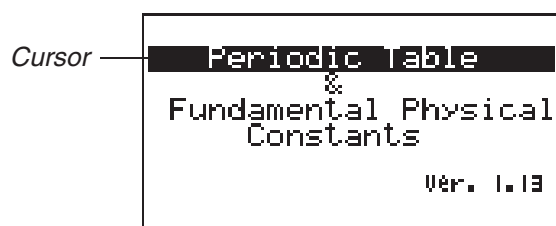
2 Starting Up Physium

1. From the Main Menu, enter the **PHYSIUM** mode.



PHYSIUM mode

2. This displays an initial screen like the one shown below.



3. Use \blacktriangle and \blacktriangledown to move the highlighting and select the type of information you want (Periodic Table or Fundamental Physical Constants).
4. Press $\boxed{\text{EXE}}$ to display the information you selected in step 3 (Periodic Table or Fundamental Physical Constants).

3 Periodic Table

■ Periodic Table Screen

Period Number

	1	2	3	4	5	6	7
1	■						
2	Li	Be					
3	Na	Mg					
4	K	Ca	Sc	Ti	V	Cr	Mn
5	Rb	Sr	Y	Zr	Nb	Mo	Tc
6	Cs	Ba	L*	Hf	Ta	W	Re
	MINI						DETAIL

Group Number
Cursor

- Group numbers run along top of the screen, while period numbers run down the left.
- Lanthanoids are indicated as L*, while actinoids are indicated as A*.
- ¹¹³Uut, ¹¹⁵Uup, ¹¹⁷Uus, and ¹¹⁸Uuo are indicated as **.
- Use ▲, ▼, ◀, and ▶ to move the cursor around the screen.

F1 (MINI)..... Displays the periodic table MINI screen.

F6 (DETAIL) (or **EXE**) Displays a dialog box with details about the element where the cursor is located. If L* or A* is selected, pressing this key displays the transition element screen.

EXIT Returns to the Physium initial screen.

● Details Dialog Box

Atomic Symbol

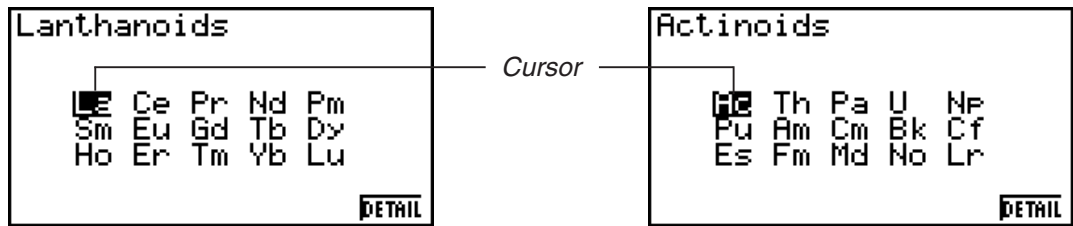
	1	2	3	4	5	6	7
Atomic Number	1	4	Be				
	2	Beryllium					
	3	Metal					
	4	Alkaline Earth					
Atomic Weight	5	Weight: 9.0121831					
	6						
		MINI					DETAIL

Element Name
Attributes

- The details dialog box shows the element atomic number, atomic symbol, element name, atomic weight, and attributes for the element that was selected on the periodic table screen.
- Brackets ([]) indicate atomic weight of a well-known isotope. The element names for these isotopes are followed by an asterisk (*).

EXIT (or **EXE** or **AC/ON**) Closes the dialog box.

• Lanthanoids Screen and Actinoids Screen

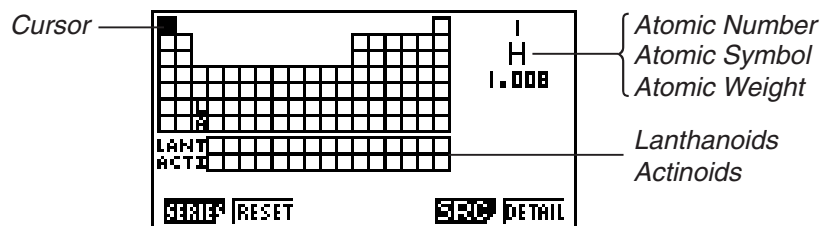


- Pressing **F6** (DETAIL) (or **EXE**) while L* or A* is selected on the periodic table screen will display the transition element screen.
- Use **▲**, **▼**, **◀**, and **▶** to move the cursor around the screen.

F6 (DETAIL) (or **EXE**) Displays a dialog box with details about the element where the cursor is located.

EXIT Returns to the periodic table screen.

■ Mini Table Screen



- The mini table screen shows a compressed version of the periodic table, with cells that represent each element.
- The information in the upper right corner of the screen shows the atomic number, atomic symbol, and atomic weight for the element where the cursor is currently located in the table.
- Lanthanoids and actinoids are shown below the periodic table (LANT, ACTI).
- When a lanthanoid is selected, the information in the upper right shows the corresponding atomic number (57 to 71), “Lant.” for the atomic symbol, and blank for the atomic weight. When an actinoid is selected, the information shows the corresponding atomic number (89 to 103), “Acti.” for the atomic symbol, and blank for the atomic weight.
- Use **▲**, **▼**, **◀**, and **▶** to move the cursor around the screen.

- F1** (SERIES) **F1** (METAL) Highlights the cells of elements whose metallicity makes them metals.
- F2** (TRANS) Highlights the cells of elements whose metallicity makes them transition elements.
- F3** (A-MET) Highlights the cells of elements that are in the Alkali Metals category.
- F4** (A-EAR) Highlights the cells of elements that are in the Alkaline Earth Metal category.
- F5** (HALGN) Highlights the cells of elements that are in the Halogens category.
- F6** (▷) **F1** (n-GAS)..... Highlights the cells of elements that are in the Noble Gases category.
- F6** (▷) **F2** (n-METL)..... Highlights the cells of elements whose metallicity makes them non-metals.
- F6** (▷) **F3** (R-EAR) Highlights the cells of elements that are in the Rare Earth category.
- F2** (RESET) Clears highlighting from the mini table screen.
- F5** (SRC) **F1** (NAME) Displays a dialog box to search for an element name.
- F2** (SYMBL) Displays a dialog box to search for an atomic symbol.
- F3** (No.) Displays a dialog box to search for an atomic number.
- F4** (WEIGH) Displays a dialog box to search for an atomic weight.
- F6** (DETAIL) (or **EXE**) Displays a dialog box with details about the element where the cursor is located. Note that the dialog box does not appear when a lanthanoid or actinoid is selected.
- EXIT** Returns to the periodic table screen.

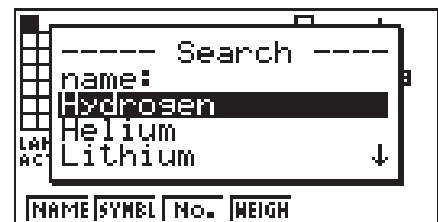
• Mini Table Details Dialog Box

- The mini table details dialog box is the same as the details dialog box described under “Details Dialog Box” on page 3-1.
- The mini table details dialog box shows details for the element where the cursor is currently located on the mini table. Note that the cursor cannot be used to select a LANT. or ACTI. cell.

EXIT (or **EXE** or **AC/ON**) Closes the dialog box.

• Searching for an Element Name

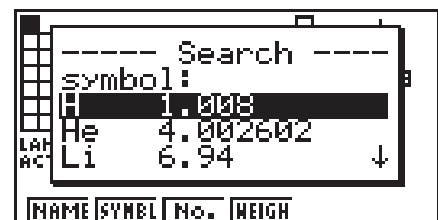
1. On the mini table screen, press **F5** (SRC) and then **F1** (NAME).
 - This will display an element name search dialog box.



2. Enter up to nine characters for the element name you want to search for.
 - The screen will change to show all elements whose names start with the character(s) you input.
3. Use **▲** and **▼** to select the element name you want.
4. Press **EXE** to return to the mini table screen, with the cursor located at the element you selected in step 3.
 - “Nothing” will appear on the screen if no element name corresponds to the character(s) you input.
 - To close the dialog box and return to the mini table screen without searching for anything, press **EXIT**.

• Searching for an Atomic Symbol

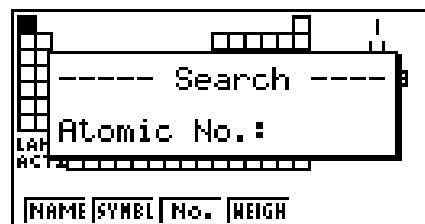
1. On the mini table screen, press **F5** (SRC) and then **F2** (SYMBL).
 - This will display an atomic symbol search dialog box.



2. Enter up to nine characters for the atomic symbol you want to search for.
 - The screen will change to show all elements whose atomic symbols start with the character(s) you input.
3. Use ▲ and ▼ to select the atomic symbol you want.
4. Press [EXE] to return to the mini table screen, with the cursor located at the element you selected in step 3.
 - “Nothing” will appear on the screen if no element name corresponds to the character(s) you input.
 - To close the dialog box and return to the mini table screen without searching for anything, press [EXIT].

• Atomic Number Search

1. On the mini table screen, press [F5] (SRC) and then [F3] (No.).
 - This will display an atomic number search dialog box.

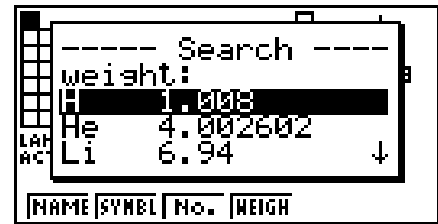


2. Enter up to three digits for the atomic number you want to search for.
3. Press [EXE] to return to the mini table screen, with the cursor located at the element that corresponds to the atomic number you input in step 2.
 - If there is no element that corresponds to the atomic number you input, pressing [EXE] will return to the mini table screen with the cursor at the same location it was when you started this procedure.
 - To close the dialog box and return to the mini table screen without searching for anything, press [EXIT].

• Atomic Weight Search

1. On the mini table screen, press **[F5]** (SRC) and then **[F4]** (WEIGH).

- This will display an atomic weight search dialog box.



2. Enter a value up to nine digits long (including numbers and decimal point) for the atomic weight you want to search for.

- The screen will change to show all elements whose atomic weight corresponds to the numbers you input.

3. Use **▲** and **▼** to select the atomic weight you want.

4. Press **[EXE]** to return to the mini table screen, with the cursor located at the element you selected in step 3.

- “Nothing” will appear on the screen if no atomic weight corresponds to the number(s) you input.
- To close the dialog box and return to the mini table screen without searching for anything, press **[EXIT]**.

4 Fundamental Physical Constants

■ Category Selection Screen

```
Physical Constants
1: Universal
2: Electromagnetic
3: Atomic & nuclear
4: Physico-chemical
5: Adopted values
0: My Drawer
```

- 1 (Universal) Displays a list of universal physical constants.
- 2 (Electromagnetic) Displays a list of electromagnetic physical constants.
- 3 (Atomic & nuclear) Displays a list of atomic and nuclear physical constants.
- 4 (Physico-chemical) Displays a list of physico-chemical physical constants.
- 5 (Adopted values) Displays a list of adopted values physical constants.
- 0 (My Drawer) Displays the “My Drawer” screen.
- EXIT Returns to the Physium initial screen.

■ Built-in Constants

- The following is a list of the constants that this application includes as built-in fundamental physical constants.

Category	Quantity	Value
Universal	c : speed of light in vacuum	299792458
	μ_0 : magnetic constant	1.2566370614E-6
	ϵ_0 : electric constant	8.854187817E-12
	Z_0 : characteristic impedance of vacuum	376.730313461
	G : Newtonian constant of gravitation	6.67408E-11
	h : Planck constant	6.626070040E-34
	\hbar : Planck constant over 2π $\frac{h}{2\pi}$	1.054571800E-34
	m_p : Planck mass	2.176470E-8
	l_p : Planck length	1.616229E-35
	t_p : Planck time	5.39116E-44
Electromagnetic	e : elementary charge	1.6021766208E-19
	ϕ_0 : magnetic flux quantum	2.067833831E-15
	G_0 : conductance quantum	7.7480917310E-5
	K_J : Josephson constant	483597.8525E9
	R_K : von Klitzing constant	25812.8074555
	μ_B : Bohr magneton	927.4009994E-26
	μ_N : nuclear magneton	5.050783699E-27

Category	Quantity	Value
Atomic & nuclear	α : fine-structure constant	7.2973525664E-3
	R_∞ : Rydberg constant	10973731.568508
	a_0 : Bohr radius	0.52917721067E-10
	m_e : electron mass	9.10938356E-31
	μ_e : electron magnetic moment	-928.4764620E-26
	m_μ : muon mass	1.883531594E-28
	μ_μ : muon magnetic moment	-4.49044826E-26
	m_τ : tau mass	3.16747E-27
	m_p : proton mass	1.672621898E-27
	μ_p : proton magnetic moment	1.4106067873E-26
	m_n : neutron mass	1.674927471E-27
	μ_n : neutron magnetic moment	-0.96623650E-26
Physico-chemical	N_A : Avogadro constant	6.022140857E23
	m_u : atomic mass constant	1.660539040E-27
	F : Faraday constant	96485.33289
	R : molar gas constant	8.3144598
	k : Boltzmann constant	1.38064852E-23
	V_m : molar volume of ideal gas (273.15 K, 100 kPa)	22.710947E-3
	σ : Stefan-Boltzmann constant	5.670367E-8
Adopted values	K_{J-90} : conventional value of Josephson constant	483597.9
	R_{K-90} : conventional value of von Klitzing constant	25812.807
	g_n : standard acceleration of gravity	9.80665

■ Constant List Screen

```

----- Universal -----
C = 299792458
P0 = 1.25663706E-6
ε0 = 8.8541878E-12
Z0 = 376.7303135
G = 6.67408E-11
h = 6.62607E-34 ↓
[EDIT] [STORE] [DETAIL] [KEEP] [INIT] [A•INIT]

```

- The constant list screen shows each constant in the format: Symbol = Value.
- Use ▲ and ▼ to move the highlighting to the constant you want to select.
 - [F1] (EDIT) Enters the editing mode for editing the currently selected constant. The editing mode is also entered automatically whenever a number key is pressed while a constant is selected.
 - [F2] (STORE) Stores the currently selected constant in Alpha memory.
 - [F3] (DETAIL) (or [EXE]) Displays a dialog box with details about the currently selected constant.
 - [F4] (KEEP) Stores the currently selected constant in My Drawer.
 - Press [F4] (KEEP) causes the message “Complete!” to appear on the display. Press [EXIT] to clear the message from the screen.
 - [F5] (INIT) Returns the currently selected constant to its initial default value.
 - [F6] (A•INIT) Returns all constants to their initial default values.
 - [EXIT] Returns to the category selection screen.

■ My Drawer Screen

```

----- My Drawer -----
C = 299792458
P0 = 1.25663706E-6
ε0 = 8.8541878E-12
[STORE] [DETAIL] [DEL]

```

- Pressing [F4] (KEEP) while the constant list screen is on the display will save the currently selected constant to the My Drawer screen. The My Drawer screen displays constants in the sequence they are saved.
 - [F2] (STORE) Stores the currently selected constant in Alpha memory.
 - [F3] (DETAIL) (or [EXE]) Displays a dialog box with details about the currently selected constant.
 - [F6] (DEL) Deletes the currently selected constant.
 - [EXIT] Returns to the category selection screen.

■ Editing a Constant

1. Select the constant you want to edit, and then press **[F1]** (EDIT).

- This enters the editing mode.

```

----- Universal -----
C      =299792458
F0    =1.25663706E-6
N0    =8.8541878E-12
M0    =376.7303135
G0    =6.67408E-11
h      =6.62607E-34      ↓

```

2. Edit the constant as desired.

3. Press **[EXE]**.

- This saves the edited version of the constant.
- Even if you input more than 15 digits for a constant value, only the 15 most significant digits are saved.
- A Syntax ERROR occurs if the resulting constant is the wrong format.
- An Ma ERROR occurs if the resulting constant is mathematically incorrect or illegal.

■ Saving a Constant to Alpha Memory

1. Select the constant you want to save in Alpha memory and then press **[F2]** (STORE).

- This displays the “Store Alpha Mem.” dialog box.

```

----- Universal -----
C      =299792458
F0    =1.25663706E-6
N0    =8.8541878E-12
M0    =376.7303135
G0    =6.67408E-11
h      =6.62607E-34      ↓

```

Store Alpha Mem.
[A~Z]:

```

----- Universal -----
C      =299792458
F0    =1.25663706E-6
N0    =8.8541878E-12
M0    =376.7303135
G0    =6.67408E-11
h      =6.62607E-34      ↓

```

[EDIT] [STORE] [DETAIL] [KEEP] [INIT] [A~Z]

2. Enter a letter that represents the Alpha memory where you want to store the constant.

3. Press **[EXE]**.

- Now when you recall the applicable Alpha memory in the **RUN • MAT** mode, the calculator will input the constant stored there.

■ Details Dialog Box

- Selecting a constant and pressing **[F3]** (DETAIL) or **[EXE]** will display a dialog box with details about the selected constant.
- The details dialog box shows the constant name, symbol, and unit.



■ Returning All Constants to their Initial Default Values

- While the constant list screen is on the display, press **[F6]** (A•INIT) to display the Init All dialog box.



- Press **[F1]** (Yes) to return all of the constants to their initial default values.

5 Calling Physium Functions from an eActivity

You can call Physium functions from an eActivity by including a “Physium strip” in the eActivity file.

This section explains how to insert a Physium strip into an eActivity file, and how to use inserted a Physium strip. For details about eActivity operations, see “Chapter 10 eActivity” in the manual that comes with the calculator.

■ Inserting a Physium Strip into an eActivity File

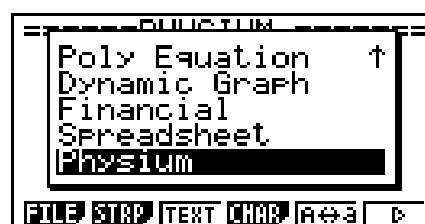
The following procedure assumes that the eActivity file into which you want to insert the Physium strip is already open.

● To insert a Physium Strip into an eActivity file

1. On the eActivity workspace screen, move the cursor the location where you want to insert the Physium strip.
2. Press **F2** (STRP).
 - This will display a dialog box with a list of insertable strips.



3. Use **▲** and **▼** to move the highlighting to the Physium strip you want to insert.



4. Press **EXE**.
 - The strip is inserted above the line or the strip where the cursor is currently located.



5. Enter up to 16 characters for the strip title.
6. Press **[EXE]** to assign the title to the strip.



- This will highlight the strip.
- You can execute the strip here by pressing **[EXE]**. For details about operations that are required when you execute a strip, see “Calling a Physium Function from a Physium Strip” below.

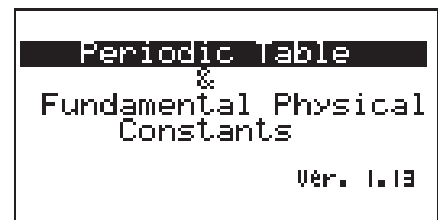
■ Calling a Physium Function from a Physium Strip

This section explains operations for Physium strip that can be inserted into an eActivity file. The following procedure assumes that the applicable Physium strip has already been inserted into an eActivity file that is currently open.

1. On the eActivity workspace screen, use the **▲** and **▼** keys to move the highlighting to the Physium strip.



2. Press **[EXE]**.
 - This launches the Physium and displays the initial screen.



3. Perform the procedure under “Starting Up Physium” (page 2-1) from step 3.
4. To return to the eActivity workspace screen, press **[SHIFT]** **[→]** (**🔒**).

6 Precautions

- You can save screen captures of Physium screens and dialog box. For details, see “1-9 Using Screen Capture” in the manual that comes with the calculator.
- Note that the Catalog Function described in the manual that comes with the calculator is not supported by Physium.
- The atomic weights in this application are based on those recommended by 2017 IUPAC (International Union of Pure and Applied Chemistry).
- The physical constants in this application are based on those recommended by the 2014 CODATA.
- A scientific constant in this application may be slightly different depending on the year it is presented or books it is presented. Refer to appropriate information that fits your purpose before use.
- The classification of rare earth elements may be slightly different from those in typical textbooks or magazines in the USA.
- The classification of transition elements may be slightly different from those in typical textbooks or magazines in the USA.

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