

Faculty of MECHANICAL ENGINEERING

Department of TECHNICAL THERMODYNAMICS

Property Library for the Industrial Formulation IAPWS-IF97 for Water and Steam

FluidTI with LibIF97 for TI Nspire[™] CX TI Nspire[™] CAS TI Nspire[™] CX CAS

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Software for the Industrial-Formulation IAPWS-IF97 for Water and Steam

FluidTl

for the TI Nspire[™] CX, TI Nspire[™] CAS and TI Nspire[™] CAS CX Pocket Calculators

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For further pocket calculator software see the following link:

www.steamtables-pocket-calculators.com

For steam tables and further property libraries for $Excel^{\ensuremath{\mathbb{R}}}$ and $Mathcad^{\ensuremath{\mathbb{R}}}$ see the following link:

www.international-steam-tables.com

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1. Property Functions for Water and Steam

The FluidTI program for the TI Nspire[™] CAS Pocket Calculator contains the following functions for the calculation of thermodynamic properties of water and steam:

Functional Dependence	Name in FluidTl	Property or Function	Unit
$p_{\rm S} = f(t)$	ps_t_97	Saturation pressure	MPa
$t_{\rm s} = f(p)$	ts_p_97	Saturation temperature	°C
v = f(p, t, x)	v_ptx_97	Specific volume	m ³ /kg
h = f(p, t, x)	h_ptx_97	Specific enthalpy	kJ/kg
s = f(p, t, x)	s_ptx_97	Specific entropy	kJ/(kg ⁺ K)
t = f(p, h)	t_ph_97	Backward function: temperature from pressure and enthalpy	°C
x = f(p, h)	x_ph_97	Backward function: vapor fraction from pressure and enthalpy	kg/kg
t = f(p, s)	t_ps_97	Backward function: temperature from pressure and entropy	°C
x = f(p,s)	x_ps_97	Backward function: vapor fraction from pressure and entropy	kg/kg
$\eta = f(p, t, x)$	eta_ptx_97	Dynamic viscosity	Pa s = kg/(m s)
$\lambda = f(p, t, x)$	lambda_ptx_97	Thermal conductivity	W/(m [·] K)
$c_p = f(p, t, x)$	cp_ptx_97	Specific isobaric heat capacity	kJ/(kg ⁺ K)

Units: p in MPa

t in °C

x in kg saturated steam / kg wet steam

Range of Validity: IAPWS-IF97 regions 1 and 2, including wet steam

Pressure: from 0.000611 MPa up to 100 MPa Temperature: from 0 °C up to 800 °C

Comment on the vapor fraction x and calculations for wet steam

The subprograms automatically deal with the wet steam region. For this purpose the following specifications for the vapor fraction *x* are to be regarded:

If the state point to be calculated is located within the single-phase regions (of liquid or superheated steam) enter either no input value, or -1 for the value of x. In this case, the backward functions will result in x = -1.

If the state point to be calculated is located within the two-phase region (wet steam), values between 0 and 1 have to be entered for the value of x (x = 0 for saturated liquid, x = 1 for saturated vapor). The backward functions result in values for x that range from 0 to 1.

In the case of wet steam it is adequate to put in either the value given for t and p = -1 (or no input value for p), or the given value for p and t = -1 (or no input value for t), as well as the value for x that ranges from 0 to 1.

If you enter p and t and x when calculating wet steam, the program considers p and t to meet the vapor-pressure curve. If this does not apply, an error message is displayed for the selected function to be calculated.

IAPWS-F97 wet steam region:

t = 0 °C ... t = 373.946 °C p = 0.000611 MPa ... p = 22.064 MPa

Please note.

If the calculation results in -1, values have been entered outside the range of validity of the IAPWS-IF97 or they do not define a state point. In this case, an error message is displayed.

2. Range of Validity and the FluidTl Program Structure

The International Association for the Properties of Water and Steam (IAPWS) issued the IAPWS-IF97 Industrial Formulation for Thermodynamic Properties of Water and Steam in 1997. This standard must be applied worldwide in final warranty calculations for power plants which use water and steam as working fluid.

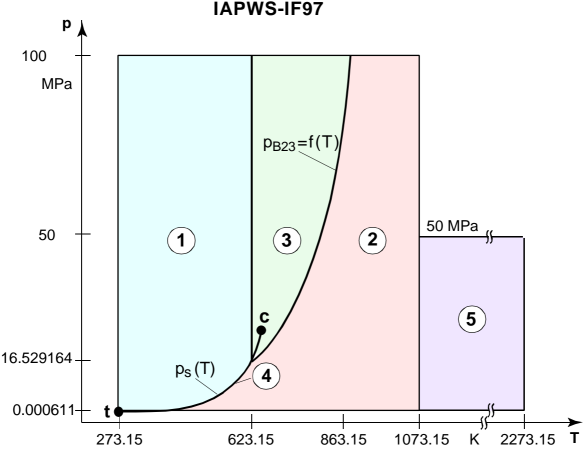
Figure 1 shows the range of validity of the equation set of the Industrial Formulation, fully named

"IAPWS Industrial Formulation 1997 for the Thermodynamic Properties of Water and Steam",

abbreviated

"IAPWS-IF97".

The IAPWS-IF97 range of state includes temperatures from 0 °C up to 800 °C at pressures from 0.000611 up to 100 MPa and temperatures up to 2000 °C at pressures up to 50 MPa.





Internally, the entire range of validity is subdivided into five calculation regions in which the appropriate constitutive equations apply (cf. Fig. 1). Those are described in detail in the official IAPWS [1] release and in the publications by *Wagner et al.* [2] and [3].

The FluidTl version at hand is valid for the regions 1, 2, and the wet steam region (region 4) up to a pressure of 16.529164 MPa (cf. Figure1). Their connection to the calculation equations is established according to the given quantities.

3. Application of FluidTl

3.1 Installation of FluidTl onto the Tl Nspire[™] Pocket Calculators

In order to run the FluidTl software on your pocket calculator, load the program from your computer into the calculator, using a special link program and the appropriate link cable.

The link program is available at service partners of Texas Instruments[®], as well as the link cable, or can be downloaded from the TI web site:

http://education.ti.com/en/us/software/details/en/79881F1D740D4A92ADB342087887BE55/tinspirecas_pc_full.

The following description is valid for the

TI-Nspire[™] CAS Student Software

link program which should have already been installed onto your computer.

(For another link program used for the file transfer see the corresponding user's guide or online-help.)

1. Insert the FluidTI CD into your CD-ROM drive. The CD contains the file listed in the following table:

TI	TI-Nspire™	TI-Nspire™	TI-Nspire™
Model	CAS	CX	CAS CX
Group File	Water.tns	Water.tns	Water.tns

2. Connect the TI calculator to the PC using the USB cable.

3. Data Transfer with the TI-Nspire[™] CAS Student Software

Make sure the TI calculator is switched on.

- Start the TI-Nspire[™] CAS Student Software program on your computer.
- The "Welcome Screen" pops up, which can be closed.
- Click the "Content Explorer" in the "Documents Toolbox".

In this toolbox there are now two windows. The upper one displays your Computer and the other the connected handhelds.

- In the "Computer" window, you can now search the "water.tns" file to copy it by rightclicking.
- After you have selected the folder "MyLib" in the lower window, insert the file by rightclicking and paste.

The last step before the program can be used, is to refresh the libraries. You do this in the main screen, after you opened "My Documents" press the menu button and select "Refresh Libraries".

3.2 Example: Calculation of h = f(p, t, x) using FluidTl

The specific enthalpy *h* as a function of pressure *p*, temperature *t* and vapor fraction *x* can be calculated for the Industrial Formulation IAPWS-IF97 on the TI NspireTM. Please carry out the following steps:

- If the "Calculator" Page is active, press the "catalog"-button and choose the library by clicking the library-symbol and select the water file.
- Now, choose "h_ptx_97" and press the <ENTER> key.
- The command line "water\h_ptx_97() is displayed. Press <ENTER>.
 The selected sub program of FluidTI is displayed:

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water\h_ptx	_97()	<
		.
S	pecific enthalpy h=f(p,T,x)	
	ок	
2		0/99

- The following three menus request the input of parameters:

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т [°С]: 1		
x [kg/kg]: 1		
h = 2502.73	p [MPa]:	
save as:		
	OK Cancel	Done
water\h_ptx_97	0	
Specific enth	alpy h=f(p,T,x)	×
		1/99
< <u>191</u> ►	*Unsaved 🗢	4 <mark>1</mark> 🔀
water\h_ptx_	97()	
Specific ent	т [*с]:	
p [MPa]: 10		
	OK Cancel	
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		0/99
₹ 1.1 ►	*Unsaved 🗢	1 🕅 🔀
water\h_ptx_	97()	
	5 4050	
Specific er		
p [MPa]: 1 .	x [kg/kg]:	
T [°C]: 400	OK Cancel	
		0/99

- Enter the given value for "p in MPa" into the appropriate window. Please consider the IAPWS-IF97 range of validity:

p = 0.000611 MPa ... 100 MPa.

 \rightarrow E.g.: Enter the value 10 and move the cursor to the next input field.

 Enter the value given for "t in °C" into the appropriate window. Please consider the IAPWS-IF97 range of validity:

t = 0 °C ... 800 °C.

 \rightarrow E.g.: Enter the value 400, and move the cursor to the next input field.

- Now, the value for the vapor fraction *x* in (kg saturated steam/kg wet steam) is to be entered into the appropriate window. Consider the following specifications:

If the state point to be calculated is located within the single-phase regions (of liquid or superheated steam) enter either no input value, or -1 for the value of x. In this case, the backward functions will result in x = -1.

If the state point to be calculated is located within the two-phase region (wet steam), values between 0 and 1 have to be entered for the value of x (x = 0 for saturated liquid, x = 1 for saturated vapor). The backward functions result in values for x that range from 0 to 1.

In the case of wet steam it is adequate to put in either the value given for t and p = -1 (or no input value for p), or the given value for p and t = -1 (or no input value for t), as well as the value for x that ranges from 0 to 1.

If you enter p and t and x when calculating wet steam, the program considers p and t to meet the vapor-pressure curve. If this does not apply, an error message is displayed for the selected function to be calculated.

IAPWS-IF97 wet steam region:

t = 0 °C ... t = 373.946 °C p = 0.000611 MPa ... p = 22.064 MPa

→ Press <ENTER> only, because in the example the state point is located in the singlephase region.

Now the calculation starts.

- After the calculation has been finished the result for *h* in kJ/kg is displayed:

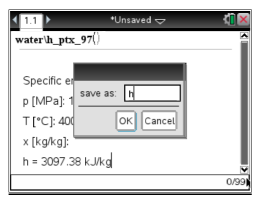
◀ 1.1 ▶	*Unsaved 🗢	
water\h_ptx_ Specific en p [MPa]: 10 T [°C]: 400 x [kg/kg]:	97() h = 3097.38 kJ/kg	
		0/99

 \rightarrow In the example, the function calculation results in 3097.38 kJ/kg.

- The calculation of h = f(p, t, x) is now complete.

The calculated value for h can be saved as a variable and used in other calculations with the pocket computer.

 \rightarrow E.g.: Enter the variable name "h" into the "save as:" window and press <ENTER> twice:



Note.

For the variable any name can be chosen, except for those which start with the ω (Omega)-symbol and system variables (cf. TI Handbook).

In general, variables beginning with the ω -symbol must not be used within the FLUIDTI programs.

3.3 Uninstalling FluidTl

Please carry out the following steps:

1. Open the menu "My Documents" in the main menu.

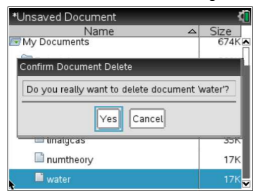
The following menu will be displayed:

*Unsaved Document	.0	ব্
Name		Size
Unsaved Document		
🖻 My Documents		674K
🗇 Beispiele		390K
📴 Bilder		ок
🖻 Examples		110K
🗇 MyLib		175K
2		

2. Select "MyLib" and press enter.

*Unsaved Document	4
Name	△ Size
🖅 My Documents	674K
🖻 Beispiele	390K
🗇 Bilder	ок
🗇 Examples	217K
🗁 MyLib	69K
🗋 linalgcas	35K
numtheory	17K
📫 water	17K

3. Press the "del" button after selecting the water file and confirm the following query.



FluidTI has been removed successfully.

4. Program Documentation

Saturation Pressure $p_s = f(t)$

Name in FluidTI: ps_t_97

Input

t - Temperature t in °C

Output

ps(t) - Saturation pressure p_s in MPa

Range of validity

from $t_{\rm t} = 0$ °C up to $t_{\rm c} = 373.946$ °C

Response on faulty input values

Error message "Out of Range!" for the following input values:

t < 0 °C or *t* > 373.946 °C

Saturation Temperature $t_s = f(p)$

Name in FluidTI: ts_p_97

Input

p - Pressure of *p* in MPa

Output

ts(p) - Saturation temperature t_s in °C

Range of validity

from $p_{t} = 0.000611$ MPa up to p = 22.064 MPa

Response on faulty input values

Error message "Out of Range!" for the following input values:

p < 0.000611 MPa or *p* > 22.064 MPa

Specific Volume v = f(p, t, x)

Name in FluidTI: v_ptx_97

Input

- **p** Pressure *p* in MPa
- t Temperature t in °C
- **x** Vapor fraction *x* in (kg saturated steam)/(kg wet steam)

Output

v(p,t,x) - Specific volume v in m³/kg

Range of validity

Liquid region:	p= p _s (t)100 MPa at 0 °C350 °C
-	0.000611 MPap=p _s (t) at 0 °C350 °C 0.000611 MPap ₂₃ (t) = p(s=5.2 kJ/(kg K)) at 350 °C590 °C 0.000611 MPa100 MPa at 590 °C800 °C

Comment on the vapor fraction x and calculations for wet steam

The subprograms automatically deal with the wet steam region. For this purpose the following specifications for the vapor fraction *x* are to be regarded:

- If the state point to be calculated is located within the single-phase region (of liquid or superheated steam) enter the value x = -1. Pressure *p* and temperature *t* are given.
- When calculating wet steam the value of x to be entered ranges from 0 to 1 (in case of boiling liquid: x = 0; in case of saturated vapor: x = 1).

In the case of wet steam it is adequate to put in either the value given for t and p = -1, or the given value for p and t = -1, as well as the value for x which ranges from 0 to 1.

If you enter p and t and x when calculating wet steam, the program considers p and t to meet the vapor-pressure curve. If this does not apply, the selected function to be calculated results in -1.

Wet steam region:t = 0 °C ... 350 °C p = 0.000611 MPa ... 16.5292 MPa

Response on faulty input values

Error message "Out of Range!" for the following input values:

Single-phase region: the parameters entered are located outside the above mentioned (x = -1) range of validity

Wet steam region:

($0 \le x \le 1$) at t = -1 and p > 16.5292 MPa or p < 0.000611 MPa or at p > 16.5292 MPa or p < 0.000611 MPa and $t > 350^{\circ}$ C or $t < 0^{\circ}$ C at $| t - t_{s}(p) | > 0.1$ K

Specific Enthalpy h = f(p, t, x)

Name in FluidTI: h_ptx_97

Input

- \mathbf{p} Pressure p in MPa
- t Temperature t in °C
- **x** Vapor fraction *x* in (kg saturated steam)/(kg wet steam)

Output

h(p,t,x) - Specific enthalpy h in kJ/kg

Range of validity

Liquid region:	p= p _s (t)100 MPa at 0 °C350 °C
Steam region:	0.000611 MPap=p _s (t) at 0 °C350 °C
	0.000611 MPap ₂₃ (t) = $p(s=5.2 \text{ kJ/(kg K)})$ at 350 °C 590 °C
	0.000611 MPa100 MPa at 590 °C 800 °C

Comment on the vapor fraction x and calculations for wet steam

The subprograms automatically deal with the wet steam region. For this purpose the following specifications for the vapor fraction *x* are to be regarded:

- If the state point to be calculated is located within the single-phase region (of liquid or superheated steam) enter the value x = -1. Pressure *p* and temperature *t* are given.
- When calculating wet steam the value of *x* to be entered ranges from 0 to 1 (in case of boiling liquid: x = 0; in case of saturated vapor: x = 1).

In the case of wet steam it is adequate to put in either the value given for t and p = -1, or the given value for p and t = -1, as well as the value for x which ranges from 0 to 1.

If you enter p and t and x when calculating wet steam, the program considers p and t to meet the vapor-pressure curve. If this does not apply, the selected function to be calculated results in -1.

Wet steam region: $t = 0 \ ^{\circ}C \dots 350 \ ^{\circ}C$

p = 0.000611 MPa ... 16.5292 MPa

Response on faulty input values

Error message "Out of Range!" for the following input values:

Single-phase region: (<i>x</i> = -1)	the parameters entered are located outside the above mentioned range of validity
Wet steam region: $(0 \le x \le 1)$	at $p = -1$ and $t > 350^{\circ}$ C or $t < 0^{\circ}$ C or at $t = -1$ and $p > 16.5292$ MPa or $p < 0.000611$ MPa or at $p > 16.5292$ MPa or $p < 0.000611$ MPa and $t > 350^{\circ}$ C or $t < 0^{\circ}$ C at $ t - t_{s}(p) > 0.1$ K

Specific Entropy s = f(p, t, x)

Name in FluidTI: s_ptx_97

Input

- ${\bf p}$ Pressure p in MPa
- t Temperature t in °C
- **x** Vapor fraction *x* in (kg saturated steam)/(kg wet steam)

Output

s(p,t,x) - Specific entropy s in kJ/(kg K)

Range of validity

Liquid region:	p= p _s (t)100 MPa for 0 °C350 °C
Steam region:	0.000611 MPap=p _s (t) for 0 °C350 °C
	0.000611 MPap ₂₃ (t) = $p(s=5.2 \text{ kJ/(kg K)})$ for 350 °C 590 °C
	0.000611 MPa100 MPa for 590 °C 800 °C

Comment on the vapor fraction x and calculations for wet steam

The subprograms automatically deal with the wet steam region. For this purpose the following specifications for the vapor fraction *x* are to be regarded:

- If the state point to be calculated is located within the single-phase region (of liquid or superheated steam) enter the value x = -1. Pressure p and temperature t are given.
- When calculating wet steam the value of *x* to be entered ranges from 0 to 1 (in case of boiling liquid: x = 0; in case of saturated vapor: x = 1).

In the case of wet steam it is adequate to put in either the value given for t and p = -1, or the given value for p and t = -1, as well as the value for x which ranges from 0 to 1.

If you enter p and t and x when calculating wet steam, the program considers p and t to meet the vapor-pressure curve. If this does not apply, the selected function to be calculated results in -1.

Wet steam region:t = 0 °C ... 350 °C

p = 0.000611 MPa ... 16.5292 MPa

Response on faulty input values

Error message "Out of Range!" for the following input values:

Single-phase region: the parameters entered are located outside the above mentioned (x = -1) range of validity

Wet steam region:at p = -1 and $t > 350^{\circ}$ C or $t < 0^{\circ}$ C or $(0 \le x \le 1)$ at t = -1 and p > 16.5292 MPa or p < 0.000611 MPa orat p > 16.5292 MPa or p < 0.000611 MPaand $t > 350^{\circ}$ C or $t < 0^{\circ}$ Cat $| t - t_s(p) | > 0.1$ K

Backward Function: Temperature t = f(p,h)

Name in FluidTI: t_ph_97

Input

p - Pressure *p* in MPa

h - Specific enthalpy h in kJ/kg

Output

t(p,h) - Temperature t in °C

Range of validity

Liquid region:	IAPWS-IF97 region 1 (Figure 1)
Steam region:	IAPWS-IF97 region 2 (Figure 1)
Wet steam region:	p = 0.000611 16.5292 MPa and h'(p) < h < h"(p)

Comment on the calculations for wet steam

The subprogram automatically calculates the wet steam region. That is, the program checks, with reference to the given values of p and h, whether the state point to be calculated is located in the single-phase region (liquid or steam) or the wet steam region. The calculation is carried out for the appropriate region.

Response on faulty input values

Error message "Out of Range!" for the following input values:

Liquid region:	at values of <i>p</i> and <i>h</i> beyond the IAPWS-IF97 region 1 (Figure 1)
Steam region:	at values of <i>p</i> and <i>h</i> beyond the IAPWS-IF97 region 2 (Figure 1)
Wet steam region:	at values of $p > 16.5292$ MPa or $p < 0.000611$ MPa

Backward Function: Temperature t = f(p,s)

Name in FluidTI: t_ps_97

Input

p - Pressure *p* in MPa

s - Specific entropy s in kJ/(kg K)

Output

t(p,s) - Temperature t in °C

Range of validity

Liquid region:	IAPWS-IF97 region 1 (Figure 1)
Steam region:	IAPWS-IF97 region 2 (Figure 1)
Wet steam region:	p = 0.000611 16.5292 MPa

Comment on the calculations for wet steam

The subprogram automatically calculates the wet steam region. That is, the program checks, with reference to the given values of p and h, whether the state point to be calculated is located in the single-phase region (liquid or steam) or the wet steam region. The calculation is carried out for the appropriate region.

Response on faulty input values

Error message "Out of Range!" for the following input values:

Liquid region:	at values of <i>p</i> and <i>s</i> beyond the IAPWS-IF97 region 1 (Figure 1)
Steam region:	at values of <i>p</i> and <i>s</i> beyond the IAPWS-IF97 region 2 (Figure 1)
Wet steam region:	at values of $p > 16.5292$ MPa or $p < 0.000611$ MPa

Backward Function: Vapor Fraction x = f(p,h)

Name in FluidTI: x_ph_97

Input

- **p** Pressure *p* in MPa
- h Specific enthalpy h in kJ/kg

Output

x(p,h) - Vapor fraction *x* in (kg saturated steam)/(kg wet steam)

Range of validity

Liquid region:	IAPWS-IF97 region 1 (Figure 1)
Steam region:	IAPWS-IF97 region 2 (Figure 1)
Wet steam region:	p = 0.000611 16.5292 MPa and h'(p) < h < h"(p)

Comment on the calculations for wet steam

The subprogram automatically calculates the wet steam region. That is, the program checks, with reference to the given values of p and h, whether the state point to be calculated is located in the single-phase region (liquid or steam) or the wet steam region. In case of wet steam the value of x is calculated. If the state point to be calculated is located in the single-phase region the vapor fraction result is x = -1.

Response on faulty input values

If the state point to be calculated is located in the single-phase region the result x(p,h) = -1 is displayed for the following input values:

p > 16.5292 MPa or *h* < *h*′(*p*) or *h* > *h*″(*p*)

Error message "Out of Range!" for the following input values: p < 0.000611 MPa or p > 100 MPa

Backward function: Vapor Fraction x = f(p,s)

Name in FluidTI: x_ps_97

Input

p - Pressure *p* in MPa

s - Specific entropy s in kJ/(kg K)

Output

x(p,s) - Vapor fraction x in (kg saturated steam)/(kg wet steam)

Range of validity

Liquid region:	IAPWS-IF97 region 1 (Figure 1)
Steam region:	IAPWS-IF97 region 2 (Figure 1)
Wet steam region:	p = 0.000611 16.5292 MPa and s'(p) < s < s"(p)

Comment on the calculations for wet steam

The subprogram automatically calculates the wet steam region. That is, the program checks, with reference to the given values of p and h, whether the state point to be calculated is located in the single-phase region (liquid or steam) or the wet steam region. In case of wet steam the value of x is calculated. If the state point to be calculated is located in the single-phase region the vapor fraction result is x = -1.

Response on faulty input values

If the state point to be calculated is located in the single-phase region the result x(p,h) = -1 occurs for the following input values:

p > 16.5292 MPa or s < s'(p) or s > s''(p)

Error message "Out of Range!" for the following input values: p < 0.000611 MPa or p > 100 MPa

Dynamic Viscosity $\eta = f(p, t, x)$

Name in FluidTI: eta_ptx_97

Input

- **p** Pressure *p* in MPa
- **t** Temperature t in °C
- **x** Vapor fraction *x* in (kg saturated steam)/(kg wet steam)

Output

 η (**p**,**t**,**x**) - Dynamic viscosity η in MPa s

Range of validity

Liquid region:	p= p _s (t)100 MPa for 0 °C350 °C
Steam region:	0.000611 MPap=p _s (t) for 0 °C350 °C
	0.000611 MPap ₂₃ (t) = $p(s=5.2 \text{ kJ/(kg K)})$ for 350 °C 590 °C
	0.000611 MPa100 MPa for 590 °C 800 °C

Comment on the vapor fraction x and calculations for wet steam

The subprograms automatically deal with the wet steam region. For this purpose the following specifications for the vapor fraction *x* are to be regarded:

- If the state point to be calculated is located within the single-phase region (of liquid or superheated steam) enter the value x = 1. Pressure *p* and temperature *t* are given.
- When calculating wet steam the value of *x* to be entered ranges from 0 to 1 (in case of boiling liquid: x = 0; in case of saturated vapor: x = 1).

In the case of wet steam it is adequate to put in either the value given for t and p = -1, or the given value for p and t = -1, as well as the value for x which ranges from 0 to 1.

If you enter p and t and x when calculating wet steam, the program considers p and t to meet the vapor-pressure curve. If this does not apply, the selected function to be calculated results in -1.

Saturated liquid and saturated vapor:

t = 0 °C ... 350 °C p = 0.000611 MPa ... 16.5292 MPa

Response on faulty input values

Error message "Out of Range!" for the following input values:

Single-phase region: the parameters
(x = -1)entered are located outside the above mentionedSaturated liquid and saturated vapor:
(x = 0 or x = 1)at p = -1 and $t > 350^{\circ}$ C or $t < 0^{\circ}$ C or
at t = -1 and p > 16.5292 MPa or p < 0.000611 MPa or
at p > 16.5292 MPa or p < 0.000611 MPa

and
$$t > 350^{\circ}$$
 C or $t < 0^{\circ}$ C
at $| t-t_{s}(p) | > 0.1$ K

References: [5], internal calculation of ρ or v: [1], [2], [3]

Thermal Conductivity $\lambda = f(p, t, x)$

Name in FluidTI: lambda_ptx_97

Input

- ${\bf p}$ Pressure p in MPa
- ${\bf t}$ Temperature t in °C
- **x** Vapor fraction *x* in (kg saturated steam)/(kg wet steam)

Output

lam(p,t,x) - Thermal conductivity λ in W/(m K)

Range of validity

Liquid region:	p= p _s (t)100 MPa for 0 °C350 °C
Steam region:	0.000611 MPap=p _s (t) for 0 °C350 °C
	0.000611 MPap ₂₃ (t) = p(s=5.2 kJ/(kg K)) for 350 °C 590 °C
	0.000611 MPa100 MPa for 590 °C 800 °C

Comment on the vapor fraction x and calculations for wet steam

The subprograms automatically deal with the wet steam region. For this purpose the following specifications for the vapor fraction *x* are to be regarded:

- If the state point to be calculated is located within the single-phase region (of liquid or superheated steam) enter the value x = 1. Pressure p and temperature t are given.
- When calculating wet steam the value of *x* to be entered ranges from 0 to 1 (in case of boiling liquid: x = 0; in case of saturated vapor: x = 1).

In the case of wet steam it is adequate to put in either the value given for t and p = -1, or the given value for p and t = -1, as well as the value for x which ranges from 0 to 1.

If you enter p and t and x when calculating wet steam, the program considers p and t to meet the vapor-pressure curve. If this does not apply, the selected function to be calculated results in -1.

Saturated liquid and saturated vapor: t = 0 °C ... 350 °C

p = 0.000611 MPa ... 16.5292 MPa

Response on faulty input values

Error message "Out of Range!" for the following input values:

Single-phase region: the parameters entered are located outside the above mentioned (x = -1) range of validity

Saturated liquid and saturated vapor: (x = 0 or x = 1) at p = -1 and $t > 350^{\circ}$ C or $t < 0^{\circ}$ C or at t = -1 and p > 16.5292 MPa or p < 0.000611 MPa or at p > 16.5292 MPa or p < 0.000611 MPa and $t > 350^{\circ}$ C or $t < 0^{\circ}$ C at $| t - t_{s}(p) | > 0.1$ K

References: [5], internal calculation of ρ or v: [1], [2], [3]

Specific Isobaric Heat Capacity $c_p = f(p, t, x)$

Name in FluidTI: cp_ptx_97

Input

- **p** Pressure *p* in MPa
- t Temperature t in °C
- **x** Vapor fraction *x* in (kg saturated steam)/(kg wet steam)

Output

cp(p,t,x) - Specific isobaric heat capacity cp in kJ/kg K

Range of validity

Liquid region:	p= p _s (t)100 MPa at 0 °C350 °C
Steam region:	0.000611 MPap=p _s (t) at 0 °C350 °C
	0.000611 MPap ₂₃ (t) = p(s=5.2 kJ/(kg K) at 350 °C 590 °C
	0.000611 MPa100 MPa at 590 °C 800 °C

Comment on the vapor fraction x and calculations for wet steam

The subprograms automatically deal with the wet steam region. For this purpose the following specifications for the vapor fraction *x* are to be regarded:

- If the state point to be calculated is located within the single-phase region (of liquid or superheated steam) enter the value x = -1. Pressure *p* and temperature *t* are given.
- When calculating wet steam the value of *x* to be entered ranges from 0 to 1 (in case of boiling liquid: x = 0; in case of saturated vapor: x = 1).

In the case of wet steam it is adequate to put in either the value given for t and p = -1, or the given value for p and t = -1, as well as the value for x which ranges from 0 to 1.

If you enter p and t and x when calculating wet steam, the program considers p and t to meet the vapor-pressure curve. If this does not apply, the selected function to be calculated results in -1.

Saturated liquid and saturated vapor: $t = 0 \ ^{\circ}C \dots 350 \ ^{\circ}C$ $p = 0.000611 \text{ MPa} \dots 16.5292 \text{ MPa}$

Response on faulty input values

Error message "Out of Range!" for the following input values:

Single-phase region: the parameters entered are located outside the above mentioned (x = -1) range of validity

Saturated liquid and saturated vapor: (x = 0 or x = 1) at p = -1 and $t > 350^{\circ}$ C or $t < 0^{\circ}$ C or at t = -1 and p > 16.5292 MPa or p < 0.000611 MPa or at p > 16.5292 MPa or p < 0.000611 MPa and $t > 350^{\circ}$ C or $t < 0^{\circ}$ C at $|t-t_{s}(p)| > 0.1$ K

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