

ABB MEASUREMENT & ANALYTICS | FUNCTION REFERENCE MANUAL

Spirit^{IT} Flow-X High accuracy flow computers



Certified flow calculations Flow and batch calculations Worksheet functions

Measurement made easy

Flow-X/P with Flow-X/M module

Introduction

Welcome to the exciting world of Spirit^{IT} Flow-X! This document describes the spreadsheet functions for the Spirit^{IT} Flow-X series and background information on related standards and calculation methods used in the industry for quality and quantity measurement of hydrocarbon and other type of fluids.

There are three reference manuals:

- Volume I This Installation manual, with the installation instructions.
- Volume II The Operation and Configuration manual. This manual consists of a general part and one of the following application-specific parts:
 - IIA Operation and configuration
 - IIB Gas Metric application
 - IIC Liquid Metric application
 - IID Gas US customary units application
- IIE Liquid US customary units application
 Volume III The manuals for solutions that exceed our standard applications. This volume consists of 1 part:

For more information

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1 Introduction

Flow-X function library

The Spirit^{IT} Flow-X series of flow computer uses Microsoft Excel as its configuration environment. Each Flow-X application consists of a single Excel workbook that contains one or more worksheets.

Flow-X functions are configured as regular Excel functions. By using the output of one function as an input (argument) in another function a complete calculation scheme can be made. Functions can be defined on multiple sheets in order to organize the application.

API Petroleum Measurement Tables

History

The first version of the API Petroleum Measurement Tables was published in **1952**. In those days measurement readings were taken manually and the tables were used to convert the observed density or gravity at the observed temperature to the value at the reference temperature. So the table values were the actual standard.

The 1952 Tables consists of 58 tables containing all kind of correction and conversion factors used in the measurement of hydrocarbon liquids. Each table deals with a particular conversion of units, correction of density, or correction of volume. The 1952 tables that have to do with the conversion of density and volume are: 5, 6, 23, 24, 53 and 54.

Table 5, 6, 23 and 24 convert density or volume to or from to a reference temperature of 60° F, while tables 53 and 54 refer to 15°C.

In **1980** a complete new set of tables was published together with computer routines to allow electronic devices to automatically calculate the volume conversion factors and API gravity / (relative) density at the reference temperature. Back then most electronic devices were not capable of performing double-precision floating point calculations, so the standard prescribed all kind of rounding and truncating rules to make sure that the calculations would always provide the same result. For the 1980 version the calculation procedures are the standard rather than the table values.

In the 1980 version, which is also referred to as **API-2540**, the tables are divided into 3 product groups and a letter designation was used to distinguish between the sub-tables. "A" was used for crude oil, "B" for refined products and "C" for special applications. The 1980 tables, however, did not cover the LPGs and NGLs density ranges and the 1952 Tables were left valid for these products. Furthermore, the lubricating oil tables (designated as "D") were not complete at the time of the printing in 1980 and were released two years later. As opposed to the A, B and C tables no implementation procedures were defined for the D tables.

In 1988 the Institute of Petroleum released its Paper No. 3 with tables 59 and 60 that are based on a reference temperature of 20°C.

This resulted in the following Petroleum Measurement Tables dealing with the conversion of volume and density to and from a reference temperature.

Number	Title
5	API Gravity Reduction to 60°F
6	Reduction of Volume to 60°F Against API Gravity at 60°F
23	Reduction of Observed Specific Gravity to Specific Gravity 60/60°F
24	Reduction of Volume to 60o F Against Specific Gravity 60/60°F
53	Reduction of Observed Density to Density at 15°C
54	Reduction of Volume to 15°C Against Density at 15°C
59	Reduction of Observed Density to Density at 20°C
60	Reduction of Volume to 20°C Against Density at 20°C

In **2004** the API MPMS 11.1 1980 tables were superseded by a new set of tables primarily for the following reasons:

- API 11.1:2004 includes the correction for both temperature and pressure in one and the same algorithm
- Taken into account the progress in electronics (and for other reasons) the complex truncating and rounding rules were abandoned. Instead the calculation procedures use double-precision floating point math. The input and output values are still rounded in order to obtain consistent results.
- The convergence methods for the correction of observed density to base density have been improved.
- On-line density measurement by densitometers became common practice, requiring the pressure and temperature correction to be incorporated in one ands the same procedure
- The tables are extended in both temperature and density to cover lower temperatures and higher densities.
- The previous standard used a significant digit format which resulted in 4 or 5 decimal places depending on whether the observed temperature was above or below the reference temperature. The new standard prescribes 5 decimal places if or both cases.
- The IP paper No. 3 tables were added to accommodate conversion to 20°C.

Tables for lubricating oils including the implementation procedures are now part of the standard.

Volume correction for pressure

The API MPMS 11.1:1980 Tables only cover the correction for temperature. The correction for pressure was published in API MPMS standards 11.2.1 and 11.2.2.

The correction for pressure is to the atmospheric pressure or, for products within the lower density range, to the equilibrium vapor pressure.

To calculate the equilibrium vapor pressure an Addendum was added to API MPMS 11.2.2. This addendum is also known as **GPA TP-15** (1988). In September 2007 the addendum was replaced by a new API standard 11.2.5 and at the same time GPA TP-15 (1988) was updated with a new 2007 revision.

NGL and LPG tables

For NGL and LPG products volume correction tables 24E and 23E (at 60 °F) were published in **GPA TP-25** (1988), so the letter 'E" was used to distinguish the tables from the related API MPMS A, B, C and D tables.

GPA TP-25 has been superseded by **GPA TP-27** / API MPMS 11.2.4 (2007), which includes tables 53E, 54E, 59E and 60E to convert to 15°C and 20°C as well. All text from TP-25 is included without technical change, so TP-25 is still viable for conversion to and from 60 °F.

Overview of hydrocarbon liquid conversion standards

- ASTM-IP Petroleum Measurement Tables, Historical Edition, 1952
- API MPMS Chapter 11.1 1980* (Temperature VCFs for Generalized Crude Oils, Refined Products, and Lubricating Oils): Historical; Published in 14 separate volumes

Also known as

- API Standard 2540 (API-2540)
- ASTM D1250
- IP 200
- In 1982 chapters XIII and XIV were published containing tables 5D, 6D, 53D and 54D for lubricating oils.
- API MPMS Chapter 11.1 2004 (Temperature & Pressure VCFs for Generalized Crude Oils, Refined Products and Lube Oils)
- API MPMS Chapter 11.2.1- 1984 (Compressibility Factors for Hydrocarbons: 0-90°API): Historical: now incorporated into Chapter 11.1-2004
- API MPMS Chapter 11.2.1M- 1984 (Compressibility Factors for Hydrocarbons: 638-1074 kg/m3): Historical: now incorporated into Chapter 11.1-2004
- API MPMS Chapter 11.2.2 1984 (Compressibility Factors for Hydrocarbons: 0.350-0.637 Relative Density and -50°F to 140°F)
- API MPMS Chapter 11.2.2M 1986 (Compressibility Factors for Hydrocarbons: 350-637 kg/m3 Density (15°C) and -46°C to 60°C)

Overview of the functions

The following table lists the volume conversion functions for hydrocarbon liquids as provided by the Flow-X series of flow computer.

Function	Temperature correction	Pressure correction	Input	Output
ASTM-IP Petroleum Mea	surement Tables 1952 - American Ed	ition		
API_Table23 (1952)	Table 23 - Specific Gravity Reduc	tion to 60 °F	SG (T)	SG (60°F)
API_Table24 (1952)	Table 24 - Volume Reduction to	50 °F	SG (60°F)	Ctl
Crude Oils, Refined Proc	ducts and Lubricating Oils (API MPMS	11.1:1980 / API-2540)		
API_Table5 (1980)	API 11.1:1980 Tables 5A, 5B and 5D	API 11.2.1:1984	°API (T, P)	°API (60°F, Pe)
API_Table6 (1980)	API 11.1:1980 Tables 6A, 6B and 6D	API 11.2.1:1984	°API (60°F, Pe)	°API (T, P)
API_Table23 (1980)	API 11.1:1980 Tables 23A and 23B	API 11.2.1:1984	RD (T, P)	RD (60°F, Pe)
API_Table24 (1980)	API 11.1:1980 Tables 24A and 24B	API 11.2.1:1984	RD (60°F, Pe)	RD (T, P)
API_Table53 (1980)	API 11.1:1980 Tables 53A, 53B and 53D	API 11.2.1M:1984	Density (T, P)	Density (15°C, Pe)
API_Table54 (1980)	API 11.1:1980 Tables 54A, 54B and 54D	API 11.2.1M:1984	Density (15°C, Pe)	Density (T, P)
Crude Oils, Refined Proc	ducts and Lubricating Oils (API MPMS	11.1:2004)		
API_Table5 (2004)	API 11.1:2004	API 11.1:2004	°API (T, P)	°API (60°F, 0 psig)
API Table6 (2004)	API 11.1:2004	API 11.1:2004	°API (60°F, 0 psig)	°API (T, P)

- API MPMS Chapter 11.2.2A 1984 (Addendum to Correlation of Vapor Pressure Correction for NGL): Superseded by Chapter 11.2.5
- API Publication/GPA TP-25/ASTM Publication (Temperature Correction for the volume of Light Hydrocarbons – Tables 24E and 23E: Superseded by API MPMS Chapter 11.2.4 GPA TP-25 was published in 1998 and replaced the 1952 tables 23, 24 for Light Hydrocarbon Liquids and GPA Technical Publication TP-16, which were previously used for volumetric measurement of LPG.
- API MPMS Chapter 11.2.4 2007 / GPA TP-27 / ASTM Publication (Temperature Correction for the Volume of NGL and LPG – Tables 23E, 24E, 53E, 54E, 59E, 60E): Supersedes GPA TP-25
- API MPMS Chapter 11.2.5 2007 / GPA TP-15 / ASTM Publication (A Simplified Vapor Pressure Correlation for Commercial NGLs): Supersedes Addendum to Chapter 11.2.2 (11.2.2A)
- IP No. 3 1988 (Energy Institute (formerly Institute of Petroleum), Petroleum Measurement Paper No 3 Computer Implementation Procedures for Correcting Densities and Volumes to 20 C. Superseded by IP No.3 - 1997
- IP No. 3 1997 (Energy Institute (formerly Institute of Petroleum), Petroleum Measurement Paper No 3 Computer Implementation Procedures for Correcting Densities and Volumes to 20 C. Supersedes IP No.3 - 1988
- ISO 91-1 1982 Petroleum measurement tables Part 1: Tables based on reference temperatures of 15 °C and 60 °F.
 Superseded by ISO 91-1 1992.
- ISO 91-1 1992 Petroleum measurement tables Part 1: Tables based on reference temperatures of 15 °C and 60 °F. Supersedes ISO 91-1 1982.
- ISO 91-2 1991 Petroleum measurement tables Part 2: Tables based on reference temperatures of 20 °C
- OIML R 63 1994 Petroleum measurement tables

Function	Temperature correction	Pressure correction	Input	Output
API_Table23 (2004)	API 11.1:2004	API 11.1:2004	RD (T, P)	RD (60°F, 0 psig)
API_Table24 (2004)	API 11.1:2004	API 11.1:2004	RD (60°F, 0 psig)	RD (T, P)
API_Table53 (2004)	API 11.1:2004	API 11.1:2004	Density (T, P)	Density (15°C, 0 bar(g))
API_Table54 (2004)	API 11.1:2004	API 11.1:2004	Density (15°C, 0 bar(g))	Density (T, P)
API_Table59 (2004)	API 11.1:2004	API 11.1:2004	Density (T, P)	Density (20°C, 0 bar(g))
API_Table60 (2004)	API 11.1:2004	API 11.1:2004	Density (20°C, 0 bar(g))	Density (T, P)
API_Table6C (2004)	API 11.1:2004	Not applicable	Thermal expansion coefficient at 60°F	Ctl
NGL and LPG (API 11.2.4)				
API_Table23E	API 11.2.4: 2007	API 11.2.2:1986	RD (T, P)	RD (60°F, Pe)
	Table 23E	GPA TP-15:1988		
		GPA TP-15:2007		
API_Table24E	API 11.2.4: 2007	API 11.2.2:1986	RD (60°F, Pe)	RD (T, P)
	Table 24E	GPA TP-15		
API_Table53E	API 11.2.4: 2007	API 11.2.2:1986	Density (T, P)	Density (15°C, Pe)
	Table 53E	GPA TP-15		
API Table54E	API 11.2.4: 2007	API 11.2.2:1986	Density (15°C, Pe)	Density (T, P)
-	Table 53E	GPA TP-15		
API_Table59E	API 11.2.4: 2007	API 11.2.2M:1986	Density (T, P)	Density (20°C, Pe)
	Table 59E	GPA TP-15		
API_Table60E	API 11.2.4: 2007	API 11.2.2M:1986	Density (20°C, Pe)	Density (T, P)
	Table 60E	GPA TP-15		

Hydrometer Correction

The API MPMS 11.1 1980 Standard (API-2540) assumes that the API gravity or relative density is observed with a glass hydrometer. Therefore a correction may be applied for the change of volume of the glass hydrometer with temperature.

The hydrometer correction applies for tables 5A, 5B, 23A, 23B, 53A and 53B.

The 2004 standard does not include a correction for a glass hydrometer.

API-2540 Boundaries

API MPMS 11.1:1980 (API 2540) is based on published data that lie within the so-called 'Data' range. The other table values were obtained from extrapolation and lie within the 'Extrapolated' range. It is recommended not to use API-2540 outside the 'Data' and 'Extrapolated' ranges.

For the lubricating oil tables no difference is made between data that is table values that are based on published data and table values that are determined by extrapolation.

Table 24D*

Range	API Gravity	Relative Density	Density	Temperature	Temperature
	[∘API]	[-]	[kg/m³]	[°F]	[°C]
Data Range	040	1.0760 0.8250	1075.0 824.0	0250	-18120
	40 50	0.8250 0.7795	824.0 778.5	0200	-1890
	50 55	0.7795 0.7585	778.5 758.0	0150	-1860
Extrapolated Range	040	1.0760 0.8250	1075.0 824.0	250300	120150
	4050	0.8250 0.7795	824.0 778.5	200 250	90125
	50 55	0.7795 0.7585	778.5 758.0	150 200	6095
	55 100	0.7585 0.6110	758.0 610.5	0200	-1895
Applies for:	Table 5A	Table 23A	Table 53A	Table 5A	Table 53A
	Table 6A	Table 24A	Table 54A	Table 6A	Table 54A
				Table 23A	
				Table 24A	
Range	API Gravity	Relative Density	Density	Temperature	Temperature
	[°API]	[-]	[kg/m³]	[°F]	[°C]
Data Range	040	1.0760 0.8250	1075.0 824.0	0250	-18120
	4050	0.8250 0.7795	824.0 778.5	0200	-1890
	50 85	0.7795 0.6535	778.5 653.0	0150	-1860
Extrapolated Range	040	1.0760 0.8250	1075.0 824.0	250300	120150
	4050	0.8250 0.7795	824.0 778.5	200250	90125
	50 85	0.7795 0.6535	778.5 653.0	150200	6095
Applies for:	Table 5B	Table 23B	Table 53B	Table 5B	Table 53B
	Table 6B	Table 24B	Table 54B	Table 6B	Table 54B
				Table 23B	
				Table 24B	
Range	API Gravity	Relative Density	Density	Temperature	Temperature
	[°API]	[-]	[kg/m ³]	[°F]	[°C]
Data Range	-1045	0.81.165	8001164	0300	-20+150
Applies for:	Table 5D	Table 23D*	Table 53D	Table 5D	Table 53D
	Table 6D	Table 24D*	Table 54D	Table 6D	Table 54D
				Table 23D*	

* Values derived from Table 5D/6D

API-2540 - Rounding and truncating rules

For each table API Standard 2540 specifies an explicit 'Calculation Procedure' that includes the rounding and truncating of all the input, intermediate and output values. The 'Calculation Procedure' is considered to be the standard rather than the table values or a set of equations. The function provides the option to either apply the full API rounding and truncating requirements or to perform the calculation procedure without any rounding and truncating being applied.

For tables 6A, 6B, 24A, 24B and 54A and 54B the standard makes a distinction between computational and table values for the calculated VCF. The table values are always rounded to 4 decimal places, Whereas the computational values has 4 decimal places when the VFC >=1 and 5 decimal places when the VCF < 1.

When API rounding is enabled the convergence limit is set to the limit value as specified in the standard. When the API rounding is disabled the convergence limit is set to 0.00001 kg/m3 to obtain highest precision.

API-11.1:2004 Limits

Range	Density	Temperature	Pressure	
Crude Oil	610.61163.5 kg/m3 @ 60°F	-58302 °F	01500 psig	
	10010 API @ 60°F	-50150 °C	0103.4 bar(g)	
	0.611201.16464 RD @ 60°F			
	611.161163.79 kg/m3 @ 15°C			
	606.121161.15 kg/m3 @ 20°C			
Refined products	610.61163.5 kg/m3 @ 60°F	-58302 °F	01500 psig	
	10010 API @ 60°F	-50150 °C	0103.4 bar(g)	
	0.611201.16464 RD @ 60°F			
	611.161163.86 kg/m3 @ 15°C			
	606.121160.62 kg/m3 @ 20°C			
Lubricating oils	800.91163.5 kg/m3 @ 60°F	-58302 °F	01500 psig	
	4510 API @ 60°F	-50150 °C	0103.4 bar(g)	
	0.801681.1646 RD @ 60°F			
	801.251163.85 kg/m3 @ 15°C			
	798.111160.71 kg/m3 @ 20°C			

API constants in US customary units

For the tables in <u>US customary units</u> the following constants apply (both for the 1980 and the 2004 tables):

Product	API Table	ко	K1	K2
Crude oil	А	341.0957	0.0	0.0
Gasoline	В	192.4571	0.2438	0.0
Transition area	В	1489.0670	0.0	-0.00186840
Jet fuels	В	330.3010	0.0	0.0
Fuel oils	В	103.8720	0.2701	0.0
Lubricating oils	D	0.0	0.34878	0.0

API constants in metric units

For the tables in <u>metric units</u> the following constants apply (both for the 1980 and the 2004 tables):

Product	API Table	К0	K1	K2
Crude oil	А	613.9723	0.0	0.0
Gasoline	В	346.4228	0.4388	0.0
Transition area	В	2680.3206	0.0	-0.00336312
Jet fuels	В	594.5418	0.0	0.0
Fuel oils	В	186.9696	0.4862	0.0
Lubricating oils	D	0.0	0.6278	0.0

2 Spirit^{IT} Gas and liquid functions

This chapter lists all available Spirit^{IT} functions for gas and liquid calculations in alphabetical order. These functions are both available in the Flow-X flow computer software and in the Flow-Xpert offline calculation software.

fxAGA3_C

Description

The function calculates the mass flow rate for Orifice pressure differential flow devices according to the AGA-3 standard for orifice meters with flange taps.

Compliance

- AGA Report No. 3 Orifice Metering Measurement of fluid flow by means of pressure differential devices, 1992
- API Manual of Petroleum Measurement Standards, Chapter 14 Natural Gas Fluids Measurement, Section 3 - Concentric Square-edged Orifice Meters 1992.
- Orifice Metering of Natural Gas and Other Related Hydrocarbon Fluids – Concentric, Square-edged Orifice Meters, Part 1: General Equations and Uncertainty Guidelines, fourth edition 2012.

Function inputs

	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Differential Pressure	Differential pressure over the primary flow device measured at	inH2O @ 60°F		01000	0
	the up- and downstream pressure tappings, which need to be in				
	the positions as specified in the standard				
Pressure	Down- or upstream pressure value of the fluid at metering	psia		030000	0
	conditions				
Temperature	Down- or upstream temperature of the fluid at metering	°F		-4002000	0
	conditions				
Density	Down or upstream density of the fluid at metering conditions	lbm/ft3		0200	0
Dynamic Viscosity	Dynamic viscosity of the fluid	lbm/ft.s	DYNVIS	010	6.9e-6
Isentropic Exponent	Also referred to as κ (kappa). For an ideal gas this coefficient is	-	KAPPA	010	1.3
	equal to the ratio of the specific heat capacity at constant				
	pressure to the specific heat at constant volume.				
	This ratio is commonly used when the real value is unknown.				
Pipe Diameter	Internal diameter of the pipe at reference temperature	inches	PIPEDIAM	0100	0
Pipe Expansion factor	The thermal expansion coefficient of the pipe material	1/°F	PIPEEXPF	01	6.2e-6
Pipe Reference	The reference temperature that corresponds to the 'Pipe	°F	PIPEREFT	-4002000	68
temperature	diameter' input value				
Orifice Diameter	Orifice diameter at reference temperature	inches	ORIFDIAM	0100	0
Orifice Expansion factor	The thermal expansion coefficient of the orifice material	1/°F	ORIFEXPF	01	9.25e-6
	Typical values are:	-/ ·	0.00	0.112	5.200 0
Orifice Reference	The reference temperature that corresponds to the 'Orifice	°F	ORIFREFT	-4002000	68
Temperature	diameter' input value	•	On the t	1002000	00
Pressure Location	1: Upstream tapping	-	PRESLOC		1
	Input 'Pressure' represents the pressure at the upstream		TRESECC		-
	pressure tapping (p_1) .				
	Since the absolute pressure is usually measured at the				
	upstream tapping this is the most common setting.				
	2: Downstream tapping				
	Input 'Pressure' represents the pressure at the downstream				
	tapping (p ₂).				
Temperature Location	1: Upstream tapping	-	TEMPLOC		3
	Input 'Temperature' represents the upstream temperature (t_1) .				5
	2: Downstream tapping				
	Input 'Temperature represents the temperature at the				
	downstream tapping (t_2).				
	3: Recovered pressure				
	•				
	Input 'Temperature' represents the downstream temperature				
	Input 'Temperature' represents the downstream temperature at a location Where the pressure has fully recovered (t ₃).				
	Input 'Temperature' represents the downstream temperature at a location Where the pressure has fully recovered (t ₃). Since temperature measurement is usually downstream of the				
	Input 'Temperature' represents the downstream temperature at a location Where the pressure has fully recovered (t ₃).				
Femperature Correction	Input 'Temperature' represents the downstream temperature at a location Where the pressure has fully recovered (t ₃). Since temperature measurement is usually downstream of the flow device this is the most common setting.	_	TEMPCOR		1
Temperature Correction	Input 'Temperature' represents the downstream temperature at a location Where the pressure has fully recovered (t ₃). Since temperature measurement is usually downstream of the flow device this is the most common setting. 1: Use $(1-\kappa)/\kappa$	-	TEMPCOR		1
Temperature Correction	$\label{eq:linear} \begin{array}{l} \mbox{Input 'Temperature' represents the downstream temperature at a location Where the pressure has fully recovered (t_3). Since temperature measurement is usually downstream of the flow device this is the most common setting. \\ \hline 1: Use (1-\kappa)/\kappa \\ \mbox{Isentropic expansion using } (1-\kappa)/\kappa \mbox{ as the temperature referral} \end{array}$	-	TEMPCOR		1
Temperature Correction	$\label{eq:linear_linear} \begin{array}{l} \mbox{Input 'Temperature' represents the downstream temperature at a location Where the pressure has fully recovered (t_3). Since temperature measurement is usually downstream of the flow device this is the most common setting. \\ \hline 1: Use (1-\kappa)/\kappa \\ \mbox{Isentropic expansion using (1-\kappa)/\kappa as the temperature referral exponent} \end{array}$	-	TEMPCOR		1
Temperature Correction	 Input 'Temperature' represents the downstream temperature at a location Where the pressure has fully recovered (t₃). Since temperature measurement is usually downstream of the flow device this is the most common setting. 1: Use (1-κ)/κ Isentropic expansion using (1-κ)/κ as the temperature referral exponent 2: Use temperature exponent 	-	TEMPCOR		1
Temperature Correction	 Input 'Temperature' represents the downstream temperature at a location Where the pressure has fully recovered (t₃). Since temperature measurement is usually downstream of the flow device this is the most common setting. 1: Use (1-κ)/κ Isentropic expansion using (1-κ)/κ as the temperature referral exponent 2: Use temperature exponent Isentropic expansion using input 'Temperature Exponent' as 	-	TEMPCOR		1
	 Input 'Temperature' represents the downstream temperature at a location Where the pressure has fully recovered (t₃). Since temperature measurement is usually downstream of the flow device this is the most common setting. 1: Use (1-κ)/κ Isentropic expansion using (1-κ)/κ as the temperature referral exponent 2: Use temperature exponent Isentropic expansion using input 'Temperature Exponent' as the temperature referral exponent [-] 	-			
Temperature Correction	 Input 'Temperature' represents the downstream temperature at a location Where the pressure has fully recovered (t₃). Since temperature measurement is usually downstream of the flow device this is the most common setting. 1: Use (1-κ)/κ Isentropic expansion using (1-κ)/κ as the temperature referral exponent 2: Use temperature exponent Isentropic expansion using input 'Temperature Exponent' as 	-	TEMPCOR		1

Function inputs	Remark	EU	SW tag	Range	Default
	used.				
	For more details refer to section 'Temperature correction'.				
Density Location	This parameter specifies if and how the density should be	-	DENSLOC		0
	corrected from downstream to upstream conditions.				
	1: Upstream tapping				
	Input 'Density' represents the density at the upstream				
	pressure tapping (ρ_1).				
	2: Downstream tapping				
	Input 'Density' represents the density at the downstream				
	tapping (ρ₂).				
	3: Recovered pressure				
	Input 'Density' represents the density downstream at a				
	location Where the pressure has fully recovered (ρ_3).				
Density Exponent.	This factor is used when density correction is enabled. The	-	DENSEXP		0
	formula 1/ κ will be used when the input value is set to 0, else the				
	input value will be used.				
	For more details refer to section function 'ISO5167- Orifice'				
	'Density correction'.				
Fluid	The type of fluid being measured	-	FLUID		0
	1: Gas				
	2: Liquid				
	For liquid the expansion factor is set to 1, i.e. the fluid is				
	considered to be incompressible.				
Drain hole	When input is > 0 then an additional correction on the orifice	in	DRAIN	0 100	0
	diameter will be applied to account for the drain hole, as				
	explained further on.				
Fpwl	Local Gravitational Correction Factor for Deadweight Calibrators	-	FPWL	0.91.1	1
	used to calibrate differential and static pressure Instruments.				
	Directly applied on the calculated mass flow rate within each				
	iteration.				
Year Of Edition	1: Edition 1992				1
	2: Edition 2012				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOVONV	
Mass flow rate	The calculated mass flow rate	klbm/hr	MASSR		0
Beta ratio	Orifice to pipe diameter ratio at upstream temperature	-	BETA		0
Orifice diameter	At the upstream temperature	inches	ORIFUP		0
Pipe diameter	At the upstream temperature	inches	PIPEUP		0
Upstream pressure	Pressure at upstream tapping (p1)	psia	PRESUP		0
Pressure at downstream	Pressure at downstream tapping (p2)	psia	PRESDN		0
tapping					
Recovered downstream	Fully recovered downstream pressure (p3)	psia	PRESREC		0
pressure					
Upstream temperature	Temperature at upstream tapping (t1)	°F	TEMPUP		0
Temperature at downstream	Temperature at downstream tapping (t ₂)	°F	TEMPDN		0
tapping					
Downstream Temperature	'Fully recovered' downstream temperature (t3)	°F	TEMPREC		0
Upstream density	Density at upstream tapping ($ ho_1$)	lbm/ft3	DENSUP		0
Density at downstream	Pressure at downstream tapping (ρ_2)	lbm/ft3	DENSDN		0
tapping					
Downstream density	'Fully recovered' downstream density (p3)	lbm/ft3	DENSREV		0
Reynolds number	The pipe Reynolds number, i.e. the Reynolds number upstream of the	-	REYN		0
	orifice and not the one within the device throat itself)				
Discharge coefficient		-	DISCF		0
Expansion Factor		-	EXPFAC		0
Velocity of Approach		-	VOA		0
Pressure out of range	0: Pressure is in valid range	-	PRESOOR	PRESOOR	0
	1: Pressure is out of valid range				
Reynolds out of range	0: Reynolds number is in valid range	-	REYNOOR	REYNOOR	0
	1: Reynolds number is out of valid range				
Diameter out of range	0: Device and pipe diameter and Beta ratio in valid range	-	DIAMOOR	DIAMOOR	0
	1: Device diameter, pipe diameter and/or Beta ratio out of valid range				

Calculations

The calculations are in accordance with the standard.

Pressure correction

The relation between the pressure at the upstream tapping p_1 and the pressure at the downstream tapping (p_2) is as following: The relation between the pressure at the upstream tapping and the downstream tapping is as following:

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 $p_3 = p_1 - p_{LOSS}$

 $p_2 = p_1 \frac{\Delta p \cdot K_{units}}{1000}$

$$p_{LOSS} = \frac{\left(1 - \alpha \cdot \beta^2\right)}{\left(1 + \alpha \cdot \beta^2\right)} \cdot \Delta p \cdot K_{units}$$

$$\alpha = C \cdot E$$
$$E = \frac{1}{\sqrt{\left(1 - \beta^4\right)}}$$

Where:

p1	Pressure at upstream tapping	psia
p ₂	Pressure at downstream tapping	psia
р₃	Fully recovered downstream pressure	psia
Δр	Differential pressure	inH20 @ 60°F
PLOSS	Pressure loss over the meter	psi
с	Discharge coefficient as calculated by the standard	-
α	Flow coefficient	-
β	Diameter ratio at the upstream pressure and temperature	-
E	Velocity of approach factor	-
K _{units}	Unit conversion factor to convert a value expressed in 'inH2O @60°F' to the corresponding expressed in 'psi' (conversion as specified in section 'Unit Types')	-

Temperature correction

When input 'Temperature correction' is set to 1, then an isentropic expansion based on the isentropic coefficient is applied:

$$t_{1} = \left(t_{2} + 459.67\right) \cdot \left(\frac{p_{2}}{p_{1}}\right)^{\frac{1-\kappa}{\kappa}} - 459.67$$
$$t_{1} = \left(t_{3} + 459.67\right) \cdot \left(\frac{p_{3}}{p_{1}}\right)^{\frac{1-\kappa}{\kappa}} - 459.67$$

When input 'Temperature correction' is set to 2, then an isentropic expansion based on input 'Temperature exponent' is applied:

$$t_{1} = \left(t_{2} + 459.67\right) \cdot \left(\frac{p_{2}}{p_{1}}\right)^{K_{TE}} - 459.67$$
$$t_{1} = \left(t_{3} + 459.67\right) \cdot \left(\frac{p_{3}}{p_{1}}\right)^{K_{TE}} - 459.67$$

Where:

t1	Upstream temperature	°F
t2	Temperature at the downstream tapping	°F
t3	Temperature at the fully recovered downstream pressure	°F
p 1	Upstream pressure	psia

Density correction

When input 'Density exponent' = 0, then the following isentropic corrections are applied (depending on the type of Density Correction)

$$\rho_1 = \rho_2 \cdot \left(\frac{p_1}{p_2}\right)^{\frac{1}{\kappa}} \qquad \qquad \rho_1 = \rho_3 \cdot \left(\frac{p_1}{p_3}\right)^{\frac{1}{\kappa}}$$

Else the value of input 'Density Exponent' is used

 $\rho_1 = \rho_2 \cdot \left(\frac{p_1}{p_2}\right)^{K_{DE}}$

$$\rho_1 = \rho_3 \cdot \left(\frac{p_1}{p_3}\right)^{K_{DE}}$$

Where:

ρ1	Upstream density	lbm/ft3
ρz	Density at the downstream tapping	lbm/ft3
ρ3	Density at the fully recovered downstream pressure	lbm/ft3
p1	Upstream pressure	psia
p2	Pressure at the downstream tapping	psia
р₃	Fully recovered downstream pressure	psia
к	Isentropic exponent	-
Kde	Density exponent	-

fxAGA5_C

Description

The AGA 5 standard defines methods to calculate the mass and volume based calorific values at 60°F and 14.73 psia for a natural gas based on known molar fractions of the <u>non</u>-hydrocarbon gas components.

Compliance

- A.G.A. Transmission Measurement Committee Report No. 5 (Fuel gas Energy Metering) 1981
- A.G.A. Transmission Measurement Committee Report No. 5 (Fuel gas Energy Metering) 1996 (Reprinted 1999)

Function inputs	Remark	(EU	SW tag	Range	Default
Name	Option	al tag name, tag description and tag group				
Composition	Standa	rd composition as defined in section 'Standard gas	molar fraction	COMP	01	0
	compos	sition.				
	Only th	e following components are considered by the calculation:				
	N2	Nitrogen				
	CO2	Carbon dioxide				
	H2O	Water				
	H2S	Hydrogen sulfide				
	H2	Hydrogen				
	CO	Carbon monoxide				
	02	Oxygen				
	He	Helium				
	Sum of	these fractions may not exceed 1				
Specific Gravity	Molar N	lass Ratio, i.e. ratio of the molar mass of the gas and of the	-	SG	01	0
	molar n	nass of air (specified in AGA-5 as 28.9644 kg/kmol				
	(lbm/lb	omol))				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
Calorific value mass	Mass based calorific value	Btu/lbm	CV_MASS		0
Calorific value volume	Volume calorific value at 60°F and 14.73 psia	Btu/scf	CV_VOL		0

Calculations

The Energy to Mass ratio is calculated according to Section III of the standard, which contains the calculation procedure for the gas mass to energy conversion. The equations based on the 'by volume' fractional values are used (and not the equations based on the 'by weight' values).

The Energy to Volume ratio is calculated according to Section II of the standard, which contains the calculation procedure for the gas volume to energy conversion.

fxAGA8_C

Description

The compressibility and density of a gas are calculated from the composition, temperature and pressure in accordance with the 'Detail Characterization' method outlined in the AGA-8 standard, with the input and output values in **US Customary** units.

Compliance

- AGA Report No. 8, Second edition November 1992 2nd printing July 1994
- API MPMS 14.2, Second edition November 1992 2nd printing July 1994
- ISO 12213 Natural gas Calculation of compression factor — Part 2: Calculation using molar-composition analysis, 1997

Boundaries

The AGA-8 calculation has defined uncertainty bounds for gas mixtures that lie within the 'Normal range'. Also an 'Expanded range' of gas mixtures is defined for which the AGA-8 calculation has a higher uncertainty. Using the AGA-8 calculation for gas mixtures that lie outside the 'Expanded range' is not recommended.

Input value	Normal Range	Expanded Range	EU
Pressure	020000	0 20000	psia
Temperature	-200 +400	-200 +400	°F
Mole fraction of Methane	0.45 1.00	0.00 1.00	-
Mole fraction of Ethane	0.00 0.10	0.00 1.00	-
Mole fraction of Propane	0.00 0.04	0.00 0.12	-
Mole fraction of Butanes	0.00 0.01	0.00 0.06	-
Mole fraction of Pentanes	0.00 0.003	0.00 0.04	-
Mole fraction of Hexanes Plus	0.00 0.002	*	-
Mole fraction of Carbon	0.00 0.03	0.00 0.03	-
monoxide			
Mole fraction of Carbon dioxide	0.00 0.30	0.00 1.00	-
Mole fraction of Nitrogen	0.00 0.50	0.00 1.00	-
Mole fraction of Helium	0.00 0.002	0.00 0.03	-
Mole fraction of Argon	0.00 0.00	0.00 0.01	-
Mole fraction of Oxygen	0.00 0.00	0.00 0.21	-
Mole fraction of Hydrogen	0.00 0.0002	0.00 1.00	-
Sulphide			
Mole fraction of Hydrogen	0.00 0.10	0.00 1.00	-
Mole fraction of Water	0.00 0.0005	*	-

* For these components the dew point temperature is the upper limit. Limit check is ignored for reason of simplicity.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Pressure	Pressure value	psia		040000	1.01325
Temperature	Temperature value	°F		-250+800	0
Composition	Standard composition as defined in section 'Standard gas composition.	mol/mol	СОМР	01	0
neo-Pentane mode	Determines what to do when component neo-Pentane is larger than zero 1: Add to i-Pentane 2: Add to n-Pentane 3: Neglect	-	NEOC5_MODE		1

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
	4: Mole fractions do not add up to 1.0 +- 0.0001			COMPERR	
Compressibility factor		-	Z		1
Mass Density		lb/ft3	MASDENS		0
Mole Density		lbmol/ft3	MOLDENS		0
Molar Mass		lb/lbmol	MOLMASS		0
Range	0: In Normal Range All inputs are within the 'Normal Range'		RANGE		0
	 In Extended Range One or more inputs within the 'Extended Range, but none of the inputs outside the Extended rang (outputs values have higher uncertainty) 	1			
	2: Out of Range One or more inputs outside the 'Extended Range' (using the AGA8 calculation is not recommended in this case)			OOR	

Calculations

The calculations are as documented in the standard.

fxAGA8_M

Description

The compressibility and density of a gas are calculated from its composition, temperature and pressure in accordance with the 'Detail Characterization' method outlined in the AGA8 standard, with the input and output values in **metric** units.

Compliance

- AGA Report No. 8, Second edition November 1992 2nd printing July 1994
- API MPMS 14.2, Second edition November 1992 2nd printing July 1994

- ISO 12213 Natural gas Calculation of compression factor — Part 2: Calculation using molar-composition analysis, 1997
- AGA Report No. 8, Third edition April 2017

Boundaries

The AGA-8 calculation has defined uncertainty bounds for gas mixtures that lie within the 'Normal range'. Also an 'Expanded range' of gas mixtures is defined for which the AGA-8 calculation has a higher uncertainty. Using the AGA-8 calculation for gas mixtures that lie outside the 'Expanded range' is not recommended.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group)			
Pressure	Pressure value	bar(a)		02800	1.01325
Temperature	Temperature value	°C		-150+450	0
Composition	Standard composition as defined in section	mol/mol	COMP	01	0
	'Standard gas composition.				
neo-Pentane mode	Determines what to do when component neo-	-	NEOC5_MODE		1
	Pentane is larger than zero				
	1: Add to i-Pentane				
	2: Add to n-Pentane				
	3: Neglect				
Year Of Edition	1: Edition 1994				1
	2: Edition 2017				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
	4: Mole fractions do not add up to 1.0 +- 0.0001			COMPERR	
Compressibility factor		-	Z		1
Mass Density		kg/m3	MASDENS		0
Mole Density		kmol/m3	MOLDENS		0
Molar Mass		kg/kmol	MOLMASS		0
Range	0: In Normal Range All inputs are within the 'Normal Range'		RANGE		0
	1: In Extended Range One or more inputs within the 'Extended Range, but none of the inputs outside the Extended rang (outputs values have higher uncertainty)				
	2: Out of Range One or more inputs outside the 'Extended Range' (using the AGA8 calculation is not recommended in this case			OOR	

Calculations

The calculations are as documented in the standard.

fxAGA8_Gross

Description

This function calculates the compressibility factor in accordance with the AGA-8 Gross Characterization Method. Although the AGA-8 Gross Method is based on the Standard GERG Virial Equation Of State (SGERG) there are slight differences in the results.

Two different methods are specified by the standard. Method 1 takes the Pressure, Temperature, Specific Gravity (Relative Density), Carbon Dioxide content and Gross Heating Value (GHV) as inputs. Method 2 takes the same inputs except for the Nitrogen content instead of GHV.

Compliance

AGA 8, Second edition November 1992 - 2nd printing July 1994

- AGA Report No. 8, Second edition November 1992 2nd printing July 1994
- API MPMS 14.2, Second edition November 1992 2nd printing July 1994

Function inputs

Boundaries

The AGA8 standard recommends using the Gross Characterization Method only when input conditions lie within the following range. For conditions outside this range the standard recommends to use the Detailed Characterization Method.

Input value	Normal Range	EU
Temperature	32130	°F
Pressure	01200	psia
Gross heating value	475 1210	Btu/ft3
Relative density	0.554 0.87	-
Carbon dioxide	0.00 0.30	mol/mol
Nitrogen	0.00 0.50	mol/mol

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Temperature	Observed temperature	°F	-250+800	60
Pressure	Observed pressure	psia	040000	
Relative density	Relative density at the corresponding reference temperature and	-	02	0
	pressure			
RD reference temperature	Reference temperature for relative density	°F	-250+800	60
RD reference pressure	Reference pressure for relative density	psia	040000	14.73
Gross heating value	Gross heating value at the corresponding reference temperature and pressure	Btu/ft3	02500	0
GHV reference temperature	Reference temperature for gross heating value	°F	-250+800	60
GHV reference pressure	Reference pressure for gross heating value	psia	040000	14.73
Nitrogen	Nitrogen (N2) fraction	mol/mol	01	0
Carbon dioxide	Carbon dioxide (CO2) fraction	mol/mol	01	0
Method	Gross Characterization Method:	-		0
	1: GHV, Relative Density, CO2			
	2: Relative Density, CO2, N2			
	Note: For Method 1 input 'Nitrogen' is not used, while for Method 2			
	inputs 'Gross heating value', 'GHV reference temperature' and 'GHV			
	reference pressure' are not used.			

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Compressibility factor		-			1
Molar mass		lb/ lbmol			0
Density	Density at observed pressure and temperature	lb/ft3			0
Range	0: In Normal Range All components are within the range that is recommended by the standard 1: Out of Range One or more inputs are outside the recommended range		RANGE		0

Calculations

The calculations are in accordance with the standard.

fxAGA10_M

Description

The function calculates the speed of sound of a gas at the specified conditions of temperature and pressure using the formulae presented in the American Gas Association Report No 10.

Compliance

AGA Report No. 10 - Speed of Sound in Natural Gas and Other Related Hydrocarbon Gases, January 2003

Boundaries

The AGA-10 calculation has defined uncertainty bounds for gas mixtures that lie within the 'Normal range'. Also an 'Expanded range' of gas mixtures is defined for which the AGA-10 calculation has a higher uncertainty. Using the AGA-10 calculation for gas mixtures that lie outside the 'Expanded range' is not recommended.

The AGA-10 standard specifies the same limits as the AGA-8 standard. Refer to the fxAGA8 function for details on the actual limit values used by this function to set output 'Range'.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Pressure	Observed pressure	bar(a)		02000	
Temperature	Observed temperature	°C		-200+400	
Composition	Standard composition as defined in section 'Standard gas composition.	mol/mol	СОМР	01	
neo-Pentane mode	Determines what to do when component neo-Pentane is larger than zero 1: Add to i-Pentane 2: Add to n-Pentane 3: Neglect	-	NEOC5_MODE		1

Function outputs

Function outputs	Remark	EU	SW tag	Alam	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
	4: Mole fractions do not add up to 1.0 +- 0.0001			COMPOOR	
Speed of sound		m/s	SOS		0
Range	0: In Normal Range		RANGE		0
	All inputs are within the 'Normal Range'				
	1: In Extended Range				
	One or more inputs within the 'Extended Range, but none of				
	the inputs outside the Extended rang (outputs values have				
	higher uncertainty)				
	2: Out of Range			OOR	
	One or more inputs outside the 'Extended Range' (using the				
	AGA10 calculation is not recommended in this case)				

Calculations

Calculations are as documented in the standard.

fxAGA10ex_M

Description

The extended AGA 10 function provides an extensive set of gas properties at the specified conditions of temperature and pressure using the formulae presented in the American Gas Association Report No 10.

Compliance

AGA Report No. 10 - Speed of Sound in Natural Gas and Other Related Hydrocarbon Gases, January 2003

Boundaries

The AGA-10 calculation has defined uncertainty bounds for gas mixtures that lie within the 'Normal range'. Also an 'Expanded range' of gas mixtures is defined for which the AGA-10 calculation has a higher uncertainty. Using the AGA-10 calculation for gas mixtures that lie outside the 'Expanded range' is not recommended.

The AGA-10 standard specifies the same limits as the AGA-8 standard. Refer to the fxAGA8 function for details on the actual limit values used by this function to set output 'Range'.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Pressure	Observed pressure	bar(a)		02000	
Temperature	Observed temperature	°C		-200+400	
Composition	Standard composition as defined in section 'Standard gas composition.	mol/mol	COMP	01	
neo-Pentane mode	Determines what to do when component neo-Pentane is larger than zero 1: Add to i-Pentane 2: Add to n-Pentane 3: Neglect	-	NEOC5_MODE		1

Function outputs

Function outputs	Remark	EU	SW tag	Alam	Fall back
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
	4: Mole fractions do not add up to 1.0 +- 0.0001			COMPOOR	
Molecular weight		kg/kmol	MOLMASS		
Molar density at base conditions		mol/m3	MOLDENSB		
Molar density at flowing conditions		mol/m3	MOLDENSF		
Mass density at base conditions		kg/m3	MASSDENSB		
Mass density at flowing conditions		kg/m3	MASSDENSF		
Ideal gas relative density		-	IRD		
Real gas relative density		-	RRD		
Velocity of sound		m/s	SOS		
Compressibility at base conditions		-	ZB		
Compressibility at flowing conditions		-	ZF		
Supercompressibility		-	FPV		
Ideal gas specific enthalpy		kJ/kg	MASSH0		
Real gas specific enthalpy		kJ/kg	MASSH		
Real gas specific entropy		kJ/kg/K	MASSS		
Ideal gas isobaric heat capacity		kJ/kg/K	MASSCP0		
Real gas isobaric heat capacity		kJ/kg/K	MASSCP		
Real gas isochoric heat capacity		kJ/kg/K	MASSCV		
Ideal gas isobaric heat capacity		kJ/kmol/K	MOLCP0		
Real gas isobaric heat capacity		kJ/kmol/K	MOLCP		
Real gas isochoric heat capacity		kJ/kmol/K	MOLCV		
Ratio of specific heats		-	CPCV		
Isentropic exponent		-	KAPPA		
Critical flow factor		-	CRITC		
Ideal gas specific enthalpy		kJ/kmol	MOLH0		
Real gas specific enthalpy		kJ/kmol	MOLH		
Isentropic perfect gas critical flow factor		-	CI		
Isentropic real gas critical flow factor		-	CRI		
Range	0: In Normal Range		RANGE		0
	All inputs are within the 'Normal Range'				Ū.
	1: In Extended Range				
	One or more inputs within the 'Extended				
	Range, but none of the inputs outside the				
	Extended rang (outputs values have higher				
	uncertainty)				
	2: Out of Range			OOR	
	One or more inputs outside the 'Extended				
	Range' (using the AGA10 calculation is not				

Function outputs	Remark	EU	SW tag	Alam	Fall back
	recommended in this case)				

Calculations

Calculations are as documented in the standard.

fxAPI_Dens15C_1952

Description

Density (T, P) <--> Density (15°C, equilibrium pressure)

This function converts a density value at the observed temperature and pressure to the density at 15°C and the equilibrium pressure (typically 0 barg) or vice versa. The temperature conversion is according to ASTM-IP Petroleum Measurements Tables 1952 (Also known as API-1952 tables) Table 54.

Note: this function is a combination of the API 1952 Tables and API 11.2.1M. For the calculation from observed to standard conditions an iterative calculation is required. The rounding and truncating of input and intermediate values is implemented such that the example calculations as specified in both standards are exactly reproduced.

Compliance

- ASTM-IP Petroleum Measurement Tables, Metric Edition, Metric Units of Measurement, 1952
- API MPMS 11.2.1M Compressibility Factors for Hydrocarbons: 638 - 1074 Kilograms per Cubic Meter Range -First Edition, August 1984

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group		-	-	
Observed Density	Depending on the conversion method this is the Density either at the observed temperature and observed pressure or at 15 °C and the equilibrium pressure	kg/m3		01300	0
Observed temperature		°C		-100200	15
Observed pressure		bar(g)		-1150	0
API 11.2.1 rounding	 0: Disabled The calculation of the compressibility factor F is performed with full precision 1: Enabled API-MPMS 11.2.1 rounding and truncating rules are applied. The compressibility factor F is rounded to 3 decimal places as specified in the standard. 	-	API1121RND		0
Equilibrium pressure	The equilibrium pressure is considered to be 0 bar(g) for liquids which have an equilibrium pressure less than atmospheric pressure (in compliance with API MPMS 12.2 par. 12.2.5.4).	bar(g)	EQUIPRES	0150	0
Conversion method	1: From observed to standard conditions 2: From standard to observed conditions		CONVERSION		1

Function outputs

	EU	SW tag	Alarm	Fallback
0: Normal	-	STS		
1: Input argument out of range			FIOOR	1
2: Calculation error			CALCERR	
3: No convergence			NOCONV	
Depending on the conversion method this is the Density either at 15 $^\circ$ C and	kg/m3	DENS		0
the equilibrium pressure or at the observed temperature and observed				
pressure				
Volume correction factor for temperature.	-	CTL		1
Volume correction factor for pressure	-	CPL		1
Value will be rounded according to input 'API 11.2.1 rounding'				
Combined volume correction factor	-	CTPL		1
CTPL = CTL * CPL				
Compressibility factor	1/bar	F		0
With respect to the standard used for the calculation of CTL the				0
combination of input values is:				
0: In Range				
1: Out of Range			CTLOOR	
With respect to the standard used for the calculation of CPL the				0
combination of input values is:				
0: In Range				
1: Out of Range			CPLOOR	
	I: Input argument out of range 2: Calculation error 3: No convergence Depending on the conversion method this is the Density either at 15 °C and the equilibrium pressure or at the observed temperature and observed pressure Volume correction factor for temperature. Volume correction factor for pressure Value will be rounded according to input 'API 11.2.1 rounding' Combined volume correction factor CTPL = CTL * CPL Compressibility factor With respect to the standard used for the calculation of CTL the combination of input values is: 0: In Range With respect to the standard used for the calculation of CPL the combination of input values is: 0: In Range	1: Input argument out of range 2: Calculation error 3: No convergence Depending on the conversion method this is the Density either at 15 °C and the equilibrium pressure or at the observed temperature and observed pressure kg/m3 Volume correction factor for temperature. - Volume correction factor for pressure - Value will be rounded according to input 'API 11.2.1 rounding' - Combined volume correction factor - CTPL = CTL * CPL - Compressibility factor 1/bar With respect to the standard used for the calculation of CTL the combination of input values is: - 0: In Range - 1: Out of Range - With respect to the standard used for the calculation of CPL the combination of input values is: - 0: In Range -	1: Input argument out of range 2: Calculation error 3: No convergence Depending on the conversion method this is the Density either at 15 °C and the equilibrium pressure or at the observed temperature and observed pressure kg/m3 DENS Volume correction factor for temperature. - CTL Volume correction factor for pressure - CPL Value will be rounded according to input 'API 11.2.1 rounding' - CPL Combined volume correction factor - CTPL CTPL = CTL * CPL - CTPL Compressibility factor 1/bar F With respect to the standard used for the calculation of CTL the combination of input values is: 0: In Range - 1: Out of Range With respect to the standard used for the calculation of CPL the combination of input values is: - - 0: In Range - : - -	1: Input argument out of range FIOOR 2: Calculation error CALCERR 3: No convergence NOCONV Depending on the conversion method this is the Density either at 15 °C and the equilibrium pressure or at the observed temperature and observed pressure kg/m3 DENS Volume correction factor for temperature. - CTL Volume correction factor for pressure - CPL Value will be rounded according to input 'API 11.2.1 rounding' - CTPL Combined volume correction factor - CTPL CTPL = CTL * CPL - CTPL Compressibility factor 1/bar F With respect to the standard used for the calculation of CTL the combination of input values is: CTLOOR 0: In Range - CTLOOR With respect to the standard used for the calculation of CPL the combination of input values is: CTLOOR 0: In Range - CTLOOR

Calculations

The calculations depend on the conversion method.

Conversion method 1: from observed to standard conditions. The function performs the following iterative algorithm to calculate the Density at standard conditions:

- 1 At the start of the iteration the initial value for Density at [15 °C, equilibrium pressure] is set to the Observed Density. The initial CPL value is set to 1.
- 2 The CTL value is determined from the Density at [15 °C, equilibrium pressure] according to API 1952 Table 54.
- 3 The Density at [15 °C, equilibrium pressure] is calculated from the Observed Density, the new CTL value and the CPL value from the previous iteration.
- 4 The compressibility factor is calculated according to API MPMS 11.2.1M from the density at [15 °C, equilibrium pressure] and the 'Observed temperature'. If API 11.2.1M rounding is enabled then the density and temperature are rounded and the calculations are performed in accordance with the rounding and truncating rules of the standard.
- 5 The CPL value is calculated from the compressibility factor and the 'Observed pressure' and 'Equilibrium pressure' input values.
- 6 The Density at [15°C, equilibrium pressure] is calculated by dividing the Observed Density by the CTL and the new CPL value.
- 7 Steps 2 through 6 are repeated taking the Density value from step 7 as the start value for the next iteration until the absolute difference between two consecutive Density values is 0.0001.

Conversion method 2: from standard to observed conditions. The function performs straightforward calculations to determine the Density at observed conditions:

- 1 The CTL value is calculated according to API 1952 Table 54
- 2 The compressibility factor is calculated according to API MPMS 11.2.1M from the input density and temperature'. If API 11.2.1M rounding is enabled then the input density and temperature are rounded and the calculations are performed in accordance with the rounding and truncating rules of the standard.
- 3 The CPL value is calculated from the compressibility factor and the 'Observed pressure' and 'Equilibrium pressure' input values.
- 4 The output Density (at observed temperature and pressure) is calculated from the input Density and the CTL and the CPL values.

fxAPI_Dens15C_1980

Description

-

Density (T, P) <--> Density (15°C, equilibrium pressure)

This function converts a density value at the observed temperature and pressure to the density value at 15°C and the equilibrium pressure (typically 0 bar(g)) or vice versa. The temperature conversion is according to API-2540, Tables 53A/54A (Generalized Crude Oils) and 53B/54B (Refined Oil Products) and API MPMS 11.1 Chapter XIV Table 53D/54D: 1984 (Lubricating Oils), while the volume correction for pressure according to API MPMS 11.2.1M.

An iterative calculation needs to be applied to convert the observed density to the value at base conditions.

Note: this function is a combination of API2540 and API 11.2.1M. For the calculation from observed to standard conditions an iterative calculation is required. The rounding and truncating of input and intermediate values is implemented such that the example calculations as specified in both standards are exactly reproduced.

Compliance

- API MPMS 11.1 Volume X (API Standard 2540) Table 53A -Generalized Crude Oils, Correction of Observed Density to Density at 15°C - First Edition, August 1980
- API MPMS 11.1 Volume X (API Standard 2540) Table 54A -Generalized Crude Oils, Correction of Volume to 15°C against Density at 15°C- First Edition, August 1980
- API MPMS 11.1 Volume X (API Standard 2540) Table 53B -Generalized Products, Correction of Observed Density to Density at 15°C - First Edition, August 1980
- API MPMS 11.1 Volume X (API Standard 2540) Table 54B -Generalized Products, Correction of Volume to 15°C against Density at 15°F - First Edition, August 1980
- API MPMS 11.1 Volume XIV Table 53D Generalized Lubricating Oils, Correction of Observed Density to Density at 15°C - January 1982
- API MPMS 11.1 Volume XIV Table 54D Generalized Lubricating Oils, Correction of Volume to 15°C against Density at 15°F - January 1982
- API MPMS 11.2.1M Compressibility Factors for Hydrocarbons: 638 - 1074 Kilograms per Cubic Meter Range -First Edition, August 1984

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Input density	Meaning depends on the input 'Conversion method'.	kg/m3		01300	0
	'Conversion method' = 1				
	Density at the observed temperature and pressure				
	'Conversion method' = 2				
	Density at 15 °C and the equilibrium pressure.				
Observed temperature		°C		-100200	15
Observed pressure		bar(g)		-1150	0
Product	1: A - Crude Oil		PRDTYP		1
	2: B - Auto select				
	Selection based on density at 15 °C				
	3: B - Gasoline				
	4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API 2540 rounding	0: Disabled		API2540RND	-	0
	The calculations are performed with full precision and the final CTL value is				
	rounded as specified by input 'CTL decimal places'				
	1: Enabled for computational value				
	API-2540 rounding and truncating rules are applied and, in case of				
	conversion method 2 (standard to observed), the computational value for				
	CTL as specified in Table 54 is used, meaning that the CTL value has:				
	4 decimal places if CTL >=1				
	5 decimal places if CTL < 1.				
	2: Enabled for table value				
	API-2540 rounding and truncating rules are applied and, in case of				
	conversion method 2 (standard to observed), the table value for CTL as				
	specified in Table 54 meaning that the CTL value has 4 decimal places in all				
	cases				
	3: Enabled with 5 decimal places				
	API-2540 rounding and truncating rules are applied, and, in case of				
	conversion method 2 (standard to observed), the CTL value has 5 decimal				
	places in all cases.				
	Note: although not strictly in accordance with the standard, this option is				
	more commonly used than option 'Enabled for computational value'				
	Note: for conversion type 1 'From observed to standard conditions' the CTL				
	factor is rounded to 6 decimal places when input 'API 2540 rounding' > 0, as				
	in accordance with table 53.				

Function inputs	Remark	EU	SW tag	Range	Default
Hydrometer correction	Only applies for conversion method	-	HYDROCOR		0
	'1: From observed to standard conditions'				
	0: Disabled				
	1: Enabled				
API 11.2.1M rounding	0: Disabled	-	API1121RND		0
j	The calculation of the compressibility factor F is performed with full precision.				
	1: Enabled				
	API-MPMS 11.2.1M rounding and truncating rules are applied. The compressibility factor F is rounded to 3 decimal places as specified in the standard.				
Equilibrium pressure	The equilibrium pressure is considered to be 0 bar(g) for liquids which have an equilibrium pressure less than atmospheric pressure (in compliance with API MPMS 12.2 par. 12.2.5.4)	bar(g)	EQUIPRES	0150	0
Conversion method	1: From observed to standard conditions		CONVERSION		1
	2: From standard to observed conditions				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Output density	Meaning depends on the input 'Conversion method'.	kg/m3	DENS		0
	'Conversion method' = 1				
	Density at 15 °C and the equilibrium pressure.				
	'Conversion method' = 2				
	Density at the observed temperature and pressure				
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API2540 rounding"				
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API 11.2.1M rounding"				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
ко	Actual value of constant K0 used for CTL calculation	-	KO		0
K1	Actual value of constant K1 used for CTL calculation	-	K1		0
К2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor	1/°C	ALPHA		0
F	Compressibility factor	1/bar	F		0
Product	When input 'Product' is 'B - Auto select', then the output is set to the actual	-	PRDCUR		0
	selected product of tables 53B/54B (enumerative value as defined for				
	input 'Product'), else the output is set equal to input 'Product'.				
CTL calc out of range	With respect to the standard used for the calculation of CTL the				0
	combination of input values is:				
	0: In Range				
	1: Out of Range			CTLOOR	
CPL calc out of range	With respect to the standard used for the calculation of CPL the				0
	combination of input values is:				
	0: In Range				
	1: Out of Range			CPLOOR	

Calculations

The calculations depend on the conversion method.

Conversion method 1: from observed to standard conditions. The function performs the following iterative algorithm to calculate the density at reference conditions:

- 1 First the inputs are rounded in accordance with the API2540 standard, provided that API2540 rounding is enabled.
- 2 The hydrometer correction on the input density is applied, provided that this correction is enabled
- 3 At the start of the iteration the density at [15 °C, equilibrium pressure] is set equal to the observed density and the initial CPL value is set to 1.
- 4 When the type of product is set to 'B Auto select' (automatic selection of the refined product range) the KO, K1 and K2 factors are determined based on the density at [15 °C, equilibrium pressure]. The Transition area is only taken in consideration in the 2nd iteration loop, as specified in the standard.

- 5 The Alpha factor is calculated according from the density at [15 °C, equilibrium pressure] and the KO, K1 and K2 factor. If API2540 rounding is enabled, then the intermediate results are rounded or truncated as specified API-2540 Table 53.
- 6 The CTL value is calculated according to API-2540 Table 53 from the Alpha factor and the differential temperature (= observed temperature – 15°C). If API2540 rounding is enabled, then the intermediate results are rounded or truncated as specified API-2540 Table 53.
- 7 Depending on the type of API2540 rounding the calculated CTL value is rounded to 6 decimal places or not rounded at all.
- 8 The density at [15 °C, equilibrium pressure] is calculated by dividing the observed density by the new CTL value and the CPL value from the previous iteration.
- 9 The compressibility factor is calculated according to API MPMS 11.2.1M from the density at [15 °C, equilibrium pressure] and the 'Observed temperature'. If API 11.2.1M rounding is enabled then the density and temperature are rounded and

the calculations are performed in accordance with the rounding and truncating rules of the standard.

- 10 The CPL value is calculated from the compressibility factor and the 'Observed pressure' and 'Equilibrium pressure' input values.
- 11 The density at [15°C, equilibrium pressure] is calculated by dividing the observed density by CTL and the new CPL value.
- 12 If API2540 rounding is enabled then the density at [15°C, equilibrium pressure] value is rounded to 3 decimal places as specified in the standard.
- 13 Steps 4 through 12 are repeated taking the density value from step 12 as the starting value until the absolute difference between two consecutive density values is either 0.05 (or 0.07 for the transition area) or 0.000001, depending of API2540 rounding being enabled or not.
- 14 For refined products the entire iteration loop is repeated if the density at [15°C, equilibrium pressure] appears to be in a different product region than the observed input density. This is required because a different product region means different K0, K1 and K2 factors.
- 15 When API2540 rounding is enabled, the final density at [15°C, equilibrium pressure] is rounded to 1 decimal place.

Conversion method 2: from standard to observed conditions. The function performs straightforward calculations to determine the density at observed conditions:

- 1 First the inputs are rounded in accordance with the API2540 standard, provided that API2540 rounding is enabled.
- 2 When the type of product is set to 'B Auto select' (automatic selection of the refined product range) the K0, K1 and K2 factors are determined based on the input density
- 3 The Alpha factor is calculated according from the input density and the K0, K1 and K2 factor. If API2540 rounding is enabled, then the intermediate results are rounded or truncated as specified API-2540 Table 54.
- 4 The CTL value is calculated according to API-2540 Table 54 from the Alpha factor and the differential temperature (= observed temperature – 15°C If API2540 rounding is enabled, then the intermediate results are rounded or truncated as specified API-2540 Table 54.
- 5 Depending on the type of API2540 rounding the calculated CTL value is rounded to 4 or 5 decimal places or not rounded at all.
- 6 The compressibility factor is calculated according to API MPMS 11.2.1M from the input density and temperature'. If API 11.2.1M rounding is enabled then the input density and temperature are rounded and the calculations are performed in accordance with the rounding and truncating rules of the standard.
- 7 The CPL value is calculated from the compressibility factor and the 'Observed pressure' and 'Equilibrium pressure' input values.
- 8 The density at [15°C, equilibrium pressure] is calculated by multiplying the input density by the CTL and the CPL values.

fxAPI_Dens15C_NGL_LPG

Description

Density (T, P) <--> Density (15°C, Pe)

This function converts the density value at the observed temperature and pressure to the density value at 15°C and the equilibrium pressure or vice versa.

The temperature correction is according to API MPMS 11.2.4:2007 (GPA TP-27), while the pressure correction is according to API MPMS 11.2.2M:1984.

The calculation of the equilibrium pressure is according to GPA TP-15 (API MPMS 11.2.2 Addendum:1994).

Compliance

- API MPMS 11.2.4: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007
- GPA TP-27: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007
- API MPMS Chapter 11.2.2M 1986 (Compressibility Factors for Hydrocarbons: 350-637 kg/m3 Density (15°C) and -46°C to 60°C)
- API MPMS 11.2.5: A Simplified Vapor Pressure Correlation for Commercial NGLs, September 2007
- GPA TP-15: A Simplified Vapor Pressure Correlation for Commercial NGLs, September 2007 (also covers GPA TP-15 1988)
- API MPMS 11.2.2 Addendum : Compressibility Factors for Hydrocarbons: Correlation of Vapor Pressure for Commercial Natural Gas Liquids (same as GPA TP-15:1988)

Function inputs

Name	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
nput density	Depending on the conversion method this represents the density either at	-		0750	0
	the observed temperature and pressure or at 15 $^\circ C$ and the equilibrium				
	pressure				
Observed temperature	Temperature at which the density is observed	°C		-100150	15
Observed pressure	Pressure at which the density is observed	bar(a)		-1200	0
API 11.2.4 rounding	0: Disabled The calculations are performed with full precision and the output values are not rounded		API1124RND		0
	1: Enabled				
	The related values are rounded as defined in the standard				
API 11.2.2M rounding	0: Disabled The calculations are performed with full precision and the output values are not rounded 1: Enabled The related values are rounded as defined in the standard		API1122RND		0
Equilibrium pressure mode	 Use Input The value of input 'Equilibrium pressure value' is used for the calculation of CPL Second Second		EQUIPMODE		2
Equilibrium pressure value	Only used when input 'Equilibrium pressure mode' is set to 'Use input'. The value will be used for the calculation of the CPL	bar(a)	EQUIPINP		0
GPA TP-15 rounding	Only used when 'Equilibrium pressure mode is set to 'GPA TP-15' 0: Disabled Full precision (no rounding and truncating applied) 1: Enabled Rounding as defined in ' GPA TP15:1988 / API MPMS 11.2.2 Addendum':1994	-	TP15RND		0
P100 Correlation	 Only used when 'Equilibrium pressure mode is set to 'GPA TP-15' O: Disabled The standard correlation is commonly used for pure products such as propane, butane and natural gasoline. It only requires the relative density and the temperature to calculate the vapor pressure 1: Enabled The improved correlation requires the vapor pressure at 100°F (37.8 °C). This method is better suited for varied NGL mixes Where different product mixes could have the same specific gravity but different equilibrium pressures. 	-	P100CORR		0
Vapor pressure at 100°F	Only used when 'Equilibrium pressure mode is set to 'GPA TP-15' and the P100 correlation is enabled.	bar(a)	EQUIP100F	0200	0
Conversion method	1: From observed to standard conditions 2: From standard to observed conditions		CONVMETH		1

Function outputs					
Name	Remark	EU	SW tag	Alarm	Fallback
Name	Remark	EU	SW tag	Alarm	Fallba

Name	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Output density	Depending on the conversion method this represents the density	kg/m3	DENS		0
	either at 15 $^\circ$ C and the equilibrium pressure or the observed				
	temperature and pressure				
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API 11.2.4 rounding'				
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API 11.2.2M rounding'				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
F	Compressibility factor	1/bar	F		0
1	The output value will be either rounded or not depending input 'API	1/ 64	•		Ũ
	rounding'				
Equilibrium pressure	The equilibrium pressure calculated by GPA TP-15	bar(a)	EQUIPCUR		0
Equilibrium pressure	Will be set to 0 when equilibrium pressure is below atmospheric	Dai (a)	LOUFCON		0
	pressure				
CTL calc out of range	With respect to the API 11.2.4 standard the combination of input values				0
CTL calc out of range	is:				0
	0: In Range				
	1: Out of Range			CTLOOR	
	The following range checks apply:			CILOOR	
	Conversion method 1: observed -> standard				
	0.21 <= RD <= 0.74				
	with RD = Input density / 999.016/CPL				
	-46 <= T <= 93 °C				
	Table 23E reference fluid ranges				
	Conversion method 1: standard -> observed				
	351.7 <= Input density <= 687.8 kg/m3				
	-46 <= T <= 93 °C				
	Table 23E reference fluid ranges				
CPL calc out of range	With respect to API 11.2.2M the combination of input values is:				0
	0: In Range				
	1: Out of Range				
	The following range checks apply:			CPLOOR	
	350 <= Density 15 °C <= 637 kg/m3				
	-46 °C <= T <= 60 °C				
GPA TP-15 out of range	Only set when the GPA TP-15 calculation is enabled	-			0
	With respect to the GPA TP-15 standard the combination of input				
	values is:				
	0: In Range				
	1: Out of Range				
	The following range checks apply:			TP15OOR	
	For lower range:				
	0.350 <= RD60 < 0.425				
	-50 to (695.51*RD60 - 155.51) °F				
	Higher range:				
	0.425 <= RD60 <= 0.676				
	-50 to 140 °F				

Calculations

The calculations depend on the conversion method.

Conversion method 1: from observed to standard conditions. The function performs the following iterative algorithm to calculate the density at 15 $^{\circ}$ C and the equilibrium pressure.

- 1 When API 11.2.4 rounding is enabled, the input density and temperature values are rounded in accordance with the standard
- $2\,$ At the start of the iteration the density at [15 °C, equilibrium pressure] is set equal to the observed density and the CPL value is set to 1.
- 3 First the density corrected for pressure is calculated by dividing the observed density by the CPL value.
- 4 The relative density corrected for pressure is calculated from the density corrected for pressure

- 5 The relative density at [60 °F, equilibrium pressure] is calculated from the relative density corrected for pressure and the observed temperature according to Table 23E
- 6 The relative density at [15 °C, equilibrium pressure] is calculated from the relative density at [60 °F, equilibrium pressure] converted to 15 °C according to Table 24E
- 7 The density at [15 °C, equilibrium pressure] is calculated from the relative density at [15 °C, equilibrium pressure]
- 8 The CTL value is calculated by dividing the density corrected for pressure by the density at [15 °C, equilibrium pressure]
- 9 Depending on the value of input 'Equilibrium pressure mode', either value of input 'Equilibrium pressure value' is used or the equilibrium pressure (vapor pressure) is calculated according to GPA TP-15. Whether the GPA TP-15 rounding and truncation rules are applied is dictated by input 'GPA-TP15 rounding'
- 10 The compressibility factor F is calculated according to API MPMS 11.2.2M from the density at [15 °C, equilibrium pressure] and the 'Observed temperature', with, depending on

input API 11.2.2M, rounding and truncation according to the standard.

- 11 The CPL value is calculated from the compressibility factor, the equilibrium pressure and the 'Observed pressure' input value.
- 12 The new value for density at [15°C, equilibrium pressure] is calculated by dividing the observed density by the CTL and CPL values.
- 13 Steps 3 though 12 are repeated taking the density value from step 12 as the staring value until the absolute difference between two consecutive density values is less than the convergence limit.

To avoid convergence problems different convergence limits are applied, depending on the whether API 11.2.2M and/or GPA TP-15 rounding is applied:

If API 11.2.2M rounding is enabled -> Limit = 0.05 kg/m3 else if GPA TP-15 rounding is enabled -> Limit = 0.005 kg/m3 else -> Limit = 0.00001 kg/m3

14 If API 11.2.4 rounding is enabled, then the density at [15°C, equilibrium pressure] is rounded to 0.1

Conversion method 2: from standard to observed conditions. The function performs straightforward calculations to determine the density at observed conditions:

- 1 When API 11.2.4 rounding is enabled, the input density and temperature values are rounded in accordance with the standard
- 2 The CTL value and the relative density at [60 °F, equilibrium pressure] are calculated according to API MPMS 11.2.4 (GPA TP-27) Table 60E from the density at [15 °C, equilibrium pressure] and the 'Observed temperature'.
- 3 Depending on the value of input 'Equilibrium pressure mode', either value of input 'Equilibrium pressure value' is used or the equilibrium pressure (vapor pressure) is calculated according to GPA TP-15.
- 4 The compressibility factor is calculated according to API MPMS 11.2.2M from the density at [15 °C, equilibrium pressure] and the 'Observed temperature'.
- 5 The CPL value is calculated from the compressibility factor, the equilibrium pressure and the 'Observed pressure' input value.
- 6 If API 11.2.4 rounding is enabled, then the CTL value is rounded at [60°F, equilibrium pressure] is rounded to 0.00001
- 7 The density at the observed conditions is calculated by multiplying the density at [15 °C, equilibrium pressure] by the CTL value and the CPL value.

fxAPI_Dens20C_NGL_LPG

Description

Density (T, P) <--> Density (20°C, Pe)

This function converts the density value at the observed temperature and pressure to the density value at 20°C and the equilibrium pressure or vice versa.

The temperature correction is according to API MPMS 11.2.4:2007 (GPA TP-27), while the pressure correction is according to API MPMS 11.2.2M:1984.

The calculation of the equilibrium pressure is according to GPA TP-15 (API MPMS 11.2.2 Addendum:1994).

Compliance

- API MPMS 11.2.4: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007
- GPA TP-27: Temperature Correction for the Volume of NGL & LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007
- API MPMS Chapter 11.2.2M 1986 (Compressibility Factors for Hydrocarbons: 350-637 kg/m3 Density (15°C) and -46°C to 60°C)
- API MPMS 11.2.5: A Simplified Vapor Pressure Correlation for Commercial NGLs, September 2007
- GPA TP-15: A Simplified Vapor Pressure Correlation for Commercial NGLs, September 2007 (also covers GPA TP-15 1988)
- API MPMS 11.2.2 Addendum : Compressibility Factors for Hydrocarbons: Correlation of Vapor Pressure for Commercial Natural Gas Liquids (same as GPA TP-15:1988)

Function inputs

Name	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group		-		
Input density	Depending on the conversion method this represents the density either	kg/m3		0750	0
	at the observed temperature and pressure or at 20 °C and the				
	equilibrium pressure				
Observed temperature	Temperature at which the density is observed	°C		-100150	20
Observed pressure	Pressure at which the density is observed	bar(a)		-1200	0
API 11.2.4 rounding	0: Disabled		API1124RND		0
Ū.	The calculations are performed with full precision and the output				
	values are not rounded				
	1: Enabled				
	The related values are rounded as defined in the standard				
API 11.2.2M rounding	0: Disabled		API1122RND		0
/	The calculations are performed with full precision and the output				
	values are not rounded				
	1: Enabled				
	The related values are rounded as defined in the standard				
Equilibrium pressure mode	1: Use Input		EQUIPMODE		2
	The value of input 'Equilibrium pressure value' is used for the				
	calculation of CPL				
	2: GPA TP-15				
	The equilibrium pressure is calculated in accordance with GPA TP-15				
Equilibrium pressure value	Only used when input 'Equilibrium pressure mode' is set to 0.	bar(a)	EQUIPINP		0
	The value will be used for the calculation of the CPL				
GPA TP-15 rounding	0: Disabled	-	TP15RND		0
5	Full precision (no rounding and truncating applied)				
	1: Enabled				
	Rounding as defined in ' GPA TP15:1988 / API MPMS 11.2.2				
	Addendum':1994				
P100 Correlation	0: Disabled	-	P100CORR		0
	The standard correlation is commonly used for pure products such as				
	propane, butane and natural gasoline. It only requires the relative				
	density and the temperature to calculate the vapor pressure				
	1: Enabled				
	The improved correlation requires the vapor pressure at 100°F (37.8				
	°C). This method is better suited for varied NGL mixes Where				
	different product mixes could have the same specific gravity but				
	different equilibrium pressures.				
Vapor pressure at 100°F		bar(a)	EQUIP100F	0200	0
Conversion method	1: From observed to standard conditions		CONVMETH		1
	2: From standard to observed conditions				

Function outputs

Name	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		1
	1: Input argument out of range			FIOOR	
	Outputs will be set to fallback values				
	2: Calculation error			CALC	

Name	Remark	EU	SW tag	Alarm	Fallback
	Outputs will be set to fallback values				
	3: No convergence within 15 iterations			NOCONV	
	Outputs will be set to values of last iteration				
Output density	Depending on the conversion method this represents the density	kg/m3	DENS		0
	either at 20 °C and the equilibrium pressure or the observed				
	temperature and pressure				
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API 11.2.4 rounding'				
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API 11.2.2M rounding'				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
F	Compressibility factor	1/bar	F		0
	The output value will be either rounded or not depending input 'API				
	rounding'				
Equilibrium pressure	The equilibrium pressure calculated by GPA TP-15	bar(a)	EQUIPCUR		0
	Will be set to 0 when equilibrium pressure is below atmospheric				
	pressure				
CTL calc out of range	With respect to the API 11.2.4 standard the combination of input				0
	values is:				
	0: In Range				
	1: Out of Range			CTLOOR	
	The following range checks apply:				
	Conversion method 1: observed -> standard				
	0.21 <= RD <= 0.74				
	with RD = Input density / 999.016/CPL				
	-46 <= T <= 93 °C				
	Table 23E reference fluid ranges				
	Conversion method 1: standard -> observed				
	331.7 <= Input density <= 683.6 kg/m3				
	-46 <= T <= 93 °C				
	Table 23E reference fluid ranges				
CPL calc out of range	With respect to API 11.2.2M the combination of input values is:				0
-	0: In Range				
	1: Out of Range				
	The following range checks apply:			CPLOOR	
	350 <= Density 15 °C <= 637 kg/m3				
	-46 °C <= T <= 60 °C				
GPA TP-15 out of range	Only set when the GPA TP-15 calculation is enabled	-			0
	With respect to the GPA TP-15 standard the combination of input				
	values is:				
	0: In Range				
	1: Out of Range				
	The following range checks apply:			TP15OOR	
	For lower range:				
	0.350 <= RD60 < 0.425				
	-50 to (695.51*RD60 - 155.51) °F				
	Higher range:				
	0.425 <= RD60 <= 0.676				
	-50 to 140 °F				
	with RD60 being the relative density at 60°F				

Calculations

The calculations depend on the conversion method.

Conversion method 1: from observed to standard conditions. The function performs the following iterative algorithm to calculate the density at 20 $^{\circ}$ C and the equilibrium pressure.

- 1 When API 11.2.4 rounding is enabled, the input density and temperature values are rounded in accordance with the standard
- 2 At the start of the iteration the density at [20 °C, equilibrium pressure] is set equal to the observed density and the CPL value is set to 1.
- 3 First the density corrected for pressure is calculated by dividing the observed density by the CPL value.
- 4 The relative density corrected for pressure is calculated from the density corrected for pressure
- 5 The relative density at [60 °F, equilibrium pressure] is calculated from the relative density corrected for pressure and the observed temperature according to Table 23E

- 6 The relative density at [20 °C, equilibrium pressure] is calculated from the relative density at [60 °F, equilibrium pressure] converted to 20 °C according to Table 24E
- 7 The density at [20 °C, equilibrium pressure] is calculated from the relative density at [20 °C, equilibrium pressure]
- 8 The CTL value is calculated by dividing the density corrected for pressure by the density at [20 °C, equilibrium pressure]
- 9 Depending on the value of input 'Equilibrium pressure mode', either value of input 'Equilibrium pressure value' is used or the equilibrium pressure (vapor pressure) is calculated according to GPA TP-15. Whether the GPA TP-15 rounding and truncation rules are applied is dictated by input 'GPA-TP15 rounding'
- 10 API 11.2.2M requires the density at [15 °C, equilibrium pressure]. For this purpose the relative density at [15 °C, equilibrium pressure] is calculated according to Table 24E from the relative density at [60 °F, equilibrium pressure] and at 15 °C. This relative density value is then converted to the density at [15 °C, equilibrium pressure].

- 11 The compressibility factor F is calculated according to API MPMS 11.2.2M from the density at [15 °C, equilibrium pressure] and the 'Observed temperature', with, depending on input API 11.2.2M, rounding and truncation according to the standard.
- 12 The CPL value is calculated from the compressibility factor, the equilibrium pressure and the 'Observed pressure' input value.
- 13 The new value for density at [20°C, equilibrium pressure] is calculated by dividing the observed density by the CTL and CPL values.
- 14 Steps 2 though 6 are repeated taking the density value from step 6 as the staring value until the absolute difference between two consecutive density values is less than the convergence limit.

To avoid convergence problems different convergence limits are applied, depending on the whether API 11.2.2M and/or GPA TP-15 rounding is applied:

If API 11.2.2M rounding is enabled -> Limit = 0.05 kg/m3 else if GPA TP-15 rounding is enabled -> Limit = 0.005 kg/m3 else -> Limit = 0.00001 kg/m3

If API 11.2.4 rounding is enabled, then the density at [20°C, equilibrium pressure] is rounded to 0.1

Conversion method 2: from standard to observed conditions. The function performs straightforward calculations to determine the density at observed conditions:

- 1 When API 11.2.4 rounding is enabled, the input density and temperature values are rounded in accordance with the standard
- 2 The CTL value and the relative density at [60 °F, equilibrium pressure] are calculated according to API MPMS 11.2.4 (GPA TP-27) Table 54 from the density at [20 °C, equilibrium pressure] and the 'Observed temperature'.
- 3 Depending on the value of input 'Equilibrium pressure mode', either value of input 'Equilibrium pressure value' is used or the equilibrium pressure (vapor pressure) is calculated according to GPA TP-15.
- 4 API 11.2.2M requires the density at [15 °C, equilibrium pressure]. For this purpose the relative density at [15 °C, equilibrium pressure] is calculated according to Table 24E from the relative density at [60 °F, equilibrium pressure] and at 15 °C. This relative density value is then converted to the density at [15 °C, equilibrium pressure].
- 5 The compressibility factor is calculated according to API MPMS 11.2.2M from the density at [15 °C, equilibrium pressure] and the 'Observed temperature'.
- 6 The CPL value is calculated from the compressibility factor, the equilibrium pressure and the 'Observed pressure' input value.
- 7 If API 11.2.4 rounding is enabled, then the CTL value is rounded at [60°F, equilibrium pressure] is rounded to 0.00001
- 8 The density at the observed conditions is calculated by multiplying the input density by the CTL value and the CPL value.

fxAPI_Gravity60F_1952

Description

°API (T, P) <--> °API (60°F, equilibrium pressure)

This function calculates the API gravity value at the observed temperature and pressure to the API gravity value at 60°F and the equilibrium pressure (typically 0 psig) or vice versa.

The volume correction for temperature is according to 1952 API Table 5 and 6, while the volume correction for pressure is according to API MPMS 11.2.1. Note: this function is a combination of the API 1952 Tables and API 11.2.1. For the calculation from observed to standard conditions an iterative calculation is required. The rounding and truncating of input and intermediate values is implemented such that the example calculations as specified in both standards are exactly reproduced.

Compliance

- ASTM-IP Petroleum Measurement Tables, American Edition, United States Units of Measurement, 1952
- API MPMS 11.2.1 Compressibility Factors for Hydrocarbons:
 0 90°API Gravity Range First Edition, August 1984

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Input API gravity	Depending of the conversion method this represents the API gravity	°API		-20120	0
	at either the observed temperature and pressure or at 60 $^\circ$ F and the				
	equilibrium pressure				
Observed temperature	Temperature at which the API gravity is observed	°F		-100400	60
Observed pressure	Pressure at which the API gravity is observed	psig		-102000	0
API 11.2.1 rounding	0: Disabled	-	API1121RND		0
	The calculation of the compressibility factor F is performed with				
	full precision				
	1: Enabled				
	API-MPMS 11.2.1 rounding and truncating rules are applied. The				
	compressibility factor F is rounded to 3 decimal places as specified				
	in the standard.				
Equilibrium pressure	The equilibrium pressure is considered to be 0 psig for liquids which	psig	EQUIPRES	02000	0
	have an equilibrium pressure less than atmospheric pressure (in				
	compliance with API MPMS 12.2 par. 12.2.5.4)				
Conversion method	1: From observed to standard conditions		CONVERSION		1
	2: From standard to observed conditions				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALC	
	3: No convergence			NOCONV	
Output API gravity	Depending of the conversion method this represents the API gravity at either	°API	API		0
	at 60 °F and the equilibrium pressure or the observed temperature and				
	pressure				
CTL	Volume correction factor for temperature.	-	CTL		1
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API 11.2.1 rounding'"				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
F	Compressibility factor	1/psi	F		0
CTL calc out of range	With respect to the standard used for the calculation of CTL the combination				0
5	of input values is:				
	0: In Range				
	1: Out of Range			CTLOOR	
CPL calc out of range	With respect to the standard used for the calculation of CPL the combination				0
_	of input values is:				
	0: In Range				
	1: Out of Range			CPLOOR	

Calculations

The calculations depend on the conversion method.

Conversion method 1: from observed to standard conditions. The function performs the following iterative algorithm to calculate the API Gravity at standard conditions:

- At the start of the iteration the initial value for API Gravity at [60 °F, equilibrium pressure] is set to the Observed API Gravity. The initial CPL value is set to 1.
- 2 The CTL value is determined from the API Gravity at [60 °F, equilibrium pressure] according to API 1952 Table 6.
- 3 The API Gravity at [60 °F, equilibrium pressure] is calculated from the Observed API gravity, the new CTL value and the CPL value from the previous iteration.
- 4 Because API 11.2.1 requires the API gravity value at 60 °F, the API gravity at [60 °F, equilibrium pressure] is calculated from the API gravity at [60 °F, equilibrium pressure].
- 5 The compressibility factor is calculated according to API MPMS 11.2.1 from the API gravity at [60 °F, equilibrium pressure] and the 'Observed temperature'. If API 11.2.1 rounding is enabled then the API gravity and temperature are rounded and the calculations are performed in accordance with the rounding and truncating rules of the standard.
- 6 The CPL value is calculated from the compressibility factor and the 'Observed pressure' and 'Equilibrium pressure' input values.
- 7 The API Gravity at [60°F, equilibrium pressure] is calculated by dividing the Observed API Gravity by the CTL and the new CPL value.
- 8 Steps 2 through 7 are repeated taking the API gravity value from step 7 as the start value for the next iteration until the absolute difference between two consecutive API gravity values is 0.01.

Conversion method 2: from standard to observed conditions. The function performs straightforward calculations to determine the API Gravity at observed conditions:

- 1 The CTL value is calculated according to API 1952 Table 6
- 2 Because API 11.2.1 requires the API gravity value at 60 °F, the API gravity at 60 °F is calculated from the 'Input API Gravity'.
- 3 The compressibility factor is calculated according to API MPMS 11.2.1 from the API gravity and the 'Observed temperature'. If API 11.2.1 rounding is enabled then the input density and temperature are rounded and the calculations are performed in accordance with the rounding and truncating rules of the standard.
- 4 The CPL value is calculated from the compressibility factor and the 'Observed pressure' and 'Equilibrium pressure' input values.
- 5 The output API Gravity (at observed temperature and pressure) is calculated from the input API Gravity and the CTL and the CPL values.

fxAPI_Gravity60F_1980

Description

°API (T, P) <--> °API (60°F, equilibrium pressure)

This function calculates the API gravity value at the observed temperature and pressure to the API gravity value at 60°F and the equilibrium pressure (typically 0 psig) or vice versa. The volume correction for temperature is according to API-2540, Tables 5/6A (Generalized Crude Oils) and 5/6B (Refined Oil Products) and API MPMS 11.1 Chapter XIII Table 5D: 1984 (Lubricating Oils), while the volume correction for pressure according to API MPMS 11.2.1.

Note: this function is a combination of API2540 and API 11.2.1. For the calculation from observed to standard conditions an iterative calculation is required. The rounding and truncating of input and intermediate values is implemented such that the example calculations as specified in both standards are exactly reproduced.

Compliance

- API MPMS 11.1 Volume X (API Standard 2540) Table 5A -Generalized Crude Oils, Correction of Observed API Gravity to API Gravity at 60°F - First Edition, August 1980
- API MPMS 11.1 Volume X (API Standard 2540) Table 5B -Generalized Products, Correction of Observed API Gravity to API Gravity at 60°F- First Edition, August 1980
- API MPMS 11.1 Volume XIII Table 5D Generalized Lubricating Oils, Correction of Observed API Gravity to API Gravity at 60°F - January 1982
- API MPMS 11.1 Volume X (API Standard 2540) Table 6A -Generalized Crude Oils, Correction of Volume to 60°F against API Gravity at 60°F - First Edition, August 1980
- API MPMS 11.1 Volume X (API Standard 2540) Table 6B -Generalized Products, Correction of Volume to 60°F against API Gravity at 60°F - First Edition, August 1980
- API MPMS 11.1 Volume XIII Table 6D Generalized Lubricating Oils, Correction of Volume to 60°F against API Gravity at 60°F F - January 1982
- API MPMS 11.2.1 Compressibility Factors for Hydrocarbons: 0 - 90°API Gravity Range - First Edition, August 1984

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Input API gravity	Depending of the conversion method this represents the API gravity at	°API		-20120	0
	either the observed temperature and pressure or at 60 $^\circ F$ and the				
	equilibrium pressure				
Observed temperature	Temperature at which the API gravity is observed	°F		-50400	60
Observed pressure	Pressure at which the API gravity is observed	psig		-102000	0
Product	1: A - Crude Oil		PRDTYP	-	1
	2: B - Auto select				
	Selection based on °API at 60 °F				
	3: B - Gasoline				
	4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API-2540 rounding	0: Disabled		API2540RND	-	0
	The calculations are performed with full precision and the final CTL value				
	is rounded as specified by input 'CTL decimal places'				
	1: Enabled for computational value				
	API-2540 rounding and truncating rules are applied and, in case of				
	conversion method 2 (standard to observed), the computational value				
	for CTL as specified in Table 6 is used, meaning that the CTL value has:				
	4 decimal places if CTL >=1				
	5 decimal places if CTL < 1.				
	2: Enabled for table value				
	API-2540 rounding and truncating rules are applied and, in case of				
	conversion method 2 (standard to observed), the table value for CTL as				
	specified in Table 6 meaning that the CTL value has 4 decimal places in				
	all cases				
	3: Enabled with 5 decimal places				
	API-2540 rounding and truncating rules are applied, and, in case of				
	conversion method 2 (standard to observed), the CTL value has 5				
	decimal places in all cases.				
	Note: although not strictly in accordance with the standard, this option				
	is more commonly used than option 'Enabled for computational value'				
	Note: for conversion type 1 'From observed to standard conditions' the				
	CTL factor is rounded to 6 decimal places when input 'API 2540 rounding'				
	> 0, as in accordance with table 5.				
Hydrometer correction	Only applies for conversion method	-	HYDROCOR		0
,	'1: From observed to standard conditions'				
	0: Disabled				
	1: Enabled				

Function inputs	Remark	EU	SW tag	Range	Default
API 11.2.1 rounding	 0: Disabled The calculation of the compressibility factor F is performed with full precision 1: Enabled API-MPMS 11.2.1 rounding and truncating rules are applied. The compressibility factor F is rounded to 3 decimal places as specified in the standard. 	-	API1121RND		0
Equilibrium pressure	The equilibrium pressure is considered to be 0 psig for liquids which have an equilibrium pressure less than atmospheric pressure (in compliance with API MPMS 12.2 par. 12.2.5.4)	psig	EQUIPRES	02000	0
Conversion method	1: From observed to standard conditions 2: From standard to observed conditions		CONVERSION		1

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALC	
	3: No convergence			NOCONV	
Output API gravity	Depending of the conversion method this represents the API gravity at either at	°API	API		0
	60 °F and the equilibrium pressure or the observed temperature and pressure				
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API2540 rounding'				
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API 11.2.1 rounding'"				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
ко	Actual value of constant K0 used for CTL calculation	-	К0		0
K1	Actual value of constant K1 used for CTL calculation	-	K1		0
K2	Actual value of constant K2 used for CTL calculation	-	К2		0
Alpha	Thermal expansion factor	1/°F	ALPHA		0
F	Compressibility factor	1/psi	F		0
Product	When input 'Product' is 'B - Auto select', then the output is set to the actual	-	PRDCUR		0
	selected product of table 5B / 6B (enumerative value as defined for input				
	'Product'), else the output is set equal to input 'Product'.				
CTL calc out of range	With respect to the standard used for the calculation of CTL the combination of				0
_	input values is:				
	0: In Range				
	1: Out of Range			CTLOOR	
CPL calc out of range	With respect to the standard used for the calculation of CPL the combination of				0
2	input values is:				
	0: In Range				
	1: Out of Range			CPLOOR	

Calculations

The calculations depend on the conversion method.

Conversion method 1: from observed to standard conditions. The function performs the following iterative algorithm to calculate the API gravity at standard conditions:

- 1 First the inputs are rounded in accordance with the API2540 standard, provided that API2540 rounding is enabled.
- 2 The observed density [kg/m3] is calculated from the observed API gravity
- 3 The hydrometer correction on the observed density is applied, provided that this correction is enabled
- 4 At the start of the iteration the initial value for density and API gravity at [60 °F, equilibrium pressure] is set to respectively the observed density and the observed API gravity. The initial CPL value is set to 1.
- 5 When the type of product is set to 'B Auto select' (automatic selection of the refined product range) the K0, K1 and K2 factors are determined based on the API gravity at [60 °F, equilibrium pressure]. The Transition area is only taken in consideration in the 2nd iteration loop, as specified in the standard.
- 6 The Alpha factor is calculated according from the density at [60 °C, equilibrium pressure] and the K0, K1 and K2 factor. If

API2540 rounding is enabled, then the intermediate results are rounded or truncated as specified API-2540 Table 5.

- 7 The CTL value is calculated according to API-2540 Table 5 from the Alpha factor and the differential temperature (= observed temperature – 60°F). If API2540 rounding is enabled, then the intermediate results are rounded or truncated as specified API-2540 Table 5.
- 8 Depending on the type of API2540 rounding the calculated CTL value is rounded to 6 decimal places or not rounded at all.
- 9 The density at [60 °F, equilibrium pressure] is calculated by dividing the observed density by the new CTL value and the CPL value from the previous iteration.
- 10 The API gravity at [60 °F, equilibrium pressure] is calculated from the density at [60 °F, equilibrium pressure]
- 11 The compressibility factor is calculated according to API MPMS 11.2.1 from the API gravity at [60 °F, equilibrium pressure] and the 'Observed temperature'. If API 11.2.1 rounding is enabled then the API gravity and temperature are rounded and the calculations are performed in accordance with the rounding and truncating rules of the standard.
- 12 The CPL value is calculated from the compressibility factor and the 'Observed pressure' and 'Equilibrium pressure' input values.

- 13 The density at [60°F, equilibrium pressure] is calculated by dividing the observed density by CTL and the new CPL value.
- 14 If API2540 rounding is enabled then the density at [60°F, equilibrium pressure] value is rounded to 3 decimal places as specified in the standard.
- 15 The API gravity at [60 °F, equilibrium pressure] is calculated from the density at [60 °F, equilibrium pressure]
- 16 If API2540 rounding is enabled then the API gravity at [60°F, equilibrium pressure] value is rounded to 1 decimal place as specified in the standard.
- 17 Steps 5 through 16 are repeated taking the density value from step 14 as the start value for the next iteration until the absolute difference between two consecutive density values is either 0.05 (or 0.07 for the transition area) or 0.000001, depending of API2540 rounding being enabled or not.
- 18 For refined products the entire iteration loop is repeated if the API gravity at [60°F, equilibrium pressure] appears to be in a different product region than the observed API gravity. This is required because a different product region means different K0, K1 and K2 factors.

Conversion method 2: from standard to observed conditions. The function performs straightforward calculations to determine the API gravity at observed conditions:

- 1 First the inputs are rounded in accordance with the API2540 standard, provided that API2540 rounding is enabled.
- 2 The density at [60°F, equilibrium pressure] is calculated from the input API gravity
- 3 When the type of product is set to 'B Auto select' (automatic selection of the refined product range) the K0, K1 and K2 factors are determined based on the input API gravity
- 4 The Alpha factor is calculated according from the density at [60°F, equilibrium pressure] and the K0, K1 and K2 factor. If API2540 rounding is enabled, then the intermediate results are rounded or truncated as specified API-2540 Table 6.
- 5 The CTL value is calculated according to API-2540 Table 6 from the Alpha factor and the differential temperature (= observed temperature – 60°F). If API2540 rounding is enabled, then the intermediate results are rounded or truncated as specified API-2540 Table 6.
- 6 Depending on the type of API2540 rounding the calculated CTL value is rounded to 4 or 5 decimal places or not rounded at all.
- 7 The compressibility factor is calculated according to API MPMS 11.2.1 from the input density and temperature'. If API 11.2.1 rounding is enabled then the input density and temperature are rounded and the calculations are performed in accordance with the rounding and truncating rules of the standard.
- 8 The CPL value is calculated from the compressibility factor and the 'Observed pressure' and 'Equilibrium pressure' input values.
- 9 The API gravity at observed temperature and pressure is calculated from the input API gravity and the CTL and the CPL values.

fxAPI_MPMS_11_2_1

Description

The API MPMS 11.2.1 standard consists of a printed table that contains compressibility factors to correct hydrocarbon volumes under pressure to the corresponding volumes at the equilibrium pressure for the metered temperature.

The table contains compressibility factors related to meter temperature and API gravity at 60°F.

From the compressibility factor the volume correction for pressure is calculated according to API MPMS 12.2.

Compliance

- API MPMS 11.2.1 Compressibility Factors for Hydrocarbons: 0 - 90°API Gravity Range - First Edition, August 1984
- API MPMS 12.2 Calculation of Liquid Petroleum Quantities Measured by Turbine or Displacement Meters

Boundaries

API MPMS 11.2.1 defines the following limits on the input values:

- 0 to 90 °API
- -20 to +200 °F
- 0 to 1500 psig.

API Rounding

The actual standard is the printed table. It also includes the 'Calculation Procedure' to obtain the table values based on the rounding and truncating of all input, intermediate and output values.

The function provides the option to either output the table value (including the <u>full</u> API rounding and truncating requirements) or to perform the calculation procedure without any rounding and truncating being applied.

Function inputs

Name	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
API60	API gravity at 60°F	°API		0120	0
Observed Temperature		°F		-50400	60
Observed Pressure		psig		-102000	0
Equilibrium Pressure	The equilibrium pressure is considered to be 0 psig for liquids which have an equilibrium pressure less than atmospheric pressure (in compliance with API MPMS 12.2 par. 12.2.5.4)	psig	EQUIPRES	02000	0
API 11.2.1 rounding	 0: Disabled The calculation of the compressibility factor F is performed with full precision. 1: Enabled API-MPMS 11.2.1 rounding and truncating rules are applied. Th compressibility factor F is rounded to 3 decimal places as specified in the standard. 	e	APIROUND	-	0

Function outputs

Name	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALERR	
CPL	Volume correction factor for pressure	-	CPL		1
	Note: to achieve compliance with API MPMS 12.2 the CPL value				
	needs to be rounded to 4 decimal places.				
F	Compressibility factor	1/psi	F		0
	The output value will rounded according to input 'API 11.2.1				
	rounding'				
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
	0: In Range				
	1: Out of Range				

Calculations

The calculations are in either full or partial compliance with the standards, depending on whether API rounding is enabled or not and on the actual number of decimal places for the CPL value (API MPMS 12.2 defines 4 decimal places for the CPL value). The CPL value is calculated as follows (in compliance with API MPMS 12.2):

$$CPL = \frac{1}{1 - F \cdot (Po - Pe)}$$

Else
$$CPL = \frac{1}{1 - F \cdot Po}$$

With:

CPL	Volume correction factor for pressure	-
F	Compressibility factor	1/psi
Ро	Observed pressure	psig
Pe	Equilibrium pressure	psig

fxAPI_MPMS_11_2_1M

Description

The API MPMS 11.2.1M standard consists of a printed table that contains compressibility factors to correct hydrocarbon volumes under pressure to the corresponding volumes at the equilibrium pressure for the metered temperature.

The table contains compressibility factors related to meter temperature and density at 15°C.

This metric standard corresponds with API MPMS 11.2.1 (the customary version)

Compliance

- 11.2.1M Compressibility Factors for Hydrocarbons: 638 -1074 Kilograms per Cubic Meter Range - First Edition, August 1984
- API MPMS 12.2 Calculation of Liquid Petroleum Quantities Measured by Turbine or Displacement Meters

Function inputs

Data Limits

API MPMS 11.2.1M defines the following limits on the input values:

- 638 to 1074 kg/m3
- -30 to 90 °C
- 0 to 103 bar(g).

It is advised not to use the standard outside these limits.

Name	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Density at 15°C		kg/m3		01300	0
Observed temperature		°C		-100200	15
Observed pressure		bar(g)		-1150	0
Equilibrium pressure	The equilibrium pressure is considered to be 0 bar(g) for liquids which have an equilibrium pressure less than atmospheric pressure (in compliance with API MPMS 12.2 par. 12.2.5.4)	bar(g)	EQUIPRES		0
API 11.2.1M rounding	 0: Disabled The calculation of the compressibility factor F is performed with full precision Enabled API-MPMS 11.2.1M rounding and truncating rules are applied. The compressibility factor F is rounded in accordance with the standard. 	-	APIROUND		0

Function outputs

Name	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALC	
CPL	Volume correction factor for pressure	-	CPL		1
	Note: to achieve compliance with API MPMS 12.2 the CPL value needs to				
	be rounded to 4 decimal places.				
F	Compressibility factor	1/bar	F		0
	The output value will be either rounded or not depending input 'API				
	11.2.1M rounding '				
Calculation out of range	With respect to the standard the input values are:	-			0
	0: In Range				
	1: Out of Range			OOR	

Calculations

The calculations are in either full or partial compliance with the standards, depending on whether API rounding is enabled or not and on the actual number of decimal places for the CPL value (API MPMS 12.2 defines 4 decimal places for the CPL value). The CPL value is calculated as follows (in compliance with API MPMS 12.2):

If Pe > 0 then

$$CPL = \frac{1}{1 - F \cdot (Po - Pe)}$$

Else

$$CPL = \frac{1}{1 - F \cdot Po}$$

With:

CPL	Volume correction factor for pressure	-
F	Compressibility factor	1/bar
Ро	Observed pressure	bar(g)
Pe	Equilibrium pressure	bar(g)

fxAPI_MPMS_11_2_2

Description

The API MPMS 11.2.2 standard consists of a printed table that contains compressibility factors to correct hydrocarbon volumes under pressure to the corresponding volumes at the equilibrium pressure for the metered temperature.

The table contains compressibility factors related to meter temperature and the relative density at 60°F.

Compliance

 11.2.2 - Compressibility Factors for Hydrocarbons: 0.350 -0.637 Relative Density (60°F/60°F) and -50°F to 140°F Metering Temperature - Second Edition, October 1986

Boundaries

API MPMS 11.2.2 defines the following limits on the input values:

- 0.350 to 0.637 (relative density)
- -50 to 140 °F
- 0 to 2200 psig

Pseudo-critical temperature check

The standard specifies that the observed temperature must be less than the **pseudo-critical** temperature

Observed Temperature [Rankin] < Pseudo Critical Temperature * 0.96

With:

Pseudo Critical Temperature [Rankin] = 621.418 - 822.686 * RD60 + 1737.86 * RD60^2

Observed Temperature [Rankin] = Observed Temperature [°F] + 459.7;

API Rounding

The actual standard is the printed table. Also included is a 'Calculation Procedure' that illustrates how to obtain the table values including all required rounding and truncating of the input, intermediate and output values.

The function provides the option to either output the table value (including all the <u>full</u> API rounding and truncating requirements) or to perform the calculation procedure without any rounding and truncating being applied.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Relative density at 60°F	Relative density at 60°F	-		00.75	0
Observed Temperature		°F		-100300	60
Observed Pressure		psig		-102500	0
Equilibrium Pressure	The equilibrium pressure is considered to be 0 psig for liquids which have an equilibrium pressure less than atmospheric pressure (in compliance with API MPMS 12.2 par. 12.2.5.4)	psig	EQUIPRES	02500	0
API 11.2.2 rounding	 0: Disabled The calculation of the compressibility factor F and CPL is performed with full precision. 1: Enabled API-MPMS 11.2.2 rounding and truncating rules are applied. The compressibility factor F is rounded to 8 decimal places with a maximum of 4 significant digits as specified in the standard. The CPL value is rounded to 4 decimal places in compliance with the standard 		APIROUND	-	0

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API 11.2.2 rounding'				
F	Compressibility factor	1/psi	F		0
	The output value will be either rounded or not depending input 'API				
	11.2.2. rounding'				
Range	With respect to the standard the input values are:	-		OOR	0
	0: In Range				
	1: Out of Range				
	This includes the boundaries for relative density and temperature and	d			
	also the check of the pseudo-critical temperature.				

Calculations

The calculations are in either full or partial compliance with the standard, depending on the selected type of API rounding.

fxAPI_MPMS_11_2_2M

Description

The API MPMS 11.2.2M standard consists of a printed table that contains compressibility factors to correct hydrocarbon volumes under pressure to the corresponding volumes at the equilibrium pressure for the metered temperature.

The table contains compressibility factors related to meter temperature and the density at 15°C.

This metric standard corresponds with API MPMS 11.2.2 (the U.S. customary version)

Compliance

 11.2.2M - Compressibility Factors for Hydrocarbons: 350 -637 Kilograms per Cubic Meter Density (15°C) and -46°C to 60 °C Metering Temperature - First Edition, October 1986

Boundaries

API MPMS 11.2.2M defines the following limits on the input values:

- 350 to 637 kg/m3
- -46 to 60 °C
- 0 to 152 bar(g)

Also the check on the **pseudo-critical** temperature as defined for fxAPI_MPMS_11_2_2 is applied.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Density at 15°C		kg/m3		0750	0
Observed Temperature		°C		-100150	60
Observed Pressure		bar(g)		-1200	0
Equilibrium Pressure	The equilibrium pressure is considered to be 0 psig for liquids which have an equilibrium pressure less than atmospheric pressure (in compliance with API MPMS 12.2 par. 12.2.5.4)	bar(g)	EQUIPRES	0200	0
API 11.2.2M rounding	 0: Disabled The calculation of the compressibility factor F and CPL is performed with full precision. 1: Enabled API-MPMS 11.2.2M rounding and truncating rules are applied. The compressibility factor F is rounded in accordance with the standard. 		APIROUND	-	0

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API 11.2.2M rounding'				
F	Compressibility factor	1/bar	F		0
	The output value will be either rounded or not depending input				
	'API 11.2.2M rounding'				
Calculation out of range	With respect to the standard the input values are:	-			0
	0: In Range				
	1: Out of Range			OOR	

Calculations

The calculations are in either full or partial compliance with the standard, depending on the selected type of API rounding.

fxAPI_MPMS_11_3_2_1

Description

The function calculates the density of Ethylene (C2H4, also called Ethene) based on the API MPMS 11.3.2.1 Equation Of State in **USC units**. This API chapter is also known as API Standard 2565.

References

• Supplement to API MPMS 11.3.1.1 (API Std. 1565) Ethylene Density, 1974, Reaffirmed 1993.

Function inputs

Boundaries

The equation of state is valid from $65...165\ ^\circ F$ and $200\ ...\ 2100$ psia.

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Temperature		°F	-100300	0
Pressure		psia	03000	0
API rounding	0: Disabled	-		1
	The calculations are performed with full precision. A convergence limit of 1e-10			
	lbm/ft3 will be applied for the iterative calculations.			
	1: Enabled			
	The calculations are performed in full compliance with the standard. The input,			
	intermediate and output values are rounded as specified and also the iteration			
	limit as specified in the standard is used to achieve convergence.			

Function outputs

Function outputs	Remark	EU	Fallback
Status	0: Normal		
	1: Input argument out of range		
	2: Calculation error		
	3: No convergence		
Density		lb/ft3	0
Compressibility		-	0
Equilibrium pressure	Equilibrium pressure at the observed temperature.	psia	0
	This property is not currently calculated, as it is not defined in the standard how to do this.		
Range	With respect to the standard the inputs are:	-	0
	0: In Range		
	1: Out of Range		

Calculations

The calculations are in compliance with the standard.

fxAPI_MPMS_11_3_3_2

Description

The API MPMS 11.3.3.2 standard consists of a table with the density values (lbm/ft3) of propylene liquid as a function of pressure and temperature. Also part of the standard is the Calculation Procedure to obtain the table values.

Compliance

• API MPMS 11.3.3.2 Propylene Compressibility Tables, 1974, Reaffirmed 1997.

Function inputs

Boundaries

The Calculation Procedure of API MPMS 11.3.3.2 defines the following limits on the input values:

- 30 to 165 °F
- 0 to 1600 psig

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Observed temperature		°F	0200	60
Observed pressure		psia	02000	0
API rounding	0: Disabled	-		1
	The calculations are performed with full precision. A convergence limit of 1e-10 lbm/ft3 will be applied for the iterative calculations. 1: Enabled The calculated density is rounded to 5 decimal places (same as table			
	values). A convergence limit of 5e-6 lbm/ft3 will be applied as defined in the standard.			

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Density	At the observed pressure and temperature	lbm/ft3	DENS		0
CTPL	Volume correction factor for temperature and pressure (also	-	CTPL		1
	referred to as the compressibility factor), equals the density at				
	the observed conditions of pressure and temperature value				
	divided by 32.6058 lbm/scf.				
	The value of 32.6058 lbm/scf is specified in the Calculation				
	Procedure of the standard as the propylene standard density at				
	60 °F and the corresponding vapor pressure.				
Equilibrium pressure	Equilibrium pressure at the observed temperature.	psia	EQUIPRES		0
	Also referred to as vapor pressure or saturated pressure				
Calculation out of range	With respect to the standard the input values are:	-			0
	0: In Range				
	1: Out of Range			OOR	

Calculations

The calculations are in full or partial compliance with the standard depending on input 'API rounding'.

fxAPI_RD60F_1980

Description

Relative Density (T, P) <--> Relative Density (60°F, equilibrium pressure)

This function converts a relative density value at the observed temperature and pressure to the relative density at 60°F and the equilibrium pressure (typically 0 psig) or vice versa.

The temperature conversion is according to API-2540, Tables 23A/24A (Generalized Crude Oils) and 23B/24B (Refined Oil Products), while the volume correction for pressure according to API MPMS 11.2.1.

In 1982 API published tables 5D, 6D, 53D and 54D for lubricating oil products as part of API MPMS 11.1. Although tables 23D and 24D are not covered in an official API standard the Flow-X series of flow computer supports tables 23D and 24D as well by combining the calculation of tables 23A/B and 24A/B with the KO and K1 constants published in the other tables for lubricating oils.

Note: this function is a combination of API2540 and API 11.2.1. For the calculation from observed to standard conditions an iterative calculation is required. The rounding and truncating of input and intermediate values is implemented such that the example calculations as specified in both standards are exactly reproduced.

Compliance

- API MPMS 11.1 Volume X (API Standard 2540) Table 23A -Generalized Crude Oils, Correction of Observed Relative Density to Relative Density at 60/60°F - First Edition, August 1980
- API MPMS 11.1 Volume X (API Standard 2540) Table 23B -Generalized Products, Correction of Observed Relative Density to Relative Density at 60/60°F - First Edition, August 1980
- API MPMS 11.1 Volume X (API Standard 2540) Table 24A -Generalized Crude Oils, Correction of Volume to 60°F against Relative Density at 60/60°F - First Edition, August 1980
- API MPMS 11.1 Volume X (API Standard 2540) Table 24B -Generalized Products Correction of Volume to 60°F against Relative Density at 60/60°F - First Edition, August 1980
- API MPMS 11.2.1 Compressibility Factors for Hydrocarbons: 0 - 90°API Gravity Range - First Edition, August 1984

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Input relative density	Depending on the conversion method this is the relative density either at	-		01.3	0
	the observed temperature and observed pressure or at 60 °F and the				
	equilibrium pressure				
Observed temperature		°F		-100400	60
Observed pressure		psig		-102000	0
Product	1: A - Crude Oil		PRDTYP	-	1
	2: B - Auto select				
	Selection based on relative density at 60 °F				
	3: B - Gasoline				
	4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API-2540 rounding	0: Disabled		API2540RND	-	0
	The calculations are performed with full precision and the final CTL value				
	is rounded as specified by input 'CTL decimal places'				
	1: Enabled for computational value				
	API-2540 rounding and truncating rules are applied and, in case of				
	conversion method 2 (standard to observed), the computational value for				
	CTL as specified in Table 24 is used, meaning that the CTL value has:				
	4 decimal places if CTL >=1				
	5 decimal places if CTL < 1.				
	2: Enabled for table value				
	API-2540 rounding and truncating rules are applied and, in case of				
	conversion method 2 (standard to observed), the table value for CTL as				
	specified in Table 24 meaning that the CTL value has 4 decimal places in				
	all cases				
	3: Enabled with 5 decimal places				
	API-2540 rounding and truncating rules are applied, and, in case of				
	conversion method 2 (standard to observed), the CTL value has 5 decimal				
	places in all cases.				
	Note: although not strictly in accordance with the standard, this option is				
	more commonly used than option 'Enabled for computational value'				
	Note: for conversion type 1 'From observed to standard conditions' the CTL				

Function inputs	Remark	EU	SW tag	Range	Default
	factor is rounded to 6 decimal places when input 'API 2540 rounding' > 0, as				
	in accordance with table 23.				
Hydrometer correction	Only applies for conversion method	-	HYDROCOR		0
	'1: From observed to standard conditions'				
	0: Disabled				
	1: Enabled				
API 11.2.1 rounding	0: Disabled	-	API1121RND		0
	The calculation of the compressibility factor F is performed with full precision				
	1: Enabled				
	API-MPMS 11.2.1 rounding and truncating rules are applied. The				
	compressibility factor F is rounded to 3 decimal places as specified in the				
	standard.				
Equilibrium pressure	The equilibrium pressure is considered to be 0 psig for liquids which have an	psig	EQUIPRES	02000	0
	equilibrium pressure less than atmospheric pressure (in compliance with				
	API MPMS 12.2 par. 12.2.5.4)				
Conversion method	1: From observed to standard conditions		CONVERSION		1
	2: From standard to observed conditions				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	1
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Output relative	Depending on the conversion method this is the relative density either at 60 $^\circ$ F	-	RD		0
density	and the equilibrium pressure or at the observed temperature and observed				
	pressure				
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API2540 rounding'				
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API 11.2.1 rounding'				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
КО	Actual value of constant K0 used for CTL calculation	-	К0		0
К1	Actual value of constant K1 used for CTL calculation	-	K1		0
K2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor	1/°F	ALPHA		0
F	Compressibility factor	1/psi	F		0
Product	When input 'Product' is 'B - Auto select', then the output is set to the actual	-	PRDCUR		0
	selected product of table 23B/24B (enumerative value as defined for input				
	'Product'), else the output is set equal to input 'Product'.				
CTL calc out of range	With respect to the standard used for the calculation of CTL the combination of				0
	input values is:				
	0: In Range				
	1: Out of Range			CTLOOR	
CPL calc out of range	With respect to the standard used for the calculation of CPL the combination of				0
	input values is:				
	0: In Range				
	1: Out of Range			CPLOOR	

Calculations

The calculations depend on the conversion method.

Conversion method 1: from observed to standard conditions. The function performs the following iterative algorithm to calculate the relative density at standard conditions:

- 1 First the inputs are rounded in accordance with the API2540 standard, provided that API2540 rounding is enabled.
- 2 The observed density [kg/m3] is calculated from the observed relative density
- 3 The hydrometer correction on the observed density is applied, provided that this correction is enabled
- 4 At the start of the iteration the initial value for density and relative density at [60 °F, equilibrium pressure] is set to respectively the observed density and the observed relative density. The initial CPL value is set to 1.
- 5 When the type of product is set to 'B Auto select' (automatic selection of the refined product range) the K0, K1 and K2

factors are determined based on the relative density at [60 $^{\circ}$ F, equilibrium pressure]. The Transition area is only taken in consideration in the 2nd iteration loop, as specified in the standard.

- 6 The Alpha factor is calculated according from the density at [60 °C, equilibrium pressure] and the K0, K1 and K2 factor. If API2540 rounding is enabled, then the intermediate results are rounded or truncated as specified API-2540 Table 23.
- 7 The CTL value is calculated according to API-2540 Table 23 from the Alpha factor and the differential temperature (= observed temperature – 60°F). If API2540 rounding is enabled, then the intermediate results are rounded or truncated as specified API-2540 Table 23.
- 8 Depending on the type of API2540 rounding the calculated CTL value is rounded to 6 decimal places or not rounded at all.
- 9 The density at [60 °F, equilibrium pressure] is calculated by dividing the observed density by the new CTL value and the CPL value from the previous iteration.

- 10 The relative density at [60 °F, equilibrium pressure] is calculated from the density at [60 °F, equilibrium pressure]
- 11 Because API 11.2.1 requires the API gravity value at 60 °F, the API gravity at [60 °F, equilibrium pressure] is calculated from the density at [60 °F, equilibrium pressure].
- 12 The compressibility factor is calculated according to API MPMS 11.2.1 from the API gravity at [60 °F, equilibrium pressure] and the 'Observed temperature'. If API 11.2.1 rounding is enabled then the API gravity and temperature are rounded and the calculations are performed in accordance with the rounding and truncating rules of the standard.
- 13 The CPL value is calculated from the compressibility factor and the 'Observed pressure' and 'Equilibrium pressure' input values.
- 14 The density at [60°F, equilibrium pressure] is calculated by dividing the observed density by CTL and the new CPL value.
- 15 If API2540 rounding is enabled then the density at [60°F, equilibrium pressure] value is rounded to 3 decimal places as specified in the standard.
- 16 The relative density at [60 °F, equilibrium pressure] is calculated from the density at [60 °F, equilibrium pressure]
- 17 If API2540 rounding is enabled then the relative density at [60°F, equilibrium pressure] value is rounded to 4 decimal places as specified in the standard.
- 18 Steps 5 through 17 are repeated taking the density value from step 14 as the start value for the next iteration until the absolute difference between two consecutive density values is either 0.05 (or 0.07 for the transition area) or 0.000001, depending of API2540 rounding being enabled or not.
- 19 For refined products the entire iteration loop is repeated if the relative density at [60°F, equilibrium pressure] appears to be in a different product region than the observed relative density. This is required because a different product region means different K0, K1 and K2 factors.
- 20 If API 11.2.4 rounding is enabled, then the relative density value at [60°F, equilibrium pressure] is rounded to 0.0001

Conversion method 2: from standard to observed conditions. The function performs straightforward calculations to determine the relative density at observed conditions:

- 1 First the inputs are rounded in accordance with the API2540 standard, provided that API2540 rounding is enabled.
- 2 The density at [60°F, equilibrium pressure] is calculated from the input relative density
- 3 When the type of product is set to 'B Auto select' (automatic selection of the refined product range) the K0, K1 and K2 factors are determined based on the input relative density
- 4 The Alpha factor is calculated according from the density at [60°F, equilibrium pressure] and the K0, K1 and K2 factor. If API2540 rounding is enabled, then the intermediate results are rounded or truncated as specified API-2540 Table 24.
- 5 The CTL value is calculated according to API-2540 Table 24 from the Alpha factor and the differential temperature (= observed temperature – 60°F). If API2540 rounding is enabled, then the intermediate results are rounded or truncated as specified API-2540 Table 24.
- 6 Depending on the type of API2540 rounding the calculated CTL value is rounded to 4 or 5 decimal places or not rounded at all.
- 7 Because API 11.2.1 requires the API gravity value at 60 °F, the API gravity at [60 °F, equilibrium pressure] is calculated from the density at [60 °F, equilibrium pressure].
- 8 The compressibility factor is calculated according to API MPMS 11.2.1 from the input density and temperature'. If API 11.2.1 rounding is enabled then the input density and temperature are rounded and the calculations are performed in accordance with the rounding and truncating rules of the standard.
- 9 The CPL value is calculated from the compressibility factor and the 'Observed pressure' and 'Equilibrium pressure' input values.
- 10 The relative density at observed temperature and pressure is calculated from the input relative density and the CTL and the CPL values.

fxAPI_SG60F_1952

Description

Specific Gravity (T, P) <--> Specific Gravity (60°F, equilibrium pressure)

This function converts a specific gravity value at the observed temperature and pressure to the specific gravity at 60°F and the equilibrium pressure (typically 0 psig) or vice versa. The temperature conversion is according to ASTM-IP Petroleum Measurements Tables 1952 (Also known as API-1952 tables) Table 24. Note: this function is a combination of the API 1952 Tables and API 11.2.1. For the calculation from observed to standard conditions an iterative calculation is required. The rounding and truncating of input and intermediate values is implemented such that the example calculations as specified in both standards are exactly reproduced.

Compliance

- ASTM-IP Petroleum Measurement Tables, American Edition, United States Units of Measurement, 1952
- API MPMS 11.2.1 Compressibility Factors for Hydrocarbons: 0 - 90°API Gravity Range - First Edition, August 1984

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Input Specific Gravity	Depending on the conversion method this is the Specific Gravity either at the observed temperature and observed pressure or at 60 $^\circ{\rm F}$ and the	-		01.3	0
Observed temperature	equilibrium pressure	°F		-100400	60
Observed pressure		 psig		-102000	0
API 11.2.1 rounding	 0: Disabled The calculation of the compressibility factor F is performed with full precision 1: Enabled API-MPMS 11.2.1 rounding and truncating rules are applied. The compressibility factor F is rounded to 3 decimal places as specified in the standard. 	-	API1121RND		0
Equilibrium pressure	The equilibrium pressure is considered to be 0 psig for liquids which have an equilibrium pressure less than atmospheric pressure (in compliance with API MPMS 12.2 par. 12.2.5.4).	psig	EQUIPRES	02000	0
Conversion method	1: From observed to standard conditions 2: From standard to observed conditions		CONVERSION		1

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	1
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Output Specific Gravity	Depending on the conversion method this is the Specific Gravity either at	-	RD		0
	60 °F and the equilibrium pressure or at the observed temperature and				
	observed pressure				
CTL	Volume correction factor for temperature.	-	CTL		1
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API 11.2.1 rounding'				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
F	Compressibility factor	1/psi	F		0
CTL calc out of range	With respect to the standard used for the calculation of CTL the				0
	combination of input values is:				
	0: In Range				
	1: Out of Range			CTLOOR	
CPL calc out of range	With respect to the standard used for the calculation of CPL the				0
	combination of input values is:				
	0: In Range				
	1: Out of Range			CPLOOR	

Calculations

The calculations depend on the conversion method.

Conversion method 1: from observed to standard conditions. The function performs the following iterative algorithm to calculate the Specific Gravity at standard conditions:

- 1 At the start of the iteration the initial value for Specific Gravity at [60 °F, equilibrium pressure] is set to the Observed Specific Gravity. The initial CPL value is set to 1.
- 2 The CTL value is determined from the Specific Gravity at [60 °F, equilibrium pressure] according to API 1952 Table 24.
- 3 The Specific Gravity at [60 °F, equilibrium pressure] is calculated from the Observed specific gravity, the new CTL value and the CPL value from the previous iteration.
- 4 Because API 11.2.1 requires the API gravity value at 60 °F, the API gravity at [60 °F, equilibrium pressure] is calculated from the Specific gravity at [60 °F, equilibrium pressure].
- 5 The compressibility factor is calculated according to API MPMS 11.2.1 from the API gravity at [60 °F, equilibrium pressure] and the 'Observed temperature'. If API 11.2.1 rounding is enabled then the API gravity and temperature are rounded and the calculations are performed in accordance with the rounding and truncating rules of the standard.
- 6 The CPL value is calculated from the compressibility factor and the 'Observed pressure' and 'Equilibrium pressure' input values.
- 7 The Specific Gravity at [60°F, equilibrium pressure] is calculated by dividing the Observed Specific Gravity by the CTL and the new CPL value.
- 8 Steps 2 through 7 are repeated taking the specific gravity value from step 7 as the start value for the next iteration until the absolute difference between two consecutive specific gravity values is 0.0001.

Conversion method 2: from standard to observed conditions. The function performs straightforward calculations to determine the Specific Gravity at observed conditions:

- 1 The CTL value is calculated according to API 1952 Table 24
- 2 Because API 11.2.1 requires the API gravity value at 60 °F, the API gravity is calculated from the 'Input Specific Gravity'.
- 3 The compressibility factor is calculated according to API MPMS 11.2.1 from the API gravity and the 'Observed temperature'. If API 11.2.1 rounding is enabled then the input density and temperature are rounded and the calculations are performed in accordance with the rounding and truncating rules of the standard.
- 4 The CPL value is calculated from the compressibility factor and the 'Observed pressure' and 'Equilibrium pressure' input values.
- 5 The output Specific Gravity (at observed temperature and pressure) is calculated from the input Specific Gravity and the CTL and the CPL values.

fxAPI_RD60F_NGL_LPG

Description

Relative Density (T, P) <--> Relative Density (60°F, Pe)

This function converts the relative density value at the observed temperature and pressure to the relative density value at 60°F and the equilibrium pressure or vice versa.

The temperature correction is according to API MPMS 11.2.4:2007 (GPA TP-25 / GPA TP-27), while the pressure correction is according to API MPMS 11.2.2:1984.

The calculation of the equilibrium pressure is according to GPA TP-15 (API MPMS 11.2.2 Addendum:1994).

Compliance

- API MPMS 11.2.4: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007
- GPA TP-27: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007
- GPA TP-25: Temperature Correction for the volume of Light Hydrocarbons – Tables 24E and 23E, 1998
- API MPMS 11.2.2 Compressibility Factors for Hydrocarbons: 0.350 - 0.637 Relative Density (60°F/60°F) and -50°F to 140°F Metering Temperature - Second Edition, October 1986
- API MPMS 11.2.5: A Simplified Vapor Pressure Correlation for Commercial NGLs, September 2007
- GPA TP-15: A Simplified Vapor Pressure Correlation for Commercial NGLs, September 2007 (also covers GPA TP-15 1988)
- API MPMS 11.2.2 Addendum : Compressibility Factors for Hydrocarbons: Correlation of Vapor Pressure for Commercial Natural Gas Liquids (same as GPA TP-15:1988)

Function inputs

Name	Remark	EU	SW tag	Range	Defaul
Name	Optional tag name, tag description and tag group				
Input relative density	Depending on the conversion method this represents the relative density either at the observed temperature and pressure or at 60 °F and the equilibrium pressure	-		00.75	0
Observed temperature	Temperature at which the relative density is observed	°F		-100300	60
Observed pressure	Pressure at which the relative density is observed	psia		-102500	0
API 11.2.4 rounding	0: Disabled The calculations are performed with full precision and the output values are not rounded 1: Enabled The input and output values are rounded as defined in the standard		API1124RND		0
API 11.2.2 rounding	0: Disabled The calculations are performed with full precision and the output values are not rounded 1: Enabled The input and output values are rounded as defined in the standard		API1122RND		0
Equilibrium pressure	1: Use Input		EQUIPMODE		2
mode	The value of input 'Equilibrium pressure value' is used for the calculation of CPL 2: GPA TP-15 The equilibrium pressure is calculated in accordance with GPA TP-15 Other equilibrium to the with the second		FOUNDING		
Equilibrium pressure value	Only used when input 'Equilibrium pressure mode' is set to 'Use input'. The value will be used for the calculation of the CPL	psia	EQUIPINP		0
GPA TP-15 rounding	0: Disabled Full precision (no rounding and truncating applied) 1: Enabled Rounding as defined in ' GPA TP15:1988 / API MPMS 11.2.2 Addendum':1994	-	TP15RND		0
P100 Correlation	 0: Disabled The standard correlation is commonly used for pure products such as propane, butane and natural gasoline. It only requires the relative density and the temperature to calculate the vapor pressure 1: Enabled The improved correlation requires the vapor pressure at 100°F (37.8 °C). This method is better suited for varied NGL mixes Where different product mixes could have the same specific gravity but different equilibrium pressures. 	-	P100CORR		0
Vapor pressure at 100°F		psia	EQUIP100F	02500	0
Conversion method	1: From observed to standard conditions 2: From standard to observed conditions		CONVMETH		1

Function outputs

Name	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		

Name	Remark	EU	SW tag	Alarm	Fallback
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Output relative	Depending on the conversion method this represents the relative density	-	DENS		Input relative
density	either at 60 °F and the equilibrium pressure or at the observed temperature				density
	and pressure				
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API 11.2.4 rounding'				
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API 11.2.2. rounding'				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
F	Compressibility factor	1/psi	F		0
	The output value will be either rounded or not depending input 'API				
	rounding'				
Equilibrium	The equilibrium pressure calculated by GPA TP-15	psia	EQUIPCUR		0
pressure	Will be set to 0 when equilibrium pressure is below atmospheric pressure	•			
CTL calc out of	With respect to the API 11.2.4 standard the combination of input values is:				0
range	0: In Range				
5	1: Out of Range				
	The following range checks apply:			CTLOOR	
	0.21 <= RD <= 0.74				
	-50.8 <= T <= 199.4 °F				
	Table 23E reference fluid ranges				
CPL calc out of	With respect to API 11.2.2M the combination of input values is:				0
range	0: In Range				
5	1: Out of Range				
	The following range checks apply:			CPLOOR	
	350 <= Density 15 °C <= 637 kg/m3				
	-46 °C <= T <= 60 °C				
	Also the check on the pseudo-critical temperature as defined for				
	fxAPI_MPMS_11_2_2 is applied.				
GPA TP-15 out of	Only set when the GPA TP-15 calculation is enabled	-			0
range	With respect to the GPA TP-15 standard the combination of input values is:				
5	0: In Range				
	1: Out of Range				
	The following range checks apply:				
	For lower range:			TP15OOR	
	0.350 <= RD60 < 0.425				
	-50 to (695.51*RD60 - 155.51) °F				
	Higher range:				
	0.425 <= RD60 <= 0.676				
	-50 to 140 °F				
	with RD60 being the relative density at 60°F				
Calculations			ty factor E is c		

Calculations

The calculations depend on the conversion method.

Conversion method 1: from observed to standard conditions. The function performs the following iterative algorithm to calculate the relative density at 60 °F and the equilibrium pressure.

- 1 When API 11.2.4 rounding is enabled, the input relative density and temperature values are rounded in accordance with the standard
- 2 At the start of the iteration the relative density at [60 °F, equilibrium pressure] is set equal to the observed relative density and the CPL value is set to 1.
- 3 First the relative density corrected for pressure is calculated by dividing the observed relative density by the CPL value.
- 4 The CTL value and the relative density at [60 °F, equilibrium pressure] is calculated from the relative density corrected for pressure and the observed temperature according to Table 23E
- 5 Depending on the value of input 'Equilibrium pressure mode', either value of input 'Equilibrium pressure value' is used or the equilibrium pressure (vapor pressure) is calculated according to GPA TP-15. Whether the GPA TP-15 rounding and truncation rules are applied is dictated by input 'GPA-TP15 rounding'

- 6 The compressibility factor F is calculated according to API MPMS 11.2.2 from the relative density at [60 °F, equilibrium pressure] and the 'Observed temperature', with, depending on input API 11.2.2, rounding and truncation according to the standard.
- 7 The CPL value is calculated from the compressibility factor, the equilibrium pressure and the 'Observed pressure' input value.
- 8 The new value for relative density at [60°F, equilibrium pressure] is calculated by dividing the observed density by the CTL and CPL values.
- 9 Steps 3 though 8 are repeated taking the density value from step 8 as the staring value until the absolute difference between two consecutive density values is less than the convergence limit.

To avoid convergence problems different convergence limits are applied, depending on the whether API 11.2.2 and/or GPA TP-15 rounding is applied:

If API 11.2.2M rounding is enabled -> Limit = 0.00005 kg/m3 else if GPA TP-15 rounding is enabled -> Limit = 0.000005 kg/m3

else -> Limit = 0.00000001 kg/m3

10 If API 11.2.4 rounding is enabled, then the relative density at [60°F, equilibrium pressure] is rounded to 0.0001

Conversion method 2: from standard to observed conditions. The function performs straightforward calculations to determine the density at observed conditions:

- 1 When API 11.2.4 rounding is enabled, the input relative density and temperature values are rounded in accordance with the standard
- 2 The CTL value is calculated according to API MPMS 11.2.4 (GPA TP-27) Table 24E from the density at [15 °C, equilibrium pressure] and the 'Observed temperature'.
- 3 Depending on the value of input 'Equilibrium pressure mode', either value of input 'Equilibrium pressure value' is used or the equilibrium pressure (vapor pressure) is calculated according to GPA TP-15.
- 4 The compressibility factor is calculated according to API MPMS 11.2.2 from the input relative density and the 'Observed temperature'.
- 5 The CPL value is calculated from the compressibility factor, the equilibrium pressure and the 'Observed pressure' input value.
- 6 If API 11.2.4 rounding is enabled, then the CTL value is rounded at [60°F, equilibrium pressure] is rounded to 0.00001
- 7 The relative density at the observed conditions is calculated by multiplying the input relative density by the CTL value and the CPL value.

fxAPI_Table5_1952

Description

°API (T) --> °API (60°F)

This function converts an API gravity value at the observed temperature to the API gravity value at 60°F in accordance with API 1952 Table 5.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Observed API	Observed API gravity	°API		-20120	0
Observed temperature	Temperature at which the API gravity is observed	°F		-100400	60

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
API at 60 °F	API gravity at 60°F	°API	API		0
Calculation out of range	With respect to the standard the input values are: 0: In Range 1: Out of Range	-		OOR	0

Calculations

The table values are the standard, so no calculations are

involved. The function performs an interpolation between the

table values that correspond to the input values.

Compliance

• ASTM-IP Petroleum Measurement Tables, American Edition, United States Units of Measurement, 1952

fxAPI_Table5_1980

Description

°API (T) --> °API (60°F)

This function converts an API gravity value at the observed temperature to the API gravity value at 60°F.

The temperature conversion is according to API MPMS 11.1:1980 (API-2540), Tables 5A (Generalized Crude Oils) and 5B (Refined Oil Products) and API MPMS 11.1 Chapter XIII Table 5D: 1984 (Lubricating Oils).

The function provides the option to correct for readings taken from a hydrometer as specified in the API-2540 standard.

Compliance

- API MPMS 11.1 Volume X (API Standard 2540) Table 5A -Generalized Crude Oils, Correction of Observed API Gravity to API Gravity at 60°F - First Edition, August 1980
- API MPMS 11.1 Volume X (API Standard 2540) Table 5B -Generalized Products, Correction of Observed API Gravity to API Gravity at 60°F- First Edition, August 1980
- API MPMS 11.1 Volume XIII Table 5D Generalized Lubricating Oils, Correction of Observed API Gravity to API Gravity at 60°F - January 1982

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Observed API	Observed API gravity	°API		-20120	0
Observed temperature	Temperature at which the API gravity is observed	°F		-100400	60
Product	1: A - Crude Oil		PRDTYP		1
	2: B - Auto select Selection based on °API at 60 °F 3: B - Gasoline 4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API2540 rounding	 0: Disabled The calculations are performed with full precision. A convergence limit of 0.000001 kg/m3 will be applied for the iterative calculations. 1: Enabled API-2540 rounding and truncating rules are applied. A convergence limit of 0.05 kg/m3 will be applied as defined in the standard. 		APIROUND		0
Hydrometer correction	0: Disabled 1: Enabled		HYDROCOR		0

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
API at 60 °F	API gravity at 60°F	°API	API		0
CTL	Volume correction factor for temperature.	-	CTL		1
ко	Actual value of constant K0 used for CTL calculation	-	К0		0
К1	Actual value of constant K1 used for CTL calculation	-	K1		0
К2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor	1/°F	ALPHA		0
Product	When input 'Product' is 'B - Auto select', then the output is set	-	PRDCUR		0
	to the actual selected product of table 5B (enumerative value				
	as defined for input 'Product'), else the output is set equal to				
	input 'Product'.				
Calculation out of range	With respect to the standard the input values are:	-			0
_	0: In Range				
	1: Out of Range			OOR	

Calculations

The calculations are in full or partial compliance with the standard depending on input 'API 2540 rounding'.

fxAPI_Table5_2004

Description

°API (T, P) --> °API (60°F, 0 psig)

This function converts an API gravity value at the observed temperature and pressure to the API gravity value at 60°F and 0 psig.

The temperature and pressure correction is according to API MPMS 11.1:2004.

An iterative calculation needs to be applied to convert the observed API gravity to the value at base conditions.

Function inputs

Note: As opposed to API-2540, the 2004 standard does not include a correction for readings taken from a hydrometer and assumes that the equilibrium pressure is below atmospheric pressure, so taking 0 psig as the base pressure.

Compliance

 API MPMS 11.1 Temperature and Pressure Volume Correction Factors for Generalized Crude Oils, Refined Products and Lubricating Oils, May 2004

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Observed API	Observed API gravity	°API		-20180	0
Observed temperature	Temperature at which the API gravity is observed	°F		-100400	60
Observed pressure	Pressure at which the API gravity is observed	psig		-102000	0
Product	1: A - Crude Oil		PRDTYP		1
	2: B - Auto select				
	Selection based on °API at 60 °F				
	3: B - Gasoline				
	4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API rounding	0: Disabled		APIROUND		0
	The calculations are performed with full precision and the final				
	CTL, CPL and CTPL values are rounded as specified by the				
	inputs 'CTL / CPL/ CTPL decimal places'				
	1: Enabled				
	The input and output values are rounded in compliance with				
	the standard. The CTL, CPL and CTPL value are rounded to 5				
	decimal places				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence within 15 iterations			NOCONV	
API at 60 °F	API gravity at 60°F and 0 psig	-	API		0
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API rounding'				
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API rounding'				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
КО	Actual value of constant K0 used for CTL calculation	-	KO		0
K1	Actual value of constant K1 used for CTL calculation	-	K1		0
К2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor	1/°F	ALPHA		0
F	Compressibility factor	1/psi	F		0
Product	When input 'Product' is 'B - Auto select', then the output is set to the	-	PRDCUR		0
	actual selected product of table B (enumerative value as defined for				
	input 'Product'), else the output is set equal to input 'Product'.				
Calculation out of	With respect to the standard the input values are:	-			0
range	0: In Range				
	1: Out of Range			OOR	

Calculations

The calculations are in either full or partial compliance with the standard, depending on whether API rounding is enabled or not.

fxAPI_Table6_1952

Description

52

°API (60°F, 0 psig) --> CTL

This function calculates the volume correction factor for temperature from the API gravity value at 60°F and the observed temperature according to API 1952 Table 6.

Compliance

ASTM-IP Petroleum Measurement Tables, American Edition, • United States Units of Measurement, 1952

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
API at 60 °F	API gravity at 60°F and the equilibrium pressure	°API		-20120	0
Observed temperature		°F		-100400	60

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALC	
CTL	Volume correction factor for temperature.	-	CTL		1
Calculation out of range	With respect to the standard the input values are:	-			0
-	0: In Range				
	1: Out of Range			OOR	

Calculations

The table values are the standard, so no calculations are involved. The function performs an interpolation between the table values that correspond to the input values.

Boundaries

Table 6 contains values for the following range:

Input value	Normal Range	EU
API Gravity at 60 °F	0100	°API
Observed temperature	0300	°F

Note that the table does not cover the full range, e.g. for an API gravity of 70 the table only specifies values between 0 .. 150 °F.

fxAPI_Table6_1980

Description

°API (60°F, 0 psig) --> CTL

This function calculates the volume correction factor for temperature from the API gravity value at 60°F and the observed temperature.

The temperature conversion is according to API-2540, Tables 6A (Generalized Crude Oils) and 6B (Refined Oil Products) and API MPMS 11.1 Chapter XIII Table 6D: 1984 (Lubricating Oils).

Compliance

- API MPMS 11.1 Volume X (API Standard 2540) Table 6A -Generalized Crude Oils, Correction of Volume to 60°F against API Gravity at 60°F - First Edition, August 1980
- API MPMS 11.1 Volume X (API Standard 2540) Table 6B -Generalized Products, Correction of Volume to 60°F against API Gravity at 60°F - First Edition, August 1980
- API MPMS 11.1 Volume XIII Table 6D Generalized Lubricating Oils, Correction of Volume to 60°F Against API Gravity at 60°F - January 1982

Function inputs & outputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
API at 60 °F	API gravity at 60°F and the equilibrium pressure	°API		-20120	0
Observed temperature		°F		-100400	60
Product	1: A - Crude Oil		PRDTYP	-	1
	2: B - Auto select				
	Selection based on °API at 60 °F				
	3: B - Gasoline				
	4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API2540 rounding	0: Disabled		APIROUND	-	0
	The calculations are performed with full precision and the final CTL value is				
	rounded as specified by input 'CTL decimal places'				
	1: Enabled for computational value				
	API-2540 rounding and truncating rules are applied and the computational				
	value for CTL as specified in the standard is used, meaning that the CTL				
	value has:				
	4 decimal places if CTL >=1				
	5 decimal places if CTL < 1.				
	2: Enabled for table value				
	API-2540 rounding and truncating rules are applied and the table value for				
	CTL as specified in the standard meaning that the CTL value has 4 decimal				
	places in all cases				
	3: Enabled with 5 decimal places				
	API-2540 rounding and truncating rules are applied, while the CTL value has				
	5 decimal places in all cases.				
	Note: although not strictly in accordance with the standard, this option is				
	more commonly used than option 'Enabled for computational value'				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALC	
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API2540 rounding'				
КО	Actual value of constant K0 used for CTL calculation	-	К0		0
К1	Actual value of constant K1 used for CTL calculation	-	K1		0
К2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor	1/°F	ALPHA		0
Product	When input 'Product' is 'B - Auto select', then the output is set to the actual selected	-	PRDCUR		0
	product of table 6B (enumerative value as defined for input 'Product'), else the				
	output is set equal to input 'Product'.				
Calculation out of	With respect to the standard the input values are:	-			0
range	0: In Range				
	1: Out of Range			OOR	

Calculations

The calculations are in full or partial compliance with the standard depending on input 'API 2540 rounding'.

fxAPI_Table6_2004

Description

54

°API (60°F, 0 psig) --> °API (T, P)

This function converts an API gravity value at 60°F and 0 psig to the API gravity value at the observed temperature and pressure. The temperature and pressure correction is according to API MPMS 11.1:2004.

Function inputs

Note: As opposed to API-2540 that the equilibrium pressure is below atmospheric pressure, so taking 0 psig as the base pressure.

Compliance

• API MPMS 11.1 Temperature and Pressure Volume Correction Factors for Generalized Crude Oils, Refined Products and Lubricating Oils, May 2004

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag				
	group				
API at 60 °F	API gravity at 60°F and 0 psig	°API		-20120	0
Observed temperature	Temperature at which the API gravity is observed	°F		-100400	60
Observed pressure	Pressure at which the API gravity is observed	psig		-102000	0
Product	1: A - Crude Oil		PRDTYP		1
	2: B - Auto select				
	Selection based on °API at 60 °F				
	3: B - Gasoline				
	4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API rounding	0: Disabled The calculations are performed with full precision		APIROUND		0
	1: Enabled				
	The input and output values are rounded in compliance with the standard. The CTL, CPL and CTPL value are rounded to 5				
	decimal places.	•			

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALC	
Observed API	API gravity at the observed temperature and pressure	-	API		0
CTL	Volume correction factor for temperature.	-	CTL		1
CPL	Volume correction factor for pressure	-	CPL		1
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
ко	Actual value of constant K0 used for CTL calculation	-	К0		0
K1	Actual value of constant K1 used for CTL calculation	-	K1		0
K2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor	1/°F	ALPHA		0
F	Compressibility factor	1/psi	F		0
Product	When input 'Product' is 'B - Auto select', then the output is set to the	-	PRDCUR		0
	actual selected product of table B (enumerative value as defined for				
	input 'Product'), else the output is set equal to input 'Product'.				
Calculation out of range	With respect to the standard the input values are:	-			0
	0: In Range				
	1: Out of Range			OOR	

Calculations

The calculations are in either full or partial compliance with the standard, depending on whether API rounding is enabled or not.

fxAPI_Table23_1952

Description

Specific Gravity (T) --> Specific Gravity (60°F)

This function converts a specific gravity value at the observed temperature to the specific gravity at 60° according to the API 1952 Table 23.

Compliance

• ASTM-IP Petroleum Measurement Tables, American Edition, United States Units of Measurement, 1952

Boundaries

Table 23 contains values for the following range:

Input value	Normal Range	EU
Observed specific gravity	0.420 1.099	
Observed temperature	0 150	°F

Note that the table does not cover the full range, e.g. for an Observed specific gravity of 0.420 the table only specifies values between 120 .. 140 $^{\circ}{\rm F}$

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Observed specific gravity	Specific gravity at the observed temperature.	-		01.3	0
Observed temperature		°F		-100400	60

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
Specific gravity 60 °F	Specific gravity at 60°F	-	RD		0
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
	0: In Range				
	1: Out of Range				

Calculations

The table values are the standard, so no calculations are involved. The function performs an interpolation between the table values that correspond to the input specific gravity and input temperature.

In case the combination of input values ('Observed specific gravity' and Observed temperature') is not covered by the table, the output 'Specific gravity at 60 °F' is set to 0 and output 'Calculation out of range' is set to 1.

fxAPI_Table23_1980

Description

Relative Density (T) --> Relative Density (60°F)

This function converts a relative density value at the observed temperature to the relative density at 60°.

The temperature conversion is according to API-2540, Tables 23A (Generalized Crude Oils) and 23B (Refined Oil Products. In 1982 API published tables 5D, 6D, 53D and 54D for lubricating oil products as part of API MPMS 11.1. Although tables 23d and 24d are not covered in an official API standard the Flow-X series of flow computer supports tables 23D and 24D as well by combining the calculation of tables 23A/B and 24A/B with the KO and K1 constants published in the other tables for lubricating oils. An iterative calculation needs to be applied to convert the observed relative density to the value at base conditions. The function provides the option to correct for readings taken from a hydrometer as specified in the API-2540 standard.

Compliance

- API MPMS 11.1 Volume X (API Standard 2540) Table 23A -Generalized Crude Oils, Correction of Observed Relative Density to Relative Density at 60/60°F - First Edition, August 1980
- API MPMS 11.1 Volume X (API Standard 2540) Table 23B -Generalized Products, Correction of Observed Relative Density to Relative Density at 60/60°F - First Edition, August 1980

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Observed relative density	Relative density at observed temperature and pressure	-		01.3	0
Observed temperature		°F		-100400	60
Product	1: A - Crude Oil		PRDTYP		1
	2: B - Auto select				
	Selection based on relative density at 60 °F				
	3: B - Gasoline				
	4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API2540 rounding	0: Disabled		APIROUND		0
	The calculations are performed with full precision.				
	A convergence limit of 0.000001 kg/m3 will be applied for the iterative				
	calculations.				
	1: Enabled				
	API-2540 rounding and truncating rules are applied.				
	A convergence limit of 0.05 kg/m3 will be applied as defined in the				
	standard.				
Hydrometer correction	0: Disabled		HYDROCOR		0
	1: Enabled				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALC	
	3: No convergence			NOCONV	
Relative density 60 °F	Relative density at 60°F and the equilibrium pressure	-	RD		0
CTL	Volume correction factor for temperature.	-	CTL		1
ко	Actual value of constant K0 used for CTL calculation	-	К0		0
K1	Actual value of constant K1 used for CTL calculation	-	K1		0
K2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor	1/°F	ALPHA		0
Product	When input 'Product' is 'B - Auto select', then the output is set to the actual	-	PRDCUR		0
	selected product of table 23B (enumerative value as defined for input				
	'Product'), else the output is set equal to input 'Product'.				
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
	0: In Range				
	1: Out of Range				

Calculations

The calculations are in full or partial compliance with the standard depending on input 'API 2540 rounding'.

fxAPI_Table23_2004

Description

Relative Density (T, P) --> Relative Density (60°F, 0 psig)

This function converts a relative density value at the observed temperature and pressure to the relative density value at 60° F and 0 psig.

The temperature and pressure correction is according to API MPMS 11.1:2004.

An iterative calculation needs to be applied to convert the observed relative density to the value at base conditions.

Function inputs

Note: As opposed to API-2540, the 2004 standard does not include a correction for readings taken from a hydrometer and assumes that the equilibrium pressure is below atmospheric pressure, so taking 0 psig as the base pressure.

Compliance

• API MPMS 11.1 Temperature and Pressure Volume Correction Factors for Generalized Crude Oils, Refined Products and Lubricating Oils, May 2004

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Observed Relative density	Relative density at the observed temperature and pressure	-		01.3	0
Observed temperature	Temperature at which the relative density is observed	°F		-100400	60
Observed pressure	Pressure at which the relative density is observed	psig		-102000	0
Product	 A - Crude Oil B - Auto select Selection based on relative density at 60 °F B - Gasoline B - Transition Area B - Jet Fuels B - Jet Fuel T - Lubricating Oil 		PRDTYP		1
API rounding	 0: Disabled The calculations are performed with full precision and the final CTL, CPL and CTPL values are rounded as specified by the inputs 'CTL / CPL/ CTPL decimal places' 1: Enabled The input and output values are rounded in compliance with the standard. The CTL, CPL and CTPL value are rounded to 5 decimal places. 		APIROUND		0

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-			1
	1: Input argument out of range				
	Outputs will be set to fallback values				
	2: Calculation error				
	Outputs will be set to fallback values				
	3: No convergence within 15 iterations				
	Outputs will be set to values of last iteration				
Relative density	Relative density at 60°F and 0 psig	-	RD		0
at 60 °F					
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API rounding'				
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API rounding'				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
КО	Actual value of constant K0 used for CTL calculation	-	KO		0
K1	Actual value of constant K1 used for CTL calculation	-	K1		0
К2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor	1/°F	ALPHA		0
F	Compressibility factor	1/psi	F		0
Product	When input 'Product' is 'B - Auto select', then the output is set	-	PRDCUR		0
	to the actual selected product of table B (enumerative value as				
	defined for input 'Product'), else the output is set equal to				
	input 'Product'.				
Calculation out of range	With respect to the standard the input values are:	-		OOR	0

Function outputs	Remark	EU	SW tag	Alarm	Fallback
	0: In Range				
	1: Out of Range				

Calculations

The calculations are in either full or partial compliance with the standard, depending on whether API rounding is enabled or not.

fxAPI_Table23E

Description

Relative Density (T) --> Relative Density (60°F)

This function converts the relative density value at the observed temperature to the corresponding relative density at 60°F. The temperature correction is according to API MPMS 11.2.4:2007 (GPA TP-25 / GPA TP-27).

Compliance

- API MPMS 11.2.4: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007
- GPA TP-27: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007
- GPA TP-25: Temperature Correction for the volume of Light Hydrocarbons Tables 24E and 23E, 1998

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag				
	group				
Observed relative density	Relative density at the observed temperature	-		00.75	0
Observed temperature	Temperature at which the relative density is observed	°F		-100300	60
API rounding	0: Disabled The calculations are performed with full precision and the output values are not rounded 1: Enabled The input and output values are rounded as defined in the standard		APIROUND		0

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Relative density at	Relative density at 60°F		RD		0
60 °F					
CTL	Volume correction factor for temperature.	-	CTL		1
Calculation out of	With respect to the standard the input values are:	-		OOR	0
range	0: In Range				
-	1: Out of Range				

Calculations

The calculations are in either full or partial compliance with the standard, depending on the selected type of API rounding.

fxAPI_Table24_1952

Description

Specific Gravity (60°F) --> CTL

This function returns the volume correction factor for temperature Ctl from the observed temperature and the specific gravity at 60° according to the API 1952 Table 24.

Compliance

• ASTM-IP Petroleum Measurement Tables, American Edition, United States Units of Measurement, 1952

Boundaries

Table 23 contains values for the following range:

Input value	Normal Range	EU
Observed specific gravity	0.500 1.100	
Observed temperature	-50 +300	°F

Note that the table does not cover the full range, e.g. for an Specific gravity at 60 °F of 0.500 the table only specifies values between -50 .. 95 °F

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Specific gravity 60 °F	Specific gravity at 60°F	-		01.3	0
Observed temperature		°F		-100400	60

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
CTL	Volume correction factor for temperature.	-	CTL		1
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
5	0: In Range				
	1: Out of Range				

Calculations

The table values are the standard, so no calculations are involved. The function performs an interpolation between the table values that correspond to the input specific gravity and input temperature.

In case the combination of input values ('Specific gravity 60 $^{\circ}$ F' and Observed temperature') is not covered by the table, the output 'CTL' is set to 1 and output 'Calculation out of range' is set to 1.

fxAPI_Table24_1980

Description

Relative Density (60°F) --> CTL

This function calculates the volume correction factor for temperature CTL from the relative density value at 60°F and the observed temperature.

The temperature conversion is according to API-2540, Tables 24A (Generalized Crude Oils) and 24B (Refined Oil Products. In 1982 API published tables 5D, 6D, 53D and 54D for lubricating oil products as part of API MPMS 11.1. Although tables 23d and 24d are not covered in an official API standard the Flow-X series of flow computer supports tables 23D and 24D as well by combining the calculation of tables 23A/B and 24A/B with the KO and K1 constants published in the other tables for lubricating oils.

Function inputs

Compliance

- API MPMS 11.1 Volume X (API Standard 2540) Table 24A -Generalized Crude Oils, Correction of Volume to 60°F against Relative Density at 60/60°F - First Edition, August 1980
- API MPMS 11.1 Volume X (API Standard 2540) Table 24B -Generalized Products, Correction of Volume to 60°F against Relative Density at 60/60°F - First Edition, August 1980

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Relative Density at 60 °F	Relative density at 60°F and the equilibrium pressure	-		01.3	0
Observed temperature		°F		-100400	60
Product	1: A - Crude Oil		PRDTYP	-	1
	2: B - Auto select				
	Selection based on relative density at 60 °F				
	3: B - Gasoline				
	4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API2540 rounding	0: Disabled		APIROUND	-	0
	The calculations are performed with full precision and the final CTL value is				
	rounded as specified by input 'CTL decimal places'				
	1: Enabled for computational value				
	API-2540 rounding and truncating rules are applied and the computational				
	value for CTL as specified in the standard is used, meaning that the CTL value				
	has:				
	4 decimal places if CTL >=1				
	5 decimal places if CTL < 1.				
	2: Enabled for table value				
	API-2540 rounding and truncating rules are applied and the table value for				
	CTL as specified in the standard meaning that the CTL value has 4 decimal				
	places in all cases				
	3: Enabled with 5 decimal places				
	API-2540 rounding and truncating rules are applied, while the CTL value has 5				
	decimal places in all cases.				
	Note: although not strictly in accordance with the standard, this option is				
	more commonly used than option 'Enabled for computational value'				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API2540 rounding'				
КО	Actual value of constant K0 used for CTL calculation	-	К0		0
К1	Actual value of constant K1 used for CTL calculation	-	K1		0
К2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor	1/°F	ALPHA		0
Product	When input 'Product' is 'B - Auto select', then the output is set to the actual	-	PRDCUR		Input
	selected product of table 24B (enumerative value as defined for input				Product
	'Product'), else the output is set equal to input 'Product'.				
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
-	0: In Range				
	1: Out of Range				

Calculations

The calculations are in full or partial compliance with the standard depending on input 'API 2540 rounding'.

The input and output values are rounded in compliance with the standard. The CTL, CPL and CTPL value are rounded to 5

fxAPI_Table24_2004

Description

Relative Density (60°F, 0 psig) --> Relative Density (T, P)

This function converts a relative density value at 60°F and 0 psig to the relative density value at the observed temperature and pressure.

decimal places.

The temperature and pressure correction is according to API MPMS 11.1:2004.

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Relative density at 60 °F	Relative density at 60°F and 0 psig	-		01.3	0
Observed temperature	Temperature at which the API gravity is observed	°F		-100400	60
Observed pressure	Pressure at which the API gravity is observed	psig		-102000	0
Product	1: A - Crude Oil		PRDTYP		1
	2: B - Auto select				
	Selection based on relative density at 60 °F				
	3: B - Gasoline				
	4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API rounding	0: Disabled		APIROUND		0
	The calculations are performed with full precision and the				
	final CTL, CPL and CTPL values are rounded as specified by				
	the inputs 'CTL / CPL/ CTPL decimal places'				
	1: Enabled				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
Observed relative	Relative density at the observed temperature and pressure	-	RD		0
density					
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API rounding'				
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API rounding'				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
КО	Actual value of constant K0 used for CTL calculation	-	KO		0
K1	Actual value of constant K1 used for CTL calculation	-	K1		0
K2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor	1/°F	ALPHA		0
F	Compressibility factor	1/psi	F		0
Product	When input 'Product' is 'B - Auto select', then the output is set to the	-	PRDCUR		0
	actual selected product of table B (enumerative value as defined for				
	input 'Product'), else the output is set equal to input 'Product'.				
Calculation out of	With respect to the standard the input values are:	-		OOR	0
range	0: In Range				
	1: Out of Range				

Calculations

The calculations are in either full or partial compliance with the standard, depending on whether API rounding is enabled or not. Note: The 2004 standard assumes that the equilibrium pressure is below atmospheric pressure, so taking 0 psig as the base pressure.

Compliance

API MPMS 11.1 Temperature and Pressure Volume Correction Factors for Generalized Crude Oils, Refined Products and Lubricating Oils, May 2004

fxAPI_Table24E

Description

Relative Density (60°F) --> CTL

This function calculates the volume correction factor for temperature from the relative density value at 60°F and the observed temperature.

The temperature correction is according to API MPMS 11.2.4:2007 (GPA TP-25 / GPA TP-27).

Function inputs

Compliance

- API MPMS 11.2.4: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007
- GPA TP-27: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007
- GPA TP-25: Temperature Correction for the volume of Light Hydrocarbons – Tables 24E and 23E, 1998

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag				
	group				
Relative density at 60 °F	Relative density at 60°F	-		00.75	0
Observed temperature	Temperature at which the relative density is observed	°F		-100300	60
API rounding	0: Disabled The calculations are performed with full precision and the output values are not rounded 1: Enabled The input and output values are rounded as defined in the standard		APIROUND		0

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error				
	3: No convergence			CALC	
CTL	Volume correction factor for temperature	-	CTL		1
	Value will be rounded according to inputs 'API rounding'				
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
2	0: In Range				
	1: Out of Range				

Calculations

The calculations are in either full or partial compliance with the standard, depending on the selected type of API rounding.

fxAPI_Table53_1952

Description

Density (T) --> Density (15°C)

This function converts a density value at the observed temperature to the density at 15°C according to the API 1952 Table 53.

Compliance

• ASTM-IP Petroleum Measurement Tables, Metric Edition, Metric Units of Measurement, 1952

Function inputs

Boundaries

Table 53 contains values for the following range:

Input value	Normal Range	EU
Observed density	420 1099	kg/m3
Observed temperature	-25 125	°C

Note that the table does not cover the full range, e.g. for an Observed specific gravity of 0.420 the table only specifies values between 45 .. 60 $^{\circ}{\rm C}$

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag	group			
Observed density	Density at the observed temperature.	kg/m3		01300	0
Observed temperature		°C		-100200	15

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
Density at 15 °C	Density at 15 °C	kg/m3	DENS15		0
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
	0: In Range				
	1: Out of Range				

Calculations

The table values are the standard, so no calculations are involved. The function performs an interpolation between the table values that correspond to the input specific gravity and input temperature.

In case the combination of input values ('Observed density' and Observed temperature') is not covered by the table, the output 'Density at 15 °C' is set to 0 and output 'Calculation out of range' is set to 1.

fxAPI_Table53_1980

Description

Density (T) --> Density (15°C)

This function converts a density value at the observed temperature to the density value at 15°C.

The temperature conversion is according to API-2540, Tables 53A (Generalized Crude Oils) and 53B (Refined Oil Products) and API MPMS 11.1 Chapter XIV Table 53D: 1984 (Lubricating Oils). The function provides the option to correct for readings taken from a hydrometer as specified in the API-2540 standard.

Compliance

- API MPMS 11.1 Volume X (API Standard 2540) Table 53A -Generalized Crude Oils, Correction of Observed Density to Density at 15°C - First Edition, August 1980
- API MPMS 11.1 Volume X (API Standard 2540) Table 53B -Generalized Products, Correction of Observed Density to Density at 15°C - First Edition, August 1980
- API MPMS 11.1 Volume XIV Table 53D Generalized Lubricating Oils, Correction of Observed Density to Density at 15°C - January 1982

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Observed density	Density at observed temperature	kg/m3		01300	0
Observed temperature		°C		-100200	15
Product	1: A - Crude Oil		PRDTYP		1
	2: B - Auto select				
	Selection based on density at 15 °C				
	3: B - Gasoline				
	4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API2540 rounding	0: Disabled		APIROUND		0
	The calculations are performed with full precision.				
	A convergence limit of 0.000001 kg/m3 will be applied for				
	the iterative calculations.				
	1: Enabled				
	API-2540 rounding and truncating rules are applied.				
	A convergence limit of 0.05 kg/m3 will be applied as defined	1			
	in the standard.				
Hydrometer correction	0: Disabled		HYDROCOR		0
	1: Enabled				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Density at 15 °C	Density at 15°C	kg/m3 (s)	DENS15		0
CTL	Volume correction factor for temperature.	-	CTL		1
К0	Actual value of constant K0 used for CTL calculation	-	КО		0
К1	Actual value of constant K1 used for CTL calculation	-	K1		0
К2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor	1/°C	ALPHA		0
Product	When input 'Product' is 'B - Auto select', then the output is	-	PRDCUR		0
	set to the actual selected product of table 53B				
	(enumerative value as defined for input 'Product'), else the				
	output is set equal to input 'Product'.				
Calculation out of	With respect to the standard the input values are:	-		OOR	0
range	0: In Range				
	1: Out of Range				

Calculations

The calculations are in full or partial compliance with the standard depending on input 'API 2540 rounding'.

fxAPI_Table53_2004

Description

Density (T, P) --> Density (15°C, 0 bar(g))

This function converts a density value at the observed temperature and pressure to the density value at 15°C and 0 bar(g).

The temperature and pressure correction is according to API MPMS 11.1:2004.

An iterative calculation needs to be applied to convert the observed density to the value at base conditions.

Function inputs and outputs

Note: As opposed to API-2540, the 2004 standard does not include a correction for readings taken from a hydrometer and assumes that the equilibrium pressure is below atmospheric pressure, so taking 0 psig as the base pressure.

Compliance

API MPMS 11.1 Temperature and Pressure Volume Correction Factors for Generalized Crude Oils, Refined Products and Lubricating Oils, May 2004

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Observed density	Density at the observed temperature and pressure	kg/m3		01300	0
Observed temperature	Temperature at which the density is observed	°C		-100200	15
Observed pressure	Pressure at which the density is observed	bar(g)		-1250	0
Product	1: A - Crude Oil		PRDTYP		1
	2: B - Auto select				
	Selection based on density at 15 °C				
	3: B - Gasoline				
	4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API rounding	0: Disabled		APIROUND		0
	The calculations are performed with full precision and the final CTL,				
	CPL and CTPL values are rounded as specified by the inputs 'CTL /				
	CPL/ CTPL decimal places'				
	1: Enabled				
	The input and output values are rounded in compliance with the				
	standard. The CTL, CPL and CTPL value are rounded to 5 decimal				
	places.				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Density at 15 °C	Density at 15°C and 0 bar(g)	-	DENS15		0
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API rounding'				
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API rounding'				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
КО	Actual value of constant K0 used for CTL calculation	-	К0		0
К1	Actual value of constant K1 used for CTL calculation	-	K1		0
К2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor at 60 °F !	1/°C	ALPHA		0
F	Compressibility factor	1/bar	F		0
Product	When input 'Product' is 'B - Auto select', then the output is set to the	-	PRDCUR		0
	actual selected product of table B (enumerative value as defined for				
	input 'Product'), else the output is set equal to input 'Product'.				
Calculation out of	With respect to the standard the input values are:	-		OOR	0
range	0: In Range				
	1: Out of Range				

Calculations

The calculations are in either full or partial compliance with the standard, depending on whether API rounding is enabled or not.

fxAPI_Table53E

Description

Density (T) --> Density (15°C)

This function converts the density value at the observed temperature to the corresponding density at 15°C.

The temperature correction is according to API MPMS 11.2.4:2007 (GPA TP-27).

Function inputs

Compliance

- API MPMS 11.2.4: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007
- GPA TP-27: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Observed density	Density at the observed temperature	kg/m3		0750	0
Observed temperature	Temperature at which the relative density is observed	°C		-100150	15
API rounding	0: Disabled The calculations are performed with full precision and the output values are not rounded 1: Enabled The input and output values are rounded as defined in the standard				0

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Density at 15°C			DENS15		
CTL	Volume correction factor for temperature.	-	CTL		1
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
	0: In Range				
	1: Out of Range				

Calculations

The calculations are in either full or partial compliance with the standard, depending on the selected type of API rounding.

fxAPI_Table54_1952

Description

Density (15°C) --> CTL

This function determines the volume correction factor for temperature CTL from the relative density value at 15°C and the observed temperature according to the API 1952 Table 54.

Compliance

• ASTM-IP Petroleum Measurement Tables, Metric Edition, Metric Units of Measurement, 1952

Boundaries

Table 54 contains values for the following range:

Input value	Normal Range	EU
Density at 15 °C	500 1100	kg/m3
Observed temperature	-50 +150	°C

Note that the table does not cover the full range, e.g. for a Density at 15 °C of 500 kg/m3the table only specifies values between -50 .. 55 °C

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Density 15 °C	Specific gravity at 60°F	Kg/m3		01300	0
Observed temperature		°C		-100200	15

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
CTL	Volume correction factor for temperature.	-	CTL		1
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
	0: In Range				
	1: Out of Range				

Calculations

The table values are the standard, so no calculations are involved. The function performs an interpolation between the table values that correspond to the input specific gravity and input temperature.

In case the combination of input values ('Specific gravity 60 $^{\circ}$ F' and Observed temperature') is not covered by the table, the output 'CTL' is set to 1 and output 'Calculation out of range' is set to 1.

fxAPI_Table54_1980

Description

Density (15°C) --> CTL

This function calculates the volume correction factor for temperature CTL from the relative density value at 15°C and the observed temperature.

The temperature conversion is according to API-2540, Tables 54A (Generalized Crude Oils) and 54B (Refined Oil Products) and API MPMS 11.1 Chapter XIV Table 54D: 1984 (Lubricating Oils).

Compliance

- API MPMS 11.1 Volume X (API Standard 2540) Table 54A -Generalized Crude Oils, Correction of Volume to 15°C against Density at 15°F - First Edition, August 1980
- API MPMS 11.1 Volume X (API Standard 2540) Table 54B -Generalized Products, Correction of Volume to 15°C against Density at 15°C - First Edition, August 1980
- API MPMS 11.1 Volume XIV Table 54D Generalized Lubricating Oils, Correction of Volume to 15°C Against Density at 15°C - January 1982

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Density at 15 °C	Density at 15°C and the equilibrium pressure	kg/m3		01300	0
Observed		°C		-100200	15
temperature					
Product	1: A - Crude Oil		PRDTYP	-	1
	2: B - Auto select				
	3: B - Gasoline				
	4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API2540 rounding	0: Disabled		APIROUND	-	0
	The calculations are performed with full precision and the final CTL value is				
	rounded as specified by input 'CTL decimal places'				
	1: Enabled for computational value				
	API-2540 rounding and truncating rules are applied and the computational				
	value for CTL as specified in the standard is used, meaning that the CTL value				
	has:				
	4 decimal places if CTL >=1				
	5 decimal places if CTL < 1.				
	2: Enabled for table value				
	API-2540 rounding and truncating rules are applied and the table value for CTL				
	as specified in the standard meaning that the CTL value has 4 decimal places in				
	all cases				
	3: Enabled with 5 decimal places				
	API-2540 rounding and truncating rules are applied, while the CTL value has 5				
	decimal places in all cases.				
	Note: although not strictly in accordance with the standard, this option is more				
	commonly used than option 'Enabled for computational value'				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API2540 rounding'				
ко	Actual value of constant K0 used for CTL calculation	-	КО		0
K1	Actual value of constant K1 used for CTL calculation	-	K1		0
К2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor	1/°C	ALPHA		
Product	When input 'Product' is 'B - Auto select', then the output is set to the	-	PRDCUR		
	actual selected product of table 54B (enumerative value as defined for				
	input 'Product'), else the output is set equal to input 'Product'.				
Calculation out of range	With respect to the standard the input values are:	-		OOR	
	0: In Range				
	1: Out of Range				

Calculations

The calculations are in full or partial compliance with the standard depending on input 'API 2540 rounding'.

fxAPI_Table54_2004

Description

Density (15°C, 0 bar(g)) --> Density (T, P)

This function converts a density value at 15°C and 0 bar(g) to the density value at the observed temperature and pressure. The temperature and pressure correction is according to API MPMS 11.1:2004. Note: The 2004 standard assumes that the equilibrium pressure is below atmospheric pressure, so taking 0 psig as the base pressure.

Compliance

• API MPMS 11.1 Temperature and Pressure Volume Correction Factors for Generalized Crude Oils, Refined Products and Lubricating Oils, May 2004

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Density at 15 °C	Density at 15°C and 0 bar(g)	kg/m3		01300	0
Observed temperature	Temperature at which the density is observed	°C		-100200	15
Observed pressure	Pressure at which the density is observed	bar(g)		-1250	0
Product	1: A - Crude Oil		PRDTYP		1
	2: B - Auto select				
	Selection based on density at 15 °C				
	3: B - Gasoline				
	4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API rounding	0: Disabled		APIROUND		0
	The calculations are performed with full precision and the final				
	CTL, CPL and CTPL values are rounded as specified by the inputs				
	'CTL / CPL/ CTPL decimal places'				
	1: Enabled				
	The input and output values are rounded in compliance with the				
	standard. The CTL, CPL and CTPL value are rounded to 5 decimal				
	places.				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence				
Observed density	Density at the observed temperature and pressure	-	DENS		0
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API rounding'				
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API rounding'				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
ко	Actual value of constant K0 used for CTL calculation	-	KO		0
K1	Actual value of constant K1 used for CTL calculation	-	K1		0
К2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor at 60 °F !	1/°C	ALPHA		0
F	Compressibility factor	1/bar	F		0
Product	When input 'Product' is 'B - Auto select', then the output is set to the actual	-	PRDCUR		0
	selected product of table B (enumerative value as defined for input 'Product'),				
	else the output is set equal to input 'Product'.				
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
	0: In Range				
	1: Out of Range				

Calculations

The calculations are in either full or partial compliance with the standard, depending on whether API rounding is enabled or not.

fxAPI_Table54E

Description

Density (15°C) --> CTL

This function calculates the volume correction factor for temperature from the relative density value at 15°C and the observed temperature.

The temperature correction is according to API MPMS 11.2.4:2007 (GPA TP-27).

Function inputs

Compliance

- API MPMS 11.2.4: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007
- GPA TP-27: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Density at 15°C		-		0750	0
Observed temperature	Temperature at which the relative density is observed	°C		-100150	15
API rounding	0: Disabled The calculations are performed with full precision and the output values are not rounded 1: Enabled The input and output values are rounded as defined in the standard				0

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		1
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence				
CTL	Volume correction factor for temperature	-	CTL		1
	Value will be rounded according to input 'API rounding'				
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
	0: In Range				
	1: Out of Range				

Calculations

The calculations are in either full or partial compliance with the standard, depending on the selected type of API rounding.

fxAPI_Table59_2004

Description

Density (T, P) --> Density (20°C, 0 bar(g))

This function converts a density value at the observed temperature and pressure to the density value at 20°C and 0 bar(g).

The temperature and pressure correction is according to API MPMS 11.1:2004.

An iterative calculation needs to be applied to convert the observed density to the value at base conditions.

Function inputs

Note: As opposed to API-2540, the 2004 standard does not include a correction for readings taken from a hydrometer and assumes that the equilibrium pressure is below atmospheric pressure, so taking 0 psig as the base pressure.

Compliance

 API MPMS 11.1 Temperature and Pressure Volume Correction Factors for Generalized Crude Oils, Refined Products and Lubricating Oils, May 2004

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Observed density	Density at the observed temperature and pressure	kg/m3		01300	0
Observed temperature	Temperature at which the density is observed	°C		-10200	20
Observed pressure	Pressure at which the density is observed	bar(g)		-1250	0
Product	1: A - Crude Oil 2: B - Auto select Selection based on density at 20 °C 3: B - Gasoline 4: B - Transition Area 5: B - Jet Fuels 6: B - Fuel Oil 7: D - Lubricating Oil		PRDTYP		1
API rounding	 0: Disabled The calculations are performed with full precision and the final CTL, CPL and CTPL values are rounded as specified by the inputs 'CTL / CPL/ CTPL decimal places' 1: Enabled The input and output values are rounded in compliance with the standard. The CTL, CPL and CTPL value are rounded to 5 decimal places. 		APIROUND		0

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Density at 20 °C	Density at 20°C and 0 bar(g)	-	DENS20		0
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API rounding'				
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API rounding'				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
КО	Actual value of constant K0 used for CTL calculation	-	KO		0
K1	Actual value of constant K1 used for CTL calculation	-	K1		0
K2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor at 60 °F !	1/°C	ALPHA		0
F	Compressibility factor	1/bar	F		0
Product	When input 'Product' is 'B - Auto select', then the output is set to the actual	-	PRDCUR		0
	selected product of table B (enumerative value as defined for input 'Product'),				
	else the output is set equal to input 'Product'.				
Calculation out of	With respect to the standard the input values are:	-		OOR	0
range	0: In Range				
	1: Out of Range				

Calculations

The calculations are in either full or partial compliance with the standard, depending on whether API rounding is enabled or not.

fxAPI_Table59E

Description

Density (T) --> Density (20°C)

This function converts the density value at the observed temperature to the corresponding density at 20°C.

The temperature correction is according to API MPMS 11.2.4:2007 (GPA TP-27).

Function inputs

Compliance

- API MPMS 11.2.4: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007
- GPA TP-27: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Observed density	Density at the observed temperature	kg/m3		0750	0
Observed temperature	Temperature at which the relative density is observed	°C		-100200	20
API rounding	0: Disabled The calculations are performed with full precision and the output values are not rounded 1: Enabled The input and output values are rounded as defined in the standard		APIROUND		0

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Density at 20°C			DENS20		0
CTL	Volume correction factor for temperature.	-	CTL		1
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
5	0: In Range				
	1: Out of Range				

Calculations

The calculations are in either full or partial compliance with the standard, depending on the selected type of API rounding.

fxAPI_Table60_2004

Description

Density (20°C, 0 bar(g)) --> Density (T, P)

This function converts a density value at 20°C and 0 bar(g) to the density value at the observed temperature and pressure. The temperature and pressure correction is according to API MPMS 11.1:2004.

Function inputs

Note: The 2004 standard assumes that the equilibrium pressure is below atmospheric pressure, so taking 0 psig as the base pressure.

Compliance

• API MPMS 11.1 Temperature and Pressure Volume Correction Factors for Generalized Crude Oils, Refined Products and Lubricating Oils, May 2004

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Density at 20 °C	Density at 20°C and 0 bar(g)	kg/m3		01300	0
Observed temperature	Temperature at which the density is observed	°C		-100200	60
Observed pressure	Pressure at which the density is observed	bar(g)		-1250	0
Product	1: A - Crude Oil		PRDTYP		1
	2: B - Auto select				
	Selection based on density at 20 °C				
	3: B - Gasoline				
	4: B - Transition Area				
	5: B - Jet Fuels				
	6: B - Fuel Oil				
	7: D - Lubricating Oil				
API rounding	0: Disabled		APIROUND		0
	The calculations are performed with full precision and the final CTL,				
	CPL and CTPL values are rounded as specified by the inputs 'CTL /				
	CPL/ CTPL decimal places'				
	1: Enabled				
	The input and output values are rounded in compliance with the				
	standard. The CTL, CPL and CTPL value are rounded to 5 decimal				
	places.				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence				
Observed density	Density at the observed temperature and pressure	-	DENS		0
CTL	Volume correction factor for temperature.	-	CTL		1
	Value will be rounded according to input 'API rounding'				
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API rounding'				
CTPL	Combined volume correction factor	-	CTPL		1
	CTPL = CTL * CPL				
К0	Actual value of constant K0 used for CTL calculation	-	К0		0
K1	Actual value of constant K1 used for CTL calculation	-	K1		0
K2	Actual value of constant K2 used for CTL calculation	-	K2		0
Alpha	Thermal expansion factor at 60 °F !	1/°C	ALPHA		0
F	Compressibility factor	1/bar	F		0
Product	When input 'Product' is 'B - Auto select', then the output is set to the actual selected	-	PRDCUR		0
	product of table B (enumerative value as defined for input 'Product'), else the output				
	is set equal to input 'Product'.				
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
	0: In Range				
	1: Out of Range				

Calculations

The calculations are in either full or partial compliance with the standard, depending on whether API rounding is enabled or not.

fxAPI_Table60E

Description

Density (20°C) --> CTL

This function calculates the volume correction factor for temperature from the relative density value at 20°C and the observed temperature.

The temperature correction is according to API MPMS 11.2.4:2007 (GPA TP-27).

Function inputs

Compliance

- API MPMS 11.2.4: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007
- GPA TP-27: Temperature Correction for the Volume of NGL and LPG Tables 23E, 24E, 53E, 54E, 59E & 60E, September 2007

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Density at 20°C		-		0750	0
Observed temperature	Temperature at which the relative density is observed	°C		-100150	20
API rounding	0: Disabled The calculations are performed with full precision and the output values are not rounded 1: Enabled The input and output values are rounded as defined in the standard		APIROUND		0

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence				
CTL	Volume correction factor for temperature	-	CTL		1
	Value will be rounded according to input 'API rounding'				
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
-	0: In Range				
	1: Out of Range				

Calculations

The calculations are in either full or partial compliance with the standard, depending on the selected type of API rounding.

fxASTM_D1550_RD60

Description

This function calculates the relative density at 60°F and 0 psig of Butadiene based on ASTM Designation 1550. The ASTM-D1550 standard specifies several tables. This function uses table 1, which contains values for the relative density at 60°F as a function of the observed relative density and the observed temperature. The compressibility factor is calculated in accordance with API MPMS 11.2.2:1984.

Compliance

- ASTM Designation: D1550 -94 (Reapproved 2005), Standard ASTM Butadiene Measurement Table 1 Reduction of observed specific gravity to specific gravity at 15.6/15. °C (60/60°F)
- API MPMS 11.2.2 Compressibility Factors for Hydrocarbons: 0.350 - 0.637 Relative Density (60°F/60°F) and -50°F to 140°F Metering Temperature - Second Edition, October 1986

Boundaries

ASTM D1550 table 1 consists of table values with a resolution of 0.005 for the relative density and 1°F for the temperature. The function applies a linear interpolation between table values based on the input values. The table specifies relative density at 60°F values that are limited to a low value of 0.621 and a high value of 0.634. When the combination of input values lies outside the range covered by the table, the function outputs either the low or high table value depending on whether the input values lay below or above the table limits.

API MPMS 11.2.2 applies for the following range:

- 0.350 to 0.637 (relative density)
- -50 to 140 °F
- 0 to 2200 psig
- . .

Function inputs

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Observed relative density		-	01	0
Observed temperature	Temperature at which the relative density is observed	°F	-100150	20
Observed pressure	Pressure at which the relative density is observed	Psig	-102500	0
API 11.2.2 rounding	0: Disabled			0
	The calculations are performed with full precision			
	1: Enabled			
	The input and intermediate values are rounded and truncated as defined in the standard			

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence				
Relative density at 60°F		-	RD60		0
F	Compressibility factor	1/psi	F		0
	The output value will be either rounded or not depending input 'API 11.2.2				
	rounding'				
CTL	Volume correction factor for temperature.	-	CTL		1
CPL	Volume correction factor for pressure	-	CPL		1
	Value will be rounded according to input 'API 11.2.2 rounding'				
ASTM D1550 calculation out	With respect to ASTM D1550 table 1 the input values are:	-	ASTM_OOR		0
of range	0: In Range				
	1: Out of Range				
API 11.2.2 calculation out of	With respect to API 11.2.2 the conditions of density, temperature and		API_OOR		
range	pressure are:				
	0: In Range				
	1: Out of Range				

Calculations

The function performs the following iterative algorithm to calculate the relative density at 60 $^\circ{\rm F}$ and 0 psig. No rounding is applied

- 1 At the start of the iteration the relative density at [60 $^\circ\text{F}$, 0 psig] is set equal to the observed relative density and the CPL value is set to 1.
- 2 The relative density corrected for pressure is calculated by dividing the observed relative density by the CPL value.
- 3 The CTL value and the relative density at [60 °F, equilibrium pressure] is calculated from the relative density corrected for pressure and the observed temperature according to ASTM D1550 Table 1
- 4 The compressibility factor F is calculated according to API MPMS 11.2.2 from the relative density at [60 °F, equilibrium pressure] and the 'Observed temperature'. Depending on setting 'API 11.2.2. rounding' the rounding and truncation rules of API 11.2.2. are applied for this step.
- 5 The CPL value is calculated from the calculated compressibility factor and the 'Observed pressure' input value.
- 6 The new value for relative density at [60°F, 0 psig] is calculated by dividing the observed density by the CTL and CPL values.
- 7 Steps 2 through 6 are repeated until the absolute difference between two consecutive relative density values is less than the convergence limit of 1e-8.

8 When no convergence is achieved in 20 iterations then the status output is set to 'No convergence' and the relative density output is set to 0.

fxASTM_D1550_Ctl

Description

This function calculates the Ctl value (VCF) of Butadiene based on the ASTM Designation 1550. The standard specifies several tables. This function uses table 2, which contains values for the volume correction factor as a function of the relative density at 60°F and the observed temperature.

Function inputs

Compliance

 ASTM Designation: D1550 -94 (Reapproved 2005), Standard ASTM Butadiene Measurement Table 2 Reduction of Observed Volume to 15.6°C (60°F) Against Specific Gravity 60/60°F

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Relative density at 60°F		-		01	0
Observed temperature	Temperature at which the relative density is observed	°F		-100150	20

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
Ctl	Volume correction factor for temperature. The output value	-	CTL		1
	will NOT be rounded.				
Calculation out of range	With respect to the standard the input values are:	-		OOR	0
	0: In Range				
	1: Out of Range				

Calculations

The Standard consists of a set of table values with a resolution of 0.001 for the relative density at 60° F and 1° F for temperature. The function applies a linear interpolation between table values based on the unrounded input values. When an input value lies outside the range of the table the function limits the input value to the minimum or maximum value of the table.

fxASTM_D4311M_09_C

Description

This function calculates the Ctl value (VCF) for Asphalt according to ASTM D4311/D4311M-09, using US Customary units.

Compliance

• ASTM Designation: D4311/D4311M-09, Standard Practice for Determining Asphalt Volume Correction to a Base Temperature

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
API gravity at 60°F		-		034.9	0
Observed temperature		°F		0500	60

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
Ctl	Volume correction factor for temperature. The output value will NOT be rounded.	-	CTL		1

Calculations

The calculations are in accordance with appendix X1. FORMULAS USED IN DETERMINING VOLUME CORRECTIONS TO A BASE TEMPERATURE of the standard.

fxASTM_D4311M_09_M

Description

This function calculates the Ctl value (VCF) for Asphalt according to ASTM D4311/D4311M-09, using metric units.

Compliance

• ASTM Designation: D4311/D4311M-09, Standard Practice for Determining Asphalt Volume Correction to a Base Temperature

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Density at 15°C		kg/m3		8001200	966
Observed temperature		°C		-25274.5	15

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
Ctl	Volume correction factor for temperature. The output value	-	CTL		1
	will NOT be rounded.				

Calculations

The calculations are in accordance with appendix X1. FORMULAS USED IN DETERMINING VOLUME CORRECTIONS TO A BASE TEMPERATURE of the standard.

fxConvertUnit

Description

This function converts a value expressed in a particular unit into the corresponding value expressed in another unit.

The input and output unit must belong category, otherwise the

conversion fails.

	Function	inputs
--	----------	--------

Function inputs	Remark	EU	SW tag	Range	Default
Name					
Input value	The value to be converted				
Input unit	Unit of the value to be converted				
	Use one of the "xu" unit constants.				
Output unit	Unit of the output value				
	Use one of the "xu" unit constants.				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Output value		<output unit=""></output>	VAL		0
Conversion failure	The conversion fails when the input unit and output unit do not belong to the same unit category (or also when the input unit and / or output unit are not a valid unit). 0: Normal 1: Failure		CONVFAIL		

fxEthylene_IUPAC_C

Description

The function calculates the compressibility factor and the density of Ethylene (C2H4, also called Ethene) based on the Equation Of State published by IUPAC and in **US customary units**.

References

• Ethylene (Ethene), International Thermodynamic Tables of the Fluid State Vol. 10 (1988), IUPAC, ISBN 0-63201-7090.

Boundaries

The limits of the tables are 104 K to 320 K (-272 .. +116 °F) for pressures up to 270 MPa (39160 psi) and 104K to 450K (-272 .. +350 °F) for pressures up to 40 MPa (5800 psi).

Function inputs				
Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Temperature		°F	-300200	0
Pressure		psia	050000	0

Function outputs

Function outputs	Remark	EU	Fallback
Status	0: Normal		
	1: Input argument out of range		
	2: Calculation error		
	3: No convergence		0
Density		lbm/ft3	0
Compressibility		-	0
Equilibrium pressure	Equilibrium pressure at the observed temperature.	psia	0
	Also referred to as vapor pressure or saturated pressure		
Range	With respect to the combination of temperature and pressure is:	-	0 0 0
	0: In Range		
	1: Out of Range		

Calculations

The calculations are in compliance with the standard.

fxEthylene_IUPAC_M

Description

The function calculates the compressibility factor and the density of Ethylene (C2H4, also called Ethene) based on the Equation Of State published by IUPAC and in **metric units**.

References

• Ethylene (Ethene), International Thermodynamic Tables of the Fluid State Vol. 10 (1988), IUPAC, ISBN 0-63201-7090.

Function inputs

Function inputs Remark EU Range Default Name Optional tag name, tag description and tag group -200..200 0 Temperature °C -200..200 0 Pressure bar(a) 3000 0

Function outputs	Remark	EU	Fallback
Status	0: Normal		
	1: Input argument out of range		
	2: Calculation error		
	3: No convergence		
Density		kg/m3	0
Compressibility		-	0
Equilibrium pressure	Equilibrium pressure at the observed temperature.	bar(a)	0
	Also referred to as vapor pressure or saturated pressure		
Range	With respect to the combination of temperature and pressure is:	-	0
	0: In Range		
	1: Out of Range		

Calculations

The calculations are in compliance with the standard.

Because the IUPAC Equation Of State specifies the calculation of the pressure from a known temperature and density iteration is required to determine the density from the input pressure. A convergence limit of 0.0005 kg/m3 is applied. A maximum of 20 iterations is applied.

Boundaries

The limits of the tables are 104 K to 320 K (-170 .. +47 °C) for pressures up to 270 MPa (2700 bar) and 104K to 450K (-170 .. +177 °C) for pressures up to 40 MPa (400 bar).

fxEthylene_NIST1045

Description

The function calculates the density of Ethylene (C2H4, also called Ethene) based on the NIST-1045 Equation Of State in **metric units**.

References

• R.D. McCarty and R.T. Jacobsen "An Equation of State for Fluid Ethylene", National Bureau of Standards (NBS), US, Technical Note 1045, 1981.

Function inputs

Boundaries

The equation of state is valid from 104 K to 400 K (-170 .. +127 $^{\circ}\text{C})$ and for pressures up to 40 MPa (400 bar).

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Temperature		°C	-200200	0
Pressure		bar(a)	500	0
Rounding	 0: Disabled The calculations are performed with full precision. A convergence limit of 1e-10 kg/m3 will be applied for the iterative calculations. 1: Enabled The calculations are performed in full compliance with the standard. The input, intermediate and output values are rounded as specified and also the iteration limit as specified in the standard is used to 	-		1
	values are rounded as specified and also the iteration limit as specified in the standard is used to achieve convergence.			

Function outputs

Function outputs	Remark	EU	Fallback	
Status	0: Normal			
	1: Input argument out of range			
	2: Calculation error			
	3: No convergence			
Density		kg/m3	0	
Compressibility		-	0	
Equilibrium pressure	Equilibrium pressure at the observed temperature.	bar(a)	0	
	Also referred to as vapor pressure or saturated pressure			
Range	With respect to the standard the inputs are:	-	0	
	0: In Range			
	1: Out of Range			

Calculations

The calculations are in compliance with the standard.

fxGasViscosity_2004

Description

The dynamic viscosity of natural gas is calculated in accordance with a method developed at the Rostock University in Germany and published in 2004 in the 'International Journal of the Thermophysics'. The estimated uncertainty of the calculated viscosity is 0.5% for natural gas (0.3% for pure methane).

Reference

 Viscosity Measurements and Predictions for Natural Gas, P. Schley, M. Jaeschke, C. Kuchenmeister and E. Vogel, International Journal of Thermophysics, Vol. 25, No. 6, November 2004 (© 2004)

Boundaries

The limits of the tables are 250 K to 450 K (-24 .. +177 °C) for pressures up to 30 MPa (300 bar).

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Density	Density value	kg/m3	02000	0
Temperature	Temperature value	°C	-200+400	0
Composition	Standard composition as defined in section 'Standard gas composition'.	mol/mol	01	0

Function outputs

Function outputs	Remark	EU	Fallback
Status	0: Normal		
	1: Input argument out of range		
	2: Calculation error		
	3: No convergence		
	4: Mole fractions do not add up to 1.0 +- 0.0001		
	6: Hardware not supported		
Dynamic viscosity		Pa.s	0
Range	With respect to the combination of temperature and pressure is:		0
	0: In Range		
	1: Out of Range		

Calculations

The calculations are in accordance with the publication, which is based on the following 12 components: methane (+ hydrogen), nitrogen (+ helium + oxygen/argon), carbon dioxide, ethane, propane, n-butane, ibutane, n-pentane (+ benzene), isentane (+ neopentane), n-hexane (hexanes + toluene), n-heptane (heptanes), and n-octane (octanes + nonanes+ C10 plus higher + xylenes).

The function uses the 'Standard gas composition' as used by most gas property functions. The 22 components of input Composition are lumped to these 12 components as defined in the following table.

Notes:

- Components 'Water', 'Hydrogen sulphide' and 'Carbon monoxide' are not supported by the publication and will therefore be neglected by the function. The lumped composition is normalized to 1 and then used in the calculation.
- The publication also mentions components benzene, toluene and xylenes, which are not part of the 'Standard gas composition'. To adhere strictly to the publication, add 'Benzene' to input 'n-Pentane', 'Toluene' to input 'n-Hexane' and 'Xylenes' to input 'n-Octane'.

Input component	Added to / neglected
Methane	Methane
Nitrogen	Nitrogen
Carbon Dioxide	Carbon Dioxide
Ethane	Ethane
Propane	Propane
Water	Neglected
Hydrogen Sulphide	Neglected
Hydrogen	Methane
Carbon Monoxide	Neglected
Oxygen	Nitrogen
i-Butane	i-Butane
n-Butane	n-Butane
i-Pentane	i-Pentane
n-Pentane	n-Pentane
n-Hexane	n-Hexane
n-Heptane	n-Heptane
n-Octane	n-Octane
n-Nonane	n-Octane
n-Decane	n-Octane
Helium	Nitrogen
Argon	Nitrogen
Neo-Pentane	Iso-Pentane

fxGERG2008_Gas

Description

The compressibility, density, speed of sound and isentropic coefficient of a gas are calculated from its composition, temperature and pressure in accordance with the GERG-2008 equation of state. The function uses an optimized routine that only applies when fluid is in the gaseous state (100% gas) and that requires limited CPU time (suitable for processing in the Flow-X flow computer).

Compliance

 The GERG-2008 Wide-Range Equation of State for Natural Gases and Other Mixtures: An Expansion of GERG-2004. Kunz, O., Wagner, W. Submitted to J. Chem. Eng. Data 57 (2012).

Boundaries

The GERG-2008 calculation has defined uncertainty bounds for fluids that lie within the 'Normal range'. Also an 'Expanded range' of gas mixtures is defined for which the calculation has a higher uncertainty. Using the calculation at conditions outside the 'Expanded range' is not recommended.

Input value	Normal Range	Expanded Range	EU
Pressure	0 35	0 70	MPa(a)
Temperature	90450	60700	К
Pressure	0 350	0 700	bar(a)
Temperature	-183 +177	-213 +426	°C
Pressure	0 5076	0 10153	psia
Temperature	-298 +350	-352 +800	°F
Mole fraction of all components	0.00 1.00	0.00 1.00	-

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Pressure	Pressure value	bar(a)		02000	1.01325
Temperature	Temperature value	°C		-200+400	0
Composition	Standard composition as defined in section 'Standard gas composition.	mol/mol	COMP	01	0
neo-Pentane mode	Determines what to do when component neo-Pentane is larger than zero 1: Add to i-Pentane 2: Add to n-Pentane 3: Neglect	-	NEOC5_MODE		1

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
	4: Mole fractions do not add up to 1.0 +- 0.0001			COMPERR	
	6: Hardware not supported				
Compressibility		-	Z		1
Mass Density		kg/m3	MASDENS		0
Molar Density		kmol/m3	MOLDENS		0
Molar Mass		kg/kmol	MOLMASS		0
Speed of Sound		m/s	SOS		0
Isentropic Exponent		-	К		0
Range	0: In Normal Range		RANGE		0
	All inputs are within the 'Normal Range'				
	1: In Extended Range				
	One or more inputs within the 'Extended Range, but none of the				
	inputs outside the Extended rang (outputs values have higher				
	uncertainty)				
	2: Out of Range				
	One or more inputs outside the 'Extended Range' (using the			OOR	
	calculation is not recommended in this case)				

Calculations

The calculations are as documented in the reference.

fxGERG2008_Flash

Description

The compressibility and density of a gas/liquid mixture are calculated from its composition, temperature and pressure in accordance with the GERG2008 standard.

Compliance

- Kunz, O., and W. Wagner. "The GERG-2008 wide-range equation of state for natural gases and other mixtures: An expansion of GERG-2004." Journal of Chemical & Engineering Data 57.11 (2012): 3032-3091.
- W. Wagner. "Description of the Software Package for the Calculation of Thermodynamic Properties from the GERG-2008 Wide-Range Equation of State for Natural Gases and Other Mixtures".

Boundaries

Function inputs

The GERG2008 calculation has defined uncertainty bounds for gas mixtures that lie within the 'Normal range'. Also an 'Expanded range' of gas mixtures is defined for which the GERG2008 calculation has a higher uncertainty. Using the GERG2008 calculation for gas mixtures that lie outside the 'Expanded range' is not recommended.

Input value	Normal Range	Expanded Range	EU
Pressure	0350	0700	bar(a)
Temperature	-183 +177	-213 +427	°C
Mole fraction of Methane	0.00 1.00	0.00 1.00	-
Mole fraction of Ethane	0.00 1.00	0.00 1.00	-
Mole fraction of Propane	0.00 1.00	0.00 1.00	-
Mole fraction of Butanes	0.00 1.00	0.00 1.00	-
Mole fraction of Pentanes	0.00 1.00	0.00 1.00	-
Mole fraction of Hexanes Plus	0.00 1.00	0.00 1.00	-
Mole fraction of Carbon monoxide	0.00 1.00	0.00 1.00	-
Mole fraction of Carbon dioxide	0.00 1.00	0.00 1.00	-
Mole fraction of Nitrogen	0.00 1.00	0.00 1.00	-
Mole fraction of Helium	0.00 1.00	0.00 1.00	-
Mole fraction of Argon	0.00 1.00	0.00 1.00	-
Mole fraction of Oxygen	0.00 1.00	0.00 1.00	-
Mole fraction of Hydrogen	0.00 1.00	0.00 1.00	-
Sulphide			
Mole fraction of Hydrogen	0.00 1.00	0.00 1.00	-
Mole fraction of Water	0.00 1.00	0.00 1.00	-

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Pressure	Pressure value	bar(a)		02000	1.01325
Temperature	Temperature value	°C		-200+400	0
Composition	Standard composition as defined in section 'Standard gas composition.	mol/mol	СОМР	01	0
neo-Pentane mode	Determines what to do when component neo-Pentane is larger than zero 1: Add to i-Pentane 2: Add to n-Pentane 3: Neglect	-	NEOC5_MODE		1

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
	4: Mole fractions do not add up to 1.0 +- 0.0001			COMPERR	
	6: Hardware not supported				
Vapor fraction			FV		0
Compressibility vapor	Compressibility of the vapor fraction of the mixture		ZV		0
Compressibility liquid	Compressibility of the liquid fraction of the mixture		ZL		0
Compressibility total	Compressibility of the mixture		Z		0
Density vapor	Density of the vapor fraction of the mixture	kg/m3	DV		0
Density liquid	Density of the liquid fraction of the mixture	kg/m3	DL		0
Density total	Density of the mixture	kg/m3	D		0

Calculations

The calculations are as documented in the standard.

fxGOST30319

Description

Thermodynamic properties of a gas are calculated from its composition, temperature and pressure in accordance with GOST30319 standard, with the input and output values in **metric** units.

Compliance

• GOST30319-2.96 - 1 July 1997

Function inputs

Boundaries

The GOST30319 calculation has the normal range 263.15K..500K, 1bar..150bar and the extended range 125K..700K, 0bar..3500bar.

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Pressure	Pressure value	bar(a)		03500	1.01325
Temperature	Temperature value	°C		-250+450	25
Composition	Standard composition as defined in section 'Standard gas	mol/mol		01	0
	composition.				
neo-Pentane mode	Determines what to do when component neo-Pentane is larger than	-			1
	zero				
	1: Add to i-Pentane				
	2: Add to n-Pentane				
	3: Neglect				
Reference pressure		bar(a)		03500	0
Reference temperature		°C		-250+450	25

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
	4: Mole fractions do not add up to 1.0 +- 0.0001			COMPERR	
Compressibility ratio	Compressibility ratio between flow and base conditions	-	ZRATIO		1
Flow compressibility	Compressibility factor at flow conditions	-	ZF		1
Base compressibility	Compressibility factor at base conditions	-	ZB		1
Critical density		kg/m3	CRITDENS		0
Reduced density	Ratio between current and critical	-	REDUDENS		1
Critical temperature		°C	CRITTEMP		0
Reduced temperature	Ratio between current and critical	-	REDUTEMP		1
Critical pressure		bar(a)	CRITPRES		0
Reduced pressure	Ratio between current and critical	-	REDUPRES		1
Pitzer factor	Denoted as Capital Omega	-	PITZER		0
Range	0: In Normal Range		OOR		0
	All inputs are within the 'Normal Range'				
	1: In Extended Range				
	One or more inputs within the 'Extended Range, but none of the				
	inputs outside the Extended range (outputs values have higher uncertainty)				
	2: Out of Range				
	One or more inputs outside the 'Extended Range' (using the			OOR	
	calculation is not recommended in this case				

Calculations

The calculations are performed according to the source code of Sick Gmbh.

fxGOST30319_AGA8

Description

Compressibility of a gas are calculated from its composition, temperature and pressure in accordance with GOST30319 standard, with the input and output values in **metric** units.

Compliance

• GOST30319-2.96, chapter AGA8 - 1 July 1997

Function inputs

Function inputs Remark EU SW tag Range Default Optional tag name, tag description and tag group Name 1.01325 0..3500 Pressure Pressure value bar(a) -250..+450 Temperature Temperature value °C 25 Composition Standard composition as defined in section 'Standard gas mol/mol 0..1 0 composition. neo-Pentane mode Determines what to do when component neo-Pentane is larger than -1 zero 1: Add to i-Pentane 2: Add to n-Pentane 3: Neglect Reference pressure bar(a) 0..3500 0 Reference temperature °C -250..+450 25

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
	4: Mole fractions do not add up to 1.0 +- 0.0001			COMPERR	
Compressibility ratio	Compressibility ratio between flow and base conditions	-	ZRATIO		1
Flow compressibility	Compressibility factor at flow conditions	-	ZF		1
Base compressibility	Compressibility factor at base conditions	-	ZB		1
Heating value		MJ/m3	HV		0

Calculations

The calculations are performed according to the source code of Sick Gmbh.

Boundaries

The GOST30319 calculation has the normal range 263.15K..500K, 1bar..150bar and the extended range 125K..700K, 0bar..3500bar.

fxGOST30319_SGERG91

Description

Thermodynamic properties of a gas are calculated from its density, speed of sound, fractions of nitrogen and carbon dioxide, temperature and pressure in accordance with GOST30319 standard, with the input and output values in **metric** units.

Compliance

- GOST30319-2.96, chapter GERG91mod (1996) •
- GOST30319-2.2015, chapter GERG91mod (2015) •

Boundaries

The 'Range' output of this function is set as follows

	1996	2015	
Pressure	1120	175	bar(a)
Temperature	250340	250350	К
Carbon dioxide	0.00 0.20	0.00 0.20	mol/mol
Nitrogen	0.00 0.15	0.00 0.20	mol/mol
Normal density	0.66 1.00	0.66 1.05	kg/m3

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group		-	-	
Pressure	Pressure value	bar(a)		02000	7
Temperature	Temperature value	°C		-200+300	25
N2	Nitrogen fraction	mol/mol		01	0.01
CO2	Carbon dioxide fraction	mol/mol		01	0.02
Normal density	Density at base conditions	kg/m3		0.661.0	
Speed of sound	Optional, needed for the calculation of molar mass	m/s			0
Edition	Year of edition	-		-	1
	1: 1997				
	2: 2015				

Description

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
Base compressibility	Compressibility factor at base conditions	-	ZB		1
Flow compressibility	Compressibility factor at flow conditions	-	ZF		1
Compressibility ratio	Compressibility ratio between flow and base	-	ZRATIO		1
	conditions				
Base density		kg/m3	BASEDENS		0
Flow density	Compressibility ratio between flow and base	kg/m3	FLOWDENS		0
	conditions				
Heating value		MJ/m3	HV		0
Molar mass	Only calculated if a valid Speed Of Sound specified	kg/kmol	MM		0
Range	The inputs are within the normal range of use		RANGE		0

Calculations

The calculations are performed according to the source code of Sick Gmbh.

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fxGPA_TP15

Description

The GPA-TP15 standard defines a generalized correlation method to determine the vapor pressure (i.e. the equilibrium pressure) for natural gas liquids (NGL).

The vapor pressure is required by the API 11.2.2 and API 11.1:2004 calculations of the CPL value for light hydrocarbon liquids that have a vapor pressure above atmospheric pressure.

References

- API MPMS 11.2.2 Addendum Compressibility Factors for Hydrocarbons: Correlation Factors of Vapor Pressure for Commercial Natural Gas Liquids - First Edition, December 1994
- GPA Technical Publication TP-15 A Simplified Vapor Pressure Correlation for Commercial NGLs - 1988
- API MPMS 11.2.5 A Simplified Vapor Pressure Correlation for Commercial NGLs - September 2007
- GPA Technical Publication TP-15 A Simplified Vapor Pressure Correlation for Commercial NGLs - September 2007
- ASTM Technical Publication [Stock No. PETROLTBL-TP15] -September, 2007

Note: the first two and the last three refer to one and the same standard. The current standard GPA-TP15 (2007) / API MPMS 11.2.5 extends the applicable range of the previous standard GPA TP-15 (1988) / API MPMS 11.2.2 Addendum 1994 while preserving the calculations and constants of the previous standard. **Function input**

Boundaries

The GPA TP-15:1988 / API MPMS 11.2.2 Addendum:1994 correlation method is valid for the following range.

- 0.490 to 0.676 (RD60)
- -50 to 140 °F

The GPA TP-15:2007 / API MPMS 11.2.5:2007 correlation method has been extended for lower density and is valid for the following ranges.

Lower range:

- 0.350 to 0.425 (RD60)
- -50 to (695.51*RD60 155.51) °F

with RD60 being the relative density at 60°F

Higher range:

- 0.425 to 0.676 (relative density)
- -50 to 140 °F

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Relative density at 60°F		-	0.3 0.75	0
Observed Temperature		°F	-100200	60
API rounding	0: Disabled	-		0
	Full precision (no rounding applied)			
	1: Enabled			
	Rounding as defined in ' GPA TP15:1988 / API MPMS 11.2.2 Addendum':1994			
P100 Correlation	0: Disabled	-		0
	The standard correlation is commonly used for pure products such as			
	propane, butane and natural gasoline. It only requires the relative density and			
	the temperature to calculate the vapor pressure			
	1: Enabled			
	The improved correlation requires the vapor pressure at 100°F (37.8 °C). This			
	method is better suited for varied NGL mixes Where different product mixes			
	could have the same specific gravity but different equilibrium pressures.			
Vapor pressure at 100°F		psia	0500	0

Function outputs					
Function outputs	Remark	EU	Fallback		
Status	0: Normal				
	1: Input argument out of range				
	2: Calculation error				
Vapor pressure	Vapor pressure at 60°F	psia	0		
Range	With respect to the 2007 standard the combination of relative density and temperature is:	-	0		
	0: In Range				
	1: Out of Range				

Calculations

The calculations are in either full or partial compliance with the standards, depending on the selected type of API rounding.

fxGPA2172_C

Description

This uses the procedure for calculating heating value, specific gravity and compressibility factor at **<u>customary</u>** (imperial) conditions from the compositional analysis of a natural gas mixture.

GPA2172 describes the calculation methods to determine the compositional properties based on the individual component values and it refers to the GPA Standard 2145 (GPA2145) standard for these individual component values.

The effect of water on the calculations is rather complicated and is accounted for with a simplified equation that is considered to be adequate for custody transfer applications.

Therefore compositional properties are calculated for the following gas compositions:

- Wet gas composition the water fraction of input "Composition' is taken as the actual water fraction.
- Dry gas composition the water fraction is set to 0 and the composition is normalized to unity.
- Saturated gas composition the water fraction is set to the value when the gas is saturated with water and the composition is normalized to unity

GPA-2172 prescribes that the most recent edition of GPA2145 used for the individual component values. The function provides the option to use the values from a specific GPA2145 standard. In order to verify the function based on the examples of GPA2172 the function provides the option to GPA2145-89 (edition 1989) as well.

Compliance

- GPA Standard 2172-96, Calculation of Gross Heating Value, Relative Density and Compressibility Factor for Natural Gas Mixtures from Compositional Analysis - 1996
- API MPMS 14.5
- ASTM D3588-98 (Reapproved 2003)
- GPA Standard 2145-89, Table of Physical Constants
- GPA Standard 2145-00, Table of Physical Constants
- GPA Standard 2145-03, Table of Physical Constants
- GPA Standard 2145-09, Table of Physical Constants
- GPA Standard 2145-16, Table of Physical Constants

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Composition	Standard composition as defined section 'Standard Gas Composition'	mol/mol	COMP	01	0
Edition	Refers to the base conditions and the editions of the GPA2145 values.	-	EDITION		2
	Note that these base conditions are used for both the density and the				
	heating value.				
	1: 60°F, 14.696 psia, GPA2145-89 (1989)				
	2: 60°F, 14.696 psia, GPA2145-00 (2000)				
	3: 60°F, 14.696 psia, GPA2145-03 (2003)				
	4: 60°F, 14.696 psia, GPA2145-09 (2009)				
	5: 60°F, 14.696 psia, GPA2145-16 (2016)				
neo-Pentane mode	Determines what to do when component neo-Pentane is larger than	-	NEOC5_MODE		1
	zero				
	1: Add to i-Pentane				
	2: Add to n-Pentane				
	3: Neglect				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: Composition error			COMPERR	
	Composition does not add up to 100% +- 0.01%				
	In case of an error the output values will be set to the				
	fallback values				
Gross Heating Value (Wet)		Btu/ft3	VOLGHV_WET		0
Molar Mass (Wet)		lbm/lbmol	MOLMASS_WET		0
Molar Mass Ratio (Wet)			ISG_WET		0
Relative Density (Wet)	Based on the compressibility of wet air	-	RRD_WET		0
Compressibility (Wet)		-	Z_DWET		0
Gross Heating Value (Wet)		Btu/lbm	MASGHV_WET		0
Net Heating Value (Wet)		Btu/ft3	VOLNHV_WET		0
Gross Heating Value (Dry)		Btu/ft3	VOLGHV_DRY		0
Molar Mass (Dry)		-	MOLMASS_DRY		0
Molar Mass Ratio (Dry)		lbm/lbmol	ISG_DRY		0
Relative Density (Dry)			RRD_DRY		0
Compressibility (Dry)		-	Z_DRY		0
Gross Heating Value (Dry)		Btu/lbm	MASGHV_DRY		0
Net Heating Value (Dry)		Btu/ft3	VOLNHV_DRY		0

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Gross Heating Value	The saturated Gross Heating Value is commonly used for	Btu/ft3	VOLGHV_SAT		0
(Saturated)	custody transfer energy calculations				
Molar Mass (Saturated)		lbm/lbmol	MOLMASS_SAT		0
Molar Mass Ratio (Saturated)	Ideal specific gravity	-	ISG_SAT		0
Relative Density (Saturated)	Based on the compressibility of saturated air	-	RRD_SAT		0
Compressibility (Saturated)		-	Z_SAT		0
Gross Heating Value		Btu/lbm	MASGHV_SAT		0
(Saturated)					
Net Heating Value		Btu/ft3	VOLNHV_SAT		0
(Saturated)					

Calculations

The calculations are as documented in the GPA-2172 standard using the GPA2145 table values. However the calculations are performed at full precision, so not with intermediate rounding as shown in the examples of the standard.

Please note that the function uses the input composition 'as is', so does <u>not</u> apply normalization to 1.

- When the water fraction input value is above the water fraction of the saturated gas then the function continues its calculations without any correction.
- GPA-2145 standard editions 2000 and 2003 do not specify properties for hydrogen, argon and carbon monoxide. The function processes these components like the other components but with all property values set to 0.

Water vapour pressure

The saturated properties are dependent on the water vapour pressure water. GPA-2172 version 1996 states a slightly different value for the water vapour pressure than the 2009 and 2014 editions. Therefore the following values are used by this function.

Edition	Value [psia]	According to
1: GPA2145-89	0.25636	GPA2172-96
2: GPA2145-00	0.25636	GPA2172-96
3: GPA2145-03	0.25636	GPA2172-96
4: GPA2145-09	0.2564	GPA2172-09/14
5: GPA2145-16	0.2564	GPA2172-09/14

fxGPA2172_M

Description

This function uses the procedure for calculating heating value, specific gravity and compressibility factor at **metric** conditions from the compositional analysis of a natural gas mixture. GPA2172 describes the calculation methods to determine the compositional properties based on the individual component values and it refers to the GPA Standard 2145 (GPA2145) standard for these individual component values.

The effect of water on the calculations is rather complicated and is accounted for with a simplified equation that is considered to be adequate for custody transfer applications.

Therefore compositional properties are calculated for the following gas compositions:

- Wet gas composition the water fraction of input "Composition' is taken as the actual water fraction.
- Dry gas composition the water mole fraction of input "Composition' is set to 0 and the composition is normalized to unity.
- Saturated gas composition the water fraction value of input "Composition' is set to the water saturated mole fraction and the composition is normalized to unity

GPA-2172 prescribes that the most recent edition of GPA2145 used for the individual component values. The function provides the option to use the values from a specific GPA2145 standard.

Compliance

- GPA Standard 2172-96, Calculation of Gross Heating Value, Relative Density and Compressibility Factor for Natural Gas Mixtures from Compositional Analysis - 1996
- API MPMS 14.5 (same as GPA2172-96)
- GPA Standard 2145-00, Table of Physical Constants
- GPA Standard 2145-03, Table of Physical Constants
- GPA Standard 2145-09, Table of Physical Constants
- GPA Standard 2145-16, Table of Physical Constants

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Composition	Standard composition as defined section 'Standard Gas	mol/mol	COMP	01	0
	Composition'.				
Edition	Refers to the base conditions and the editions of the GPA2145 values.	-	EDITION		2
	Note that the same temperature value is used for the density and				-
	heating value.				
	1: 15°C, 1.01325 bar(a), GPA2145-00				
	2: 15°C, 1.01325 bar(a), GPA2145-03				
	3: 15°C, 1.01325 bar(a), GPA2145-09				
	4: 15°C, 1.01325 bar(a), GPA2145-16				
neo-Pentane mode	Determines what to do when component neo-Pentane is larger than	-	NEOC5_MODE		1
	zero				
	1: Add to i-Pentane				
	2: Add to n-Pentane				
	3: Neglect				

Function outputs

Function inputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: Composition error			COMPERR	
	Composition does not add up to 100% +- 0.01%				
	In case of an error the output values will be set to the				
	fallback values				
Gross Heating Value (Wet)		MJ/m3	VOLGHV_WET		0
Molar Mass (Wet)		kg/kmol	MOLMASS_WET		0
Molar Mass Ratio (Wet)		-	ISG_WET		0
Relative Density (Wet)	Based on the compressibility of wet air	-	RRD_WET		0
Compressibility (Wet)		-	Z_DWET		0
Gross Heating Value (Wet)		MJ/kg	MASGHV_WET		0
Net Heating Value (Wet)		MJ/m3	VOLNHV_WET		0
Gross Heating Value (Dry)		MJ/m3	VOLGHV_DRY		0
Molar Mass (Dry)		kg/kmol	MOLMASS_DRY		0
Molar Mass Ratio (Dry)		-	ISG_DRY		0
Relative Density (Dry)		-	RRD_DRY		0
Compressibility (Dry)		-	Z_DRY		0
Gross Heating Value (Dry)		MJ/kg	MASGHV_DRY		0
Net Heating Value (Dry)		MJ/m3	VOLNHV_DRY		0

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Gross Heating Value (Saturated)	<u> </u>	MJ/m3	VOLGHV_SAT		0
	used for custody transfer energy calculations				
Molar Mass (Saturated)		kg/kmol	MOLMASS_SAT		0
Molar Mass Ratio (Saturated)	Ideal specific gravity	-	ISG_SAT		0
Relative Density (Saturated)	Based on the compressibility of saturated air	-	RRD_SAT		0
Compressibility (Saturated)		-	Z_SAT		0
Gross Heating Value (Saturated)		MJ/kg	MASGHV_SAT		0
Net Heating Value (Saturated)		MJ/m3	VOLNHV SAT		0

Calculations

The calculations are as documented in the GPA-2172 standard using the GPA2145 table values. However the calculations are performed at full precision, so not with intermediate rounding as shown in the examples of the standard.

Please note that the function uses the input composition 'as is', so does <u>not</u> apply normalization to 1.

- When the water fraction input value is above the water fraction of the saturated gas then the function continues its calculations without any correction.
- GPA-2145 standard editions 2000 and 2003 do not specify properties for hydrogen, argon and carbon monoxide. The function processes these components like the other components but with all property values set to 0.

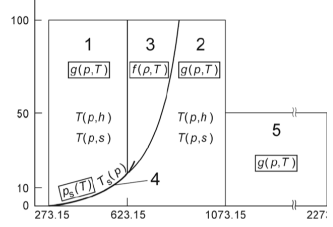
fxIAPWS_IF97_C

Description

The function calculates the density and enthalpy of steam and water according to AIPWS-IF97 in **US Customary units**.

IAPWS-IF97 defines calculations for 5 regions as shown in the picture below.

p/MPa



References

• Revised Release on the IAPWS Industrial Formulation 1997 for the Thermodynamic Properties of Water and Steam, August 2007.

Boundaries

The IAPWS Industrial Formulation 1997 consists of a set of equations for different regions which cover the following range of validity:

- 32 ≤ T ≤ 1472 °F p ≤ 14500 psia
- 1472 ≤ T ≤ 3632 °F p ≤ 7250 psia

Region	1: Water
--------	----------

Region 2: Superheated steam

Region 3: Water

Region 4: Saturation line (saturated steam / water)

Region 5: Superheated steam

Function inputs

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Temperature		°F	04000	0
Pressure		psia	015000	0
Phase	The phase (water or steam) can be calculated automatically or be set to either steam			1
	or water.			
	1: Auto-select (calculate from t and p inputs)			
	2: Steam			
	3: Water			
	If 'Steam' or 'Water' is selected, while the combination of temperature and pressure			
	indicates the opposite phase, then the function uses either the saturation pressure			
	(region 4) or the boundary pressure (intersection regions 2 and 3) instead of the input			
	pressure for its calculations.			

Function outputs

Function outputs	Remark	EU	Fallback
Status	0: Normal		
	1: Input argument out of range		
	2: Calculation error		
	3: No convergence		
Density		lbm/ft3	0
Enthalpy	Energy flow = Mass flow * Enthalpy	btu/lbm	0
Region	Actual IAPWS-IF97 region	-	0
-	0: Combination of t and p is outside the valid range		
	1: Water		
	2: Steam		
	3: Pressurized water		
	4: At the saturation line		
	5: High temperature steam (1472 ≤ T ≤ 3632 °F)		
Saturation pressure	Saturation pressure at the input temperature.	psia	0
	Note: only calculated up to the critical temperature of 647.096 K (+- 705 °F), set to 0 for		
	higher temperatures		
Dynamic viscosity	The dynamic viscosity is required for flow rate calculations based on a differential	lbm/ft.s	0

Function outputs	Remark	EU	Fallback
	pressure measurement (e.g. orifice)		
Ratio of specific heats	Equals the ratio of the specific heats cp / cv	-	0
	cp : specific heat at constant pressure		
	cv : specific heat at constant volume		
	This ratio can be used as the isentropic exponent value (also called 'kappa') when the real value is unknown.		
	The isentropic exponent is required for flow rate calculations based on a differential		
	pressure measurement (e.g. orifice)		

Calculations

The calculations are in compliance with the standard.

For regions 1, 2 and 5 the density can be calculated directly from the temperature in pressure. For region 3 an iterative calculation is required because the Equation Of State for this region calculates the pressure from a known temperature and density iteration. A convergence limit of 0.00001 kg/m3 (+- 0.000006 lbm/ft3) is applied. A maximum of 20 iterations is applied.

The other properties can be calculated directly from the temperature and pressure for all regions.

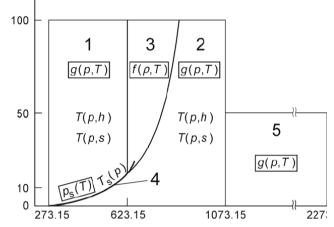
fxIAPWS_IF97_M

Description

The function calculates the density and enthalpy of steam and water according to AIPWS-IF97 in **Metric units**.

IAPWS-IF97 defines calculations for 5 regions as shown in the picture below.

p/MPa



References

• Revised Release on the IAPWS Industrial Formulation 1997 for the Thermodynamic Properties of Water and Steam, August 2007.

Boundaries

The IAPWS Industrial Formulation 1997 consists of a set of equations for different regions which cover the following range of validity:

- 0 ≤ T ≤ 800 °C p ≤ 1000 bar(a)
- 800 ≤ T ≤ 2000 °C p ≤ 500 bar(a)

Region	1:	Water
--------	----	-------

Region 2: Superheated steam

Region 3: Water

Region 4: Saturation line (saturated steam / water)

Region 5: Superheated steam

Function inputs

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Temperature		°C	-50 2200	0
Pressure		bar(a)	01100	0
Phase	The phase (water or steam) can be calculated automatically or be set to either steam or water.			1
	1: Auto-select (calculate from t and p inputs) 2: Steam 3: Water			
	If 'Steam' or 'Water' is selected, while the combination of temperature and pressure indicates the opposite phase, then the function uses either the saturation pressure (region 4) or the boundary pressure (intersection regions 2 and 3) instead of the input pressure for its calculations.			

Function outputs

Function outputs	Remark	EU	Fallback
Status	0: Normal		
	1: Input argument out of range		
	2: Calculation error		
	3: No convergence		
Density		kg/m3	0
Enthalpy	Energy flow = Mass flow * Enthalpy	MJ/kg	0
Region	Actual IAPWS-IF97 region	-	0
	0: Combination of t and p is outside the valid range		
	1: Water		
	2: Steam		
	3: Pressurized water		
	4: At the saturation line		
	5: High temperature steam (800 ≤ T ≤ 2000 °C)		
Saturation pressure	Saturation pressure at the input temperature	bar(a)	0
	Note: only calculated up to the critical temperature of 647.096 K (+- 374 °C), set to 0 for higher		
	temperatures		
Dynamic viscosity	The dynamic viscosity is required for flow rate calculations based on a differential pressure	lbm/ft.s	0
-	measurement (e.g. orifice)		
Ratio of specific heats	Equals the ratio of the specific heats cp / cv	-	0

Function outputs	Remark	EU	Fallback
	cp : specific heat at constant pressure		
	cv : specific heat at constant volume		
	This ratio can be used as the isentropic exponent value (also called 'kappa') when the real value is unknown.		
	The isentropic exponent is required for flow rate calculations based on a differential pressure measurement (e.g. orifice)		

Calculations

The calculations are in compliance with the standard.

For regions 1, 2 and 5 the density can be calculated directly from the temperature in pressure. For region 3 an iterative calculation is required because the Equation Of State for this region calculates the pressure from a known temperature and density iteration. A convergence limit of 0.00001 kg/m3 is applied. A maximum of 20 iterations is applied.

The enthalpy can be calculated directly from the temperature and pressure for all regions.

fxInterpolationCurve

Description

This function calculates the output value from a set of reference points and the actual input value. The function can take an arbitrary number of reference points.

The function is typically used for applying a calibration curve to a K factor or a meter factor (liquid) or a meter error curve (gas). Besides of the 1st calibration point (which is always used) the function will only use the calibration points (starting from the 2nd point) for which the x value is greater than the previous x value. All further points will be ignored.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Input value	Reference to the cell containing the actual input value (e.g. flow rate or pulse				
	frequency)				
Reference values	Array of reference values, assuming the following sequence:		REFVAL		
	Point 1 - Input value				
	Point 1 - Output value				
	Point 2 - Input value				
	Point 2 - Output value				
	etc				
	The array must contain an even number of values with the input values in				
	ascending order.				
	So it is required that Input 1 < Input 2 < Input 3 etc.				
	However, when an input value equals 0, then the function will not use this				
	point and all subsequent points of the array.				
Extrapolation mode	Determines whether or not extrapolation must be applied when the input		EXPMODE		
	value is outside the linearization curve. When disabled either the first or last				
	output value will be used.				
	0: Disabled				
	1: Enabled				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Output value	Interpolated value	Same as input 'Output reference 1'	VAL		0
Date/time	Latest modification date and time of the any of the		DTTM		
	reference point values				
Out of range	Input value is outside the range that is covered by the			OOR	
	reference values				

Calculations

$$y = (x - In_L) \cdot \frac{Out_H - Out_L}{In_H - In_L} + Out_L$$

Where:

х	Input value
у	Interpolated value
ln∟	Closest input reference value that is smaller than the input value
In _H	Closest input reference value that is larger than the input value
Out∟	Output reference value that corresponds with In_L
Out _H	Output reference value that corresponds with $\ensuremath{In_{H}}$

fxISO5167_ISA1932Nozzle

Description

The function calculates the mass flow rate for **ISA1932 Nozzle** pressure differential flow devices according to the ISO-5167 standard.

Compliance

- ISO-5167 1991 Measurement of fluid flow by means of pressure differential devices, 1st edition, 1991
- ISO-5167 Amd.1 : 1998(E)
- ISO-5167 2003 Measurement of fluid flow by means of pressure differential devices inserted in circular cross-section conduits running full, 2nd edition, 2003

Function inputs

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Differential Pressure	Differential pressure over the primary flow device measured at the up- and downstream	mbar		0
	pressure tappings, which need to be in the positions as specified in the standard			
Pressure	Upstream pressure value of the fluid at metering conditions	bar(a)		0
Temperature	Down- or upstream temperature of the fluid at metering conditions	°C		0
Density	Down or upstream density of the fluid at metering conditions	kg/m3		0
Dynamic Viscosity	Dynamic viscosity of the fluid	Pa.s		0
sentropic Exponent	Also referred to as κ (kappa). For an ideal gas this coefficient is equal to the ratio of the			0
	specific heat capacity at constant pressure to the specific heat at constant volume.			
	According to the ISO standard this ratio may be used, when the real value is unknown.			
Pipe Diameter	Internal diameter of the pipe at reference temperature	mm		0
Pipe Expansion factor	The thermal expansion coefficient of the pipe material	1/°C		0.0000108
Pipe Reference temperature	The reference temperature that corresponds to the 'Pipe diameter' input value	°C		20
SA1932 Nozzle Diameter	ISA1932 Nozzle diameter at reference temperature	mm		0
SA1932 Nozzle Expansion factor	The thermal expansion coefficient of the ISA1932 Nozzle material	1/°C		0.0000163
SA1932 Nozzle Reference Femperature	The reference temperature that corresponds to the 'ISA1932 Nozzle diameter' input value	°C		20
Pressure Location	1: Upstream	-		1
	Input 'Pressure' represents the pressure at the upstream pressure tapping (p1). Since the absolute pressure is usually measured at the upstream tapping this is the most common setting. 2: Downstream			-
	Input 'Pressure' represents the pressure at the downstream tapping (p $_2$).			
Temperature Location	1: Upstream	-		2
	Input 'Temperature' represents the upstream temperature (t $_1$).			
	2: Downstream			
	Input 'Temperature represents the temperature at the downstream tapping (t_2).			
	3: Recovered Input 'Temperature' represents the downstream temperature at a location Where the pressure has fully recovered (t ₃). Since temperature measurement is usually downstream of the flow device this is the most common setting.			
Femperature Correction	This parameter specifies how the temperature should be corrected from downstream to upstream conditions (or vice versa)			3
	1: $(1-\kappa)/\kappa$ Isentropic expansion using $(1-\kappa)/\kappa$ as the temperature referral exponent			
	2: Constant Isentropic expansion using input 'Temperature Exponent' as the temperature referral exponent [-].			
	Please note that this value must be < 0			
	3: Joule Thomson			
	Isenthalpic expansion using input 'Temperature Exponent' as the Joule Thomson coefficient [°C/bar].			
	This method is prescribed by ISO5167-1:2003.			
emperature Exponent	Refer to input Temperature Correction	-		0
	Unit depends on input Temperature Correction value	°C/bar		
Density Location	This parameter specifies if and how the density should be corrected from downstream to upstream conditions (or vice versa).	-		1
	1: Upstream			
	Input 'Density' represents the density at the upstream pressure tapping ($ ho_1$).			
	2: Downstream			
	Input 'Density' represents the density at the downstream tapping (ρ_2). 3: Recovered			
	Input 'Density' represents the density downstream at a location Where the pressure has fully recovered (ρ_3).			
				0
Density Exponent.	This factor is used when density correction is enabled. The formula $1/\kappa$ will be used when the input value is set to 0, else the input value will be used.	-		0
Density Exponent. Fluid	This factor is used when density correction is enabled. The formula $1/\kappa$ will be used when	-		1

Function inputs	Remark	EU	Range	Default
	2: Liquid			
Year Of Edition	1: Edition 1991	-		3
	2: Edition 1998			
	3: Edition 2003			
	Only used for calculation of pressure loss with options 1 and 2 giving the same result			

Function outputs Function outputs Remark EU Status 0: Normal (No error condition) 1: Input argument out of range 2: No convergence Mass flow rate The calculated mass flow rate tonne/h Beta ratio Nozzle to pipe ratio at upstream temperature Nozzle diameter At the upstream temperature mm Pipe diameter At the upstream temperature mm Upstream pressure Pressure at upstream tapping (p1) bar(a) Pressure at downstream tapping Pressure at downstream tapping (p₂) bar(a) Recovered downstream pressure Fully recovered downstream pressure (p₃) bar(a) Upstream temperature Temperature at upstream tapping (t1) °C Temperature at downstream tapping °C Temperature at downstream tapping (t₂) °C 'Fully recovered' downstream temperature (t₃) Downstream Temperature Upstream density Density at upstream tapping (ρ_1) kg/m3 Density at downstream tapping Pressure at downstream tapping (ρ_2) kg/m3 Downstream density 'Fully recovered' downstream density (ρ_3) kg/m3 The pipe Reynolds number (this is the Reynolds number upstream of the ISA1932 Nozzle Reynolds number and not the one within the device throat itself Discharge coefficient -**Expansion Factor** -Velocity of Approach 0: Pressure is in valid range Pressure Range _ 1: Pressure is out of valid range Reynolds Range 0: Reynolds number is in valid range _ 1: Reynolds number is out of valid range Diameter Range 0: Device and pipe diameter and Beta ratio in valid range -1: Device diameter, pipe diameter and/or Beta ratio out of valid range

fxISO5167_LongRadiusNozzle

Description

The function calculates the mass flow rate for **Long Radius Nozzle** pressure differential flow devices according to the ISO-5167 standard.

Compliance

ISO-5167 - 1991 Measurement of fluid flow by means of pressure differential devices, 1st edition, 1991 ISO-5167 Amd.1 : 1998(E) ISO-5167 - 2003 Measurement of fluid flow by means of pressure differential devices inserted in circular cross-section conduits running full, 2nd edition, 2003

Function inputs

Function inputs	Remark	EU Range	Default
Name	Optional tag name, tag description and tag group		
Differential Pressure	Differential pressure over the primary flow device measured at the up- and downstream pressure tappings, which need to be in the positions as specified in the standard	mbar	0
Pressure	Upstream pressure value of the fluid at metering conditions	bar(a)	0
Temperature	Down- or upstream temperature of the fluid at metering conditions	°C	0
Density	Down or upstream density of the fluid at metering conditions	kg/m3	0
Dynamic Viscosity	Dynamic viscosity of the fluid	Pa.s	0
Isentropic Exponent	Also referred to as κ (kappa). For an ideal gas this coefficient is equal to the ratio	-	0
	of the specific heat capacity at constant pressure to the specific heat at constant		
	volume. According to the ISO standard this ratio may be used, when the real value		
	is unknown.		
Pipe Diameter	Internal diameter of the pipe at reference temperature	mm	0
Pipe Expansion factor	The thermal expansion coefficient of the pipe material	1/°C	0.0000108
Pipe Reference temperature	The reference temperature that corresponds to the 'Pipe diameter' input value	°C	20
Long Radius Nozzle Diameter	Long Radius Nozzle diameter at reference temperature	mm	0
Long Radius Nozzle Expansion	The thermal expansion coefficient of the Long Radius Nozzle material	1/°C	0.0000163
factor			
Long Radius Nozzle Reference	The reference temperature that corresponds to the 'Long Radius Nozzle diameter'	°C	20
Temperature	input value		
Pressure Location	1: Upstream	-	1
	Input 'Pressure' represents the pressure at the upstream pressure tapping (p_1) .		
	Since the absolute pressure is usually measured at the upstream tapping this is		
	the most common setting.		
	2: Downstream		
	Input 'Pressure' represents the pressure at the downstream tapping (p ₂).		
Temperature Location	1: Upstream	-	2
	Input 'Temperature' represents the upstream temperature (t1).		
	2: Downstream		
	Input 'Temperature represents the temperature at the downstream tapping		
	(t₂). 3: Recovered		
	Input 'Temperature' represents the downstream temperature at a location		
	Where the pressure has fully recovered (t_3) .		
	Since temperature measurement is usually downstream of the flow device this		
	is the most common setting.		
Temperature Correction	This parameter specifies how the temperature should be corrected from		3
	downstream to upstream conditions (or vice versa)		5
	1: (1-κ)/κ		
	Isentropic expansion using $(1-\kappa)/\kappa$ as the temperature referral exponent		
	2: Constant		
	Isentropic expansion using input 'Temperature Exponent' as the temperature		
	referral exponent [-].		
	Please note that this value must be < 0		
	3: Joule Thomson		
	Isenthalpic expansion using input 'Temperature Exponent' as the Joule		
	Thomson coefficient [°C/bar].		
	This method is prescribed by ISO5167-1:2003.		
Temperature Exponent	Refer to input Temperature Correction	-	0
	Unit depends on input Temperature Correction value	°C/bar	
Density Location	This parameter specifies if and how the density should be corrected from	-	1
	downstream to upstream conditions (or vice versa).		
	1: Upstream		
	Input 'Density' represents the density at the upstream pressure tapping (ρ_1).		
	2: Downstream		
	Input 'Density' represents the density at the downstream tapping (ρ_2).		
	3: Recovered		
	Input 'Density' represents the density downstream at a location Where the		
	pressure has fully recovered (ρ_3).		
Density Exponent.	This factor is used when density correction is enabled. The formula $1/\kappa$ will be	-	0
	used when the input value is set to 0, else the input value will be used.		

Function inputs	Remark	EU	Range	Default
	For more details refer to section 'Density correction'.			
Fluid	The type of fluid being measured	-		1
	1: Gas			
	2: Liquid			
Year Of Edition	1: Edition 1991	-		3
	2: Edition 1998			
	3: Edition 2003			
	Only used for calculation of pressure loss, with options 1 and 2 giving the same	9		
	result			

Function outputs

Function outputs	Remark	EU
Status	0: Normal (No error condition)	-
	1: Input argument out of range	
	2: No convergence	
Mass flow rate	The calculated mass flow rate	tonne/h
Beta ratio	Nozzle to pipe ratio at upstream temperature	
Nozzle diameter	At the upstream temperature	
Pipe diameter	At the upstream temperature	
Upstream pressure	Pressure at upstream tapping (p1)	bar(a)
Pressure at downstream tapping	Pressure at downstream tapping (p2)	bar(a)
Recovered downstream pressure	Fully recovered downstream pressure (p3)	bar(a)
Upstream temperature	Temperature at upstream tapping (t1)	°C
Temperature at downstream tapping	Temperature at downstream tapping (t₂)	°C
Downstream Temperature	'Fully recovered' downstream temperature (t_3)	°C
Upstream density	Density at upstream tapping (ρ_1)	kg/m3
Density at downstream tapping	Pressure at downstream tapping (p2)	kg/m3
Downstream density	'Fully recovered' downstream density (ρ₃)	kg/m3
Reynolds number	The pipe Reynolds number (this is the Reynolds number upstream of the Long Radius	-
	Nozzle and not the one within the device throat itself	
Discharge coefficient		-
Expansion Factor		-
Velocity of Approach		
Pressure Range	0: Pressure is in valid range	-
	1: Pressure is out of valid range	
Reynolds Range	0: Reynolds number is in valid range	-
	1: Reynolds number is out of valid range	
Diameter Range	0: Device and pipe diameter and Beta ratio in valid range	-
	1: Device diameter, pipe diameter and/or Beta ratio out of valid range	

fxISO5167_Orifice

Description

The function calculates the mass flow rate for **Orifice** pressure differential flow devices according to the ISO-5167 standard and the ISO/TR 15377 technical report.

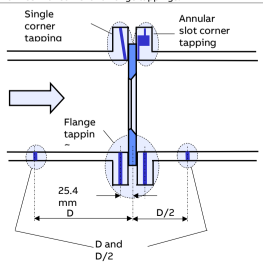
Orifices plates with the following type of pressure tappings are supported by this function:

- ISO5167 Square-edged orifice with corner tappings
- ISO5167 Square-edged orifice with flange tappings
- ISO5167 Square-edged orifice with D and D/2 tappings
- ISO/TR 15377 Quarter circle orifice with corner tappings
- ISO/TR 15377 Quarter circle orifice with flange tappings
- ISO/TR 15377 Conical entrance orifice with corner tappings

References

- ISO 5167 Measurement of fluid flow by means of pressure differential devices, 1991
- ISO 5167 Measurement of fluid flow by means of pressure differential devices, 1991
- Amendment 1 1998
- ISO 5167 Measurement of fluid flow by means of pressure differential devices, 2003
- British standard 1042: Part 1: 1964
- ISO/TR 15377:2007 Measurement of fluid flow by means of pressure differential devices – Guidelines for the specification of orifice plates, nozzle and Venturi tubes beyond the scope of ISO 5167

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Differential Pressure	Differential pressure over the primary flow device measured at the up- and downstream	mbar	010000	0
	pressure tappings, which need to be in the positions as specified in the standard			
Pressure	Down- or upstream pressure value (p1) of the fluid at metering conditions	bar (a)	02000	0
Temperature	Down- or upstream temperature of the fluid at metering conditions	°C	-2401000	0
Density	Down or upstream density of the fluid at metering conditions	kg/m3	02000	0
Dynamic Viscosity	Dynamic viscosity of the fluid	Pa.s	01	0
Isentropic Exponent	Also referred to as κ (kappa). For an ideal gas this coefficient is equal to the ratio of the	-	010	0
	specific heat capacity at constant pressure to the specific heat at constant volume.			
	According to the ISO standard this ratio may be used, when the real value is unknown.			
Pipe Diameter	Internal diameter of the pipe at reference temperature	mm	02000	0
Pipe Expansion factor	The thermal expansion coefficient of the pipe material	1/°C	01	0.0000108
Pipe Reference	The reference temperature that corresponds to the 'Pipe diameter' input value	°C	-3001000	20
temperature				
Orifice Diameter	Orifice diameter at reference temperature	mm	02000	0
Orifice Expansion factor	The thermal expansion coefficient of the orifice material	1/°C	01	0.0000163
Orifice Reference Temperature	The reference temperature that corresponds to the 'Orifice diameter' input value	°C	-3001000	20
Configuration	The location of the pressure tappings. Several configurations are permitted by the	-		2
	ISO5167 standard. Each configuration has a different calculation of the discharge			
	coefficient and of the expansion factor			
	1: ISO5167 Corner			
	2: ISO5167 D and D/2			
	3: ISO5167 Flange			
	4: ISO15377 Quarter circle (*)			
	5: ISO15377 Conical entrance			
	(*) The calculation of the discharge and expansion factor are equal for the quarter circle orifice with corner and flange tappings.			



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Function inputs	Remark	EU	Range	Default
Pressure Location	1: Upstream	-		1
	Input 'Pressure' represents the pressure at the upstream pressure tapping (p $_1$).			
	Since the absolute pressure is usually measured at the upstream tapping this is the			
	most common setting.			
	2: Downstream			
	Input 'Pressure' represents the pressure at the downstream tapping (p ₂).			
Temperature Location	1: Upstream	-		2
	Input 'Temperature' represents the upstream temperature (t ₁).			
	2: Downstream			
	Input 'Temperature represents the temperature at the downstream tapping (t_2) .			
	3: Recovered			
	Input 'Temperature' represents the downstream temperature at a location Where the			
	pressure has fully recovered (t_3).			
	Since temperature measurement is usually downstream of the flow device this is the			
	most common setting.			
emperature Correction	This parameter specifies how the temperature should be corrected from downstream to			3
	upstream conditions (or vice versa)			
	1: $(1-\kappa)/\kappa$			
	Isentropic expansion using $(1-\kappa)/\kappa$ as the temperature referral exponent 2: Constant			
	Isentropic expansion using input 'Temperature Exponent' as the temperature referral exponent [-].			
	Please note that this value must be < 0			
	3: Joule Thomson			
	Isenthalpic expansion using input 'Temperature Exponent' as the Joule Thomson			
	coefficient [°C/bar].			
	This method is prescribed by ISO5167-1:2003.			
remperature Exponent	Refer to input Temperature Correction	_		0
	Unit depends on input Temperature Correction value	°C/bar		0
Density Location	This parameter specifies if and how the density should be corrected from downstream to	-		1
Density Location	upstream conditions (or vice versa).	-		1
	1: Upstream			
	Input 'Density' represents the density at the upstream pressure tapping (ρ_1).			
	2: Downstream			
	Input 'Density' represents the density at the downstream tapping (ρ_2).			
	3: Recovered			
	Input 'Density' represents the density downstream at a location Where the pressure has			
	fully recovered (ρ_3).			
Density Exponent.	This factor is used when density correction is enabled. The formula $1/\kappa$ will be used when	-		0
sensity Exponent.	the input value is set to 0, else the input value will be used.			Ū
	For more details refer to section 'Density correction'.			
luid	The type of fluid being measured	-		1
	1: Gas			-
	2: Liquid			
'ear Of Edition	1: Edition 1991	-		3
	2: Edition 1998			5
	3: Edition 2003			
	Note: Only applicable for ISO5167. ISO/TR15377 refers to ISO5167:2003.			
Drain hole	When input value is > 0 then an additional correction on the orifice diameter will be	mm		0
	applied to account for the drain hole, as explained further on.			~

Function outputs	Remark	EU	Fallback
Status	0: Normal (No error condition)		
	1: Input argument out of range		
	2: No convergence		
Mass flow rate	The calculated mass flow rate	kg/s	0
Beta ratio	Orifice to pipe diameter ratio at upstream temperature	-	0
Orifice diameter	At the upstream temperature and optionally with a correction for the drain hole	mm	0
Pipe diameter	At the upstream temperature	mm	0
Upstream pressure	Pressure at upstream tapping (p1)	bar(a)	0
Pressure at downstream tapping	Pressure at downstream tapping (p2)	bar(a)	0
Recovered downstream pressure	Fully recovered downstream pressure (p₃)	bar(a)	0
Upstream temperature	Temperature at upstream tapping (t1)	°C	0
Temperature at downstream	Temperature at downstream tapping (t₂)	°C	0
tapping			
Downstream Temperature	'Fully recovered' downstream temperature (t ₃)	°C	0
Upstream density	Density at upstream tapping (p1)	kg/m3	0
Density at downstream tapping	Pressure at downstream tapping (p2)	kg/m3	0
Downstream density	'Fully recovered' downstream density (ρ₃)	kg/m3	0
Reynolds number	The pipe Reynolds number (this is the Reynolds number upstream of the orifice and not	-	0
-	the one within the device throat itself		

Function outputs	Remark	EU	Fallback
Discharge coefficient		-	0
Expansion Factor		-	0
Velocity of Approach		-	0
Pressure Range	0: Pressure is in valid range	-	0
	1: Pressure is out of valid range		
Reynolds Range	0: Reynolds number is in valid range	-	0
	1: Reynolds number is out of valid range		
Diameter Range	0: Device and pipe diameter and Beta ratio in valid range	-	0
-	1: Device diameter, pipe diameter and/or Beta ratio out of valid range		
Dressing compation			

Pressure correction

The relation between the pressure at the upstream tapping p_1 and the pressure at the downstream tapping (p_2) is as following:

 $p_2 = p_1 - \Delta p / 1000$

The relation between the pressure at the upstream tapping and the fully recovered pressure (p3) is as following:

$$p_3 = p_1 - p_{LOSS}$$

The calculation of P_{LOSS} is as defined in the standard.

Where:

p1	Pressure at upstream tapping	[bar(a)]
p2	Pressure at downstream tapping	[bar(a)]
р3	Fully recovered downstream pressure	[bar(a)]
Δр	Differential pressure	[mbar]
pLOSS	Pressure loss over the meter	[bar]

Temperature correction

When input 'Temperature correction' is set to 1, then an <u>isentropic</u> expansion based on the isentropic coefficient is applied:

$$t_{2} = (t_{3} + 273.15) \cdot \left(\frac{p_{3}}{p_{2}}\right)^{\frac{1-\kappa}{\kappa}} - 273.15$$
$$t_{1} = (t_{3} + 273.15) \cdot \left(\frac{p_{3}}{p_{1}}\right)^{\frac{1-\kappa}{\kappa}} - 273.15$$

When input 'Temperature correction' is set to 2, then an <u>isentropic</u> expansion based on input 'Temperature exponent' is applied:

$$t_{2} = (t_{3} + 273.15) \cdot \left(\frac{p_{3}}{p_{2}}\right)^{K_{TE}} - 273.15$$
$$t_{1} = (t_{3} + 273.15) \cdot \left(\frac{p_{3}}{p_{1}}\right)^{K_{TE}} - 273.15$$

When input 'Temperature correction' is set to 3, then an <u>isenthalpic</u> expansion based on the linear Joule Thomson correction as defined in ISO5167-1:2003, taking input 'Temperature exponent' as the Joule Thomson coefficient:

$$t_1 = t_2 + (p_1 - p_2) \cdot \mu_{JT}$$
 $t_1 = t_3 + (p_1 - p_3) \cdot \mu_{JT}$

Where:			
t1	Upstream temperature	°C	
t ₃	Downstream temperature	°C	
p1	Upstream pressure	bar(a)	
рз	Fully recovered downstream pressure	bar(a)	
κ	Isentropic exponent	-	
Кте	Temperature exponent	-	
μյτ	Joule Thomson coefficient	°C/bar	

ISO-5167 edition 2003 prescribes an <u>isenthalpic</u> expansion instead of an isentropic expansion. This can be achieved by assigning a fixed Joule Thomson coefficient to input 'Temperature Exponent'.

Note: ISO is working a method to calculate the Joule Thomson rather than using a fixed value.

Density correction

When input 'Density exponent' = 0, then the following <u>isentropic</u> corrections are applied (depending on the type of Density Correction)

$$\rho_1 = \rho_2 \cdot \left(\frac{p_1}{p_2}\right)^{\frac{1}{\kappa}} \qquad \qquad \rho_1 = \rho_3 \cdot \left(\frac{p_1}{p_3}\right)^{\frac{1}{\kappa}}$$

Else the value of input 'Density Exponent' is used

$$\rho_1 = \rho_2 \cdot \left(\frac{p_1}{p_2}\right)^{K_{DE}} \qquad \qquad \rho_1 = \rho_3 \cdot \left(\frac{p_1}{p_3}\right)^{K_{DE}}$$

Where:

ρ1	Upstream density	[kg/m3]
ρ2	Density at the downstream tapping	[kg/m3]
ρ3	Density at the fully recovered downstream pressure	[kg/m3]
p1	Upstream pressure	[bar(a)]
p2	Pressure at the downstream tapping	[bar(a)]
р3	Fully recovered downstream pressure	[bar(a)]
κ	Isentropic exponent	[-]
KDE	Density exponent	[-]

Note: In March 2007 the British DTI (Department of Trade and Industry) has recommended that the density correction method should not be based on isentropic expansion but on <u>isenthalpic</u> expansion instead.

The correction assumes that the density is measured at p_2 (downstream pressure tapping pressure) and t_3 (downstream recovered temperature). For this situation the following density correction is defined:

$$\rho_1 = \rho_M \cdot \frac{p_1 \cdot t_3 \cdot Z(p_2, t_3)}{p_2 \cdot t_1 \cdot Z(p_1, t_1)}$$

Upstream density	[kg/m3]
Measured density from the densitometer at p2 and t3	[kg/m3]
Upstream pressure	[bar(a)]
Pressure at the downstream tapping	[bar(a)]
Upstream temperature	°C
Temperature at downstream side Where pressure has fully recovered	°C
Compressibility at p1 and t1	[-]
Compressibility at p2 and t3	[-]
	Measured density from the densitometer at p2 and t3 Upstream pressure Pressure at the downstream tapping Upstream temperature Temperature at downstream side Where pressure has fully recovered Compressibility at p1 and t1

When required, this correction should be applied outside the ISO5167 function by defining an additional compressibility calculation (e.g. AGA8) at p_2 and t_3 in the application and calculating the upstream density, Where the value of p_2 is provided as an output by the ISO5167 function. Subsequently the upstream density is calculated and fed into the ISO5167 function (input 'Density') with input 'Density Correction' set to 'Upstream density'.

Correction for drain hole

When input 'Drain hole' is > 0 then the following correction factor is applied on the orifice diameter according to the British standard 1042: Part 1: 1964 and ISO/TR 15377:2007

$$C_{DH} = 1 + 0.55 * \left(\frac{d_{DH}}{d_0}\right)^2$$

CDH	Darin hole correction factor on orifice diameter	[-]
dDH	Drain hole diameter	[mm]
d0	Orifice diameter at reference temperature (i.e. input	[mm]
	'Orifice diameter')	

fxISO5167_Venturi

Description

The function calculates the mass flow rate for **classical Venturi** tube pressure differential flow devices according to the ISO-5167 standard.

Compliance

ISO-5167 - 1991 Measurement of fluid flow by means of pressure differential devices, 1st edition, 1991 ISO-5167 Amd.1 : 1998(E) ISO-5167 - 2003 Measurement of fluid flow by means of pressure differential devices inserted in circular cross-section conduits running full, 2nd edition, 2003

Function inputs

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Differential Pressure	Differential pressure over the primary flow device measured at the up- and downstream	mbar		0
	pressure tappings, which need to be in the positions as specified in the standard			
Pressure	Upstream pressure value of the fluid at metering conditions	bar(a)		0
Temperature	Down- or upstream temperature of the fluid at metering conditions	°C		0
Density	Down or upstream density of the fluid at metering conditions	kg/m3		0
Dynamic Viscosity	Dynamic viscosity of the fluid	Pa.s		0
Isentropic Exponent	Also referred to as κ (kappa). For an ideal gas this coefficient is equal to the ratio of the specific	-		0
	heat capacity at constant pressure to the specific heat at constant volume. According to the			
	ISO standard this ratio may be used, when the real value is unknown.			
Pipe Diameter	Internal diameter of the pipe at reference temperature	mm		0
Pipe Expansion factor	The thermal expansion coefficient of the pipe material	1/°C		0.0000108
Pipe Reference	The reference temperature that corresponds to the 'Pipe diameter' input value	°C		20
temperature	· · · · · · · · · · · · · · · · · · ·			
Venturi Diameter	Venturi diameter at reference temperature	mm		0
Venturi Expansion factor	The thermal expansion coefficient of the Venturi material	1/°C		0.0000163
Venturi Reference	The reference temperature that corresponds to the 'Venturi diameter' input value	°C		20
Temperature	The reference temperature that corresponds to the ventur diameter input value	C		20
Configuration	The type of classical venturi tube.			2
configuration	Three configurations are permitted by the ISO5167 standard. Each configuration has a different	-		2
	calculation of the discharge coefficient and of the expansion factor			
	1: As cast convergent section			
	2: Rough welded 3: Machined			
	4: User-defined (not according to the standard!) When 'User-defined' is selected then the input 'Discharge coefficient' will be used in the			
Durana i a anti-u	calculations instead.			
Pressure Location	1: Upstream	-		1
	Input 'Pressure' represents the pressure at the upstream pressure tapping (p1).			
	Since the absolute pressure is usually measured at the upstream tapping this is the most			
	common setting.			
	2: Downstream			
T	Input 'Pressure' represents the pressure at the downstream tapping (p2).			2
Temperature Location	1: Upstream	-		2
	Input 'Temperature' represents the upstream temperature (t1).			
	2: Downstream			
	Input 'Temperature represents the temperature at the downstream tapping (t_2).			
	3: Recovered			
	Input 'Temperature' represents the downstream temperature at a location Where the			
	pressure has fully recovered (t_3).			
	Since temperature measurement is usually downstream of the flow device this is the most			
	common setting.			3
Temperature Correction	This parameter specifies how the temperature should be corrected from downstream to			5
	upstream conditions (or vice versa)			
	1: $(1-\kappa)/\kappa$			
	Isentropic expansion using $(1-\kappa)/\kappa$ as the temperature referral exponent			
	2: Constant			
	Isentropic expansion using input 'Temperature Exponent' as the temperature referral			
	exponent [-].			
	Please note that this value must be < 0			
	3: Joule Thomson			
	Isenthalpic expansion using input 'Temperature Exponent' as the Joule Thomson coefficient			
	[°C/bar].			
	This method is prescribed by ISO5167-1:2003.			
Temperature Exponent	Refer to input Temperature Correction	-		0
	Unit depends on input Temperature Correction value	°C/bar		
Density Location	This parameter specifies if and how the density should be corrected from downstream to	-		1
	upstream conditions (or vice versa).			
	1: Upstream			
	Input 'Density' represents the density at the upstream pressure tapping (ρ_1).			
	2: Downstream			

Function inputs	Remark	EU	Range	Default
	Input 'Density' represents the density at the downstream tapping (ρ_2).			
	3: Recovered			
	Input 'Density' represents the density downstream at a location Where the pressure has fully recovered (ρ_3).			
Density Exponent.	This factor is used when density correction is enabled. The formula $1/\kappa$ will be used when the input value is set to 0, else the input value will be used.	-		0
	For more details refer to section 'Density correction'.			
Fluid	The type of fluid being measured	-		1
	1: Gas			
	2: Liquid			
Pressure Loss Mode	The method for determining the pressure loss	-		1
	1: Absolute value in mbar The value of input 'Pressure Loss Value' is taken as a value in mbar			
	2: Percentage of differential pressure			
	The value of input 'Pressure Loss Value' is taken as a percentage from input 'Differential Pressure'			
Pressure Loss Value	Value in mbar or %, depending on the 'Pressure Loss Mode'.	mbar		0
	The pressure loss over the Venturi is used to calculate the downstream fully recovered pressure.	%		
	The pressure loss equals the difference between the upstream pressure (p_1) and the fully			
	recovered downstream pressure (p3)			
	The standard prescribes that only the pressure loss that is caused by the venturi tube should be			
	included (so it should not include the pressure loss that occurred between the two pressure			
	tappings before the venturi tube was installed).			
Discharge coefficient	This value will used instead of the discharge coefficient as specified in the standard.	-		0
	Only used when input 'Configuration' is set to 'User-defined'.			

Function outputs

Function outputs	Remark	EU	Fallback
Status	0: Normal (No error condition)		
	1: Input argument out of range		
	2: No convergence		
Mass flow rate	The calculated mass flow rate	tonne/h	0
Beta ratio	Venturi to pipe diameter ratio at upstream temperature	-	0
Venturi diameter	At the upstream temperature	mm	0
Pipe diameter	At the upstream temperature	mm	0
Upstream pressure	Pressure at upstream tapping (p1)	bar(a)	0
Pressure at downstream tapping	Pressure at downstream tapping (p2)	bar(a)	0
Recovered downstream pressure	Fully recovered downstream pressure (p3)	bar(a)	0
Upstream temperature	Temperature at upstream tapping (t1)	°C	0
Temperature at downstream	Temperature at downstream tapping (t2)	°C	0
tapping			
Downstream Temperature	'Fully recovered' downstream temperature (t3)	°C	0
Upstream density	Density at upstream tapping (p1)	kg/m3	0
Density at downstream tapping	Pressure at downstream tapping (ρ_2)	kg/m3	0
Downstream density	'Fully recovered' downstream density ($ ho_3$)	kg/m3	0
Reynolds number	The pipe Reynolds number (this is the Reynolds number upstream of the Venturi and not	-	0
	the one within the device throat itself		
Discharge coefficient		-	0
Expansion Factor		-	0
Velocity of Approach			0
Pressure Range	0: Pressure is in valid range	-	0
	1: Pressure is out of valid range		
Reynolds Range	0: Reynolds number is in valid range	-	0
	1: Reynolds number is out of valid range		
Diameter Range	0: Device and pipe diameter and Beta ratio in valid range	-	0
	1: Device diameter, pipe diameter and/or Beta ratio out of valid range		

fxISO5167_VenturiNozzle

Description

The function calculates the mass flow rate for **Venturi Nozzle** pressure differential flow devices according to the ISO-5167 standard.

Compliance

ISO-5167 - 1991 Measurement of fluid flow by means of pressure differential devices, 1st edition, 1991 ISO-5167 Amd.1 : 1998(E) ISO-5167 - 2003 Measurement of fluid flow by means of pressure differential devices inserted in circular cross-section conduits running full, 2nd edition, 2003

Function inputs

Function inputs	Remark	EU	Default
Name	Optional tag name, tag description and tag group		
Differential Pressure	Differential pressure over the primary flow device measured at the up- and downstream pressure tappings, which need to be in the positions as specified in the standard	mbar	0
Pressure	Upstream pressure value of the fluid at metering conditions	bar(a)	0
Temperature	Down- or upstream temperature of the fluid at metering conditions	°C	0
Density	Down or upstream density of the fluid at metering conditions	kg/m3	0
Dynamic Viscosity	Dynamic viscosity of the fluid	Pa.s	0
sentropic Exponent	Also referred to as κ (kappa). For an ideal gas this coefficient is equal to the ratio of the specific heat		0
	capacity at constant pressure to the specific heat at constant volume. According to the ISO standard this ratio may be used, when the real value is unknown.		
Pipe Diameter	Internal diameter of the pipe at reference temperature	mm	0
Pipe Expansion factor	The thermal expansion coefficient of the pipe material	1/°C	0.0000108
Pipe Reference	The reference temperature that corresponds to the 'Pipe diameter' input value	°C	20
temperature	The reference temperature that corresponds to the ripe diameter input value	C	20
Venturi Nozzle Diameter	Venturi Nozzle diameter at reference temperature	mm	0
Venturi Nozzle Expansion	The thermal expansion coefficient of the Venturi Nozzle material	1/°C	0.0000163
actor	·		
Venturi Nozzle Reference Temperature	The reference temperature that corresponds to the 'Venturi Nozzle diameter' input value	°C	20
Pressure Location	 Input 'Pressure' represents the pressure at the upstream pressure tapping (p1). Since the absolute pressure is usually measured at the upstream tapping this is the most common setting. 	-	1
	2 Input 'Pressure' represents the pressure at the downstream tapping (p ₂).		
remperature Location	1 Input 'Temperature' represents the upstream temperature (t1).	-	2
	2 Input 'Temperature represents the temperature at the downstream tapping (t ₂).		
	 Input 'Temperature' represents the downstream temperature at a location Where the pressure has fully recovered (t₃). Since temperature measurement is usually downstream of the flow device this is the most common setting. 		
Temperature Correction	1 Isentropic expansion using $(1-\kappa)/\kappa$ as the temperature referral exponent		3
·	2 Isentropic expansion using input 'Temperature Exponent' as the temperature referral exponent [-		
	 Isenthalpic expansion using input 'Temperature Exponent' as the Joule Thomson coefficient [°C/bar]. This method is prescribed by ISO5167-1:2003. 		
Temperature Exponent	Refer to input Temperature Correction	-	0
		°C/bar	0
Density Location	This parameter specifies if and how the density should be corrected from downstream to upstream conditions.	-	1
	 Input 'Density' represents the density at the upstream pressure tapping (ρ1). 		
	 2 Input 'Density' represents the density at the downstream tapping (p1). 2 Input 'Density' represents the density at the downstream tapping (p2). 		
	3 Input 'Density' represents the density downstream at a location Where the pressure has fully		
	recovered (ρ ₃).		
Density Exponent.	This factor is used when density correction is enabled. The formula $1/\kappa$ will be used when the input value is set to 0, else the input value will be used.	-	0
	For more details refer to function 'ISO5167 - Orifice' section 'Density correction'		
Fluid	The type of fluid being measured 1: Gas	-	1
	2: Liquid		
Pressure Loss Mode	The method for determining the pressure loss	-	1
	 Absolute value in mbar The value of input 'Pressure Loss Value' is taken as a value in mbar 		
	2 Percentage of differential pressure The value of input 'Pressure Loss Value' is taken as a percentage from input 'Differential Pressure'		
Pressure Loss Value	Value in mbar or %, depending on the 'Pressure Loss Mode'.	mbar	0
	The pressure loss over the Venturi nozzle is used to calculate the downstream fully recovered	%	v
	pressure. The pressure loss equals the difference between the upstream pressure (p_1) and the fully	70	
	recovered downstream pressure (p_3) The standard prescribes that only the pressure loss that is caused by the venturi nozzle should be		
	included (so it should not include the pressure loss that occurred between the two pressure tappings		

Function inputs	Remark	EU	Default
	before the venturi nozzle was installed).		

Function outputs

Function outputs	Remark	EU	Fallback
Status	0: Normal (No error condition)		
	1: Input argument out of range		
	2: No convergence		
Mass flow rate	The calculated mass flow rate	tonne/h	0
Beta ratio	Venturi nozzle to pipe diameter ratio at upstream temperature		0
Venturi Nozzle diameter	At the upstream temperature		0
Pipe diameter	At the upstream temperature		0
Upstream pressure	Pressure at upstream tapping (p1)	bar(a)	0
Pressure at downstream tapping	Pressure at downstream tapping (p2)	bar(a)	0
Recovered downstream pressure	Fully recovered downstream pressure (p3)	bar(a)	0
Upstream temperature	Temperature at upstream tapping (t1)	°C	0
Temperature at downstream	Temperature at downstream tapping (t₂)	°C	0
tapping			
Downstream Temperature	'Fully recovered' downstream temperature (t_3)	°C	0
Upstream density	Density at upstream tapping (ρ1)	kg/m3	0
Density at downstream tapping	Pressure at downstream tapping (ρ₂)	kg/m3	0
Downstream density	'Fully recovered' downstream density (ρ_3)	kg/m3	0
Reynolds number	The pipe Reynolds number (this is the Reynolds number upstream of the Venturi	-	0
	nozzle and not the one within the device throat itself		
Discharge coefficient		-	0
Expansion Factor		-	0
Velocity of Approach			0
Pressure Range	0: Pressure is in valid range	-	0
	1: Pressure is out of valid range		
Reynolds Range	0: Reynolds number is in valid range	-	0
	1: Reynolds number is out of valid range		
Diameter Range	0: Device and pipe diameter and Beta ratio in valid range	-	0
	1: Device diameter, pipe diameter and/or Beta ratio out of valid range		

Temperature correction

When input "Temperature exponent' = 0, then an isentropic expansion is applied:

$$t_1 = (t_3 + 273.15) \cdot \left(\frac{p_3}{p_1}\right)^{\frac{1-\kappa}{\kappa}} - 273.15$$

Else the value of input 'Temperature exponent' is used:

$$t_1 = (t_3 + 273.15) \cdot \left(\frac{p_3}{p_1}\right)^{k_{TE}} - 273.15$$

t1	Upstream temperature	[°C]
t₃	Downstream temperature	[°C]
p1	Upstream pressure	[bar(a)]
р₃	Fully recovered downstream pressure	[bar(a)]
κ	Isentropic exponent	[-]
Кте	Temperature exponent	[-]

fxISO6976_1983_M

Description

ISO standard 6976 defines component properties and calculations to determine the calorific value, density and relative density for a gas composition at the specified metering and combustion reference temperatures and 1.01325 bar(a).

Compliance

 International standard, Natural Gas - Calculation of calorific values, density, relative density and Wobbe index (ISO 6976:1983)

Boundaries

ISO6976:1983 does not define limits for its input data.

Function inputs

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Composition	Standard composition as defined in section 'Standard gas composition.	mol/mol	01	0
Metering reference	Temperature used for calculating the compressibility, the density and the real	-		1
temperature	relative density values			
	1: 0 °C			
	2: 15 °C			
Calorific value reference	Temperatures used for calculating the calorific values.	-		1
temperature	1st value represents the combustion reference temperature and the 2nd value			
	the Gas volume reference temperature			
	1: 25 °C / 0 °C			
	2: 0 °C / 0 °C			
	3: 15 °C / 0 °C			
	4: 15 °C / 15 °C			
	5: 60 °F / 60 °F			

Function outputs

Function outputs	Remark	EU	Fallback	
Status	0: Normal			
	1: Input argument out of range			
	2: Calculation error			
	3: Mole fractions do not add up to 1.0 +- 0.0001			
Superior calorific value	Real value at the reference conditions of temperature and	MJ/m3	0	
	pressure			
Density	At the reference conditions of temperature and pressure	kg/m3	0	
Compressibility		-	1	
Relative density		-	0	
Molar mass		kg/kmol	0	

Calculations

Calculations are performed in accordance with the standard,

using the values as listed in the tables of the standard.

fxISO6976_1995_M

Description

ISO standard 6976 edition 1995 defines component properties and calculations to determine the calorific value, density, relative density and Wobbe index for a gas composition at the specified metering and combustion reference temperatures and 1.01325 bar(a).

Both the definitive and alternative methods of calculating the calorific value on a mass and volumetric basis are included.

Boundaries

- The valid ranges for molar fractions are as follows:
- Methane
 0.5 <= .. <= 1.0
 Nitrogen
 0.0 <= .. <= 0.3
 Ethane
 0.0 <= .. <= 0.15
- Carbon dioxide 0.0 <= .. <= 0.15
- All others 0.0 <= .. <= 0.05

Compliance

 International standard, Natural Gas - Calculation of calorific values, density, relative density and Wobbe index (ISO 6976:1995/BS7589)

Function inputs and outputs

Remark	EU	Range	Defaul
Optional tag name, tag description and tag group			
Standard composition as defined in section 'Standard gas composition.	mol/mol	01	0
The reference temperature for combustion / metering:			1
1: 15°C / 15°C			
2: 0°C / 0°C			
3: 15°C / 0°C			
· · ·	-		1
			-
	-		1
Uses the values from tables 3, 4 and 5 as listed in the standard.			
Remark	EU		Fallback
0: Normal			
1: Input argument out of range			
2: Calculation error			
3: Mole fractions do not add up to 1.0 +- 0.0001			
•	MJ/m3		0
	-, -		
•	ka/m3		0
			•
· ·	-		1
· · ·	-		1
	-		0
	- - kg/kmol		0
Real superior calorific value on mass basis at the reference conditions of temperature and	-		0
Real superior calorific value on mass basis at the reference conditions of temperature and pressure	- - kg/kmol MJ/kg		0 0 0
Real superior calorific value on mass basis at the reference conditions of temperature and pressure Real superior calorific value on mole basis at the reference conditions of temperature and	- - kg/kmol		0
Real superior calorific value on mass basis at the reference conditions of temperature and pressure Real superior calorific value on mole basis at the reference conditions of temperature and pressure	- - kg/kmol MJ/kg MJ/kmol		0 0 0 0
Real superior calorific value on mass basis at the reference conditions of temperature and pressure Real superior calorific value on mole basis at the reference conditions of temperature and	- - kg/kmol MJ/kg		0 0 0
Real superior calorific value on mass basis at the reference conditions of temperature and pressure Real superior calorific value on mole basis at the reference conditions of temperature and pressure	- - kg/kmol MJ/kg MJ/kmol		0 0 0 0
Real superior calorific value on mass basis at the reference conditions of temperature and pressure Real superior calorific value on mole basis at the reference conditions of temperature and pressure Real inferior calorific value on volume basis at the reference conditions of temperature and	- - kg/kmol MJ/kg MJ/kmol		0 0 0 0
Real superior calorific value on mass basis at the reference conditions of temperature and pressure Real superior calorific value on mole basis at the reference conditions of temperature and pressure Real inferior calorific value on volume basis at the reference conditions of temperature and pressure	- - kg/kmol MJ/kg MJ/kmol MJ/m3		0 0 0 0 0
Real superior calorific value on mass basis at the reference conditions of temperature and pressure Real superior calorific value on mole basis at the reference conditions of temperature and pressure Real inferior calorific value on volume basis at the reference conditions of temperature and pressure Real inferior calorific value on mass basis at the reference conditions of temperature and pressure	- - kg/kmol MJ/kg MJ/kmol MJ/m3		0 0 0 0 0
Real superior calorific value on mass basis at the reference conditions of temperature and pressure Real superior calorific value on mole basis at the reference conditions of temperature and pressure Real inferior calorific value on volume basis at the reference conditions of temperature and pressure Real inferior calorific value on mass basis at the reference conditions of temperature and pressure	- kg/kmol MJ/kg MJ/kmol MJ/m3 MJ/kg		0 0 0 0 0 0
Real superior calorific value on mass basis at the reference conditions of temperature and pressure Real superior calorific value on mole basis at the reference conditions of temperature and pressure Real inferior calorific value on volume basis at the reference conditions of temperature and pressure Real inferior calorific value on mass basis at the reference conditions of temperature and pressure Real inferior calorific value on mass basis at the reference conditions of temperature and pressure Real superior calorific value on mole basis at the reference conditions of temperature and pressure	- - MJ/kg MJ/kmol MJ/m3 MJ/kg MJ/kg		0 0 0 0 0 0 0
Real superior calorific value on mass basis at the reference conditions of temperature and pressure Real superior calorific value on mole basis at the reference conditions of temperature and pressure Real inferior calorific value on volume basis at the reference conditions of temperature and pressure Real inferior calorific value on mass basis at the reference conditions of temperature and pressure Real inferior calorific value on mass basis at the reference conditions of temperature and pressure Real superior calorific value on mole basis at the reference conditions of temperature and pressure	- kg/kmol MJ/kg MJ/kmol MJ/m3 MJ/kg		0 0 0 0 0 0
Real superior calorific value on mass basis at the reference conditions of temperature and pressure Real superior calorific value on mole basis at the reference conditions of temperature and pressure Real inferior calorific value on volume basis at the reference conditions of temperature and pressure Real inferior calorific value on mass basis at the reference conditions of temperature and pressure Real inferior calorific value on mass basis at the reference conditions of temperature and pressure Real superior calorific value on mole basis at the reference conditions of temperature and pressure	- - MJ/kg MJ/kmol MJ/m3 MJ/kg MJ/kg		0 0 0 0 0 0 0
	Remark Optional tag name, tag description and tag group Standard composition as defined in section 'Standard gas composition. The reference temperature for combustion / metering: 1: 15°C / 15°C 2: 0°C / 0°C 3: 15°C / 0°C 4: 25°C / 0°C 5: 20°C / 20°C 6: 25°C / 20°C 1: Calculate Calculates the molar mass from the atomic masses as defined in the note of Table 1 of the standard 2: Table Uses the values from Table 1 of the standard West the mass based calorific value from the molar based calorific values from table 3 and from the calculated molar mass values. Calculates the volume based calorific value by multiplying the molar based calorific values from table 3 by p2/R.T2 2: Alternative method Uses the values from tables 3, 4 and 5 as listed in the standard. Remark 0: Normal 1: Input argument out of range 2: Calculation error 3: Mole fractions do not add up to 1.0 +- 0.0001 Real superior calorific value on volume basis at the reference conditions of temperature and pressure	Remark EU Optional tag name, tag description and tag group standard composition as defined in section 'Standard gas composition. mol/mol The reference temperature for combustion / metering: i: 15°C / 15°C ?? 2: 0°C / 0°C 3: 15°C / 0°C ? 4: 25°C / 0°C ? ? 6: 25°C / 20°C ? ? 1: Calculate - . Calculates the molar mass from the atomic masses as defined in the note of Table 1 of the standard . 2: Table . . Uses the values from Table 1 of the standard - . Calculates the molar mass from the atomic masses as defined in the note of Table 1 of the standard . 2: Table Uses the values from Table 1 of the standard - Use the values from Table 1 of the standard - . Calculates the mass based calorific value from the molar based calorific values from table 3 and from the calculated molar mass values. . Calculates the volume based calorific value by multiplying the molar based calorific values from table 3 by p2/R.T2 . 2: Alternative method Uses the values from tables 3, 4 and 5 as listed in the standard. Normal 1. Input argument out of range .	Optional tag name, tag description and tag group standard composition as defined in section 'Standard gas composition. mol/mol 0.1 The reference temperature for combustion / metering: 1:15°C / 15°C 0°C / 0°C 3:15°C / 0°C 3: 15°C / 0°C 3:5°C / 0°C - - 4: 25°C / 0°C - - - 1: Calculate - - - Calculates the molar mass from the atomic masses as defined in the note of Table 1 of the standard - - 2: Table - - - - Uses the values from Table 1 of the standard - - - Refer to paragraph 6.1 and 7.1 of the standard - - - 1: Definitive method - - - - Calculates the walues from Table 1 of the standard - - - - 2: Table - <

Calculations

Calculations are performed in accordance with the standard.

fxISO6976ex_1995_M

Description

Function inputs

Extended version that takes the first 55 components as used in the standard plus an additional set of user-definable components.

ISO standard 6976 edition 1995 defines component properties and calculations to determine the calorific value, density, relative density and Wobbe index for a gas composition at the specified metering and combustion reference temperatures and 1.01325 bar(a).

Both the definitive and alternative methods of calculating the calorific value on a mass and volumetric basis are included.

Boundaries

The valid ranges for molar fractions are as follows:

•	Methane	0.5 <= <= 1.0
•	Nitrogen	0.0 <= <= 0.3
•	Ethane	0.0 <= <= 0.15
•	Carbon dioxide	0.0 <= <= 0.15

• All others 0.0 <= .. <= 0.05

Compliance

 International standard, Natural Gas - Calculation of calorific values, density, relative density and Wobbe index (ISO 6976:1995/BS7589)

Function inputs	Remark	EU	Default
lame	Optional tag name, tag description and tag group		
omposition	Array of mole fractions of the first 55 components as used in the standard.	mol/mol	0
	1: Methane		
	2: Ethane		
	3: Propane		
	4: n-Butane		
	5 2-Methylpropane		
	6: n-Pentane		
	7: 2-Methylbutane		
	8: 2,2-Dimethylpropane		
	9: n-Hexane		
	10: 2-Methylpentane		
	11: 3-Methylpentane		
	12: 2,2-Dimethylbutane		
	13: 2,3-Dimethylbutane		
	14: n-Heptane		
	15: n-Octane		
	16: n-Nonane		
	17: n-Decane		
	18: Ethylene		
	19: Propylene		
	20: 1-Butene		
	21: cis-2-Butene		
	22: trans-2-Butene		
	23: 2-Methylpropene		
	24: 1-Pentene		
	25: Propadiene		
	26: 1,2-Butadiene		
	27: 1,3-Butadiene		
	28: Acetylene		
	29: Cyclopentane		
	30: Methylcyclopentane		
	31: Ethylcyclopentane		
	32: Cyclohexane		
	33: Methylcyclohexane		
	34: Ethylcyclohexane		
	35: Benzene		
	36: Toluene		
	37: Ethylbenzene		
	38: o-Xylene		
	39: Methanol		
	40: Methanethiol		
	41: Hydrogen		
	42: Water		
	43: Hydrogen sulfide		
	44: Ammonia		
	45: Hydrogen cyanide		
	46: Carbon monoxide		
	47: Carbonyl sulfide		
	48: Carbon disulfide		
	49: Helium		
	50: Neon		
	51: Argon		
	52: Nitrogen		
	53: Oxygen		

Function inputs	Remark	EU	Default
	54: Carbon dioxide		
	55: Sulfur dioxide		
Reference conditions	The reference temperature for combustion / metering:		1
	1: 15°C / 15°C		
	2: 0°C / 0°C		
	3: 15°C / 0°C		
	4: 25°C / 0°C		
	5: 20°C / 20°C		
	6: 25°C / 20°C		
Molar mass table method	1: Calculate	-	1
	Calculates the molar mass from the atomic masses as defined in the note of Table 1 of the standard		
	2: Table		
	Uses the values from Table 1 of the standard		
Calorific value calculation	Refer to paragraph 6.1 and 7.1 of the standard	-	1
method	1: Definitive method		
	Calculates the mass based calorific value from the molar based calorific values from table 3 and from		
	the calculated molar mass values.		
	Calculates the volume based calorific value by multiplying the molar based calorific values from table		
	3 by p2/R.T2		
	2: Alternative method		
	Uses the values from tables 3, 4 and 5 as listed in the standard.		
Jser-defined composition	Array of mole fractions of the additional user-definable components.	mol/mol	0
	Note: The inputs for the additional components are optional inputs.		
User-defined molar mass	Array of molar masses of the additional user-definable components. This should be the molar mass	kg/kmol	0
	corresponding to the current 'Molar mass table method'.		
	Note: This array should be exactly as long as the user-defined composition array.		
Jser-defined summation	Array of summation factor values of the additional user-definable components.	-	0
factor	Note: The input values have to correspond with the metering conditions of input 'Reference conditions.		
	This array should be exactly as long as the user-defined composition array.		
Jser-defined superior	Array of Superior Calorific Values of the additional user-definable components. This should be the SCV	KJ/mol	0
calorific value (SCV)	corresponding to the current reference conditions.		
	Note: the input values have to correspond with the conditions as defined by input 'Reference conditions.		
	This array should be exactly as long as the user-definable composition array.		
User-defined inferior	Array of Inferior Calorific Values of the additional user-definable components. This should be the ICV	KJ/mol	0
calorific value (ICV)	corresponding to the current reference conditions.		
	Note: the input values have to correspond with the conditions as defined by input 'Reference conditions.		
	This array should be exactly as long as the user-definable composition array.		

Function outputs

Function outputs	Remark	EU	Fallback
Status	0: Normal		
	1: Input argument out of range		
	2: Calculation error		
	3: Mole fractions do not add up to 1.0 +- 0.0001		
Superior calorific value	Real superior calorific value on volume basis at the reference conditions of temperature	MJ/m3	0
	and pressure		
Density	At the reference conditions of temperature and pressure	kg/m3	0
Compressibility		-	1
Relative density		-	0
Molar mass		kg/kmol	0
Superior calorific value	Real superior calorific value on mass basis at the reference conditions of temperature and pressure	MJ/kg	0
Superior calorific value	Real superior calorific value on mole basis at the reference conditions of temperature and pressure	MJ/kmol	0
Inferior calorific value	Real inferior calorific value on volume basis at the reference conditions of temperature and pressure	MJ/m3	0
Inferior calorific value	Real inferior calorific value on mass basis at the reference conditions of temperature and pressure	MJ/kg	0
Inferior calorific value	Real superior calorific value on mole basis at the reference conditions of temperature and pressure	MJ/kmol	0
Wobbe index		MJ/m3	0
Data range	With respect to the ISO6976-1995 standard the combination of input values is:		-
	0: In Range		
	1: Out of Range		

Calculations

Calculations are performed in accordance with the standard with the addition of the user-definable components.

based heating values that are calculated from these input properties. Only the Definitive Method for calculating the calorific value is supported in this case.

The properties of the user-definable components are specified by the corresponding inputs, except for the volume and mass

fxISO6976_2016_M

Description

ISO standard 6976 edition 2016 defines component properties and calculations to determine the calorific value, density, relative density and Wobbe index for a gas composition at the specified metering and combustion reference temperatures and metering pressure.

Boundaries

The standard defines the following validity ranges

- 0.9 < Pressure < 1.1 bar
- Compressibility > 0.9

Compliance

• International standard, Natural Gas - Calculation of calorific values, density, relative density and Wobbe indices from composition (ISO 6976:2016(e))

Function inputs and outputs

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Composition	Standard composition as defined in section 'Standard gas composition.	mol/mol	01	0
Reference conditions	The reference temperature for combustion / metering:			1
	1: 15°C / 15°C			
	2: 0°C / 0°C			
	3: 15°C / 0°C			
	4: 25°C / 0°C			
	5: 20°C / 20°C			
	6: 25°C / 20°C			
	7. 60°F / 60°F			
Molar mass table	1: Calculate	-	12	1
method	Calculates the molar mass from the atomic masses as defined in the note of Table 1 of the			
	standard			
	2: Table			
	Uses the values from Table 1 of the standard			
Metering reference	Metering reference pressure p2. The default value is 1.01325 bar (101.325 kPa).	bar	02	1.01325
pressure				

Function outputs	Remark	EU	Fallback
Status	0: Normal		
	1: Input argument out of range		
	2: Calculation error		
	3: Mole fractions do not add up to 1.0 +- 0.0001		
Gross calorific value	Real gas gross calorific value on volume basis at the reference conditions of temperature and pressure	MJ/m3	0
Density	At the reference conditions of temperature and pressure	kg/m3	0
Compressibility		-	1
Relative density		-	0
Molar mass		kg/kmol	0
Gross calorific value	Real gas gross calorific value on mass basis at the reference conditions of temperature and pressure	MJ/kg	0
Gross calorific value	Real gas gross calorific value on mole basis at the reference conditions of temperature and pressure	MJ/kmol	0
Net calorific value	Real gas net calorific value on volume basis at the reference conditions of temperature and pressure	MJ/m3	0
Net calorific value	Real gas net calorific value on mass basis at the reference conditions of temperature and pressure	MJ/kg	0
Net calorific value	Real gas net calorific value on mole basis at the reference conditions of temperature and pressure	MJ/kmol	0
(Gross) Wobbe index		MJ/m3	0
Data range	With respect to the ISO6976-2016 standard the combination of input values is:		-
	0: In Range		
	1: Out of Range		

Calculations

Calculations are performed in accordance with the standard.

fxISO6976ex_2016_M

Description

Extended version that takes the first 60 components as used in the standard plus an additional set of user-definable components.

ISO standard 6976 edition 2016 defines component properties and calculations to determine the calorific value, density, relative density and Wobbe index for a gas composition at the specified metering and combustion reference temperatures and metering pressure.

Function inputs

Boundaries

The standard defines the following validity ranges

- 0.9 < Pressure < 1.1 bar
- Compressibility > 0.9

Compliance

 International standard, Natural Gas - Calculation of calorific values, density, relative density and (Gross) Wobbe index (ISO 6976:2016/BS7589)

Function inputs	Remark	EU	Default
ame	Optional tag name, tag description and tag group	· · ·	
omposition	Array of mole fractions of the first 55 components as used in the standard.	mol/mol	0
	1: Methane		
	2: Ethane		
	3: Propane		
	4: n-Butane		
	5 2-Methylpropane		
	6: n-Pentane		
	7: 2-Methylbutane		
	8: 2,2-Dimethylpropane		
	9: n-Hexane		
	10: 2-Methylpentane		
	11: 3-Methylpentane		
	12: 2,2-Dimethylbutane		
	13: 2,3-Dimethylbutane		
	14: n-Heptane		
	15: n-Octane		
	16: n-Nonane		
	17: n-Decane		
	18: Ethylene		
	19: Propylene		
	20: 1-Butene		
	21: cis-2-Butene		
	22: trans-2-Butene		
	23: 2-Methylpropene		
	24: 1-Pentene		
	25: Propadiene		
	26: 1,2-Butadiene		
	27: 1,3-Butadiene		
	28: Acetylene		
	29: Cyclopentane		
	30: Methylcyclopentane		
	31: Ethylcyclopentane		
	32: Cyclohexane		
	33: Methylcyclohexane		
	34: Ethylcyclohexane		
	35: Benzene		
	36: Toluene		
	37: Ethylbenzene		
	38: o-Xylene		
	39: Methanol		
	40: Methanethiol		
	41: Hydrogen		
	42: Water		
	43: Hydrogen sulfide		
	44: Ammonia		
	45: Hydrogen cyanide		
	46: Carbon monoxide		
	47: Carbonyl sulfide		
	48: Carbon disulfide		
	49: Helium		
	50: Neon		
	51: Argon		
	52: Nitrogen		
	53: Oxygen		
	54: Carbon dioxide		
	55: Sulfur dioxide		
	56: N_undecane		
	57: N_Dodecane		

Function inputs	Remark	EU	Default
	58: N_tridecane		
	59: N_pentadecane		
	60: N_pentadecane		
Reference conditions	The reference temperature for combustion / metering:		1
	1: 15°C / 15°C		
	2: 0°C / 0°C		
	3: 15°C / 0°C		
	4: 25°C / 0°C		
	5: 20°C / 20°C		
	6: 25°C / 20°C		
	7. 60°F / 60°F		
Molar mass table method	1: Calculate	-	1
	Calculates the molar mass from the atomic masses as defined in the note of Table 1 of the standard		
	2: Table		
	Uses the values from Table 1 of the standard		
Metering reference pressure	Metering reference pressure p2.	bar	1.01325
Jser-defined composition	Array of mole fractions of the additional user-definable components.	mol/mol	0
	Note: The inputs for the additional components are optional inputs.		
Jser-defined molar mass	Array of molar masses of the additional user-definable components. This should be the molar mass	kg/kmol	0
	corresponding to the current 'Molar mass table method'.	5.	
	Note: This array should be exactly as long as the user-defined composition array.		
Jser-defined summation	Array of summation factor values of the additional user-definable components.	-	0
actor	Note: The input values have to correspond with the metering conditions of input 'Reference conditions.		
	This array should be exactly as long as the user-defined composition array.		
Jser-defined gross calorific	Array of Gross Calorific Values of the additional user-definable components. This should be the CV	KJ/mol	0
value	corresponding to the current reference conditions.		
	Note: the input values have to correspond with the conditions as defined by input 'Reference conditions.		
	This array should be exactly as long as the user-definable composition array.		

Function outputs

Function outputs	Remark	EU	Fallback
Status	0: Normal		
	1: Input argument out of range		
	2: Calculation error		
	3: Mole fractions do not add up to 1.0 +- 0.0001		
Gross calorific value	Real gross calorific value on volume basis at the reference conditions of temperature and pressure	MJ/m3	0
Density	At the reference conditions of temperature and pressure	kg/m3	0
Compressibility		-	1
Relative density		-	0
Molar mass		kg/kmol	0
Gross calorific value	Real gas gross calorific value on mass basis at the reference conditions of temperature and pressure	MJ/kg	0
Gross calorific value	Real gas gross calorific value on mole basis at the reference conditions of temperature and pressure	MJ/kmol	0
Net calorific value	Real gas net calorific value on volume basis at the reference conditions of temperature and pressure	MJ/m3	0
Net calorific value	Real gas net calorific value on mass basis at the reference conditions of temperature and pressure	MJ/kg	0
Net calorific value	Real gas net calorific value on mole basis at the reference conditions of temperature and pressure	MJ/kmol	0
(Gross) Wobbe index		MJ/m3	0
Data range	With respect to the ISO6976-2016 standard the combination of input values is: 0: In Range 1: Out of Range		-

Calculations

Calculations are performed in accordance with the standard with the addition of the user-definable components. The properties of the user-definable components are specified by the corresponding inputs, except for the volume and mass based heating values that are calculated from these input properties. Only the Definitive Method for calculating the calorific value is supported in this case.

fxMR113

Description

The relative humidity, compressibility, density, speed of sound, isentropic coefficient and optionally the dynamic viscosity of a gas are calculated from its composition, absolute humidity, temperature and pressure in accordance with the GOST MR113 standard. The CPU-intensive calculation of viscosity can be optionally enabled if the (flow computer) CPU capacity allows for this.

Compliance

• The GSSSD method MR 113-03. Kozlov, Mamonov, Rogovin, Rybakov, (10/06/2003).

Boundaries

The standard defines a valid range for pressure and temperature. The function will not accept input values outside this range

The valid range is as follows

- Pressure: 1..150 bar(a)
- Temperature: 263..500 K (-10 ..+226 °C)

Note: the standard does not specify a range limitation for the components.

Function inputs

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Pressure	Flowing pressure	bar(a)	1150	1.01325
Temperature	Flowing temperature	°C	-10+226	0
Composition	Standard composition as defined in section 'Standard gas composition. Unused components must be set to 0.	mol/mol	01	0
neo-Pentane mode	Determines what to do when component neo-Pentane is larger than zero 1: Add to i-Pentane 2: Add to n-Pentane 3: Neglect	-	13	1
Absolute humidity	Absolute humidity at the humidity meter	kg/m3	01	0
Humidity pressure	Pressure value at the humidity meter	bar(a)	03500	1.01325
Humidity temperature	Temperature value at the humidity meter	°C	-250+450	0
Reference pressure	Reference pressure (base conditions)	bar(a)	03500	1.01325
Reference temperature	Reference temperature (base conditions)	°C	-250+450	0
Rounding	0: Disabled, full precision		01	0
	1: Enabled, results are rounded			
Viscosity calculation	0: Disabled		01	0
	1: Enabled			
	Note: the viscosity calculation is relative CPU-intensive.			

Function outputs

Function outputs	Remark	EU	Fallback
Status	0: Normal		
	1: Input argument out of range		
	2: Calculation error		
	3: No convergence		
Base Compressibility	At reference conditions	-	1
Flow Compressibility	At flowing conditions	-	1
Base Density	At reference conditions	kg/m3	0
Flow Density	At flowing conditions	kg/m3	0
Kappa	Isotropic exponent	-	0
Molar Mass		kg/kmol	0
Speed of Sound	At flowing conditions	m/s	0
Dew point	Water dew point, the temperature at which the water in the gas starts to	К	0
	condensate.		
Viscosity	The dynamic viscosity of the gas. The calculation is disabled by default.	Pa.s	0
Max water fraction	Maximum water fraction at flowing conditions	%mole	0
Water fraction	Actual water fraction at flowing conditions	%mole	0
Relative humidity	At flow conditions	%	0
Range	0: All input values are within the 'Normal Range'		0
-	1: One or more input values are outside the 'Normal Range		

Calculations

The calculations are as documented in the reference.

fxNX19_1962

Description

The AGA NX-19 standard describes a method to calculate the super-compressibility for natural gases and was developed in 1962.

The 1962 standard describes a standard method for calculating the super-compressibility factor that is based on the actual pressure and temperature, the specific gravity and the mole fractions of the carbon dioxide and nitrogen.

The 1962 standard also specifies 3 alternate methods, which are based on a full compositional analysis (1st alternate method), the relationship between methane and specific gravity (2nd alternate method) and the relationship between the heating value and the specific gravity

The function only performs the standard method as specified by the standard and none of the alternate methods.

Note: the definition of the specific gravity as used in the standard is that of the ratio of the density of the gas to that of air at base conditions, so the real specific gravity or real relative density.

Compliance

- AGA Par Research Project NX-19 Manual for the Determination. of the Supercompressibility Factors for Natural Gas, 1962
- Flow Measurement Engineering Handbook, Third edition, . Richard W. Miller, 1996, ISBN-0-07-042366-0

Boundaries

The following boundaries apply for the input values. Using the standard for conditions that lie outside this range will yield to a higher uncertainty and is not recommended.

-40240	°F
05000	psig
0.554 01.000	-
0.00 0.15	mol/mol
0.00 0.15	mol/mol
	05000 0.554 01.000 0.00 0.15

Function inputs

Function inputs				
Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Pressure	Observed pressure	psig	040000	0
Temperature	Observed temperature	°F	-250800	60
Specific gravity	Ratio of density of gas and density of air at the applicable reference conditions of pressure and temperature, i.e. the <u>real</u> specific gravity (real relative density).at 60°F and 14.73 psia.	-	02	0
Nitrogen	Mole fraction of nitrogen	mol/mol	01	0
Carbon dioxide	Mole fraction of carbon dioxide	mol/mol	01	0

Function outputs

Function outputs	Remark	EU	Fallback
Status	0: Normal		
	1: Input argument out of range		
	2: Calculation error		
	Note: value 1 is also reported when one of the intermediate results is out of limit	t.	
Compressibility factor	Compressibility factor at the observed temperature and pressure (Zf)	-	1
Base compressibility factor	Compressibility factor at 60 °F and 14.7 psia (Zb)	-	1
Super-compressibility factor	Super-compressibility Fpv. Refer to section Calculations for ist definition.	-	1
Range	0: All input values are within the 'Normal Range'	-	0
-	1. One or more input values are outside the 'Normal Range		

Calculations

The AGA-NX-19 standard specifies the calculations of the supercompressibility Fpv, which is defined as follows:

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Farm	_	Zb
Fpv	=	 Zf

Where:	
Fpv	Supercompressibility
Zb	Compressibility at base conditions of 60 °F and 14.7 psia
Zf	Compressibility at flowing conditions, i.e. at the input temperature and pressure

The definition of the base compressibility factor when applying the AGA-NX19:1962 method for custody transfer is given in chapter 2 of [Miller:1996].

$$Zb = \left[\sqrt{1 + \frac{0.00132}{T^{3.25}}}\right]^{-2}$$

Where T = Tadj/500 with Tadj calculated in accordance with the AGA NX-19:1962 standard

fxNX19_M

Description

The AGA NX-19 standard describes a method to calculate the super-compressibility for natural gases and was developed in 1962.

The 1962 standard describes a standard method for calculating the super-compressibility factor that is based on the actual pressure and temperature, the specific gravity and the mole fractions of the carbon dioxide and nitrogen.

The 1962 standard also specifies 3 alternate methods, which are based on a full compositional analysis (1st alternate method), the relationship between methane and specific gravity (2nd alternate method) and the relationship between the heating value and the specific gravity

The function only performs the standard method as specified by the standard and none of the alternate methods.

The function provides the option to perform the PTB G9 correction instead of the 1962 standard method. This consists of the modified NX-19 method (NX-19-mod) per Herning & Wolowsky and the additional 'BR.KORR.3H' correction for highcaloric gases (gross heating value >= 39.8 MJ/m3).

Note: the definition of the specific gravity as used in the standard is that of the ratio of the density of the gas to that of air at base conditions, so the <u>real</u> specific gravity or real relative density.

Compliance

- AGA Par Research Project NX-19 Manual for the Determination. of the Supercompressibility Factors for Natural Gas, 1962
- Berechnung von Realgasfaktoren und Kompressibilitätszahlen für Erdgas, Technische Richtlinie G9 der Physikalisch - Technische Bundesanstalt für meßgeräte für Gas (PTB), TRG 9 8/82

Boundaries

The following bounds apply for the input values. Using the standard for conditions that lie outside this range will yield to a higher uncertainty and is not recommended.

Input value	AGA-NX-19 (1962)	AGA-NX-19- mod	AGA-NX-19- mod.BR. KORR.3H	EU
Pressure	0 350	0 137.9	080	bar(a)
Temperature	-40115.6	-40115.6	030	°C
Relative density	0.554 01.000	0.5540.75	0.5540.691	-
Gross heating value	Not used	31.839.8	39.846.2	MJ/m3
Nitrogen	0.00 0.15	0.00 0.15	0.000.025	mol/mol
Carbon dioxide	0.00 0.15	0.00 0.15	0.000.07	mol/mol

Function inputs

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Pressure	Observed pressure	bar(a)	0200	1.01325
Temperature	Observed temperature	°C	-100300	0
Specific gravity	Ratio of density of gas and density of air at the applicable reference conditions of pressure and temperature, i.e. the <u>real</u> specific gravity (real relative density). If setting 'PTB G9 correction' is disabled the reference conditions are 60°F and 14.73 psia. Else the specific gravity value shall be at the applicable reference conditions of pressure and temperature.	-	02	0
Gross heating value	At the applicable reference conditions of pressure and temperature Only required when the 'PTB G9 correction' is enabled.	MJ/m3	0100	0
Nitrogen		mol/mol	01	0
Carbon dioxide		mol/mol	01	0
PTB G9 correction	Determines if the AGA-NX-19-mod / AGA-NX-19-mod.BR.KORR.3H is used instead of the AGA-NX-19-1962 standard calculation. 0: Disabled 1: Enabled	-		1

Function outputs

Function outputs	Remark	EU	Fallback
Status	0: Normal		
	1: Input argument out of range		
	2: Calculation error		
	3: No convergence		
	Note: value 1 is also reported when one of the intermediate results is out of limit.		
Compressibility factor		-	1
Range	0: All input values are within the 'Normal Range'	-	0
-	1: One or more input values are outside the 'Normal Range		

Calculations

The calculations are as specified in the standards.

fxOIML_R22_1975

Description

This function calculates the density, mass and volume based alcoholic strength of ethanol and water mixtures.

Compliance

• International Organisation of Legal metrology: International alcoholometric tables.

Function input

Function inputs	Remark	EU	Range	Default
Observed temperature		°C	-2545	7
Standard temperature for the mixture		°C	-2545	20
Standard temperature for ethanol		°C	-2545	20
Input density		kg/m3	771.93999.97	829.110
Conversion method	1: From observed to standard conditions 2: From standard to observed conditions.		12	1
Density margin.	Allow the input density to go out of the limits of the standard by this margin. E.g. Now the function allows an Input density of 769.	kg/m3		3

Function output

Function outputs	Remark	EU	Fallback
Status	0: Normal (No error condition)		
	1: Input argument out of range		
	2: No convergence		
Density		kg/m3	1
Ctl	Volume correction factor for temperature.	01	1
Mass strength	Mass based alcoholic strength.	01	1
Volume strength	Volume based alcoholic strength.	01	1
CBSW	Correction factor to calculate the volume of ethanol at the ethanol standard	01	1
	temperature from the volume of the mixture at the mixture standard temperature.		
Range status	0: Input values are in the range of the standard		
	1: Input values are outside the range of the standard		

fxSarasota_C

Description

The function calculates the density from a frequency input signal provided by a Sarasota densitometer and corrects it for temperature and pressure effects in **US customary** units.

Note: The calibration constants also need to be in US customary

units (°F, psi and lbm/ft3).

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Periodic time	In microseconds	μs		01e6	0
	Equals 1000 divided by the frequency in [Hz]				
Line temperature	Used when temperature correction is enabled	°F		-273.15+500	20
Line pressure	Used when pressure correction is enabled	psig		0200	0
Temperature correction	0: Disabled	-	TEMPCOR		1
	1: Enabled				
Pressure correction	0: Disabled	-	PRESCOR		1
	1: Enabled				
Reference temperature	Used when temperature correction is enabled	°F	REFTEMP	0100	20
Reference pressure	Used when pressure correction is enabled	psig	REFPRES	0100	0
do	Constant from calibration certificate	lbm/ft3	D0	02000	0
	Note: value required in lbm/ft3				
το	Constant from calibration certificate	μs	то	01e6	0
к	Spool calibration constant from calibration certificate	-	К	03e3	0
Temperature coefficient	Constant from calibration certificate	μs/°F	TEMPCOEF	-1e61e6	0
Pressure coefficient	Constant from calibration certificate	μs/psi	PRESCOEF	-1e61e6	0

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALC	
Corrected density	Density corrected for temperature and pressure	lbm/ft3	CORDENS		0

Calculations

The corrected density ρ_{C} is calculated by

$$\rho_{c} = d_{0} \cdot \frac{\tau - \tau_{c}}{\tau_{c}} \cdot \left(2 + K \cdot \frac{\tau - \tau_{c}}{\tau_{c}}\right)$$

$$\tau_{c} = \tau_{0} + t_{COEF} \cdot (t - t_{CAL}) + p_{COEF} \cdot (p - p_{CAL})$$

ρο	The corrected density	lbm/ft3
d٥	Obtained from the calibration certificate	lbm/ft3
το	Obtained from the calibration certificate	μs
к	Obtained from the calibration certificate	-
PCOEF	Obtained from the calibration certificate	µs/psi
tcoef	Obtained from the calibration certificate	µs/°F
t	Line temperature	°F
tcal	Reference temperature	°F
р	Line temperature	psig
PCAL	Reference pressure	psig
τc	Time periodic input corrected for temperature and	μs
	pressure	
τ	Measured time period	μS

fxSarasota_M

Description

The function calculates the density from a frequency input signal provided by a Sarasota densitometer and corrects it for temperature and pressure effects in **metric** units.

Note: Calibration constants also need to be in metric units (°C,

bar and kg/m3).

Function inputs

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Periodic time	In microseconds.Equals 1000 divided by the frequency in [Hz]	μs	01e6	0
Line temperature	Used when temperature correction is enabled	°C	-273.15+500	20
Line pressure	Used when pressure correction is enabled	bar(g)	0200	0
Temperature correction	0: Disabled	-		1
	1: Enabled			
Pressure correction	0: Disabled	-		1
	1: Enabled			
Reference temperature	Used when temperature correction is enabled	°C	0100	20
Reference pressure	Used when pressure correction is enabled	bar(g)	0100	0
do	Constant from calibration certificate	kg/m3	03000	0
το	Constant from calibration certificate	μs	01e6	0
к	Spool calibration constant from calibration certificate	-	03e3	0
Temperature coefficient	Constant from calibration certificate	μs/°C	-1e61e6	0
Pressure coefficient	Constant from calibration certificate.	μs/bar	-1e61e6	0
	Note: value required in µs/bar			

Function outputs

Function outputs	Remark	EU	SW tag	Fallback
Status	0: Normal	STS		
	1: Input argument out of range		FIOOR	
	Outputs will be set to fallback values			
	2: Calculation error		CALC	
	Outputs will be set to fallback values			
Corrected density	Density corrected for temperature and pressure	kg/m3	CORDENS	0

Calculations

The corrected density ρ_{C} is calculated by

$$\rho_{c} = d_{0} \cdot \frac{\tau - \tau_{c}}{\tau_{c}} \cdot \left(2 + K \cdot \frac{\tau - \tau_{c}}{\tau_{c}}\right)$$

$$\tau_{C} = \tau_{0} + t_{COEF} \cdot (t - t_{CAL}) + p_{COEF} \cdot (p - p_{CAL})$$

ρς	The corrected density	kg/m3
d٥	Obtained from the calibration certificate	kg/m3
το	Obtained from the calibration certificate	μs
к	Obtained from the calibration certificate	-
do	Obtained from the calibration certificate	-
P COEF	Obtained from the calibration certificate	µs/bar
tcoef	Obtained from the calibration certificate	µs/°C
t	Line temperature	°C
tcal	Reference temperature	°C
р	Line temperature	bar(g)
PCAL	Reference pressure	bar(g)
τc	Time periodic input corrected for temperature and pressure	μs
τ	The time period in μS	μs

fxSGERG_C

Description

This function performs the SGERG calculation in **USC** units.

The Standard (or Simplified) GERG TM5 1991 Virial Equation (SGERG or SGERG-88) has defines a method to calculate the Compressibility Factor (Z) for Natural Gases. The SGERG calculation is equivalent to the AGA8 Gross Characterisation Method, however .the results are slightly different

Instead of the full compositional analysis (as used by the AGA 8 Detailed Characterization method), the Gross Characterisation Method (SGERG) uses a restricted set of input variables for its equation, comprising Relative Density, Superior Calorific Value, Carbon Dioxide and Nitrogen together with pressure and temperature.

Compliance

• GERG Technical Monograph 5, Standard GERG Virial Equation, 1991

Boundaries

The standard specifies a maximum uncertainty of the compressibility factor of 0.1% provided the inputs lie in the following range.

Pressure	01740	psi(a)
Temperature	17 143	°F
Gross heating value	509 1288	Btu/ft3
Relative density	0.55 0.90	-
Carbon dioxide	0.00 0.30	mol/mol
Nitrogen	0.00 0.50	mol/mol
Hydrogen	0.00 0.10	mol/mol

Function inputs

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Pressure	Observed pressure	psia	030000	
Temperature	Observed temperature	°F	-250500	0
Relative density	At the reference conditions according to input 'Reference conditions'	-	02	0
Gross heating value	At the combustion and reference conditions according to input 'Reference conditions'	Btu/ft3	02500	0
Nitrogen		mol/mol	01	0
Carbon dioxide		mol/mol	01	0
Hydrogen		mol/mol	01	0
Method	Calculation method:	-		0
	1: All inputs are known			
	2: Unknown Nitrogen mole fraction			
	3: Unknown Carbon Dioxide mole fraction			
	4: Unknown Gross Heating Value			
	5: Unknown Relative Density			
Reference conditions	Reference conditions that correspond with the values of inputs 'Relative density' and 'Gross			1
	heating value'.			
	Combustion temp. / metering temp. / pressure			
	1: 60°F / 60 °F / 14.73 psia			
	2: 60 °F / 60 °F / 1.01592 bar			
	Note: the calculations are based on 25°C / 0°C / 1.01325 bar(a). For the other conditions			
	conversion factors are applied as specified in GERG Technical Monograph 5, Standard GERG			
	Virial Equation, 1991.			
	Refer to section 'Calculations' for more details			

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Compressibility factor		-			1
Molar mass		lb/lbmol			0
Range	0: In Normal Range All components are within the 'Normal Range'		RANGE		0
	1: In Extended Range One or more components within the 'Extended Range, but none of the components outside the Extended rang (outputs values have higher				
	uncertainty) 2: Out of Range One or more components outside the 'Extended Range' (using the AGA8 calculation is not recommended in this case)			OOR	

Calculations

The calculations are in accordance with the standard.

As specified in the standard and depending on the selected reference conditions (input 'Reference conditions') one of the following set of conversions is carried out to obtain the input values of the relative density (RD) at 0°C and 1.01325 bar(a) and the gross heating value (GHV) at 25°C and 1.01325 bar(a) combustion and 0°C and 1.01325 bar(a) metering conditions.

Input 'Reference Conditions'	Multiply input GHV with	Multiply input RD with
60 °F, 60 °F @ 14.73 psia	1.0543/26.85	1.0002
60 °F, 60 °F @ 1.01592 bar(a)	1.0543/26.86	1.0002

fxSGERG_M

Description

This function performs the SGERG calculation in metric units. The Standard (or Simplified) GERG TM5 1991 Virial Equation (SGERG or SGERG-88) has defines a method to calculate the Compressibility Factor (Z) for Natural Gases. The SGERG calculation is equivalent to the AGA8 Gross Characterisation Method, however .the results are slightly different Instead of the full compositional analysis (as used by the AGA 8 Detailed Characterization method), the Gross Characterisation Method (SGERG) uses a restricted set of input variables for its equation, comprising Relative Density, Superior Calorific Value, Carbon Dioxide and Nitrogen together with pressure and temperature.

Compliance

GERG Technical Monograph 5, Standard GERG Virial • Equation, 1991

Boundaries

The standard specifies a maximum uncertainty of the compressibility factor of 0.1% provided the inputs lie in the following range.

Pressure	0 120	bar(a)
Temperature	265335	К
Gross heating value	1948	MJ/m3
Relative density	0.55 0.90	-
Carbon dioxide	0.00 0.30	mol/mol
Nitrogen	0.00 0.50	mol/mol
Hydrogen	0.00 0.10	mol/mol

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Pressure	Observed pressure	bar(a)	02000	1.01325
Temperature	Observed temperature	°C	-200300	0
Relative density	At the reference conditions according to input 'Reference conditions'	-	02	0
Gross heating value	At the combustion and reference conditions according to input 'Reference conditions'	MJ/m3	0100	0
Nitrogen		mol/mol	01	0
Carbon dioxide		mol/mol	01	0
Hydrogen		mol/mol	01	0
Method	Calculation method:	-		0
	1: All inputs are known			
	2: Unknown Nitrogen mole fraction			
	3: Unknown Carbon Dioxide mole fraction			
	4: Unknown Gross Heating Value			
	5: Unknown Relative Density			
Reference conditions	Reference conditions that correspond with the values of inputs 'Relative density' and 'Gross heating value'.			1
	Combustion temp. / metering temp. / pressure			
	1: 25°C / 0°C / 1.01325 bar(a)			
	2: 0°C / 0 °C / 1.01325 bar(a)			
	3: 15°C / 15°C / 1.01325 bar(a)			
	Note: the calculations are based on 25° C / 0° C / 1.01325 bar(a). For the other conditions			
	conversion factors are applied as specified in GERG Technical Monograph 5, Standard GERG			
	Virial Equation, 1991.			
	Refer to section 'Calculations' for more details			

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallbac
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
	3: No convergence			NOCONV	
Compressibility factor		-			1
Molar mass		kg/kmol			0
Range	0: In Normal Range All components are within the 'Normal Range'		RANGE		0
	1: In Extended Range One or more components within the 'Extended Range, but none of the components outside the Extended rang (outputs values have higher uncertainty)				
	2: Out of Range One or more components outside the 'Extended Range' (using the AGA8 calculation OOR is not recommended in this case)				
Calculations	is not recommended in this case) combustion and 0°C and 1.01325 bar(a) metering conditions.				

Calculations

The calculations are in accordance with the standard. As specified in the standard and depending on the selected reference conditions (input 'Reference conditions') one of the following set of conversions is carried out to obtain the input values of the relative density (RD) at 0°C and 1.01325 bar(a) and the gross heating value (GHV) at 25°C and 1.01325 bar(a)

Input 'Reference Conditions'	Multiply input GHV with	Multiply input RD with
25°C / 0 °C / 1.01325 bar(a)	Not applicable	Not applicable
0 °C, 0 °C @ 1.01325 bar(a)	0.9974	Not applicable
15 °C, 15°C @ 1.01325 bar(a)	1.0543	1.0002

fxSolartron_Gas_C

Description

The function calculates the density from a frequency input signal provided by a Solartron 7810, 7811 or 7812 gas densitometer and corrects it for temperature and velocity of sound effects in **US customary** units.

The function requires that the calibration constants are based on the following units:

g/cc

- Temperature °F
- Pressure psi
- Density

Function inputs

The calculations are in accordance with documents:

- 78125010 'Solartron 7812 Gas Density Transducer Manual', 2001.
- 78125040 Rev. C, 'Micro Motion 7812 Gas Density Meter', October 2007.

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group		-	-	
Periodic time	In microseconds	μs		01e6	0
	Equals 1000 divided by the frequency in [Hz]				
Line temperature	Used when temperature correction is enabled	°F		-459.67+1000	20
Temperature correction	0: Disabled	-	TEMPCOR		1
	1: Enabled				
VOS correction	0: Disabled	-	VOSCOR		1
	1: Enabled				
Reference temperature	Used when temperature correction is enabled	°F	REFTEMP	0200	60
ко	Constant K0 from calibration certificate	-	К0	-1e91e9	
К1	Constant K1 from calibration certificate	-	K1	-1e91e9	
К2	Constant K2 from calibration certificate	-	K2	-1e91e9	
K18	Constant K18 from calibration certificate	-	K18	-1e91e9	
К19	Constant K19 from calibration certificate	-	K19	-1e91e9	
К3	Constant K3 from calibration certificate	-	K3	-1e91e9	
К4	Constant K4 from calibration certificate	-	K4	-1e91e9	
Calibration gas constant	Constant Kc from calibration certificate	-	KC	-1e91e9	
G value method	Method of determining value G, which is the ratio of Gas Specific	-	GMETHOD		1
	Gravity and the Ratio of Specific Heats				
	1: Use input 'G value'				
	2: Uses ratio of inputs				
	Uses the ratio of inputs 'Specific Gravity' and 'Ratio of Specific				
	Heats'				
G value	Value will be used when VOS correction is enabled and the G value	-	GVAL		
	method is 'Use input G value'				
Specific gravity	Value will be used when VOS correction is enabled and the G value	-	SG	02	0
	method is 'Use ratio of inputs'				
Ratio of specific heats	Value will be used when VOS correction is enabled and the G value	-	CP_CV	0.0110	0
	method is 'Use ratio of inputs'				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm Fallback
Status	0: Normal		STS	
	1: Input argument out of range			FIOOR
	2: Calculation error			CALCERR
Corrected density	Density corrected for temperature and VOS	lbm/ft3	CORDENS	0
Density corrected for	Density corrected for temperature	lbm/ft3	TCORDENS	0
temperature				
Uncorrected density	Uncorrected (indicated density	lbm/ft3	UNCDENS	0
Calaulatiana				

Calculations

Density calculations are performed in g/cc, while the function outputs are in lbm/ft3 $\,$

Where:

ρι	The uncorrected density	g/cc
ко	Obtained from the calibration certificate	-
К1	Obtained from the calibration certificate	-
К2	Obtained from the calibration certificate	-
τ	The time period in µS	μs

The uncorrected density ρ_i is calculated by $\rho_i = K0 + K1 \cdot \tau + K2 \cdot \tau^2$ The temperature corrected density ρt is calculated by

$$\rho_t = \rho_i \cdot \left[1 + K18 \cdot (t - t_R)\right] + K19 \cdot (t - t_R)$$

Where:

ρt	The density corrected for temperature	g/cc
K18	Obtained from the calibration certificate	-
К19	Obtained from the calibration certificate	-
t	The line temperature	°F
t _R	The reference temperature	°F

The density value corrected for Velocity of Sound is calculated as follows:

$$\rho_{VOS} = \rho_t \left[1 + \frac{K_3}{(\rho_t + K_4)} \cdot \left(Kc - \frac{G}{t + 273} \right) \right]$$
$$Kc = \frac{Cc}{t_c + 273}$$

ρ _{vos}	The density corrected for temperature and VOS	g/cc
КЗ	Obtained from the calibration certificate	-
К4	Obtained from the calibration certificate	-
Kc	Calibration gas constant from the calibration certificate	-
G	G value. Equals either input 'G value' or the ratio of inputs 'Specific gravity' and 'Ratio of specific heats', depending on input 'G value method'	-
t	The line temperature	°F
Cc	Specific Gravity/Ratio of specific heats of calibration gas	-
tc	Calibration temperature	°F

fxSolartron_Gas_M

Description

The function calculates the density from a frequency input signal provided by a Solartron 7810, 7811 or 7812 gas densitometer and corrects it for temperature and velocity of sound effects in metric units.

The function requires that the calibration constants are based on the following units:

•	Temperature	°C
•	Temperature	°C

- Pressure bar
- kg/m3 Density

nction inputs

Compliance

The calculations are in accordance with documents:

- 78125010 'Solartron 7812 Gas Density Transducer Manual', 2001.
- 78125040 Rev. C, 'Micro Motion 7812 Gas Density Meter', . October 2007.

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Periodic time	In microseconds	μs		01e6	0
	Equals 1000 divided by the frequency in [Hz]				
Line temperature	Used when temperature correction is enabled	°C		-273+500	20
Temperature correction	0: Disabled	-	TEMPCOR		1
	1: Enabled				
VOS correction	0: Disabled	-	VOSCOR		1
	1: Enabled				
Reference temperature	Used when temperature correction is enabled	°C	REFTEMP	0100	20
KO	Constant K0 from calibration certificate	-	KO	-1e91e9	-1.104252E+2
K1	Constant K1 from calibration certificate	-	K1	-1e91e9	-1.882012E-2
K2	Constant K2 from calibration certificate	-	K2	-1e91e9	4.749797E-4
K18	Constant K18 from calibration certificate	-	K18	-1e91e9	-1.360E-5
K19	Constant K19 from calibration certificate	-	K19	-1e91e9	8.440E-4
K3	Constant K3 from calibration certificate	-	К3	-1e91e9	354
K4	Constant K4 from calibration certificate	-	K4	-1e91e9	57.4
Calibration gas constant	Constant Kc from calibration certificate	-	КС	-1e91e9	0.00236
G value method	Method of determining value G, which is the ratio of Gas Specific	-	GMETHOD		1
	Gravity and the Ratio of Specific Heats				
	1: Use input 'G value'				
	2: Uses ratio of inputs				
	Uses the ratio of inputs 'Specific Gravity' and 'Ratio of Specific				
	Heats'				
G value	Value will be used when VOS correction is enabled and the G	-	GVAL	0100	0
	value method is 'Use input G value'				
Specific gravity	Value will be used when VOS correction is enabled and the G	-	SG	02	0
	value method is 'Use ratio of inputs'				
Ratio of specific heats	Value will be used when VOS correction is enabled and the G	-	CP_CV	010	0
	value method is 'Use ratio of inputs'				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
Corrected density	Density corrected for temperature and VOS	kg/m3	CORDENS		0
Density corrected for	Density corrected for temperature	kg/m3	TCORDENS		0
temperature					
Uncorrected density	Uncorrected (indicated density	kg/m3	UNCDENS		0
Calculations		The temperature	corrected density	ν pt is calculated	d by
The uncorrected density ρ_i is calculated by $\rho_i = K \Omega + K 1 \cdot \tau + K 2 \cdot \tau^2$		$\rho_t = \rho_i \cdot [1 + K]$	$18 \cdot (t - t_p)] + R$	$(t-t_p)$	

$$\rho_i = K0 + K1 \cdot \tau + K2 \cdot \tau^2$$

Where:				
ρι	The uncorrected density	kg/m3		
ко	Obtained from the calibration certificate	-		
К1	Obtained from the calibration certificate	-		
К2	Obtained from the calibration certificate	-		
τ	The time period in µs	μs		

The density corrected for temperature	kg/m3
Obtained from the calibration certificate	-
Obtained from the calibration certificate	-
The line temperature	°C
The reference temperature	°C
	Obtained from the calibration certificate Obtained from the calibration certificate The line temperature

The density value corrected for Velocity of Sound is calculated as follows:

$$\rho_{VOS} = \rho_t \left[1 + \frac{K_3}{(\rho_t + K_4)} \cdot \left(Kc - \frac{G}{t + 273} \right) \right]$$
$$Kc = \frac{Cc}{t_c + 273}$$

ρ _{vos}	The density corrected for temperature and VOS	kg/m3
К3	Obtained from the calibration certificate	-
K4	Obtained from the calibration certificate	-
Kc	Calibration gas constant from the calibration certificate	-
G	G value. Equals either input 'G value' or the ratio of inputs 'Specific gravity' and 'Ratio of specific heats', depending on input 'G value method'	-
t	The line temperature	°C
Cc	Specific Gravity/Ratio of specific heats of calibration gas	-
tc	Calibration temperature	°C

fxSolartron_Liquid _C

Description

The function calculates the density from a frequency input signal provided by a Solartron 7835, 7845, 7846 or 7847 liquid densitometer and corrects it for temperature, pressure and velocity of sound effects using US Customary units.

The function requires that the calibration constants are based on the following units:

•	Temperature	°F
-	remperature	

- Pressure psi -
- Density g/cc

Compliance

The calculations are in accordance with documents:

- 78355010 'Solartron 7835, 7845, 7846 & 7847 Advanced Liquid Density Transducer Manual', 2001
- 78355080, Rev. C, 'Micro Motion 7835, 7845, 7846 & 7847 . Liquid Density Transducer', October 2007

Function inputs				
Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Periodic time	In microseconds	μs	01e6	0
	Equals 1000 divided by the frequency in [Hz]			
Line temperature	Used when temperature correction is enabled	°F	-459.67+100	0
Line pressure	Used when pressure correction is enabled	psig	03000	0
Temperature correction	0: Disabled	-		1
	1: Enabled			
Pressure correction	0: Disabled	-		1
	1: Enabled			
VOS correction	0: Disabled	-		2
	1: Based on Kr / Kj Constants			
	Solartron manual edition 1985			
	2: Based on VOS value			
	Solartron manual edition 1996 and later			
Reference temperature	Used when temperature correction is enabled	°F	0200	60
Reference pressure	Used when pressure correction is enabled	psig	01500	0
ко	Constant from calibration certificate	-	-1e91e9	
K1	Constant from calibration certificate	-	-1e91e9	
K2	Constant from calibration certificate	-	-1e91e9	
K18	Constant from calibration certificate	-	-1e91e9	
K19	Constant from calibration certificate	-	-1e91e9	
K20A	Constant from calibration certificate	-	-1e91e9	
K20B	Constant from calibration certificate	-	-1e91e9	
K21A	Constant from calibration certificate	-	-1e91e9	
K21B	Constant from calibration certificate	-	-1e91e9	
Kr	Used when VOS method is set to 'Based on Kr / Kj Constants'.	-		
	Constant needs to be obtained from Solartron			
Kj	Used when VOS method is set to 'Based on Kr / Kj Constants'.	lbm/ft3		
2	Constant needs to be obtained from Solartron	- ,		
Liquid VOS	Velocity of sound of liquid	ft/s		
• • •	Used when VOS method is set to 'Based on VOS value'	-, -		

Function outputs

Function outputs	Remark			EU	SW tag	Alarm	Fallback
Status	0: Normal				STS		
	1: Input argument out of range					FIOOR	
	2: Calculation error					CALCERR	
Corrected density	Density corrected for temperature,	pressure	and VOS	lbm/ft3	CORDENS		0
Density corrected for temperature				lbm/ft3	TCORDENS		0
Density corrected for temperature and	d			lbm/ft3	PTCORDENS		0
pressure							
Uncorrected density				lbm/ft3	UNCDENS		0
Calculations		Whe	re:				
		- ρi	The indicat	ed density			g/cc
Density calculations are performed in g/cc and m/s, while the function inputs and outputs are in lbm/ft3 and ft/s		KO	Obtained f	rom the calibra	tion certificate		-
		K1	Obtained from the calibration certificate				-
		K2	Obtained f	rom the calibra	tion certificate		-
			The time pe	eriod in µS			μs

The indicated density ρ_i is calculated by $\rho_i = K0 + K1 \cdot \tau + K2 \cdot \tau^2$

$$\rho_t = \left[\rho_i \cdot \left[1 + K18 \cdot (t - t_R)\right] + K19 \cdot (t - t_R)\right]$$

Where:

ρt	The density corrected for temperature	g/cc
K18	Obtained from the calibration certificate	-
K19	Obtained from the calibration certificate	-
t	The line temperature	°F
t _R	The reference temperature	°F

The pressure and temperature corrected density ρ_{PT} is calculated by

$$\rho_{pt} = \left[\rho_t \cdot \left[1 + K20 \cdot \left(p - p_{REF}\right)\right] + K21 \cdot \left(p - p_{REF}\right)\right]$$

$$K20 = K20A + K20B \cdot (p - p_{REF})$$

$$K21 = K21A + K21B \cdot \left(p - p_{REF}\right)$$

Where:

ρ_{pt}	The density corrected for pressure and temperature	g/cc
K20A	Obtained from the calibration certificate	-
K20B	Obtained from the calibration certificate	-
K21A	Obtained from the calibration certificate	-
K21B	Obtained from the calibration certificate	-
р	The line pressure	psig
p _R	The reference pressure	psig

When 'VOS Correction' is set to '<u>Based on Kr and Kj Constants</u>' the following correction for velocity of sound is applied:

 $\rho_{VOS} = \rho_{pt} + K_{r.} \cdot (\rho_{pt} - K_j)^3$

When 'VOS Correction' is set to '<u>Based on VOS value</u>', the following correction for velocity of sound is applied:

$$\rho_{VOS} = \rho_{pt} \cdot \left[1 + \frac{1.4e^6}{1000 \cdot \rho_{pt} + 1.4} \cdot \left(\frac{1}{V_c^2} + \frac{1}{V_A^2} \right) \right]$$

When 0.3 <= ρ_{pt} <= 1.1: $V_C = (100 + 1455 \cdot \rho_{pt})$

When 1.1 < ρ_{pt} <= 1.6: $V_C = 2690 - 900 \cdot \rho_{pt}$

Else (ρ_{pt} < 0.3 or ρ_{pt} > 1.4) the VOS of sound correction is not performed.

ρvos	The density corrected for temperature	g/cc
Kr	Constant obtained from Solartron	-
Kj	Constant obtained from Solartron	g/cc
Vc	Calibration VOS	m/s
VA	Liquid VOS	m/s

fxSolartron_Liquid_M

The function calculates the density from a frequency input signal provided by a Solartron 7835, 7845, 7846 or 7847 liquid densitometer and corrects it for temperature, pressure and velocity of sound effects using metric units.

The function requires that the calibration constants are based on the following units:

°C

- Temperature •
 - Pressure bar
- Density kg/m3 •

. .

Compliance

The calculations are in accordance with documents:

- 78355010 'Solartron 7835, 7845, 7846 & 7847 Advanced • Liquid Density Transducer Manual', 2001
- 78355080, Rev. C, 'Micro Motion 7835, 7845, 7846 & 7847 ٠ Liquid Density Transducer', October 2007

Function inputs				
Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group			
Periodic time	In microseconds	μs	01e6	0
	Equals 1000 divided by the frequency in [Hz]			
Line temperature	Used when temperature correction is enabled	°C	-273.15+500	0
Line pressure	Used when pressure correction is enabled	bar(g)	0200	0
Temperature correction	0: Disabled	-		1
	1: Enabled			
Pressure correction	0: Disabled	-		1
	1: Enabled			
VOS correction	0: Disabled	-		2
	1: Based on Kr / Kj Constants			
	Solartron manual edition 1985			
	2: Based on VOS value			
	Solartron manual edition 1996 and later			
Reference temperature	Used when temperature correction is enabled	°C	0100	20
Reference pressure	Used when pressure correction is enabled	bar(g)	0100	0
ко	Constant from calibration certificate	-	-1e91e9	1.7418E2
К1	Constant from calibration certificate	-	-1e91e9	-1.10493e0
К2	Constant from calibration certificate	-	-1e91e9	3.703268e-4
К18	Constant from calibration certificate	-	-1e91e9	-6.415e-4
К19	Constant from calibration certificate	-	-1e91e9	-5.674e-1
K20A	Constant from calibration certificate	-	-1e91e9	2.888e-4
К20В	Constant from calibration certificate	-	-1e91e9	-5.581e-6
K21A	Constant from calibration certificate	-	-1e91e9	-4.467e-1
K21B	Constant from calibration certificate	-	-1e91e9	-8.633e-3
Kr	Used when VOS method is set to 'Based on Kr / Kj Constants'.	-	-1+1	0.0
	Constant needs to be obtained from Solartron			
Kj	Used when VOS method is set to 'Based on Kr / Kj Constants'.	kg/m3	02000	0.0
-	Constant needs to be obtained from Solartron			
Liquid VOS	Velocity of sound of liquid	m/s	02000	0.0
	Used when VOS method is set to ' Based on VOS value'			

Function outputs

Function outputs	Remark		EU	SW tag	Alarm	Fallback
Status	0: Normal			STS		
	1: Input argument out of range				FIOOR	
	2: Calculation error				CALCERR	
Corrected density	Density corrected for temperature, pressure and V	'OS	kg/m3	CORDENS		0
	(provided that particular correction is enabled)					
Density corrected for			kg/m3	TCORDENS		0
temperature						
Density corrected for			kg/m3	PTCORDENS		0
temperature and pressure						
Uncorrected density			kg/m3	UNCDENS		0
Calculations		Wher	e:			
The indicated density p	is calculated by	ρι	The indicated of	density		kg/m3
$\rho_i = K0 + K1 \cdot \tau + K2 \cdot \tau^2$		ко	Obtained from the calibration certificate		cate	-
$p_i = K0 + K1 \cdot t +$	K 2 · t	К1	Obtained from the calibration certificate		cate	-
		К2	Obtained from the calibration certificate		cate	-
		τ	The time perio	d in µS		μs

$$\rho_t = \left[\rho_i \cdot \left[1 + K18 \cdot (t - t_R)\right] + K19 \cdot (t - t_R)\right]$$

Where:

ρt	The density corrected for temperature	kg/m3
K18	Obtained from the calibration certificate	-
K19	Obtained from the calibration certificate	-
t	The line temperature	°C
t _R	The reference temperature	°C

The pressure and temperature corrected density ρ_{PT} is calculated by

$$\rho_{pt} = \left[\rho_t \cdot \left[1 + K20 \cdot \left(p - p_{REF}\right)\right] + K21 \cdot \left(p - p_{REF}\right)\right]$$

$$K20 = K20A + K20B \cdot (p - p_{REF})$$

$$K21 = K21A + K21B \cdot (p - p_{REF})$$

Where:

ρ_{pt}	The density corrected for pressure and temperature	kg/m3
K20A	Obtained from the calibration certificate	-
K20B	Obtained from the calibration certificate	-
K21A	Obtained from the calibration certificate	-
K21B	Obtained from the calibration certificate	-
р	The line pressure	bar(g)
p _R	The reference pressure	bar(g)

When 'VOS Correction' is set to '<u>Based on Kr and Kj Constants</u>' the following correction for velocity of sound is applied:

$$\rho_{VOS} = \rho_{pt} + K_{r.} \cdot (\rho_{pt} - K_j)^3$$

When 'VOS Correction' is set to '<u>Based on VOS value</u>', the following correction for velocity of sound is applied:

$$\rho_{VOS} = \rho_{pt} \cdot \left[1 + \frac{1.4e^6}{\rho_{pt} + 1400} \cdot \left(\frac{1}{V_c^2} - \frac{1}{V_A^2} \right) \right]$$

When 300 <= ρ_{pt} <= 1100: $V_C = 100 + 1.455 \cdot \rho_{pt}$

When 1100 < ρ_{pt} <= 1600: $V_c = 2690 - 0.9 \cdot \rho_{pt}$

Else (ρ_{pt} < 300 or ρ_{pt} > 1600) the VOS of sound correction is not performed.

Where	:	
ρvos	The density corrected for temperature	kg/m3
Kr	Constant obtained from Solartron	-
Kj	Constant obtained from Solartron	kg/m3
Vc	Calibration VOS	m/s
VA	Liquid VOS	m/s

fxSolartron_SG

Description

The function calculates the specific gravity from a frequency input signal provided by a Solartron 3096 or 3098 Specific Gravity transducer.

Compliance

The calculations are in accordance with documents:

- 30985020 'Solartron 3098 Gas Specific Gravity Transducer Manual', 2001
- 30985020, Rev. B, 'Micro Motion 3098 Gas Specific Gravity Meter', October 2007

Function input

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Periodic time	In microseconds	μs		01e6	0
	Equals 1000 divided by the frequency in [Hz]				
КО	Constant from calibration certificate	-	К0	-100000	-11.952
K2	Constant from calibration certificate	-	K2	01	4.719593e-5

Function output

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALCERR	
Specific Gravity	Ratio of the molecular weight of the gas (mixture) to that of the molecular	-	SG		0
	weight of dry air (i.e. the ideal Specific Gravity).				

-

-

-

μs

Calculations

Where:

τ κο

К2

The specific gravity is calculated by

Obtained from the calibration certificate

Obtained from the calibration certificate

 $SG = K0 + K2 \cdot \tau^2$

Specific gravity

Periodic time

If the calibration certificate contains factors Gx, Gy, tx and ty then K2 must be calculated as follows:

$$K2 = \frac{Gx - Gy}{t_x^2 - t_y^2}$$

Gx	Specific gravity of calibration gas x	-
Gy	Specific gravity of calibration gas y	-
τχ	Periodic time of calibration gas x	μs
τу	Periodic time of calibration gas y	μs

fxUGC_C

Description

The function calculates the density from a frequency input signal provided by a UGC densitometer and corrects it for temperature and pressure effects in **US customary** units.

Note: Calibration constants also need to be in US customary units.

Function input

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Periodic time	In microseconds	μs		01e6	
	Equals 1000 divided by the frequency in [Hz]				
Line temperature	Used when temperature correction is enabled	°F		-459.67+1000	
Line pressure	Used when pressure correction is enabled	psig		03000	
Temperature correction	0: Disabled	-	TEMPCOR		1
	1: Enabled				
Pressure correction	0: Disabled	-	PRESCOR		1
	1: Enabled				
Reference temperature	Used when temperature correction is enabled	°F	REFTEMP	0200	60
Reference pressure	Used when pressure correction is enabled	psig	REFPRES	01500	0
ко	Constant K0 from calibration certificate	-	KO	-1e91e9	
К1	Constant K1 from calibration certificate	-	K1	-1e91e9	
К2	Constant K2 from calibration certificate	-	K2	-1e91e9	
КТ1	Constant KT1 from calibration certificate		KT1	-1e91e9	
КТ2	Constant KT2 from calibration certificate		KT2	-1e91e9	
ктз	Constant KT3 from calibration certificate		KT3	-1e91e9	
KP1	Constant KP1 from calibration certificate		KP1	-1e91e9	
KP2	Constant KP2 from calibration certificate		KP2	-1e91e9	
КРЗ	Constant KP3 from calibration certificate		KP3	-1e91e9	

Function output

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal (No error condition)		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALC	
Corrected density	Density corrected for temperature and pressure	lbm/ft3	CORDENS		0
Uncorrected density	Uncorrected (indicated) density	lbm/ft3	UNCDENS		0

Calculations

The uncorrected density $\rho_{\rm i}$ is calculated by

 $\rho_i = K0 + K1 \cdot \tau + K2 \cdot \tau^2$

Whe	re:	
ρι	The uncorrected density	lbm/ft3
ко	Obtained from the calibration certificate	-
К1	Obtained from the calibration certificate	-
К2	Obtained from the calibration certificate	-
τ	The time period in µS	μs

The temperature and pressure corrected density ρt is calculated by

$$\rho_{i} = \rho_{i} + \left[K_{p_{1}} + K_{p_{2}} \cdot \rho_{i} + K_{p_{3}} \cdot \rho_{i}^{2}\right] \cdot \left(p - p_{R}\right) + \left[K_{T_{1}} + K_{T_{2}} \cdot \rho_{i} + K_{T_{3}} \cdot \rho_{i}^{2}\right] \cdot \left(t - t_{R}\right)$$
We are:

ρt	The density corrected for temperature and pressure	lbm/ft3
Кр1	Obtained from the calibration certificate	-
K _{P2}	Obtained from the calibration certificate	-
Крз	Obtained from the calibration certificate	-
Кті	Obtained from the calibration certificate	-
К т2	Obtained from the calibration certificate	-
Ктз	Obtained from the calibration certificate	-
t	The line temperature	°F
tr	The reference temperature	°F
р	The line pressure	psig
p _R	The reference pressure	psig

fxUGC_M

Description

The function calculates the density from a frequency input signal provided by a UGC densitometer and corrects it for temperature and pressure effects in **metric** units. Calibration constants also need to be in metric units.

Function input

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Periodic time	In microseconds	μs		01e6	
	Equals 1000 divided by the frequency in [Hz]				
Line temperature	Used when temperature correction is enabled	°C		-273.15+500	
Line pressure	Used when pressure correction is enabled	bar(g)		0200	
Temperature correction	0: Disabled	-	TEMPCOR		1
	1: Enabled				
Pressure correction	0: Disabled	-	PRESCOR		1
	1: Enabled				
Reference temperature	Used when temperature correction is enabled	°C	REFTEMP	0100	20
Reference pressure	Used when pressure correction is enabled	bar(g)	REFPRES	0100	0
ко	Constant K0 from calibration certificate	-	K0	-1e91e9	
K1	Constant K1 from calibration certificate	-	K1	-1e91e9	
K2	Constant K2 from calibration certificate	-	K2	-1e91e9	
KT1	Constant KT1 from calibration certificate		KT1	-1e91e9	
KT2	Constant KT2 from calibration certificate		KT2	-1e91e9	
ктз	Constant KT3 from calibration certificate		КТЗ	-1e91e9	
KP1	Constant KP1 from calibration certificate		KP1	-1e91e9	
KP2	Constant KP2 from calibration certificate		KP2	-1e91e9	
KP3	Constant KP3 from calibration certificate		KP3	-1e91e9	

Function output

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal (No error condition)		STS		
	1: Input argument out of range			FIOOR	
	2: Calculation error			CALC	
Corrected density	Density corrected for temperature and pressure	kg/m3	CORDENS		0
Uncorrected density	Uncorrected (indicated) density	kg/m3	UNCDENS		0

Calculations

The uncorrected density ρ_i is calculated by

$$\rho_i = K0 + K1 \cdot \tau + K2 \cdot \tau^2$$

The temperature and pressure corrected density pt is calculated

by

$$\rho_{i} = \rho_{i} + \left[K_{p_{1}} + K_{p_{2}} \cdot \rho_{i} + K_{p_{3}} \cdot \rho_{i}^{2}\right] \cdot (p - p_{R}) + \left[K_{T_{1}} + K_{T_{2}} \cdot \rho_{i} + K_{T_{3}} \cdot \rho_{i}^{2}\right] \cdot (t - t_{R})$$

Whe	re:	
ρι	The uncorrected density	kg/m3
ко	Obtained from the calibration certificate	-
К1	Obtained from the calibration certificate	-
К2	Obtained from the calibration certificate	-
τ	The time period in µS	μs

ρt	The density corrected for temperature and pressure	kg/m3
K _{P1}	Obtained from the calibration certificate	-
K _{P2}	Obtained from the calibration certificate	-
Крз	Obtained from the calibration certificate	-
Кті	Obtained from the calibration certificate	-
Ктг	Obtained from the calibration certificate	-
Ктз	Obtained from the calibration certificate	-
t	The line temperature	°C
t _R	The reference temperature	°C
р	The line pressure	bar(g)
P R	The reference pressure	bar(g)

fxVCone_C

Description

This function calculates the mass flow rate for a measured differential pressure over a McCrometer V-Cone meter in <u>U.S.</u> <u>customary</u> units.

The calculation, as specified by the meter supplier, is essentially a modified ISO 5167 flow rate calculation. As opposed to ISO-5167 the discharge coefficient is a function input. Because the discharge coefficient is a function of Reynolds number an optional calibration correction needs to be applied outside this function.

Function inputs

Compliance

• McCrometer: Flow Calculations for the V-Cone Flow meter Literature part #24509-54 Rev 3.1/02-05 2005

Function inputs	Remark	EU	Range	Default
Name	Optional tag name, tag description and tag group		-	
Differential Pressure	Differential pressure over the V-Cone device measured at the up- and downstream pressure taps	inH2O @ 60°F	0100	0
Pressure	Upstream pressure value of the fluid at metering conditions	psia	030000	0
Temperature	Down- or upstream temperature of the fluid at metering conditions	°F	-4002000	0
Density	Down or upstream density of the fluid at metering conditions	lbm/ft3	0200	0
Dynamic Viscosity	Dynamic viscosity of the fluid	lbm/ft.s	010	6.9e-6
Isentropic Exponent	Also referred to as κ (kappa). For an ideal gas this coefficient is equal to the ratio		010	1.3
	of the specific heat capacity at constant pressure to the specific heat at constant volume. According to the ISO standard this ratio may be used, when the real value is unknown.			
Pipe Diameter	Internal diameter of the pipe at reference temperature	inches	0100	0
Pipe Expansion factor	The thermal expansion coefficient of the pipe material	1/°F	01e-4	6.2e-6
Pipe Reference	The reference temperature that corresponds to the 'Pipe diameter' input value	°F	-4002000	68
temperature				
Cone Diameter	Cone diameter at reference temperature	inches	0 'Pipe Diameter'	0
Cone Expansion factor	The thermal expansion coefficient of the Cone material	1/°F	01e-4	9.25e-6
Cone Reference	The reference temperature that corresponds to the 'Cone diameter' input value	°F	-4002000	68
Temperature	provide the second s			
Configuration	The type of McCrometer V-Cone meter. This setting is used to select the appropriate equation for determination of the	-		1
	gas expansion factor as specified by McCrometer 1: Standard V-Cone 2: Wafer-Cone Note of input 'Fluid' is set to 'Liquid', then this input is not used (because the			
Pressure Location	expansion factor is set to 1) 1: Upstream			1
	 Input 'Pressure' represents the pressure at the upstream pressure tapping (p1). Since the absolute pressure is usually measured at the upstream tapping this is the most common setting. 2: Downstream Input 'Pressure' represents the pressure at the downstream tapping (p2). 			
Temperature Location	1: Upstream	-		2
	 Input 'Temperature' represents the upstream temperature (t₁). 2: Downstream Input 'Temperature represents the temperature at the downstream tapping (t₂). 3: Recovered Input 'Temperature' represents the downstream temperature at a location Where the pressure has fully recovered (t₃). Since temperature measurement is usually downstream of the flow device 			
	this is the most common setting.			
Temperature Correction	This parameter specifies if and how the temperature should be corrected from downstream to upstream conditions (or vice versa) 1: $(1-\kappa)/\kappa$			1
	Isentropic expansion using $(1-\kappa)/\kappa$ as the temperature referral exponent			
	 Constant Isentropic expansion using input 'Temperature Exponent' as the temperature referral exponent [-] 			
	3: Joule Thomson Isenthalpic expansion using input 'Temperature Exponent' as the Joule Thomson coefficient [°F/psi]. This method is prescribed by ISO5167-1:2003.			
Temperature Exponent	Refer to input Temperature Correction	- °E (nci		0
Donaity Location	Unit depends on input Temperature Correction value	°F/psi		1
Density Location	This parameter specifies if and how the density should be corrected from downstream to upstream conditions (or vice versa). 1: Upstream	-		1

Function inputs	Remark	EU	Range	Default
	Input 'Density' represents the density at the upstream pressure tapping (ρ_1).			
	2: Downstream			
	Input 'Density' represents the density at the downstream tapping (ρ_2).			
	3: Recovered			
	Input 'Density' represents the density downstream at a location Where the pressure has fully recovered ($ ho_3$).			
Density Exponent.	This factor is used when density correction is enabled. The formula $1/\kappa$ will be	-		0
	used when the input value is set to 0, else the input value will be used.			
	For more details refer to section 'Density correction'.			
Fluid	The type of fluid being measured	-		1
	1: Gas			
	2: Liquid			
Discharge coefficient	The McCrometer reference document states that the discharge coefficient is a	-	02	0.85
	function of Reynolds number. A calibration correction needs to be implemented			
	through an additional function and input 'Discharge coefficient' needs to be			
	linked to the corresponding output of this additional function.			

Function outputs

Function outputs	Remark	EU	Fallback	
Status	0: Normal (No error condition)			
	1: Input argument out of range			
	2: No convergence			
Mass flow rate	The calculated mass flow rate	klbm/hr	0	
Beta ratio	Cone to pipe diameter ratio at upstream temperature	-	Input Cone diameter / Input Pipe diameter	
Cone diameter	At the upstream temperature	inches	Input Cone diameter	
Pipe diameter	At the upstream temperature	inches	Input Pipe diameter	
Upstream pressure	Pressure at upstream tapping (p1)	psia	Input Pressure	
Pressure at downstream tapping	Pressure at downstream tapping (p2)	psia	Input Pressure	
Recovered downstream pressure	Fully recovered downstream pressure (p3)	psia	Input Pressure	
Upstream temperature	Temperature at upstream tapping (t1)	°F	Input Temperature	
Temperature at downstream	Temperature at downstream tapping (t_2)	°F	Input Temperature	
tapping				
Downstream Temperature	'Fully recovered' downstream temperature (t3)	°F	Input Temperature	
Upstream density	Density at upstream tapping (ρ_1)	lbm/ft3	Input Density	
Density at downstream	Pressure at downstream tapping (ρ₂)	lbm/ft3	Input Density	
tapping				
Downstream density	'Fully recovered' downstream density (ρ3)	lbm/ft3	Input Density	
Reynolds number		-	0	
Discharge coefficient	Same as input value	-		
Expansion Factor		-	0	
Velocity	Pipeline velocity	Ft/s	0	
Expansion Factor Range	The McCrometer reference document states that for gas applications the expansion	-	0	
	factor should not get below 0.84			
	0: Expansion factor is in valid range			
	1: Expansion factor is out of valid range			

Calculations

The flow calculation is as specified in the McCrometer reference document.

The downstream to upstream correction (and vice versa) for pressure, temperature and density are as specified for function 'ASME MFC-3M Orifice'.

fxVCone_M

Description

This function calculates the mass flow rate for a measured differential pressure over a McCrometer V-Cone meter in <u>metric</u> units.

The calculation, as specified by the meter supplier, is essentially a modified ISO 5167 flow rate calculation. As opposed to ISO-5167 the discharge coefficient is a function input. Because the discharge coefficient is a function of Reynolds number an optional calibration correction needs to be applied outside this function.

Compliance

• McCrometer: Flow Calculations for the V-Cone Flow meter Literature part #24509-54 Rev 3.1/02-05 2005

Function inputs	Remark	EU	Range	Default
ame	Optional tag name, tag description and tag group			
oifferential Pressure	Differential pressure over the V-Cone device measured at the up- and downstream	mbar	02000	0
	pressure taps			
ressure	Upstream pressure value of the fluid at metering conditions	bar(a)	02000	0
emperature	Down- or upstream temperature of the fluid at metering conditions	°C	-2401000	0
Density	Down or upstream density of the fluid at metering conditions	kg/m3	02000	0
Dynamic Viscosity	Dynamic viscosity of the fluid	Pa.s	01	0.01115
lsentropic Exponent	Also referred to as κ (kappa). For an ideal gas this coefficient is equal to the ratio	-	02	0
	of the specific heat capacity at constant pressure to the specific heat at constant			
	volume. According to the ISO standard this ratio may be used, when the real value			
	is unknown.			
lipe Diameter	Internal diameter of the pipe at reference temperature	mm	02000	0
ipe Expansion factor	The thermal expansion coefficient of the pipe material	1/°C	01	0.0000108
ipe Reference temperature	The reference temperature that corresponds to the 'Pipe diameter' input value	°C	-2401000	20
Cone Diameter	Cone diameter at reference temperature	mm	0	-
one Expansion factor	The thermal expansion coefficient of the Cone material	1/°C	240 1000	0.0000163
Cone Reference Temperature	The reference temperature that corresponds to the 'Cone diameter' input value	°C	-240 1000	20
Configuration	The type of McCrometer V-Cone meter.	-		1
	This setting is used to select the appropriate equation for determination of the			
	gas expansion factor as specified by McCrometer 1: Standard V-Cone			
	2: Wafer-Cone			
	Note of input 'Fluid' is set to 'Liquid', then this input is not used (because the			
	expansion factor is set to 1)			
Pressure Location	1: Upstream			1
Pressure Location	Input 'Pressure' represents the pressure at the upstream pressure tapping (p1).			1
	Since the absolute pressure is usually measured at the upstream tapping this is			
	the most common setting.			
	2: Downstream			
	Input 'Pressure' represents the pressure at the downstream tapping (p_2).			
emperature Location	1: Upstream	-		2
	Input 'Temperature' represents the upstream temperature (t_1).			
	2: Downstream			
	Input 'Temperature represents the temperature at the downstream tapping (t_2) .			
	3: Recovered			
	Input 'Temperature' represents the downstream temperature at a location			
	Where the pressure has fully recovered (t_3).			
	Since temperature measurement is usually downstream of the flow device this			
	is the most common setting.			
Temperature Correction	This parameter specifies if and how the temperature should be corrected from			1
	downstream to upstream conditions (or vice versa)			
	1: (1-κ)/κ			
	Isentropic expansion using (1- κ)/ κ as the temperature referral exponent			
	2: Constant			
	Isentropic expansion using input 'Temperature Exponent' as the temperature			
	referral exponent [-]			
	3: Joule Thomson			
	Isenthalpic expansion using input 'Temperature Exponent' as the Joule			
	Thomson coefficient [°C/bar].			
	This method is prescribed by ISO5167-1:2003.			
emperature Exponent	Refer to input Temperature Correction	- °C /h		0
	Unit depends on input Temperature Correction value	°C/bar		
Density Location	This parameter specifies if and how the density should be corrected from	-		1
	downstream to upstream conditions (or vice versa).			
	1: Upstream Input 'Density' represents the density at the upstream pressure tapping (ρ_1).			
	2: Downstream			
	Input 'Density' represents the density at the downstream tapping (ρ_2).			
	3: Recovered			
	Input 'Density' represents the density downstream at a location Where the			
	pressure has fully recovered (ρ_3).			
ensity Exponent.	This factor is used when density correction is enabled. The formula $1/\kappa$ will be used	-		0
Density Exponent.	when the input value is set to 0, else the input value will be used.			0
	For more details refer to section 'Density correction'.			
Fluid	The type of fluid being measured	-		1
	1: Gas			-
	2: Liquid			
Discharge coefficient	The discharge coefficient of the meter as specified by the manufacturer.	-	02	0.85
	The McCrometer reference document states that the discharge coefficient is a		•	0.00
	function of Reynolds number. It is advised that the meter is calibrated across the			
	range of Reynold numbers for which the meter is used.			
	The resulting correction curve can be implemented through function			
	fxInterolationCurve. Input 'Discharge coefficient' needs to be linked to the			

Function outputs

Function outputs	Remark	EU	Fallback
Status	0: Normal (No error condition)		
	1: Input argument out of range		
	2: No convergence		
Mass flow rate	The calculated mass flow rate	tonne/h	0
Beta ratio	Cone to pipe diameter ratio at upstream temperature	-	Input Cone diameter /
			Input Pipe diameter
Cone diameter	At the upstream temperature	mm	Input Cone diameter
Pipe diameter	At the upstream temperature	mm	Input Pipe diameter
Upstream pressure	Pressure at upstream tapping (p1)	bar(a)	Input Pressure
Pressure at downstream tapping	Pressure at downstream tapping (p2)	bar(a)	Input Pressure
Recovered downstream pressure	Fully recovered downstream pressure (p3)	bar(a)	Input Pressure
Upstream temperature	Temperature at upstream tapping (t1)	°C	Input Temperature
Temperature at downstream tapping	Temperature at downstream tapping (t2)	°C	Input Temperature
Downstream Temperature	'Fully recovered' downstream temperature (t ₃)	°C	Input Temperature
Upstream density	Density at upstream tapping ($ ho_1$)	kg/m3	Input Density
Density at downstream tapping	Pressure at downstream tapping (ρ_2)	kg/m3	Input Density
Downstream density	'Fully recovered' downstream density (ρ_3)	kg/m3	Input Density
Reynolds number		-	0
Discharge coefficient	Same as input value	-	
Expansion Factor		-	0
Velocity	Pipeline velocity	m/s	0
Expansion Factor Range	The McCrometer reference document states that for gas applications the expansion factor should not get below 0.84	-	0
	0: Expansion factor is in valid range		
	1: Expansion factor is out of valid range		

Calculations

The flow calculation is as specified in the McCrometer reference document.

The downstream to upstream correction (and vice versa) for pressure, temperature and density are as specified for function 'ISO 5167 Orifice'.

3 Flow-X flow computer functions

This chapter lists all available general Spirit^{IT} Flow-X functions. These functions are only available in the Flow-X flow computer software.

fx2CellSelection

Description

The function selects between 2 input cells (e.g. differential pressure cells) based on the actual measured value and the failure status of each cell.

The function can handle the following type of cell range configurations:

- Lo-Hi
- Hi Hi

Where 'Lo' means low range, 'Mid' mid range and 'Hi' high range.

Function inputs	Remark	EU	SW tag	Range	Default
Name					
Cell A value	Input value as percentage of span of cell A				
lame iell A value I iell A status I iell B value I iell B status I iell B status I iange type I iange I	Input status of cell A				
	0: Normal				
	<> 0 : Failure				
Cell B value	Input value as percentage of span of cell B				
Cell B status	Input status of cell B				
	0: Normal				
	<> 0 : Failure				
Range type	For a description of the functionality refer to adjacent section 'Logic'		RNGTYP		
	1: Lo Hi				
	Cell A at low range				
	Cell B at high range				
	2: Hi Hi				
	Cell A and B at same range				
Auto switchback	For a description of the functionality refer to adjacent section 'Logic'				
	0: Disabled				
	1: Enabled				
Switch-up percentage	Switch-up value expressed as percentage of span of the lower range	-	SWUPPERC	0100	95
Switch-down percentage	Switch-down value expressed as percentage of span of the lower range	-	SWDNPERC	0100	90

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
Selected cell number	1: Cell 1		SELNR		1
	2: Cell 2				
Selected cell status	0: Normal		SELSTS		0
	1: Failure				

Logic

The function will switch from one cell to another at the following conditions:

Range type = 'Lo Hi'

When cell A is currently selected

- Select cell B when cell A value is above or equal to the switchup percentage of its range and cell B is healthy.
- Select cell B when cell A fails while cell B is healthy

When cell B is currently selected

- Select cell A when cell A value is below or equal to the switchdown percentage of its range and cell A is healthy
- Select cell A when cell B fails and cell A is healthy

Range type = 'Hi Hi'

When cell A is currently selected

• Select cell B when cell A value fails and cell B is healthy

When cell B is currently selected

- Select cell A when cell A is healthy and 'Auto switchback' is enabled
- Select cell A when cell B fails and cell A is healthy.

fx3CellSelection

Description

The function selects between 3 input cells (typically differential pressure cells) based on the actual measured value and the failure status of each cell.

The function can handle the following type of cell range configurations:

- Lo Mid Hi
- Lo Hi Hi
- Hi Hi Hi

Where 'Lo' means low range, 'Mid' mid range and 'Hi' high range.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name					
Cell A value	Input value as percentage of span of cell A				
Cell A status	Input status of cell A				
	0: Normal				
	<> 0 : Failure				
Cell B value	Input value as percentage of span of cell B				
Cell B status	Input status of cell B				
	0: Normal				
	<> 0 : Failure				
Cell C value	Input value as percentage of span of cell C				
Cell C status	Input status of cell C				
	0: Normal				
	<> 0 : Failure				
Range type	For a description of the functionality refer to adjacent section		RNGTYP		
	'Logic'				
	1: Lo Mid Hi				
	Cell A at low range				
	Cell B at mid range				
	Cell C at high range				
	2: Lo Hi Hi				
	Cell A at low range				
	Cell B and C at high range				
	3: Hi Hi Hi				
	Cell A, B and C at same range				
Auto switchback	For a description of the functionality refer to adjacent section				
	'Logic'				
	0: Disabled				
	1: Enabled	s of cell B as percentage of span of cell C s of cell C ie iption of the functionality refer to adjacent section NNGTYP low range mid range high range low range d C at high range and C at same range iption of the functionality refer to adjacent section ralue expressed as percentage of span of the lower ralue expressed as percentage of span of the lower ralue expressed as percentage of span of the lower ralue expressed as percentage of span of the lower ralue expressed as percentage of span of the lower ralue expressed as percentage of span of the lower ralue expressed as percentage of span of the - SWDNPERC			
Switch-up percentage	Switch-up value expressed as percentage of span of the lower	-	SWUPPERC	0100	95
	range				
	Does not apply for selection type 'Hi Hi Hi'				
Switch-down percentage	Switch-down value expressed as percentage of span of the	-	SWDNPERC	0100	90
	lower range				
	Does not apply for selection type 'Hi Hi Hi'				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			FIOOR	
Selected cell number	1: Cell 1		SELNR		1
	2: Cell 2				
	3: Cell 3				
Selected cell status	0: Normal		SELSTS		0
	1: Failure				

Logic

The function will switch from one cell to another at the following conditions:

Range type = 'Lo Mid Hi'

When cell A is currently selected

- Select cell B when cell A value is above or equal to the switchup percentage of its range and cell B is healthy.
- Select cell B when cell A fails while cell B is healthy
- Select cell C when cell A and cell B fail and cell C is healthy

When cell B is currently selected

- Select cell C when cell B value is above or equal to the switch-up percentage of its range and cell C is healthy
- Select cell A when cell A value is below or equal to the switchdown percentage of its range and cell A is healthy
- Select cell A when cell B fails while cell A is healthy
- Select cell C when cell B and cell A fail and cell C is healthy

When cell C is currently selected

- Select cell B when cell B value is below or equal to the switchdown percentage of its range and cell B is healthy
- Select cell B when cell C fails while cell B is healthy
- Select cell A when cell C and cell B fail and cell A is healthy

Range type = 'Lo Hi Hi'

When cell A is currently selected

- Select cell B when cell A value is above or equal to the switchup percentage of its range and cell B is healthy.
- Select cell C when cell A value is above or equal to the switch-up percentage of its range and cell B fails and cell C is healthy.
- Select cell B when cell A fails while cell B is healthy
- Select cell C when cell A and cell B fail and cell C is healthy

When cell B is currently selected

- Select cell A when cell A value is below or equal to the switchdown percentage of its range and cell A is healthy
- Select cell C when cell B fails while cell C is healthy
- Select cell A when cell B and cell C fail and cell A is healthy

When cell C is currently selected

- Select cell A when cell A value is below or equal to the switchdown percentage of its range and cell A is healthy
- Select cell B when cell B is healthy and 'Auto switchback' is enabled
- Select cell A when cell C and cell B fail and cell A is healthy

Range type = 'Hi Hi Hi'

When cell A is currently selected

- Select cell B when cell A value fails and cell B is healthy
- Select cell C when cell A and cell B fail and cell C is healthy

When cell B is currently selected

- Select cell A when cell A is healthy and 'Auto switchback' is enabled
- Select cell A when cell B fails and cell A is healthy
- Select cell C when cell B and A fail and cell C is healthy

When cell C is currently selected

- Select cell A when cell A is healthy and 'Auto switchback' is enabled
- Select cell B when cell B is healthy and cell A fails and 'Auto switchback' is enabled
- Select cell A when cell C fails and cell A is healthy
- Select cell B when cell C and A fail and cell B is healthy

fxBatchFWA

Description

The function calculates a **f**low-**w**eighted **a**verage (FWA) for a batch.

A batch can be any batch type of process, such as product loading, meter proving or transmitter validation. The function weights the input value with a flow increment and updates the average accordingly. The flow increment is provided by either a 'fxTotalizerDelta' or a 'xTotalizerRate' function.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name used for tag-prefix and retentive storage.				
Input value	Value to be averaged	Same as linked cell		-1e111e11	
Enabled	0: Disabled		EN		
	1: Enabled				
Increment	Flow increment with which the input value is weighed.	Same as linked cell		01e11	
	Must refer to the corresponding output from a 'TotalizerRate' or				
	'TotalizerDelta' function				
	Negative values will be ignored.				
nput value inabled ncrement Reset command	Trigger to reset the batch.				
	At a batch reset the current average is stored in the previous				
	value and the current value is reset to 0				
Identification	Batch identification.		ID		
	Can be any string of maximum 255 characters long.				
	If multiple identifications need to be stored for future referral,				
	e.g. the batch number, the ship name and the nomination				
	number, then the individual strings should be concatenated with				
	an "." character in-between.				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Current average	Average calculated over the current batch.	Same as input 'Input value'	СВ		0
Previous average	Average of the previous batch.	Same as input 'Input value'	PB		0

fxBatchHistData

Description

The function retrieves historical 'batch' data from the flow computer persistent memory.

A 'batch' can be any batch type of process, such as product loading, meter proving or transmitter validation. The function retrieves one or more historical values for the specified function instance. The function instance must be one of the following function types:

- fxBatchFWA
- fxBatchLatch
- fxBatchStore
- fxBatchTotal
- fxBatchTWA
- fxBatchWatch

The function instance is referred to by its name (i.e. the 1st argument of the referred function).

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name used for tag-prefix and retentive storage.				
First ID	Optional.		FIRSTID		
	Batch identification of the first historical batch for which the value				
	has to be retrieved.				
Last ID	Optional.		LASTID		
	Batch identification of the last historical batch for which the value				
	has to be retrieved.				
Sequence	Sequence in which the retrieved values must be copied to the				1
	function outputs.				
	1: Ascending order (Value 1 contains oldest value)				
	2: Descending order (Value 1 contains newest value)				

Function outputs	Remark	EU	SW tag	Alarm	Fallback	
Number of values	Number of historical values that was retrieved from the flow computer memory according to the input criteria		ACTSIZE			
Value 1	The 1st retrieved historical data value		1			
Value 2	The 2nd retrieved historical data value		2			
etc.						

fxBatchLatch

Description

The function latches a value at every batch reset.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name used for tag-prefix and retentive storage.				
Input value	Value to be latched	Same as linked cell			
Latch command *	Trigger to latch the value				
Reset command *	Trigger to reset the batch.				
	At every batch reset the last latched value is stored in the				
	previous latch output value and the current latch output				
	value is reset to 0				
Identification	Batch identification.		ID		
	Can be any string of maximum 255 characters long.				

Note: When the latch and reset commands are given at the same time, then the current value becomes the 'Previous latch' output value and the current latch is reset to 0.

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Current latch	Value that is latched since the last batch reset.	Same as input 'Input value'	СВ		0
	Is reset to 0 at every batch reset.				
Previous latch	Value that was latched during the previous batch. If no value was latched during the previous batch, then the value is set to 0.	Same as input 'Input value'	РВ		0

fxBatchMax

Description

The function determines the maximum for a particular input value over a batch.

A batch can be any batch type of process, such as product

loading, meter proving or transmitter validation.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name used for tag-prefix and retentive storage.				
Input value	Value for which the maximum has to be determined	Same as linked cell		-1e111e11	
nput value Enabled Reset command	0: Disabled		EN		
	1: Enabled				
Reset command	Trigger to reset the batch.				
	At a batch reset the current average is stored in the previo	us			
	value and the current value is reset to 0				
dentification	Batch identification.		ID		
	Can be any string of maximum 255 characters long.				
	If multiple identifications need to be stored for future				
	referral, e.g. the batch number, the ship name and the				
	nomination number, then the individual strings should be				
	concatenated with an "." character in-between.				

Function outputs	Remark	EU	SW tag	Alarm	Fallback	
Current minimum	Minimum over the current batch.	Same as input 'Input value'	СВ		0	
Previous minimum	Minimum over the previous batch.	Same as input 'Input value'	PB		0	

fxBatchMin

Description

The function determines the minimum for a particular input value over a batch.

A batch can be any batch type of process, such as product

loading, meter proving or transmitter validation.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name used for tag-prefix and retentive storage.				
Input value	Value for which the minimum has to be determined	Same as linked cell		-1e111e11	
Enabled	0: Disabled		EN		
	1: Enabled				
Reset command	Trigger to reset the batch.				
	At a batch reset the current average is stored in the				
	previous value and the current value is reset to 0				
Identification	Batch identification.		ID		
	Can be any string of maximum 255 characters long.				
	If multiple identifications need to be stored for future				
	referral, e.g. the batch number, the ship name and the				
	nomination number, then the individual strings				
	should be concatenated with an "." character in-				
	between.				

Function outputs	Remark	EU	SW tag	Alarm	Fallback	
Current minimum	Minimum over the current batch.	Same as input 'Input value'	СВ		0	
Previous minimum	Minimum over the previous batch.	Same as input 'Input value'	PB		0	

fxBatchTotal

Description

The function accumulates a flow increment into a batch total. At every batch reset the current batch total is stored into the previous value and the current value is reset to 0. The flow increment originates from a 'TotalizerRate' or

'TotalizerDelta' function.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name used for tag-prefix and retentive storage.				
Increment	Increment value to be added to the batch total.	Same as linked cel		01e11	
	Negative values will be ignored, so the batch total will not decrease.				
Enabled	0: Disabled		EN		
	1: Enabled				
Identification	Batch identification.		ID		
	Can be any string of maximum 255 characters long.				
	If multiple identifications need to be stored for future referral, e.g. the				
	batch number, the ship name and the nomination number, then the				
	individual strings should be concatenated with an "." character in-				
	between.				
Rollover value	The batch total will be reset to 0 when it reaches the rollover value	Same as input	ROVAL	01e15	1e12
		Increment			
Decimal places	Defines the number of decimal places for the current and previous total		DECPLS	-110	-1
	output values.				
	-1 means full precision (no rounding applied)				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Current total	Accumulated total for the current batch, so since the last batch reset.	Same as input 'Increment'	СВ		0
Previous total	Accumulated total for the previous batch	Same as input 'Increment'	PB		0
Rollover flag	Flag indicating a rollover to 0. 0: Off 1: On Note: stays 'On' for one calculation cycle only)	0		ROALM	

fxBatchTWA

Description

The function calculates a time-**w**eighted **a**verage (TWA) for a batch. At a batch reset the current average is stored in the previous value and the current value is reset to 0. The function weights the input value with the time (in fact the actual calculation cycle time) and updates the average

accordingly.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name used for tag-prefix and retentive storage.				
Input value	Value to be averaged	Same as linked cell		-1e111e11	
Enabled	0: Disabled		EN		
	1: Enabled				
Reset command	Trigger to reset the batch.				
	At a batch reset the current latch is stored in the previous value and the				
	current latch is reset to 0				
Identification	Batch identification.		ID		
	Can be any string of maximum 255 characters long.				
	If multiple identifications need to be stored for future referral, e.g. the				
	batch number, the ship name and the nomination number, then the				
	individual strings should be concatenated with an "." character in-				
	between.				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Current average	Average calculated over the current batch.	Same as input 'Input value'	СВ		0
Previous average	Average of the previous batch.	Same as input 'Input value'	PB		0

fxBatchWatch

Description

The function 'remembers' that a condition has been valid during a batch.

A typical example is a transmitter that was overridden with a

keypad value.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name used for tag-prefix and retentive storage.				
Condition	0: Condition is not valid				
	<>0: Condition is valid				
nabled	0: Disabled	EN			
	1: Enabled				
Reset command	Trigger to reset the batch.				
	At a batch reset the current watched value is stored in the previous value and the				
	current watched value is reset to 0				
Identification	Batch identification.		ID		
	Can be any string of maximum 255 characters long.				
	If multiple identifications need to be stored for future referral, e.g. the batch				
	number, the ship name and the nomination number, then the individual strings				
	should be concatenated with an "." character in-between.				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Current watch	Indicates whether or not the condition has been valid during the current batch:		СВ		0
	0: Not valid				
	1: Valid				
Previous watch	Indicates whether or not the condition has been valid during the previous batch:		PB		0
	0: Not valid				
	1: Valid				

fxDeviationAlarm

Description

The function watches the deviation, or the difference or discrepancy, between two values and generates an alarm when the deviation exceeds the specified limit.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag				
	group				
Input value 1	Must be linked to another cell	Same as linked cell		-1e111e11	0
Input value 2	Must be linked to another cell	Must be same as for input value 1		-1e111e11	0
Deviation type	Determines whether the absolute or the relative		DEVTYP		1
	difference needs to be checked. The unit of the				
	deviation limit will be in accordance.				
	1: Absolute				
	2: Relative				
Deviation limit	The unit depends on the 'Deviation type'	Absolute: Same as input value 1	DEVLIM	01e11	0
		Relative : %			
Enabled	Enabled or disabled the alarm			True or false	True
Alarm type	1: Alarm				1
	2: Warning				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Deviation alarm	0: Normal	-	DEVALM		
	1: Alarm			DEVALM	

Logic

• A deviation alarm is raised when the absolute difference between the two values is greater than the 'Discrepancy limit'.

Deviation type = 2 (Relative)

• A deviation alarm is raised when the absolute difference between the two values divided by the minimum of the two values times 100 % is greater than the 'Discrepancy limit'.

Deviation type = 1 (Absolute)

fxGenerateReport

Description

This function generates prints and stores a report.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Report definition				
	Must be the name of the report definition (Flow-Xpress, section Reports).				
Event	Event to generate the report.				
	Event occurs when value changes from zero to non-zero (or from FALSE to TRUE).				
Identifier	Optional report file name, defined as a string				
	When defined the Identifier is used as the report file name .				
	When left empty, the UniqueMethod setting as defined for the report definition is used				
	for the report file name (Flow-Xpress, section Reports).				
Printer	Optional printer.				
	Must be the name of one of the printers that are defined in Flow-Xpress .				
	When defined this printer is used instead of the printer that is assigned to the report				
	template (Flow-Xpress , section Reports).				
	When left empty the printer that is assigned to the report template is used.				
Number of copies	Number of copies to print.				1
	This setting is ignored when no printer is defined.				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range or in conflict			FIOOR	
	2: Latest report could not be generated			RPTERR	

fxIndex

Description

The 'fxIndex' provides the same functionality as the Excel Index function with the addition of the creation of tag names.

A spreadsheet cell that contains a 'fxIndex' function obtains the tag name as defined by its 'Name' input with the tag inheriting all properties including the value and units from the referred tag. When the referred cell contains a writable tag (i.e. a value and no function) than the cell with the 'fxIndex' function also represents a writable tag with the same properties.

When the referred cell contains a tag that represents one or more alarms, then the same alarms are created for the cell with the 'fxIndex' function.

The 'fxIndex' is especially useful for setting up generic (template) applications as illustrated by the following examples:

- For each of the 6 analog inputs the application contains one 'Analog input' function that generates tag names with prefix, "AIN1_", "AIN2_etc. Also more meaningful tag names such as "..PT" (pressure transmitter), ".._TT" are used in the application. When changing the high alarm limit for the pressure transmitter (e.g. through an OPC interface) it makes more sense to address the tag as "..PT_HISCALE" instead of "..AIN3_HISCALE" (assuming AIN 3 being used for the pressure transmitter). This can be achieved by using the 'fxIndex' function for the cell that represents the '..PT_HISCALE' tag.
- In some cases one and the same input signal is used for multiple process variables that are defined in the generic application. E.g. when the generic application assumes a prover inlet temperature input signal as well as a prover outlet temperature signal (and has corresponding tag names), while there is only a temperature transmitter in the prover loop, then the input tags of both signals can refer to the same "AIN" signal by using the 'fxIndex' function.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Tag name for the cell that contains the function				
Reference	Reference to one or more cell ranges				1
Row number	Optional				1
	Number of the row in reference from which to return a reference				
Column number	Optional.				1
	Number of the column in reference from which to return a reference				
Area number	Optional				1
	Selects a range in reference from which to return the intersection of 'Row				
	number' and 'Column number'. The first area selected or entered is numbered				
	1, the second is 2, and so on.				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Value	The cell obtains a tag name that consist of the prefix as defined by input 'Name' with the addition of the suffix as was generated for the tag that is being referred to. The same applies for an alarm if one has been defined for the referred tag.	Same as referred cell	See Remark		

fxKeypadFallback

Description

This function provides a generic interface to any input signal, such as a pressure, temperature, density or flow input. It provides the option to override the 'live' value with a keypad value and to fallback to a specific value when the input value is faulty.

When the input signal fails, the in-use value may fall back to the last good value, the keypad value or additionally a separate fallback value. The fallback value allows the user to define a fixed value (e.g. 0) that is independent from the current keypad value.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Input status	Status of he input signal		INPSTS		
	0: Normal				
	<> 0: Failure				
	Must be linked to the Status output of the related input function				
Input value	Value of the input signal		INPVAL		
	Must be linked to the (scaled) value output of the related input function.				
Fallback type	Determines what to do when input fails		FBTYP		
	1: Use last good value				
	2: Use fallback value				
	3: Use keypad value				
	4: Use measured				
Fallback value	Used when output 'Input status' becomes 'Faulty' and 'Fallback type' is set to	Same as	FBVAL		
	'Use fallback value'	input			
Keypad mode	Forces the usage of the keypad value		KPMOD	-	0
	0: Disabled				
	1: Enabled				
Keypad value	Used when output 'Input status' becomes 'Faulty' and 'Fallback type' is set to	Same as	KPVAL		0
	'Use fallback value'	input			

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Argument out of range			FIOOR	
	2: Keypad			FAILLG	
	3: Fail Last Good			FAILKP	
	4: Fail Keypad			FAILFB	
	5: Fail Fallback			KEYPAD	
	6: Fail Measured			FAILMS	
	Only for status 'Function Input argument out of range' the output				
	vales will revert to the corresponding fallback value.				
n-use value		Same as input	CUR		0

fxKeypadFallbackArray

Description

This function provides a generic interface to an array of input values, typically a gas composition. It provides the option to override the 'live' values with keypad values and to fallback to specific values when the set of input values is faulty. When the input signals fail, the in-use values may fall back to the last good values, the keypad values or additionally separate fallback values. The fallback values allow the user to define a fixed values (e.g. 0) that is independent from the current keypad values.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Input status	Status of the input signals		INPSTS		
	0: Normal				
	<> 0: Failure				
Input values	Array if input values		INPVAL		
Fallback type	Determines what to do when input fails		FBTYP		
	1: Use last good value				
	2: Use fallback value				
	3: Use keypad value				
	4: Use measured				
Fallback value	Array of fallback values.	Same as input	FBVAL		
	Used when output 'Input status' becomes 'Faulty' and 'Fallback				
	type' is set to 'Use fallback value'				
Keypad mode	Forces the usage of the keypad value		KPMOD	-	0
	0: Disabled				
	1: Enabled				
Keypad value	Array of keypad values.	Same as input	KPVAL		0
	Used when output 'Input status' becomes 'Faulty' and 'Fallback				
	type' is set to 'Use fallback value'				

Function outputs	Remark	EU	SW tag	Alarm	Fallback	
Status	0: Normal		STS			
	1: Argument out of range			FIOOR		
	2: Keypad			FAILLG		
	3: Fail Last Good			FAILKP		
	4: Fail Keypad			FAILFB		
	5: Fail Fallback			KEYPAD		
	6: Fail Measured			FAILMS		
	Only for status 'Function Input argument					
	out of range' the output vales will revert					
	to the corresponding fallback value.					
In-use value	Array of in-use values	Same as input	CUR		0	

fxLatch

Description

The 'fxLatch' function provides generic latching functionality.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Tag name for the cell that contains the 'fxLatchValue' function.				
Latch trigger	Trigger to latch the input value				
Input value	Value to be latched. May be a constant a formula or a reference to another cell or tag.				
Reset trigger	Optional				
	Trigger to resets the latched value to the reset value				
Reset value	Optional				
	Reset value (default 0). May be a constant or a formula.				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Latched value	The most recent latched value	Same as of input	Input Name		
	Note: this value is persistent and will be reloaded upon startup	Input Value			

fxLimitAlarm

Description

The function applies alarm limits on any value.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name				
Input value	Must be linked to another cell	Same as linked cell	INPVAL	-1e111e11	0
Low limit value		Same as Input value	LLIM	-1e111e11	-1e11
High limit value		Same as Input value	HLIM	Low limit value 1e11	-1e11
Low low limit value		Same as Input value	LLLIM	-1e11 Low limit value	1e11
High high limit value		Same as Input value	HHLIM	High limit value 1e11	1e11
Deadband		Same as Input value		01e11	0
Enabled	Enables or disables the alarm			True or false	True
Warning behavior	Determines the warning behavior when the lo and hi limit are violated: 1: No warnings, just alarms 2: Warn on lo limit 3: Warn on hi limit 4: Warn on lo and hi limit				1

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Alarm status	0: Normal	-	ALMSTS		
	1: Low alarm			LALM	
	2: High alarm			HALM	
	3: Low low alarm			LLALM	
	4: High high alarm			HHALM	

Logic

The order of priority in setting the alarm status output is as follows:

- 1 If process value is below the 'Low low limit value' then the status becomes '3: Low low alarm'.
- 2 Else if process value is above the 'High high limit value' then the status becomes '4: High high alarm'.
- 3 Else if process value is below the 'Low limit value' then the status becomes '1: Low alarm'.
- 4 Else if process value is above the 'High limit value' then the status becomes '2: High alarm' and the 'High high alarm' is raised
- 5 Else the status becomes '0: Normal'.

fxName

Description

The 'fxName' function creates a string that defines the prefix, description and group for any function that generates tags,

except for function fxTag.

Function outputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name for the tag				
Description	Optional description for the tag				<empty></empty>
Group	Optional Group for the tag, including optional parent groups.				<empty></empty>
	The parent group must proceed the child group and be separated by the '\'				
	character.				
	E.g. "Meter setup\Meter data" defines that the tag belongs to group 'Meter data',				
	which is a subgroup of group "Meter setup".				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
	String containing the name, description and group for the tag(s) to be				
	created.				

fxPeriodFWA

Description

The function calculates a **f**low-**w**eighted **a**verage (FWA) for a particular period.

The function weights the input value with a flow increment and

updates the average accordingly. The flow increment is provided

by a 'TotalizerDelta' or 'TotalizerRate' function.

At the end of the period the current average is stored in the

previous value and the current value is reset to 0.

Function inputs

Function inputs	Remark		EU	SW tag	Range	Default
Name	Name us	ed for tag prefix and retentive storage.				
Input value	Value to	be averaged	Same as linked cell			
Enabled	0:	Disabled		EN		
	<> 0:	Enabled				
Increment	Flow inc	rement with which the input value is weighed.	Same as linked cell			
	Must ref	er to the corresponding output from a 'TotalizerRate' or				
	'Totalize	rDelta' function				
	Negative	values will be ignored.				
Period type	Type of I	beriod:		TYP		
	1: Secon	t de la constante de				
	2: Minute					
	3: Hour					
	4: Day					
	5: Week					
	6: Month					
	7: Quarte	er				
	8: Year					
Period count	Number	of periods (e.g. 5 minutes, 8 hours)		CNT	11e11	
Period start	Absolute	start date and time of the period. This will be used as the		START	<datetime></datetime>	
	referenc	e point to calculate the next period rollover from.				
	The valu	e may be defined in the past or the future. The next rollover				
	period w time).	ill be calculated accordingly (so forwards or backwards in				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Current average	Average calculated over the current period.	Same as input 'Input value'	CUR		0
Previous average	Average of the previous period.	Same as input 'Input value'	PRV		0
Pre-previous average	Average of the pre-previous period (i.e. the period before the previous period).	Same as input 'Input value'	PPRV		0

fxPeriodLatch

Description

The function latches a value at the end of a repeating period of time.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name used for tag prefix and retentive storage.				
Input value	Value to be latched	Same as linked cell		Not applied	
Period type	Type of period:		TYP		
	1: Second				
	2: Minute				
	3: Hour				
	4: Day				
	5: Week				
	6: Month				
	7: Quarter				
	8: Year				
Period count	Number of periods (e.g. 5 minutes, 8 hours)		CNT	11e11	
Period start	Absolute start date and time of the period. This will be used		START	<datetime></datetime>	
	as the reference point to calculate the next period rollover				
	from.				
	The value may be defined in the past or the future. The next				
	rollover period will be calculated accordingly (so forwards or				
	backwards in time).				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Current latch	Value that is latched at start of the current period (end of previous period)	Same as input 'Input value'	CUR		0
Previous latch	Value that is latched at the start of the previous period	Same as input 'Input value'	PRV		0

fxPeriodMax

Description

The function determines a maximum value over a particular

period.

Function inputs

Function inputs	Remark		EU	SW tag	Range	Default
Name	Name us	ed for tag prefix and retentive storage.				
Input value	Value fo	r which the maximum has to be determined	Same as linke	d		
			cell			
Enabled	0:	Disabled		EN		
	<> 0:	Enabled				
Period type	Type of I	period:		TYP		
	1: Secon	d				
	2: Minute	e				
	3: Hour					
	4: Day					
	5: Week					
	6: Month	1				
	7: Quarte	er				
	8: Year					
Period count	Number	of periods (e.g. 5 minutes, 8 hours)		CNT	11e11	
Period start	Absolute	e start date and time of the period. This will be used as the		START	<datetime></datetime>	
	referenc	e point to calculate the next period rollover from.				
	The valu	e may be defined in the past or the future. The next rollover				
	period w	ill be calculated accordingly (so forwards or backwards in tim	ie).			

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Current maximum	Maximum over the current period.	Same as input 'Input value'	CUR		0
Previous maximum	Maximum over the previous period.	Same as input 'Input value'	PRV		0
Pre-previous maximum	Maximum over the pre-previous period (period before the previous period).	Same as input 'Input value'	PPRV		0

fxPeriodMin

Description

The function determines a minimum value over a particular period.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name used for tag prefix and retentive storage.				
Input value	Value for which the minimum has to be determined	Same as			
-		linked cell			
Enabled	0: Disabled		EN		
	<> 0: Enabled				
Period type	Type of period:		TYP		
	1: Second				
	2: Minute				
	3: Hour				
	4: Day				
	5: Week				
	6: Month				
	7: Quarter				
	8: Year				
Period count	Number of periods (e.g. 5 minutes, 8 hours)		CNT	11e11	
Period start	Absolute start date and time of the period. This will be used as the		START	<datetime></datetime>	
	reference point to calculate the next period rollover from.				
	The value may be defined in the past or the future. The next rollover				
	period will be calculated accordingly (so forwards or backwards in time	e).			

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Current minimum	Minimum over the current period.	Same as input 'Input value'	CUR		0
Previous minimum	Minimum over the previous period.	Same as input 'Input value'	PRV		0
Pre-previous maximum	Minimum over the pre-previous period (i.e. period before the previous period).	Same as input 'Input value'	PPRV		0

fxPeriodTotal

Description

The function accumulates a flow increment into a period total. At the end of the period the current total is stored into the previous value and the current value is reset to 0.

The flow increment originates from a 'TotalizerRate' or

'TotalizerDelta' function.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name used for tag prefix and retentive storage.				
Increment	Increment value to be added to the period total.	Same as linked		01e11	
	Negative values will be ignored, so the period total will not decrease.	cell			
Enabled	0: Disabled		EN		
	1: Enabled				
Period type	Type of period:		ТҮР		
	1: Second				
	2: Minute				
	3: Hour				
	4: Day				
	5: Week				
	6: Month				
	7: Quarter				
	8: Year				
Period count	Number of periods (e.g. 5 minutes, 8 hours)		CNT	11e11	
Period start	Absolute start date and time of the period. This will be used as the		START	<datetime></datetime>	
	reference point to calculate the next period rollover from.				
	The value may be defined in the past or the future. The next rollover				
	period will be calculated accordingly (so forwards or backwards in time).				
Rollover value	The period total will be reset to 0 when it reaches the rollover value	Same as input	ROVAL	01e15	1e12
		Increment			
Decimal places	Defines the number of decimal places for the current and previous total		DECPLS	-110	-1
	output values.				
	-1 means full precision (no rounding applied)				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Current total	Accumulated total for the current period	Same as input 'Increment'	CUR		0
Previous total	Accumulated total for the previous period	Same as input 'Increment'	PRV		0
Rollover flag	Flag indicating a rollover to 0. 0: Off 1: On Note: stays 'On' for one calculation cycle only).			ROALM	
Pre-previous total	Accumulated total for the pre-previous period (i.e. the period before the previous period	Same as input 'Increment'	PPRV		0

fxPeriodTWA

Description

The function calculates a **t**ime-**w**eighted **a**verage (TWA) for a particular period. At the end of a period the current average is stored in the previous value and the current value is reset to 0. The function weights the input value with the time (in fact the actual calculation cycle time) and updates the average accordingly.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name used for tag prefix and retentive storage.		_		
Input value	Value to be averaged	Same as linked cell		-1e111e11	
Enabled	0: Disabled		EN		
	1: Enabled				
Period type	Type of period:		TYP		
	1: Second				
	2: Minute				
	3: Hour				
	4: Day				
	5: Week				
	6: Month				
	7: Quarter				
	8: Year				
Period count	Number of periods (e.g. 5 minutes, 8 hours)		CNT	11e11	
Period start	Absolute start date and time of the period. This will be used as		START	<datetime></datetime>	
	the reference point to calculate the next period rollover from.				
	The value may be defined in the past or the future. The next				
	rollover period will be calculated accordingly (so forwards or				
	backwards in time).				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Current average	Average calculated over the current period	Same as input 'Input value'	CUR		0
Previous average	Average of the previous period	Same as input 'Input value'	PRV		0
Pre-previous average	Average of the pre-previous period (i.e. the period	Same as input 'Input value'	PPRV		0
Pre-previous average	Average of the pre-previous period (i.e. the period before the previous period)	Same as input 'Input value'	PPRV		0

fxPeriodWatch

Description

The function 'remembers' that a condition has been valid during

a period of time.

A typical example is a transmitter that was overridden with a

keypad value.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name used for tag-prefix and retentive storage.				
Condition	The condition to be watched				
	0: Condition is not valid				
	<>0: Condition is valid				
Enabled	0: Disabled		EN		
	1: Enabled				
Period type	Type of period:		TYP		
	1: Second				
	2: Minute				
	3: Hour				
	4: Day				
	5: Week				
	6: Month				
	7: Quarter				
	8: Year				
Period count	Number of periods (e.g. 5 minutes, 8 hours)		CNT	11e11	
Period start	Absolute start date and time of the period. This will be used as the reference point		START	<datetime></datetime>	
	to calculate the next period rollover from.				
	The value may be defined in the past or the future. The next rollover period will be				
	calculated accordingly (so forwards or backwards in time).				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Current watch	Indicates whether or not the condition has been valid during the current		CUR		0
	period:				
	0: Not valid				
	1: Valid				
Previous watch	Indicates whether or not the condition has been valid during the previous		PRV		0
	period:				
	0: Not valid				
	1: Valid				
Pre-previous watch	Indicates whether or not the condition has been valid during the pre-		PPRV		0
	previous period (period before the previous period):				
	0: Not valid				
	1: Valid				

fxPID

Description

PID control is a generic method to control a process variable by means of a feedback control loop and is widely used in the industry.

A PID controller adjusts its control output by applying a Proportional, Integral and Derivative algorithm based on the error between the measured process variable and the desired setpoint.

The Proportional part of the algorithm determines the reaction to the current error. The Integral part reacts to the recent errors accumulated over a sliding time window, while the Derivative part reacts to the change rate of the error. The 3 actions are added up by applying individual weigh factors and the sum is used to adjust a control device, e.g. the position of a control valve.

Note: In flow measurement systems typically only PI control is applied, so the derivative action is disabled.

The Flow-X PID function provides several features for enhanced PID control such as:

• In <u>cascade control</u> there are two PID control loops arranged with one loop controlling the set point of the other loop. Within the outer loop the primary physical parameter is controlled, such as fluid level or velocity. The inner loop reads the output of the outer loop as its set point and usually controls a more rapid changing parameter such as flow rate or acceleration.

- For systems with a slow responsiveness to disturbances or setpoint changes <u>feed forward control</u> may be beneficial.
 Besides of the closed PID loop an open feed-forward loop is added that reacts immediately to a change in process or setpoint value.
- The function provides the option for <u>bumpless transfers</u> between auto and manual mode and vice versa. The actual process value is copied into the required setpoint value while manual mode is enabled (<u>PV tracking</u>). When reverting to auto mode the process will stay on the current process value. For the same reason the actual output % is copied to the manual output % while Auto mode is enabled.
- When the control output reaches its limit (e.g. control valve is fully opened) there is the risk for wind-up of the integral part, because the error will continue to be integrated. This results in the integral part to become very large, so the error must have the opposite value for a long time before the control loop returns to normal. In order to avoid this windup (i.e. achieve <u>anti-windup</u>) the function compensates the integral part when the control output has reached its limit.
- To avoid that a change in setpoint value will result in an impulse in the control signal the function provides the feature to define a maximum for the setpoint <u>clamp rate</u> and the control output <u>slew rate</u>. The setpoint clamp rate causes the setpoint to change gradually until it has caught up with the required value. The slew rate directly limits the rate of change of the control output.

Function inputs	Remark	EU	SW tag	Range	Defaul
Name	Optional tag name, tag description and tag group				
Process value	This represents the actual process value that is being controlled	EU			
Setpoint value	The control loop will try to achieve this input value provided that both the 'Manual mode' and	EU			
	'Cascade mode' are disabled.				
Proportional gain	Proportional gain factor	-			
Integral gain	Integral gain factor	s			
	The value 0 disables the integral part of the PID algorithm				
Derivative gain	Derivative gain factor	s			
	The value 0 disables the derivative part of the PID algorithm				
Low scale value	Process / setpoint value that corresponds to 0% of the control output	EU			
High scale value	Process / setpoint value that corresponds to 100% of the control output	EU			
Reverse	Selects the direct or reverse action of control				0
	0: Forward -> Error = (PV - SP)				
	1: Reverse -> Error = (SP - PV)				
Manual mode	0: Disabled				0
	<> 0: Enabled				
	When this input is enabled the 'Control output %' is set to input 'Manual output %'.				
	When this input is disabled the PID algorithm is applied and either the 'Setpoint value' or Cascade				
	value' is used depending on the 'Cascade mode'.				
Manual output %	The control output % will be set this value when 'Manual mode' is enabled	%			0
Upwards SP clamp	The setpoint will not be allowed to increase faster than this limit	EU/s			0
rate	Enter 0 disable this feature				
Downwards SP clamp	The setpoint will not be allowed to decrease faster than this limit	EU/s			0
rate					
Upwards OP slew rate	The control output % will not be allowed to increase faster than this limit	%/s			0
Downwards OP slew	The control output % will not be allowed to decrease faster than this limit	%/s			0
rate					
Low limit value	The control output % will not be allowed to go below this limit	%			0
High limit value	The control output % will not be allowed to go above this limit	%			100
Manual at startup	Forces manual mode at restart of flow computer				0
	0: Disabled				
	1: Enabled				
Bumpless transfer	When this input is enabled bump-less transfers between auto and manual mode and vice versa				1

Function inputs	Remark	EU	SW tag	Range	Default
	will be performed.				
	When enabled and when the mode changes from manual to auto, input 'Setpoint value' will be set				
	to the scaled value that corresponds with the current control output %.				
	When the mode changes from auto to manual, then input 'Manual output %' will be set to the				
	current output %.				
	0: Disabled				
	<> 0: Enabled				
Permissive flag	When the Permissive flag is not set the output is forced to the 'Idle output %'				1
	This input can be used for user-defined logic				
	0: Disabled				
	<> 0 : Enabled				
Idle output %	Value used for control output when the PID permissive flag is not set				0
Cascade mode	0: Disabled				0
	<>0 : Enabled				
	When this input is enabled while Manual mode is disabled, the PID algorithm is applied using the				
	'Cascade input %' (after scaling) as the set point value.				
Cascade input %	The control loop will try to achieve this input value (after scaling) provided that 'Manual mode' is	%			0
	disabled and 'Cascade mode' is enabled				
	Must be linked to the output 'Control output" of the primary PID controller.				
Tracking mode	0: Disabled				0
Ū.	<> 0: Enabled				
	This output is meant for cascade control. If this function acts as the primary (Master) PID				
	controller in a cascade configuration, this input needs to be connected to output 'Tracking mode'				
	of the secondary (Slave) PID function.				
	This input tells this function that the secondary (Slave) PID function is not using its Cascade input,				
	but its Manual output % or Setpoint value instead. This allows the primary PID function to track				
	the secondary process or setpoint value enabling a bumpless transfer between modes.				
	0: Disabled				
	<> 0 : Enabled				
Tracking value	This output is meant for cascade control. If this function acts as the primary (Master) PID	%			0
	controller in a cascade configuration, this input needs to be connected to output 'Tracking value'				
	of the secondary (Slave) PID function.				
	The value represents the process or setpoint value of the secondary (Slave) PID function as				
	percentage of scale.				
Feed forward	Value is directly added to the control output	%			0
	The advantage of feed forward control is that corrective action is taken for a change in a				
	disturbance input before it affects the controlled parameter.				

Function outputs

Function outputs	Remark			EU	SW tag	Alarm	Fallback	
Control output	The actual output value as percentage of scale that sl	hall be used for	actual control.	%				
	Equals the required control output including the slew	rate and min/	max limitations.					
Setpoint value	The actual setpoint in-use (may differ from the requir	ed setpoint be	cause of the SP clamp rate)	EU				
Tracking mode	0: Disabled							
	<> 0: Enabled							
	This output is meant for cascade control. If this funct		, ,					
	in a cascade configuration, this input needs to be cor	nnected to outp	out 'Tracking mode' of the					
	primary (Master) PID function							
	This output tells the primary (Master) PID function th		5					
	Cascade input, but the Manual output % or Setpoint							
	function to track the secondary process or setpoint v modes.	alue enabling a	bumpless transfer between					
	This output is meant for cascade control. If this funct			0/				
Tracking value	in a cascade configuration, this input needs to be cor		, ,	90				
	primary (Master) PID function	mected to outp	but Tracking value of the					
	The value depends on the Manual and Cascade mode							
	If Manual mode is enabled, this output equals the per		le of input 'Process value', else					
	this output equals the percentage of scale of input 'S		·····, ····, ····, ····, ····,					
Error	Current error	· ·						
P	Current proportional part							
1	Current integral part							
D	Current derivative part							
v	Required control output as percentage of scale (refer	to section cal	ulations). This output is for	%				
	information only and shall not be used for actual cont	trol						
Logic		e _{PRV}	error in previous cycle [El	U]				
Symbols		Δu	control output deviation	value	e [%]			
PV _{CUR} process	value in current cycle [EU]	V _{CUR}	required control output v	alue	in curren [.]	t cycle [%]	
PV _{PREV} process	value in previous cycle [EU]	V _{PRV}	required control output value in previous cycle					
SP _{REQ} required	setpoint value [EU]	U _{CUR}	actual control output val	ue in	current c	ycle [%]		

 $\mathbf{u}_{\mathsf{MAX}}$

 $\mathsf{SP}_{\mathsf{CUR}}$ in-use setpoint value in current cycle [EU] actual control output value in previous cycle [%] U_{PRV} SP_{PRV} in-use setpoint value in previous cycle [EU] low limit for control output [%] U_{MIN} high limit for control output [%]

- Δt calculation cycle time [s]
- K_P Proportional gain factor
- K₁ Integral gain factor
- K_D Derivation gain factor
- P Proportional part of current cycle
- I_{CUR} Integral part of current cycle
- I_{PRV} Integral part of previous cycle
- D Derivative part of current cycle

Control output logic

The logic for the control output depends on the current manual and tracking modes.

Manual mode = Enabled

Set current output equal to manual input value
 v_{CUR} = 'Manual output %'

Manual mode = Disabled AND Tracking mode = Enabled

- Set current output equal to tracking input value
 - v_{CUR} = 'Tracking value' (input)

Manual mode = Disabled AND Tracking mode = Disabled

Determine the current setpoint

If cascade mode enabled then

SP_{CUR} = Cascade input value * (High scale value - Low scale value) + Low scale value

Else

 $SP_{CUR} = SP_{REQ}$

• Check if the current setpoint needs to be gradually ramped up or down to the required setpoint:

If (SP_{CUR} - SP_{PRV}) > (Upwards SP clamp rate * Δt) then

 $SP_{CUR} = SP_{PRV} + (Upwards SP clamp rate * \Delta t)$ Else if (SP_{PRV} - SP_{CUR}) > (Downwards SP clamp rate * Δt) then

 $SP_{CUR} = SP_{PRV}$ - (Downwards SP clamp rate * Δt)

Calculate the current error:

```
If Control direction = Forward then
e<sub>CUR</sub> = SP<sub>CUR</sub> – PV<sub>CUR</sub>
```

Else

 $e_{CUR} = PV_{CUR} - SP_{CUR}$

• Calculate the Proportional part:

 $P = K_P * e_{CUR}$

- Calculate the Integral part:

 I_{CUR} = I_{PRV} + Ki * Δt * (e_{CUR} + (u_{PRV} v_{PRV})*(High scale value Low scale value)/100)
 Note: the latter part is required to avoid anti-windup.
- Calculate the Derivative part: D = Kd / Δt * (e_{CUR} - e_{PRV})
- Calculate the required control output:
 v_{CUR} = [P + I_{CUR} + D Low scale value] / [High scale vale
 Low scale value]
 - Check if change in control output is within the slew rate If v_{CUR} - u_{PRV} > Upwards slew rate * Δt **then** u_{CUR} = u_{PRV} + (Upwards slew rate * Δt) Else if Δu < - (Downwards slew rate * Δt) **then** u_{CUR} = u_{PRV} - (Downwards slew rate * Δt) Else

 $u_{CUR} = v_{CUR}$

Check if new control output is outside its limits
 If u_{CUR} > u_{MAX} then
 u_{CUR} = u_{MAX}

Else If $u_{CUR} < u_{MIN}$ then $u_{CUR} = u_{MIN}$

Bumpless transfer logic

If bumpless transfer is enabled, then the following logic is applied.

Setpoint tracking

IF Manual mode = Enabled OR Tracking mode = Enabled SP_{REQ} = PV_{CUR}

ELSE IF Cascade mode = Enabled

 $SP_{REQ} = SP_{CUR}$

Manual output tracking

IF Manual mode = Disabled 'Manual output %' = u_{CUR}

Tracking mode and value Outputs 'Tracking mode' and 'Tracking value' are set as follows:

Tracking mode

'Tracking mode' = ('Manual mode' = Enabled) OR ('Cascade mode' = Disabled)

Tracking value

IF 'Manual mode' = Enabled 'Tracking value' = (PV_{CUR} * - Low scale value) / (High scale value - Low scale value) * 100

IF 'Manual mode' = Disabled

'Tracking value' = (SP_{CUR} * - Low scale value) / (High scale value - Low scale value) * 100

fxROCAlarm

Description

The function checks if a value does not change its value at a rate that is higher than a specific limit ('rate of change').

Function inputs

Function inputs	Remark	EU	SW tag	Range	Defaul
Name	Optional tag name, tag description and tag	I			
	group				
Input value	Must be linked to another cell	Same as linked cell		-1e111e11	0
Rate of change limit	The unit depends on the 'Deviation type'	Absolute: Same as input value 1 / s	ROCLIM	01e11	0
		Relative : % /s			
Enabled	Enables or disabled the alarm			True or false	True
Alarm type	1: Alarm				1
	2: Warning				

Function outputs

			Alarm	Fallback	
Rate of change alarm 0: Norm	d -	ROCALM	ROCALM		
1: Alarm					

Logic

A 'Rate of change alarm' is raised when the absolute difference between two consecutive values divided by the calculation cycle time in seconds is more than the limit.

fxSetOnChange

Description

The 'fxSetOnChange function sets a tag or cell to a specific value

whenever another value changes

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Module	Number of the Flow-X/M module.			-18	
	-1 : local module				
	18 : module 1 through 8				
Target	The cell or tag that has to be set. This must be a direct reference to a cell.				
Value	Value to be assigned. May be a constant a formula or a reference to another cell or tag.				
Change	A change if this value will set the Target to the Value. This must be a direct reference to				
	a cell.				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Date and time	Date and time that the most recent change has occurred				

fxSetOnCondition

Description

The 'fxSetOnCondition' function sets a tag or cell to a specific

value whenever a condition is true.

It is a generic function that is especially useful for implementing

logic for controlling output signals (e.g. valve commands) and

state machines (e.g. prove sequences).

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Module	Number of the Flow-X/M module.			-18	
	-1 : local module				
	18 : module 1 through 8				
Target	The cell or tag that has to be set. This must be a direct reference to a cell.				
Value	Value to be assigned. May be a constant a formula or a reference to another cell or tag.				
Condition	Boolean expression.				
	When the expression outcome is TRUE (<> 0), then the target is set to the value.				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Date and time	Date and time that the most recent event has occurred				

fxSetOnEvent

Description

The 'fxSetOnEvent' function sets a tag to a specific value whenever an event occurs.

It is a generic function that is especially useful for implementing logic for controlling output signals (e.g. valve commands) and state machines (e.g. prove sequences).

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Module	Number of the Flow-X/M module.			-18	-1
	-1 : local module				
	18 : module 1 through 8				
Target	The cell or tag that has to be set. This must be a direct reference to a cell.				
Value	Value to be assigned. May be a constant a formula or a reference to another cell or tag.				
Event	Boolean expression.				
	When the expression outcome <u>changes</u> from FALSE to TRUE (or from 0 to <> 0), then				
	the target is set to the value.				
Condition	Optional condition that needs to be valid while the event occurs.				TRUE
	If the condition is not valid, then teh target will not be set to the value.				

Function outputs Remark EU SW tag Alarm Fallback Date and time Date and time that the most recent event has occurred EV SW tag Alarm Fallback

fxSetIndexOnChange

Description

The 'fxSetIndexOnChange function sets one tag from an array of tags to a specific value whenever another value changes.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Module	Number of the Flow-X/M module.			-18	-1
	-1 : local module				
	18 : module 1 through 8				
Index	The index number of the target tag to be set				
Value	Value to be assigned. May be a constant a formula or a reference to another cell or				
	tag.				
Change	A change if this value will set the Target to the Value. This must be a direct				
	reference to a cell.				
Target 1	The tag that has to be set when the index number is 1. This must be a direct				
	reference to a cell.				
Target 2	The tag that has to be set when the index number is 2. This must be a direct				
	reference to a cell.				
etc.					

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Date and time	Date and time that the most recent change has occurred				

fxSetIndexOnCondition

Description

The 'fxSetIndexOnCondition' function sets one tag from an array of tags to a specific value whenever a condition is true. It is a generic function that is especially useful for implementing logic for controlling output signals (e.g. valve commands) and state machines (e.g. prove sequences).

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Module	Number of the Flow-X/M module.			-18	-1
	-1 : local module				
	18 : module 1 through 8				
Index	The index number of the target tag to be set				
Value	Value to be assigned. May be a constant a formula or a reference to another				
	cell or tag.				
Condition	Boolean expression.				
	When the expression outcome <u>is</u> TRUE (<> 0), then the target is set to the				
	value.				
Target 1	The tag that has to be set when the index number is 1. This must be a direct				
	reference to a cell.				
Target 2	The tag that has to be set when the index number is 2. This must be a direct				
	reference to a cell.				
etc.					

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Date and time	Date and time that the most recent event has occurred				

fxSetIndexOnEvent

Description

The 'fxSetIndexOnEvent' function sets one tag from an array of tags to a specific value whenever an event occurs. It is a generic function that is especially useful for implementing

logic for controlling output signals (e.g. valve commands) and

state machines (e.g. prove sequences).

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Module	Number of the Flow-X/M module.			-18	-1
	-1 : local module				
	18 : module 1 through 8				
Index	The index number of the target tag to be set				
Value	Value to be assigned. May be a constant a formula or a reference to another cell or				
	tag.				
Event	Boolean expression.				
	When the expression outcome <u>changes</u> from FALSE to TRUE (or from 0 to <> 0),				
	then the target is set to the value.				
Condition	Condition that needs to be valid while the event occurs.				
	If the condition is not valid, then the target will not be set to the value.				
Target 1	The tag that has to be set when the index number is 1. This must be a direct				
	reference to a cell.				
Target 2	The tag that has to be set when the index number is 2. This must be a direct				
	reference to a cell.				
etc.					

	-				
Function outputs	Remark	EU	SW tag	Alarm	Fallback
Date and time	Date and time that the most recent event has occurred				

fxStatusAlarm

Description

The 'fxStatusAlarm' alarm function generates an alarm that can be triggered by a boolean condition.

Function input

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name for the tag				
Condition	A boolean indicating whether the alarm should be triggered				
Enable	Enables or disables the alarm			True or false	True
Alarm type	1: Alarm				1
	2: Warning				

fxTag

Description

The 'fxTag' function creates a tag for the cell that contains the 'fxTag' function.

Function input

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name for the tag				
Description	Description for the tag				
Value	May contain a value or a formula.				
	In case of a formula or a reference to a cell or another tag, the tag becomes				
	read-only.				
	On the other hand when it is a value the tag is writable and the specified value is				
	considered at the initial value.				
Unit	Defines an engineering unit, enumeration date/time, or a special data type, e.g.				
	'xt_bool', 'xu_kg_s' or 'xe_period').				
Write level	Applies for writeable tags only.				
	Security access level that is required to write a new value to the tag. Only applies				
	when input 'Value' contains a value and not a formula.				
	When not defined (i.e. function argument is left empty) the tag is only internally				
	writable by a spreadsheet function but not externally writable through the				
	display or communications				
Retentive	Applies for writable tags only				
	Defines whether or not the value needs to be 'remembered' (retentive).				
	When not defined (i.e. function argument is left empty) the tag is retentive				
	provided that the tag is writable and AutoReset is not enabled				
AutoReset	Applies for writeable tags only.				
	Automatically resets the tag to its initial value after it has been written to.				
Minimum	Minimum value that is accepted when the tag is externally writable.				
	May be left empty, in which case no minimum check is applied				
Maximum	Maximum value that is accepted when the tag is externally writable				
	May be left empty, in which case no maximum check is applied				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Value	Depends on setting 'Mode'	Input Unit	Input Name		
	Mode = 'This Cell'	-			
	Cell shows the value or the result of the formula that is defined for input				
	'Value'				
	Mode = 'Referred Cell'				
	Cell returns TRUE when the 'fxTag' function evaluates successfully or				
	FALSE otherwise				

fxTimer

Description

The 'fxTimer' function provides generic timer functionality.

Function input

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Start	Trigger to start the timer.		START		
	Sets output Running to 1 and starts accumulation of the actual waited time, provided that				
	the Enable condition is true (<> 0)				
	If the timer is already running, then it will be restarted, i.e. the Wait Time will be set to 0 and				
	Elapsed status will be reset to False (in case it was True)				
leset	Trigger to resets the timer.		RST		
	Sets outputs Elapsed, Running and Wait time to 0				
Limit	Time-out period for the timer.	sec	LIM		
	When the actual wait time is larger than the limit output Elapsed is set to 1.				
	The limit value is expressed in seconds and may contain a fractional part				
	The actual wait time however will be a multifold of the flow computer cycle time.				
	When the limit value does not match an exact number of cycles, then the actual limit value				
	will be rounded upwards to match the 'next' number of cycles. E.g. when the flow computer				
	cycle time is 250 ms and the Limit is set to 3.15 sec, the actual limit value being used will be				
	3.25 sec.				
Enable	Condition that controls the accumulation of actual waited time.		EN		
	0: Disabled				
	1: Enabled				
	When disabled the actual wait time will be frozen until the timer is enabled again.				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Elapsed	Flag that indicates that timer has timed out, i.e. the actual wait time is larger than the limit		ELAP		
Running	Flag that indicates that the timer is running, i.e. has been started and not been reset yet, irrespective of the Enable condition		RUN		
Wait time	Time accumulated since the latest start and while being enabled. When this time reached the limit, the Elapsed output is set to 1. Time will remain accumulated even when the timer has elapsed.	sec	TIM		

fxTotalizerDelta

Description

The function accumulates a **flow increment** into a cumulative (eternal) total.

Besides of the cumulative total the function also outputs the flow increment that represents the increase in flow quantity in the last calculation cycle. This increment value serves as an input for related batch and period flow-weighted averaging and totalization functions.

Function input

Note: As opposed to the flow increment input value, the flow increment output value is set to 0 when the totalization is disabled and has the proper units and is therefore better suited as input for other functions.

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group and retentive storage.				
Flow increment	Actual flow increment to be accumulated.	Defined by		01e11	
	Negative values will be ignored, so the cumulative total will not decrease.	'Input unit'			
Enabled	Dictates whether the flow accumulation is enabled or not. When disabled the		EN		
	cumulative total will not be updated and the increment will be set to 0.				
	0: Disabled				
	1: Enabled				
Input unit	Unit of input 'Flow increment'.				
Output unit	Unit to be used for total and increment.				
	Refer to the next section 'Unit conversion' for more information.				
	Changing the unit will only be possible when the cumulative total value equals 0.				
Rollover value	The cumulative total will be reset to 0 when it reaches the rollover value	Defined by	ROVAL	01e15	1e12
		'Output unit'			
Decimal places	Defines the number of decimal places for the total and increment output values.		DECPLS	-110	-1
	-1 means full precision (no rounding applied)				
Reset	This should be used with great care!				0
	Command to reset the cumulative total to 0				
	0: No reset				
	1: Reset				

Function output

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Cumulative total	Total quantity accumulated so far since the last rollover or	Defined by 'Input unit'	CUM		
	reset				
Increment	Increment in last calculation cycle	Defined by 'Input unit'	INCR		
Rollover flag	Flag indicating a rollover to 0.			ROALM	
	0: Off				
	1: On				
	Note: stays 'On' for one cycle only.				
Reset flag	Flag indicating a reset to 0.			RESET	
	0: Off				
	1: On				
	Note: stays 'On' for one cycle only				
Recoverable reload	Flag that indicates that at startup only 2 of the 3 copies were			RTOTERR	
error	equal and that that value is used as the initial total.				
	0: Off				
	1: On				
	Note: stays 'On' for one cycle only				
Fatal reload error	Flag that indicates that at startup all 3 copies were different			FTOTERR	
	and the total was reset to 0				
	0: Off				
	1: On				
	Note: stays 'On' for one cycle only				

Unit conversion

The function will automatically apply the required conversion based on the unit of the flow increment input value (Input unit) and the selected 'Output unit' for the flow quantity.

E.g. when flow rate input is in 'scf' the output unit is 'MMscf' then the following conversions are applied:

1 The flow increment input value is converted to the corresponding base unit, which is 'sm3' for a 'Volume' unit

2 The flow increment is converted from 'sm3' to 'MMscf' before the cumulative total and increment output values are updated.

Similar conversions are applied for flow increments that are expressed in mass, volume, normal volume and energy units.

fxTotalizerRate

Description

The function accumulates a **flow rate** into a cumulative (eternal) total.

Besides of the cumulative total the function also outputs the flow increment that represents the increase in flow quantity in the last calculation cycle. This increment value serves as an input for related batch and period flow-weighted averaging and totalization functions.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group and retentive storage.				
Flow rate	Actual flow rate to be accumulated.	Defined by 'Input		01e11	
	Negative values will be ignored, so the cumulative total will not decrease.	unit'			
Enabled	Dictates whether the flow accumulation is enabled or not. When disabled the		EN		
	cumulative total will not be updated and the increment will be set to 0.				
	0: Disabled				
	1: Enabled				
Input unit	Unit of input 'Flow rate.				
Output unit	Unit to be used for total and increment.				
	Refer to the next section 'Unit conversion' for more information.				
	Changing the unit will only be possible when the cumulative total value equals 0.				
Rollover value	The cumulative total will be reset to 0 when it reaches the rollover value	Defined by	ROVAL	01e15	1e12
		'Output unit'			
Decimal places	Defines the number of decimal places for the total and increment output values.		DECPLS	-110	-1
	-1 means full precision (no rounding applied)				
Reset	This should be used with great care!				0
	Command to reset the cumulative total to 0				
	0: No reset				
	1: Reset				

Function output

Function outputs	Remark	EU	SW tag	Alarm	Default
Cumulative total	Total quantity accumulated so far since the last rollover or reset	Defined by Input unit'	CUM		
Increment	Increment in last calculation cycle	Defined by 'Input unit'	INCR		
Rollover flag	Flag indicating a rollover to 0.			ROALM	
	0: Off				
	1: On				
	Note: stays 'On' for one cycle only.				
Reset flag	Flag indicating a reset to 0.			RESET	
	0: Off				
	1: On				
	Note: stays 'On' for one cycle only				
Reload warning	Flag that indicates that at startup only 2 of the 3 copies were equal			RLWARN	
	and that that value is used as the initial total.				
	0: Off				
	1: On				
	Note: stays 'On' for one cycle only				
Reload error	Flag that indicates that at startup all 3 copies were different and			RLERR	
	the total was reset to 0				
	0: Off				
	1: On				
	Note: stays 'On' for one cycle only				

Unit conversion

The function will automatically apply the required conversion based on the unit of the flow rate input value (Input unit) and the selected 'Output unit' for the flow quantity.

E.g. when flow rate input is in 'kg/hr' the selected unit is 'tonne' then the following conversions are applied:

- 1. The flow rate value is converted to the corresponding base unit, which is 'kg/s' for a 'Mass per Time' unit
- The flow increment over the last calculation cycle is calculated from the 'kg/s' value and the actual calculation cycle time, resulting in a value expressed in 'kg' (i.e. the base unit for 'Mass').
- The flow increment is converted from 'kg' to 'tonne' before the cumulative total and increment output values are updated.

Similar conversions are applied for flow rates that are expressed in volume, standard volume, normal volume and energy units.

fxWatchUpdate

Description

This function raises a flag whenever a value has been updated in the latest calculation cycle.

It is a generic function that can be used for any purpose, e.g. to report the number of times that a gas chromatograph has sent updates of the gas composition.

Use function fxTotalizerDelta to accumulate the number of times the flag has been raised.

Function input

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				<empty></empty>
Value	The value that needs to be checked for updates		VAL		0
Neglect zeros	Controls if the value 0 has to be considered as an update or not.		NGLZERO		0
	0: Disabled				
	The value 0 is also an update				
	1: Enabled				
	The value 0 is not considered as an update				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
UpdateFlag	Update flag		UPDATEFLAG		0
	0: Value has not changed				
	1: Value has not changed				
	Flag is automatically cleared (set to 0) at next cycle				

4 Flow-X IO Functions

This chapter lists all available Flow-X IO functions. Please refer to the pinouts for your specific model of flow computer in the installation manual provided with the Spirit Flow Suite software.

fxAnalogInput

Description

Each flow module supports a maximum of 6 analog input signals. The first 2 signals can be used as either a mA/VDC input or as a RTD input.

The Analog input function is used for mA and VDC inputs. For RTD inputs refer to function 'RTD Input'.

Analog signals are sampled at a rate of about 15 Hz. Every calculation cycle the samples are averaged and the average is scaled to a value in engineering units.

The output value can be equal to either the last sample or the average of the samples of the last calculation cycle. The average is either the arithmetic mean or the Root of the Mean of the Squares (RMS), Where the latter is meant for a differential pressure signal of a primary flow device (e.g. an orifice plate) **Function inputs**

The input signal is considered to be faulty when the input circuitry has an open or a short circuit or when the measured value is outside a configurable range.

Note: Function fxKeypadfallback provides the option to force the analog input value to a specific fallback value in case it should fail. It also provides the option to force the input value to a keypad value e.g. upon user request.

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Channel number			CHAN	16	1
Input type	1: 4-20 mA		INPTYP		2
	2: 0-20 mA				
	3: 1-5 VDC				
	4: 0-5 VDC				
Averaging type	1: Arithmetic mean		AVGTYP		1
5 5 51	2: Root Mean Square				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Function Input argument out of range			FIOOR	
Percentage value	Actual percentage of scale value	%	PERC		0

fxAnalogOutput

Description

This function configures a single 4-20 mA output channel on the local Flow-X module. Use function 'fxSetAnalogOutput' to control the actual output signal.

The output current is either set directly or gradually changed to the required set point based on the specified filter method.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional Optional tag name, tag description and tag group				<empty></empty>
Channel number		14	CHAN		0
Filter setpoint	015		FILSP		0
	0: No filtering				
	1: Fastest filter				
	15: Slowest filter				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Function Input argument out of range			FIOOR	
Percentage value	The actual (and filtered) output value expressed as a percentage	%	PERC		0

fxSetAnalogOutput

Description

This function writes an analog value to an output. Make sure that the analog output has been set up by calling fxAnalogOutput first.

Name	Optional Optional tag name, tag description and tag group				< Empty >
	description and tag group				<empty></empty>
Module	Not used, set to -1			-116	-1
Channel number		14	CHAN		0
Filter setpoint	015		FILSP		0
	0: No filtering				
	1: Fastest filter				
	15: Slowest filter				

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Function Input argument out of range			FIOOR	

fxDigitalInput

Description

The Digital Input function processes a single digital input and outputs either the actual input signal or the latched input signal.

Function inputs

Function inputs	Remark	EU	SW tag	Default
Name	Optional tag name, tag description and tag group			
Channel number	116		CHAN	1
Logic	1: Positive		LOGIC	1
	2: Negative			
Mode	1: Actual		MODE	1
	2: Latched			
Threshold	Each digital input has 2 threshold levels that determine whether the signal is considered		THRESHOLD	1
	to be either high (above the threshold) or low (below the threshold).			
	The threshold levels are as follows (all relative to signal ground):			
	Channels 1 through 8:			
	1: + 1.25 Volts			
	2: + 12 Volts			
	Channels 9 through 16:			
	1: + 3.75 Volts			
	2: + 12 Volts			

Function outputs

Function outputs	Remark	SW tag	Alarm	EU
Status	0: Normal	STS		-
	1: Input argument out of range		FIOOR	
Signal State	0: Off	SIGSTATE		-
	1: On			
	Meaning depends on the input Logic and Mode, refer to the table below.			

Calculations

The following table summarizes the relationship between the input signal and the output 'State', depending on the inputs 'Logic and 'Mode'.

Logic	Mode	Signal state = Off	Signal State = On
Positive	Actual	Actual signal is low	Actual signal is high
Positive	Latched	Signal has not been high during last calculation cycle	Signal has been high during last calculation cycle
Negative	Actual	Actual signal is high	Actual signal is low
Negative	Latched	Signal has not been low during last calculation cycle	Signal has been low during last calculation cycle

fxDigitalOutput

Description

The Digital Output function configures a single digital output on the local module. Use function 'fxSetDigitalOutput' to control the actual output signal.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name for output 'Signal State'				
Channel number	One of the 16 digital I/O channels.		CHAN	116	1
Logic	1: Positive		LOGIC		1
	2: Negative				
Delay	Period of time that the control signal must be high (> 0) without interruption	ms	DELAY	>= 0	0
	before the output will be activated.				
	The value 0 disables the delay function				
	If the control signal becomes 0 before the time has elapsed, then the output				
	signal will not be activated.				
Minimum activation	Minimum period of time that the signal will remain activated.	ms	MINACTITM	>= 0	0
time	After the minimum activation time has elapsed the output signal will remain				
	activated until the control value becomes 0.				

Function outputs

Function outputs	Remark	SW tag	Alarm	EU	Fallback
Status	0: Normal	STS		-	0
	1: Input argument out of range or conflict		FIOOR		
Signal State	0: Not activated	SIGSTATE		-	0
	1: Activated				

Calculations

The following table summarizes the relationship between the output signal, the input 'Logic and the control value as set by function 'fxSetDigitalOutput'.

fxSetDigitalOutput

Description

This function writes a value to a configured digital output.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Name for output 'Signal State'				
Module	Unused, set to -1			-116	-1
Channel number	One of the 16 digital i/o channels.			116	
Setpoint	Logic is positive:				
	0: Output is activated				
	<>0: Output is not activated				
	Logic is negative:				
	0: Output is not activated				
	<>0: Output is activated				

Function outputs	Remark	SW tag	Alarm	EU	Fallback
Status	0: Normal	STS		-	0
	1: Input argument out of range or conflict		FIOOR		

fxFrequencyOutput

Description

This function provides a programmable output frequency on one of the digital pins. The frequency as well as the duty cycle are user programmable. When using the frequency output to configure a dual pulse system, channel 2 can be configured.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name					
Index	Index of the frequency output			14	
Channel number	Output channel number, the digital IO pin that is used.			016	
Channel number 2	Output channel number, the digital IO pin that is used.			016	0

Function outputs	Remark	SW tag	Alarm	EU	Fallback
Status	0: Normal	STS		-	1
	1: Input argument out of range or conflict		FIOOR		

fxSetFrequencyOutput

Description

This function starts a programmed frequency on the digital pin, set by fxFrequencyOutput. In case of a dual pulse via frequency out, here the phase shift between the pins can be configured.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name					
Module	Not used, set to -1			-116	-1
Index	Index of the frequency output			14	
Frequency	Output frequency (Hz)	Hz		0.01 10000	
Duty cycle	Duty cycle	%		0100	
Enabled	Enables (<>0) or disables (0) the output signal.				
Phase shift	Phase shift between dual pulse channel A and B expressed as the ratio of the time delay between the last A and B and the time between two A pulses. A typical phase difference is 90° corresponding to a value of 0.25. The shift is limited to [0°180°).			00.5	0

Function outputs	Remark	SW tag	Alarm	EU	Fallback
Status	0: Normal	STS		-	1
	1: Input argument out of range or conflict		FIOOR		
Actual frequency	Actual frequency, which can deviate from the programmed	ACTFREQ		Hz	0
	frequency because of hardware limitations.				
Actual duty cycle		ACTDC		%	0

fxDoubleChronometry

Description

The function provides double chronometry measurement for meter proving and calibration purposes with a resolution of 100 nanoseconds.

The function monitors a (dual) pulse input signal provided by the meter under test and by one or more digital signals that represent the start and stop of the measurement.

The digital inputs used for start and stop of the measurement are typically connected to the sphere detector switches of a pipe prover or the piston detectors of a compact prover.

The start / stop signal may also be any other type of signal, e.g. a digital output from another flow computer. This allows for flexible master meter configurations in which a single prove measurement is performed by two separate flow computers, one that monitors the meter under test and one that monitors the reference or master meter.

By using multiple Double Chronometry functions multiple prove measurements can be performed at the same time by one and the same flow module.

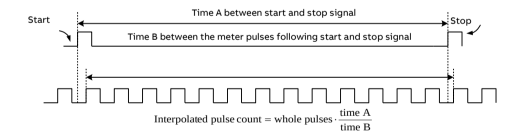
Some examples of simultaneous prove measurements are:

• For a prover with 2 begin and 2 end detector switches the flow meter can be proved against all 4 the calibrated prover volumes at the same time.

- With 2 detectors at one end of the prover (A and B) and 2 at the other end (C and D) there are 4 possible detector combinations A-C, A-D, B-C and B-D, resulting in four available prover volumes. By defining 4 Prove measurement functions all 4 prove volumes can be used to prove the meter by a single prove sequence. The operator selects the primary detector combination that will be used for calculating the new K factor or meter factor. The result of the 4 detector combinations are compared to each other and an alarm is raised in case of discrepancy.
- Meters that operate in series can all be proved at the same time.
- For meters that provide more than flow signal all signals can be proved at the same time
- For provers with a range of calibrated prover volumes to accommodate proving of meters of different capacities, multiple prover volumes can be processed at the same time.

Combinations of these scenarios are possible as well.

The double chronometry method comprises the measurement of the time between the start and stop signals and the time between the two meter pulses that immediately follow the start and stop signals. The flow module then determines the interpolated number of meter pulses that correspond to the measurement start and stop signals as follows:



Function inputs and outputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Index	Index number		INDEX	14	1
Flow input channel number	Number of the digital input channel		FLOWCHAN		
Start DI channel number	Channel number of the digital input that is to be used as the Start signal		STRTCHAN		0
Start DI logic	1: Positive 2: Negative		STRTLOGIC	12	
Stop DI channel number	Channel number of the digital input that is to be used as the Stop signal. Note: the same digital input may be used for both Start and Stop of measurement		STOPCHAN		0
Stop DI logic	1: Positive 2: Negative		STOPLOGIC	12	
Start minimum delay time *	After the Reset command has been given the function will wait for at least this delay time before considering the activation of the Start digital input	sec	STRTMINDLY	01e9	0
Start maximum delay time *	After the Reset command has been given the function will wait for no longer than this delay time before the Start digital input must have been activated 0 disables this delay check	sec	STRTMAXDLY	01e9	0
Stop minimum delay time *	After the Reset command has been given the function will wait for at least this delay time before considering the activation of the Stop digital input. O disables this delay check Note: The stop minimum delay time must be defined when a common digital input is used as both the Start and Stop signals	sec	STOPMINDLY	01e9	0

Function inputs	Remark		EU	SW tag	Range	Default
Stop maximum delay	After th	e Reset command has been given the function will wait for no longer	sec	STOPMAXDLY	01e9	0
time *	than thi	s delay time before the Stop digital input must have been activated.				
	0 disabl	es this delay check				
Number of pulses per	Applies	for flow meter that provide a direct pulse signal (typically turbine and		PLSPERREV	11e9	1
revolution	Positive	Displacement (PD) meters)				
	To exclu	Ide the influence of geometrical imperfections of the primary metering				
	device t	he function may be forced to accumulate a number of pulses that				
	corresp	onds to a whole number of flow meter revolutions.				
	E.g. sup	pose that at the stop signal 6754 pulses have been accumulated and				
	that cor	responding turbine meter provides 20 pulses for each turbine revolution	n,			
	the fund	tion will accumulate another 6 pulses before the measurement is				
	stopped	1.				
Start	0	No action		START		0
	<> 0	Starts the double chronometry				
	If the do	puble chronometry function is already running, then it will be restarted.				
Reset	0	No action		RST		
	<> 0	Resets all outputs to 0 or, where applicable, FALSE.				

Function outputs	Remark	SW tag	Alarm	EU
Status	0: Normal	STS		-
	1: Input argument out of range or conflict		FIOOR	
Interpolated pulse count	Equals :	INPOLCNT		
	Whole pulse count * Time A / Time B			
	According to API requirements the interpolated pulse count should be used when less than 10000 pulses are acquired			
Whole pulse count	The number of meter pulses that were acquired within the measurement period (time B).	WHOLECNT		
	Could be used instead of interpolated pulses when more than 10000 pulses are acquired	WHOLECHT		
	Note: Whole pulse count = the decimal part of Interpolated pulse count			
Time between start and	Time period between start and stop signal	INPOLTIM		sec
stop				
Time between pulses	Time between the meter pulses immediately following the start and stop signals	WHOLETIM		sec
In Progress	Status bit that indicates that the measurement is in progress	INPRG		
Completed	Prove measurement has been completed	COMPL		
Start time-out	Prove measurement has been aborted - Start signal time-out	STRTTO		
Stop time-out	Aborted - Stop signal time-out	STOPTO		

fxPulseInput

Description

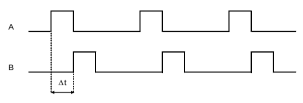
The 'fxPulseInput' function is meant for a flow meter that provides a single or a dual pulse output signal. Each flow module supports either 1 single or 1 dual pulse input.

A dual pulse signal is a set of two pulse signals ('pulse trains') A and B that originate from the same flow meter. The two pulse trains are similar but shifted in phase (typically 90°).

The primary purpose of the dual signal is to allow for **pulse integrity checking**. Added or missing pulses on either pulse train are detected and corrected for and simultaneous noise pulses are rejected.

The function provides detailed information on the raw, corrected and bad pulses for both channels and for both the forward and reverse flow direction.

The phase shifted pulse train signal also allows for automatic detection of <u>flow direction</u>. Each A pulse is followed by a B pulse within a time period (Δ t) in case the flow runs in the forward direction. In case the flow runs in the reverse direction, the opposite is the case, i.e. each B pulse is followed by an A pulse within the same time period Δ t.



Channel B lags channel A

The function also provides the option to output the raw pulse signals, which is useful in case a separate flow computer is used for proving purposes. The proving flow computer takes the pulse output from the flow computer that processes the meter on prove to perform prove measurements including double chronometry if required. The prover output signal is generated at 20 MHz, the same frequency at which the raw pulse input signals are sampled.

Compliance

ISO 6551:1982, Petroleum liquids and gases -- Fidelity and security of dynamic measurement -- Cabled transmission of electric and/or electronic pulsed data

Note: The Flow/X series of flow computers provides Level A pulse security as defined in ISO 6551, which means that bad pulses are not only detected (level B) but also corrected for.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Index	Index number. Always 1.		INDEX	11	1
Channel A	Input channel number for pulse A		CHANA	016	0
	0 disables the entire function				
Channel B	Input channel number for pulse B		CHANB	016	Input Channel A + 1
	Enter 0 in a case of single pulse input				
Pulse fidelity threshold	All pulse fidelity checking will be disabled when the corrected	Hz	FRQTHD	01e5	0
	pulse frequency is below the 'Pulse fidelity threshold'				
	Enter a 0 to disable this functionality (the default value)				
	Note: In FPGA later then v13, pulse fidelity checking is always				
	enabled				
Error pulses limit	When the total number of missing pulses, added and		ERRLIM	01e99	0
	simultaneous pulses for either channel becomes larger than				
	this value, the status becomes 'Error Pulses'				
	The value 0 disables the error pulses limit check.				
Missing pulses limit	When the total number of missing pulses on channel A or B	-	MISLIM	01e99	0
	becomes larger than this value, the status becomes 'Missing				
	Pulses Channel x' (with x either A or B)				
	The value 0 disables the missing pulses limit check.				
Added pulses limit	When the total number of added pulses on channel A or B	-	ADDLIM	01e99	0
	becomes larger than this value, the status becomes 'Added				
	Pulses Channel x' (with x either A or B)				
	The value 0 disables the added pulses limit check.				
Simultaneous pulses	When the total number of simultaneous pulses on both	-	SIMLIM	01e99	0
limit	channels becomes larger than this value, the status becomes				
	Simultaneous Pulses '				
	The value 0 disables the simultaneous pulses limit check.				
Good pulse reset limit	When the number of good pulses since the last 'bad' pulse has	-	RSTLIM	01e99	0
	reached this value, all the bad pulse count and alarms will be				
	reset automatically.				
	The value 0 disables the automatic reset function.				
Bad pulse reset	When the value changes the bad pulse count and alarms are	-	RSTCMD	Any value	0
command	reset. Can be used reset the bad pulses manually or				
	automatically e.g. at a every new batch.				
Error rate limit	When the difference in frequency between the two raw pulse	%	ERRLIM	0100	0
	trains is larger than this limit within the last calculation cycle,				
	the status becomes 'Pulse Rate Error'				
	The value 0 disables the error rate limit check				

Function inputs	Remark	EU	SW tag	Range	Default
Pulse A output channel	Number of digital I/O channel that is used to output the raw A	-	POCHANA		0
	pulses.				
	0:Not used				
	1:Digital I/O channel 1				
	16:Digital I/O channel 16				
Pulse B output channel	Number of digital I/O channel that is used to output the raw B	-	POCHANA		0
	pulses.				
	0:Not used				
	1:Digital I/O channel 1				
	16:Digital I/O channel 16				
Pulse A output mode	Used to switch the pulse A output on and off.		POMODA		0
	0: Pulse output A is disabled				
	<> 0: Pulse output A is enabled				
Pulse B output mode	Used to switch the pulse B output on and off.		POMODA		0
	0: Pulse output B is disabled				
	<> 0: Pulse output B is enabled				

Function outputs	Remark	EU	SW tag	Alarm	Fallbac
Status	0: Normal	-	STS		
	1: Input argument out of range			FIOOR	
	2: No A pulses (while B pulses)			NOPLSA	
	3: No B pulses (while A pulses)			NOPLSB	
	4: Missing pulses channel A			MISPLSA	
	5: Missing pulses channel B			MISPLSB	
	6: Added pulses channel A			ADDPLSA	
	7: Added pulses channel B			ADDPLSB	
	8: Simultaneous pulses			SIMPLS	
	9: Pulse rate error			ERRRATE	
	10: Low frequency (above 0 and below cut-off)			LOFRQ	
				•	
	11: Error pulses			ERRPLS	
	Note: during normal operation status 'Low frequency' occurs for a relative				
	short time whenever the flow starts or stops. In order to avoid				
	unnecessary alarms the corresponding alarm delay time shall be defined				
	accordingly (default 5 sec)				
Corrected pulse increment	Number of good pulses within the last calculation cycle	-		CORINC	
Corrected frequency	Frequency that corresponds to the last corrected pulse increment within the last calculation cycle	Hz		CORFRQ	
Error pulse increment	Number of bad pulses within the last calculation cycle			ERRINC	
Error pulse frequency	Frequency that corresponds to the last error pulse increment within the	Hz		ERRFRQ	
	last calculation cycle			-	
Flow direction	0: Forward	-		FLOWDIR	
	1: Reverse			LOUDIN	
Raw pulse increment	Number of raw pulses on channel A within the last calculation cycle			RAWINCA	
channel A	Number of haw pulses of channel A within the last calculation cycle			NAWINCA	
	Number of your pulses on abound D within the last calculation and			DAMINICD	
Raw pulse increment	Number of raw pulses on channel B within the last calculation cycle			RAWINCB	
channel B					
Missing pulse count channel	Depends on FPGA version:	-		MISCNTA	
A	• FPGA v13 and earlier: Total missing pulse count channel A since the				
	last reset				
	 FPGA following v13: Total number of B pulses while channel A was 				
	determined to be missing since the last reset				
Missing pulse count channel	Depends on FPGA version:	-		MISCNTB	
В	• FPGA v13 and earlier: Total missing pulse count channel B since the				
	last reset				
	• FPGA following v13: Total number of A pulses while channel B was				
	determined to be missing since the last reset				
Added pulse count channel	Depends on FPGA version:	-		ADDCNTA	
Α	• FPGA v13 and earlier: Total added pulse count channel A since the				
	last reset				
	 FPGA following v13: Total A pulses without preceding B pulse since 				
	the last reset, when missing channel is not determined				
Added pulse count channel	Depends on FPGA version:			ADDCNTB	
B	•	-		ADDCIVID	
D					
	last reset				
	• FPGA following v13: Total B pulses without preceding A pulse since				
	the last reset, when missing channel is not determined				
Simultaneous pulses	Total simultaneous pulse count since the last reset			SIMPLSF	
Phase difference	Last measured phase difference between A and B pulse expressed as the	-		PHASEDIF	
	ratio of the time delay between the last A and B and the time between two				
	A pulses. A typical phase difference is 90° corresponding to a value of				
	ratio of the time delay between the last A and B and the time between two				

fxResetPulseInputErrors

Description

This function resets the error pulse counters (error, added, missing and simultaneous pulses) of the pulse input of one of

the modules part of the same configuration.

It is required that the pulse input of the specified module is

configured by function 'fxPulseInput'.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				<empty></empty>
Module	Unused, set to 0			016	0
Index	Index number as defined in the corresponding fxPulseInput function			01	
	0 disables the function				
Reset	0: No action				
	<> 0 All error counters are reset to 0				

Function inputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range or in conflict			FIOOR	

xPulseOutput

Description

This function is used to feed pulses to an electro-mechanical (E/M) counter. There are 4 pulse output units, to be selected by the index parameter. The function uses a reservoir to accumulate the pulses. On one hand the number of pulses that need to be added to the reservoir is calculated from inputs 'Increment' and 'Significance factor'. On the other hand pulses are taken from the reservoir and fed to the E/M counter at a rate that will not exceed the specified output rate.

This can also be used to configure a dual pulse situation, for this case you can configure a second channel and the delay it has on the primary channel.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Index	Index number 14.		INDEX	14	1
Channel	Digital I/O channel number		CHAN	016	0
	0 disables the function				
Significance factor	Factor that specifies the relation between the Increment value and the	-	SIGFCT	01e99	1
	number of output pulses.				
	E.g. a value of 100 means that 1 pulse is generated whenever 100				
	increment units have been accumulated.				
Pulse width	Time that each output pulse remains active (high) in millisecond	ms	PLSWID	01e6	20
	Restriction (pulse duty cycle is 50%):				
	Pulse width <= 1000 / (2* max. output rate)				
Max. output rate	Maximum pulse output rate.	Hz	MAXFREQ	01e6	40
	When output pulses are generated at a frequency higher than the				
	maximum output rate, the superfluous pulses will be accumulated in the				
	pulse reservoir.				
	Restriction when pulse duty cycle should not exceed 50%:				
	Max. output rate <= 1000 / (2* Pulse width)				
Reservoir alarm limit	Alarm limit for the number of pulses in the reservoir buffer	-	RSVLIM	01e99	1e9
Channel 2	Digital I/O channel number		CHAN	016	0
	0 disables the function				
Delay	Delay between dual pulse channel A and B in milliseconds	ms		01e99	0

Function outputs

Function outputs	Remark		SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range or in conflict			FIOOR	
Pulse reservoir	Number of pulses that are accumulated in the reservoir.	-	RSVPLS		0
Reservoir alarm status	With respect to the 'Reservoir alarm limit' the number of pulses in		RSVSTS		0
	the reservoir is:				
	0: Within limit				
	1: Out of limit			RSVALM	

Calculations

Every calculation cycle a number of pulses is added to the reservoir that is equal to input value 'Increment' divided by input value Significance factor.

fxSetPulseOutput

Adds pulses to the reservoir, to be output on the next cycle. Takes the 'significance factor' into account, see fxPulseOutput.

Description

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default	
Name	Optional tag name, tag description and tag group					
module	Not used, set to -1116					
index	Index of the pulse output			04		
	0 disables the function					
Increment	Number of pulses to be added to the reservoir					
Reset command	0 is normal, 1 is no hardware state changed				0	

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal	-	STS		
	1: Input argument out of range or in conflict			FIOOR	

fxPT100Input

Description

Each flow module provides 2 inputs for **Pt-100** sensors. A Pt-100 sensor is a Platinum Resistance Thermometer (**PRT**) element with a resistance of 100 Ω at 0°C. 'Pt' is the symbol for platinum.

Resistance thermometers as such are also referred to as Resistance Temperature Detectors (**RTD**).

The resistance changes linearly with temperature. For Pt-100 elements the linearity coefficient is 0.385 Ohm/°C for European elements and 0.392 Ohm/°C for American elements

Function inputs

For each sample the temperature value is obtained from the measured Ohms by means of the standard RTD quadratic equation.

The input signal is considered to be faulty when the input circuitry has an open or a short circuit or when the measured value is outside its range (as defined in table below).

Function inputs	Remark	EU	Default
Name	Optional tag name, tag description and tag group		
Channel number		12	1
Input Type	1: European, 0.00385 Ω/ Ω /°C	12 1	1
	As per DIN 43760, BS1905,IEC751		
	Range - 200+850 °C		
	2: American, 0.00392 Ω/ Ω /°C		
	Range - 100+457 °C		

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback	
Status	Status output		STS			
	0: Normal					
	1: Input argument out of range			FIOOR		
	2: Under range failure			(*)		
	3: Over range failure			(*)		
	4: Open circuit			(*)		
	5: Short circuit			(*)		
Value	Temperature in degrees Celsius	°C	VAL		0	

(*) Note that no alarm is generated for this status output value. This is to avoid an unnecessary alarm in case the input is not used

fxRTDInput

Description

Reads the PT100 input as a 'raw' Ohm number. Use fxPT100Table to convert to degrees Celsius.

Function inputs

Function inputs	Remark	EU	Default
Name	Optional tag name, tag description and tag group		
Channel number		12	

Function outputs	Remark	EU	SW tag	Alarm	Fallback	
Status	Status output		STS			
	0: Normal					
	1: Input argument out of range			FIOOR		
	2: Under range failure			(*)		
	3: Over range failure			(*)		
	4: Open circuit			(*)		
	5: Short circuit			(*)		
Value		Ohm			0	

fxPT100Table

Description

This function converts a resistance value (Ohm) into a temperature value (°C) according to the Pt-100 conversion tables. A Pt-100 sensor is a Platinum Resistance Thermometer (**PRT**) element with a resistance of 100 Ω at 0°C. 'Pt' is the symbol for platinum.

Resistance thermometers as such are also referred to as Resistance Temperature Detectors (**RTD**).

The resistance changes linearly with temperature. For Pt-100 elements the linearity coefficient is 0.385 Ohm/°C for European elements and 0.392 Ohm/°C for American elements For each sample the temperature value is obtained from the measured Ohms by means of the standard RTD quadratic equation.

The input signal is considered to be faulty when the input circuitry has an open or a short circuit or when the measured value is outside its range (as defined in table below).

Function inputs

Function inputs	Remark	EU	Default
Name	Optional tag name, tag description and tag group		
Ohm	Measured temperature in Ohms	Ohm	
nm put Type	1: European, 0.00385 Ω/ Ω /°C		1
	As per DIN 43760, BS1905,IEC751		
	Range - 200+850 °C		
	2: American, 0.00392 Ω/ Ω /°C		
	Range - 100+457 °C		

Function outputs	Remark	EU	SW tag	Alarm	Fallback	
Value		°C			0	

fxTimePeriodInput

Description

This function measures the time period between two pulses with a high resolution (100 nanoseconds) and is typically used for densitometer inputs. The measurement is performed continuously in the background and the function returns the average value since the previous calculation cycle. The measured time period value can be linked to a subsequent densitometer-specific (Solartron, Sarasota, UGC) function that calculates the density at the densitometer conditions. Abnormal measurements are filtered out and alarmed for.

Function inputs

Function inputs	Remark	EU	SW tag	Range	Default
Name	Optional tag name, tag description and tag group				
Channel	Channel number for the signal	-	CHAN	116	0
Time period differential limit	Maximum allowable difference in microseconds.	μs			
	When the time period between two consecutive pulses differs more				
	than this limit from the previous time period, the reading is				
	considered to be abnormal.				
	Following an abnormal reading there must be 3 consecutive readings				
	within the limit before the time period value is considered normal				
	again.				
	When no 3 consecutive readings within the limit are available in the				
	last 5 readings then the input signal is considered to be invalid.				
	Resolution of the limit value is 100 nanoseconds.				
Minimum stable time	Minimum time for high- and low-times of a pulse to be considered	μs		010000	30
	valid.				

Function outputs

Function outputs	Remark	EU	SW tag	Alarm	Fallback
Status	0: Normal		STS		
	1: Input argument out of range			INPERR	
	Outputs will be set to fallback values				
	2: No valid measurement			(*)	
	Outputs will be set to fallback values				
Time period	Average (arithmetic mean) in microseconds of the last calculation	μs	TIME		0
	cycle				
Frequency	Average (arithmetic mean) of the last calculation cycle	Hz	FREQ		0
Curcount	Current counter value		CURCNT		0
Curtime	Current time value	sec	CURTIME		0
Oldcount	Previous counter value		PRVCNT		0
Oldtime	Previous time value	sec	PRVTIME		0

(*) Note that no alarm is generated for this status output value. This is to avoid an unnecessary alarm in case the input is not used

5 Reference

Unit Types

Type of unit	Description	Unit	Convert to	Excel constant	Multiply by	Conversion
Acceleration	meters per second squared	m/s2		xu_m_s2		
	kilometers per second squared	km/s2	m/s2	xu_km_s2	1.0 E+03	Exact
	inch per second squared	in/s2	m/s2	xu_in_s2	2.54 E-02	Exact
	foot per second squared	ft/s2	m/s2	xu_ft_s2	3.048 E-01	Exact
Area	square meter	m2		xu_m2	by 1.0 E+03 2.54 E-02	
	square millimeter	mm2	m2	xu_mm2		Exact
	square centimeter	cm2	m2	xu_cm2		Exact
	square kilometer	km2	m2	xu_km2		Exact
	square inch	in2	m2	xu_in2		Exact
	square foot	ft2	m2	xu_ft2	9.290304 E-02	Exact
Dynamic Viscosity	pascal second	Pa.s		xu_Pa.s		
	poise	poise	Pa.s	xu_poise		Exact
	centipoise	cP	Pa.s	xu_cP		Exact
	kilogram force second per square meter	kgf.s/m2	Pa.s	xu_kgf.s_m2		Exact
	pound-mass per foot second	lbm/ft.s	Pa.s	xu_lbm_ft.s	0.45359237 / 0.3048	Exact
Energy	joules	J		xu_J		
	kilojoules	kJ	J	xu_kJ		Exact
	megajoules	MJ	J	xu_MJ	1.0 E+06	Exact
	gigajoules	GJ	J	xu_GJ		Exact
	terajoules	TJ	J	xu_TJ		Exact
	watt hour	W.h	J			Exact
	kilowatt hour	kW.h	J		3.6 E+06	Exact
	watt second	W.s	J		1	Exact
	British thermal unit	Btu	J		1.05505585262 E+03	Exact
	kilo British thermal unit	kBtu	J		1.05505585262 E+06	Exact
	million British thermal unit	MMBtu	J		1.05505585262 E+09	Exact
	calorie	cal	J		4.1868	Exact
	kilocalorie	kcal	J		4.1868 E+03	Exact
	megacalorie	Mcal	J		4.1868 E+09	Exact
	decatherm	dT	J		1.05505585262 E+09	Exact
nergy per Mass	joule per kilogram	J/kg				
	kilojoule per kilogram	kJ/kg	J/kg		1.0 E+03	Exact
	megajoule per kilogram	MJ/kg	J/kg		1.0 E+06	Exact
	British thermal unit per pound (avoirdupois)	Btu/lbm	J/kg		2.32601 E+03	Exact
	kilo British thermal unit per pound (avoirdupois)	kBtu/lbm	J/kg		2.32601 E+06	Exact
	calorie per kilogram	cal/kg	J/kg		4.1868	Exact
	kilocalorie per kilogram	kcal/kg	J/kg			Exact
	million calorie per kilogram	MMcal/kg	J/kg			Exact
nergy per Mole	joules per mole	J/mol	57 Kg		4.1000 E . 00	Exact
incigy per hole	kilojoules per mole	kJ/mol	J/mol		1.0 E+03	Exact
	megajoules per mole	MJ/mol	J/mol			Exact
	kilojoules per kilomole	kJ/kmol	J/mol			Exact
	megajoules per kilomole	MJ/kmol	J/mol			Exact
	British thermal unit per pound	Btu/lbmol	J/mol			Exact
	mole kilo British thermal unit per	kBtu/lbmol	J/mol			Exact
	pound mole	cal/mol	J/mol			
	calorie per mole	kcal/mol	J/mol			Exact Exact
	kilocalorie per mole megacalorie per mole	Mcal/mol	J/mol			
noray por Timo	· ·	1	J/IIIII		+.1000 E+U0	Exact
inergy per Time	joules per second	J/s M1/br	1/c		(10/2600) E+06	Evact
	megajoules per hour	MJ/hr	J/s			Exact
	gigajoules per hour	GJ/hr M1/day	J/s			Exact
	megajoules per day	MJ/day	J/s			Exact
	gigajoules per day	GJ/day	J/s		· · · ·	Exact
	kilo British thermal unit per hour	kBtu/hr	J/s			Exact
	million British thermal unit per hour	MMBtu/hr	J/s			Exact
	kilo British thermal unit per day	kBtu/d	J/s		E+06	Exact
	million British thermal unit per day	MMBtu/d	J/s		(1.05505585262 / 86400) E+09	Exact
	mega calorie per hour	Mcal/hr	J/mol		(4.1868/3600) E+06	Exact
	giga calorie per hour	Gcal/hr	J/mol		(4.1868/3600) E+09	Exact

Type of unit	Description	Unit	Convert to	Excel constant	Multiply by	Conversion
	million calorie per day	Mcal/d	J/mol		(4.1868/86400) E+06	Exact
	giga calorie per day	Gcal/d	J/mol		(4.1868/86400) E+09	Exact
Energy per Volume	joules per cubic meter	J/m3				
	kilojoules per cubic meter	kJ/m3	J/m3		1.0 E+03	Exact
	megajoules per cubic meter	MJ/m3	J/m3		1.0 E+06	Exact
	British thermal unit per cubic foot	Btu/ft3	J/m3		(1.05505585262 / 0.02831685) E+03	Exact
	kilo British thermal unit per cubic foot	kBtu/ft3	J/m3		(1.05505585262 / 0.02831685) E+06	Exact
	calorie per cubic meter	cal/m3	J/mol		4.1868	Exact
	kilocalorie per cubic meter	kcal/m3	J/mol		4.1868 E+03	Exact
inergy per Standard Volume	joules per standard cubic meter	J/sm3				
	kilojoules per standard cubic meter	kJ/sm3	J/sm3		1.0 E+03	Exact
	megajoules per standard cubic meter	MJ/sm3	J/sm3		1.0 E+06	Exact
	British thermal unit per standard cubic foot	Btu/scf	J/sm3		(1.05505585262 / 0.02831685) E+03	Exact
	kilo British thermal unit per standard cubic foot	kBtu/scf	J/sm3		(1.05505585262 / 0.02831685) E+06	Exact
	calorie per standard cubic meter	cal/sm3	J/sm3		4.1868	Exact
	kilocalorie per standard cubic meter	kcal/sm3	J/sm3		4.1868 E+03	Exact
Energy per Normal /olume	joules per standard cubic meter	J/m3(n)				
	kilojoules per standard cubic meter	kJ/m3(n)	J/m3(n)	xu_J/m3n	1.0 E+03	Exact
	megajoules per standard cubic meter	MJ/m3(n)	J/m3(n)		1.0 E+06	Exact
	calorie per standard cubic meter	cal/m3(n)	J/m3(n)		4.1868	Exact
	kilocalorie per standard cubic meter	kcal/m3(n)	J/m3(n)		4.1868 E+03	Exact
actor	scaling value	Decimal				
	percent	%	Decimal		1.0 E-02	Exact
	parts per million	ppm	Decimal		1.0 E-06	Exact
orce	Newton	N				
	kilogram-force	kgf	N		9.80665	Exact
	pound-force	lbf	N		4.4482216152605	Exact
requency	Hertz	Hz				
leat Capacity per 1ass	Joule per kilogram per degree Celsius	J/kg.°C		xu_J_kg.degC		
	Joule per kilogram per degree Celsius	kJ/kg.°C	J/kg.°C		1.0 E+03	Exact
	British thermal unit per pound (avoirdupois) per degree Fahrenheit	Btu/lbm.°F	J/kg. °C		4186.8	Exact
Heat Capacity per Mole	Joule per mole per degree Celsius	J/mol.°C				
	kilo Joule per kilo mole per degree Celsius	kJ/kmol.°C	J/mol.°C		1	Exact
	British thermal unit per mole per degree Fahrenheit	Btu/lbmol.°F	J/mol.°C		2.326E+03 / 1.8	Exact
Kinematic Viscosity	square meter per second	m2/s				
	square millimeter per second	mm2/s	m2/s		1.0 E-06	Exact
	centistokes	cSt	m2/s		1.0 E-06	Exact
	stokes	St	m2/s		1.0 E-04	Exact
ength	meter	m				
	centimeter	cm	m		1.0 E-02	Exact
	millimeter	mm	m		1.0 E-03	Exact
	kilometer	km	m		1.0 E+03	Exact
	micron	μ	m		1.0 E-06	Exact
	foot	ft	m		3.048 E-01	Exact
ength per	inch meter per degree Celsius	in m/°C	m		2.54 E-02	Exact
emperature	meter per dograa Eabranhait	m/°E	m/°C		1.9	Evact
	meter per degree Fahrenheit centimeter per degree Celsius	m/°F cm/°C	m/°C m/°C		1.8 1.0 E-02	Exact Exact
	centimeter per degree	cm/°F	m/°C m/°C		1.8 E-02	Exact
	Fahrenheit	mm /°C	m /°C		1.0.5.02	Evact
	millimeter per degree Celsius millimeter per degree	mm/°C mm/°F	m/°C m/°C		1.0 E-03 1.8 E-03	Exact Exact
					TO F-00	LAGL

Type of unit	Description	Unit	Convert to	Excel constant	Multiply by	Conversion
	feet per degree Celsius	ft/°C	 m/°C	constant	3.048 E-01	Exact
	feet per degree Fahrenheit	ft/°F	m/°C		5.4864 E-01	Exact
	inches per degree Celsius	in/°C	m/°C		2.54 E-02	Exact
	inches per degree Fahrenheit	in/°F	m/°C		4.572 E-02	Exact
ass	kilogram	kg	iii/ C		4.572 2-02	LACC
1855			ka		10502	Evact
	gram	g	kg		1.0 E-03	Exact
	milligram	mg	kg		1.0 E-06	Exact
	pound mass (avoirdupois)	lbm	kg		4.5359237 E-01	Exact
	kilopound mass (avoirdupois)	klbm	kg		4.5359237 E+02	Exact
	Million pound mass (avoirdupois)	Mlbm	kg		4.5359237 E+05	Exact
	metric ton	tonne	kg		1.0 E+03	Exact
	short ton (equals 2000 lb, also	short ton	kg		9.0718474 E+02	Exact
	called tonUS) long ton (equals 2240 lb, also	long ton	kg		1.016046909 E+03	Exact
	called tonUK)		ĸy		1.010040305 2103	Exact
lass per Mass	mass fraction	mass/mass				
	mass percentage	%mass	mass/mass		1.0 E-02	Exact
ass per Mole	kilograms per mole	kg/mol				
	kilograms per kilomole	kg/kmol	kg/mol		1.0 E-03	Exact
	grams per mole	g/mol	kg/mol		1.0 E-03	Exact
	pound per pound mole	lbm/lbmol	kg/mol		1.0 E-03	Exact
	kilopound per kilopound mole	klbm/klbmol	kg/mol		1.0 E-03	Exact
lace por Dules	kilograms per pulse	,	kg/ IIIOI		1.0 2 05	LAULI
ass per Pulse		kg/pulse				
	grams per pulse	g/pulse	kg/pulse		1.0 E-03	Exact
	pounds mass per pulse	lbm/pulse	kg/pulse		4.535924 E-01	Exact
ass per Time	kilogram per second	kg/s				
	kilogram per hour	kg/hr	kg/s		(1.0/3600)	Exact
	ton (metric) per hour	tonne/hr	kg/s		(1.0/3600) E+03	Exact
	ton (short) per hour	short ton/hr	kg/s		(9.071847/3600) E+02	Exact
	ton (long) per hour	long ton/hr	kg/s		(1.016046909/3600) E+03	Exact
			-		· · · ·	
	pound mass (avoirdupois) per second	lbm/s	kg/s		4.535924 E-01	Exact
	pound mass (avoirdupois) per hour	lbm/hr	kg/s		(4.535924/3600) E-01	Exact
	kilopound mass (avoirdupois) per hour	klbm/hr	kg/s		(4.535924/3600) E+02	Exact
	kilogram per day	kg/d	kg/s		(1.0/86400)	Exact
	ton (metric) per day	tonne/d	kg/s		(1.0/86400) E+03	Exact
	ton (short) per day	short ton/d	kg/s		(9.071847/86400) E+02	Exact
	ton (long) per day	long ton/d	kg/s		(1.016046909/86400) E+03	Exact
	pound mass (avoirdupois) per day	lbm/d	kg/s		(4.535924/86400) E-01	Exact
	kilopound mass (avoirdupois) per day	klbm/d	kg/s		(4.535924/86400) E+02	Exact
1ass per Volume	kilogram per cubic meter	kg/m3				
	gram per cubic centimeter	g/cm3	kg/m3		1.0 E03	Exact
	pound per cubic foot	lbm/ft3	kg/m3		1.601846337 E+01	Exact
ass per Pulse		-				
	API gravity ⁽¹⁾	°API	kg/m3		ρ _{H2O,60°F} * 141.5 / (°API+131.5) (2)	Exact ⁽³⁾
	Specific Gravity at 60 degrees Fahrenheit ⁽¹⁾	SG @ 60°F	kg/m3		ρ _{H20,60°F} ⁽²⁾	Exact ⁽³⁾
	Relative Density at 60 degrees Fahrenheit ⁽¹⁾	RD @ 60°F	kg/m3		ρ _{H2O,60°F} ⁽²⁾	Exact ⁽³⁾
	⁽¹⁾ This conversion only applies					
	when conversion is to /from the					
	'Mass per Volume' value at 60 °F					
	$^{(2)}$ $\rho_{H20,60^{\circ}F}$ the density of water					
	at 60 °F is a global setting with					
	a default value of 999.012					
	kg/m3					
	⁽²⁾ The conversion is exact,					
	however the resulting value is an approximation because of					
	Р H2O,60°F					
	kilogram per standard cubic	kg/sm3				
Mass per Standard	meter				1.0 E03	Exact
•	meter gram per standard cubic	g/scm3	kg/sm3		1.0 205	LAUCE
1ass per Standard /olume		g/scm3	kg/sm3		1.0 203	Exact
•	gram per standard cubic	g/scm3 lbm/scf	kg/sm3 kg/sm3		1.601846337 E+01	Exact
olume	gram per standard cubic centimeter pound per standard cubic foot kilogram per normal cubic					
	gram per standard cubic centimeter pound per standard cubic foot	lbm/scf				

Type of unit	Description	Unit	Convert to	Excel constant	Multiply by	Conversion
	centimeter					
lole	mole	mol				
	<u> </u>	lune el			105.02	E
						Exact
	•	Ibmol	mol		4.5359237 E-01	Exact
	5					
	•					
	-				4 5250227 5 02	Event
			moi		4.5359237 E+02	Exact
lole per Mole		•				
			mole/mole		1.0 E-02	Exact
Iole per Volume	•	•				
	mole per cubic centimeter	•			1.0 E+6	Exact
	mole per litre	•	-			Exact
	kilomole per cubic meter	,	mol/m ³		1.0 E+3	Exact
	kilomole per cubic centimeter	kmol/cm³	mol/m³		1.0 E+9	Exact
	kilomole per litre	kmol/l	mol/m³		1.0 E+6	Exact
	kilomole per cubic feet	kmol/ft³	mol/m³		1 / 28.31685 E -03	Exact
Power	watt	W				
	kilowatt	kW	W		1.0 E+03	Exact
	megawatt	MW	W		1.0 E+06	Exact
	gigawatt	GW	W		1.0 E+09	Exact
Pressure		Pa				
centimeter mol Mole mole mol A mole resembles 6.0251 x 10^23 molecules of a substance, a standard number of molecules known as Avogadro's number. kmol mol 1.0 E+03 pound mole kmol mol 4.5359237 E- In English units, the pound-mass (lbm) is the standard unit of mass. In order to use the same molecular weights as those listed on the periodic chart, the pound-mol, or lbm-mole) is defined mole/mole 4.5359237 E- Mole per Mole mole rection mole/mole 4.5359237 E- Mole per Mole nole rouse the same molecular weights as those listed on the periodic chart, the pound-mol, or lbm-mole) is defined 4.5359237 E- Mole per Mole mole fraction mole/mole 4.5359237 E- Mole per Mole mole mole or 1.0 E-02 Mole per Lobic meter mol/mole 4.5359237 E- Mole per Cubic meter mol/mole 1.0 E-02 Mole per Cubic meter mol/mole 4.5359237 E- Mole per Cubic centimeter mol/mole 1.0 E+02 Mole per Cubic centimeter mol/mole 1.0 E+02 Mole per cubic cent						
	kilo pascal	kPa	Pa		1.0 E+03	Exact
	•					Exact
	<u> </u>		-			Exact
	5	Kgr/ chile	i u		5.00003 2.04	Exact
		lhf/ft2	Pa		47 8803	Exact
		•				Exact
			Fa		0894.70	Exact
		nei	Pa		6894 76	Exact
		p31	īα		0004.10	Exact
		bar	Pa		1 0 E+05	Exact
						Exact
			-			Exact
	2	ттнд	Pa		133.322387415	Exact
			D -		0.00000	E
						Exact
	¥				•	Approximate
						Exact
	· · · ·					Approximate
	,,,		Pa			Approximate
	inch of water, convential	inH2O con	Pa		249.08891	Exact
	inch of water @ 39.2°F (4°C)	inH2O @ 39.2°F	Pa		249.082	Approximate
	inch of water @ 60°F	inH2O @ 60°F	Pa		248.84	Approximate
	inch of water @ 68°F	inH2O @ 68°F	Pa			Approximate
Pressure (absolute)		-				
	•		Pa(a)		1.0 E+03	Exact
	•					Exact
	· · · ·		/			
		bar(a)	Pa(a)		1 0 F+05	Exact
						Exact
		ппппуа	ra(a)		133.32238/415	Exact
		mmU20-			0.80665	Eve at
		mmH2Oa	Pa(a)		9.8005	Exact
		mm1120- 0 0005			240.04/25.4	A
		mmH2Oa @ 60°F	Pa(a)		248.84/25.4	Approximate
		•	- / \			
	2	ınHga con	Pa(a)		3386.38864	Exact
		• • •				
		inHga @ 32°F	Pa(a)		3386.38	Approximate
	inch of mercury @ 60°F absolute	inHga @ 60°F	Pa(a)		3376.85	Approximate
	inch of water, convential	inH2Oa con	Pa(a)		249.08891	Exact

Type of unit	Description	Unit	Convert to	Excel constant	Multiply by	Conversion
	inch of water @ 39.2°F (4°C) absolute	inH2Oa @ 39.2°F	Pa(a)		249.082	Approximate
	inch of water @ 60°F absolute	inH2Oa @ 60°F	Pa(a)		248.84	Approximate
	inch of water @ 68°F absolute	inH2Oa @ 68°F	Pa(a)		248.64107	Approximate
Pressure (gauge)	pascal gauge					
	kilo pascal gauge					Exact
	pound-force per square inch (psi) gauge	psig	Pa(g)		6894.76	Exact
	bar gauge	bar(g)	Pa(g)		1 0 F+05	Exact
	millibar gauge	-	-			Exact
	millimeter of mercury,				133.322387415	Exact
	convential gauge millimeter of water, convential	mmH2Og	Pa(g)		9.80665	Exact
	gauge millimeter of water @ 60°F	mmH2Og @ 60°F	Pa(g)		248.84/25.4	Approximate
	gauge inch of mercury, convential	inHgg con	Pa(g)		3386.38864	Exact
	gauge					
	inch of mercury @ 32°F (0°C) gauge					Approximate
	inch of mercury @ 60°F gauge					Approximate
	inch of water, convential gauge inch of water @ 39.2°F (4°C) gauge	inH2Og con inH2Og @ 39.2°F	Pa(g) Pa(g)		249.08891 249.082	Exact Approximate
	inch of water @ 60°F gauge	inH2Og @ 60°F	Pa(g)		248.84	Approximate
Pressure inverse	per pascal	•	1 /D-		10502	Eve -t
	per kilo pascal	•	•			Exact
	per Mega pascal	1 -	,			Event
	(psi)	1/psi	1/Pa		1/6894.76	Exact
	per bar	1/bar	1/Pa		1.0 E-05	Exact
ressure per Mass	pascals per kilogram	•	_,			
	kilopascals per kilogram		Pa/kg		1.0 E+03	Exact
	megapascals per kilogram	MPa/kg	Pa/kg		1.0 E+06	Exact
	pounds mass (avoirdupois) per	psi/kg	Pa/kg		6894.76	Exact
	square inch per kilogram					
	bar per per kilogram	-	Pa/kg		1.0 E+05	Exact
Pulses per Mass	pulses per kilogram	· -			105.00	
	pulses per gram	1 / 5				Exact
Pulsos por Volumo	(avoirdupois)		puises/kg		1/0.4555924	Exact
Pulses per Volume	· · ·		pulsos /m2		105.06	Exact
	pulses per cubic centimeter		· · ·			Exact
	pulses per cubic inch		· ·			Exact
	pulses per cubic freet					Exact
Temperature	Kelvin		puises/115		1,0.0203100	Exact
	degree Celsius		К		T[K] = t[°C] + 273.15	Exact
	degree Fahrenheit					Exact
	Rankine	R	К		T[K] = T[R]/1.8	Exact
Femperature nverse	per Kelvin	to constant by ater @ 30-7" int20a @ 30-7" Pa(a) 249.082 Adviter @ 60" ater @ 60" absolute int20a @ 60" Pa(a) 248.84107 Adviter @ 60" ater @ 60" absolute int20a @ 60" Pa(a) 248.84107 Adviter @ 60" algange Pa(a) 10.6"+03 E algange Pa(a) 10.6"+03 E ge bar(g) Pa(a) 10.6"+03 E ge corr per square inch psig 10.6"+02 E ge bar(g) Pa(a) 10.6"+03 E ge or formeruny, mmH2Og Pa(a) 3386.38864 E er of water @ 60"F mmH2Og @ 60"F Pa(a) 3386.38864 E nercuny @ 60"F mH2Og @ 60"F Pa(a) 248.844 Ad atter, Gorvential inH3@ @ 32*F Pa(a) 240.08891 E atter @ 60"F gauge inH2Og @ 60"F Pa(a) 240.08891 E atter, Gorvential intPa 10.6"-06				
	per degree Celsius		· ·			Exact
	per degree Fahrenheit	•				Exact
	per Rankine		1/K		1.8	Exact
lemperature per Pressure	degree Celsius per bar (Joule-Thomson coefficient)	-	20.4		10(000) ==	
Fime	cocond	· · · · · · · · · · · · · · · · · · ·	°C/bar		1.8/6894.76	Exact
Гіme	second milli second		<u> </u>		10502	Evact
	milli second micro second					Exact
	nano second	· · · · · · · · · · · · · · · · · · ·				Exact Exact
	minute					Exact
	hour					Exact
	day					Exact
/elocity	meters per second		~			
	kilometers per second		m/s		1.0 E+03	Exact
	•					Exact
	Kilometers ber nour				· · · · · · · · = ·	
	foot per second	-	m/s		3.048 E-01	Exact
/olume		ft/s	m/s		3.048 E-01	Exact
Volume	foot per second	ft/s	m/s m3		3.048 E-01 1.0 E-06	Exact Exact

Type of unit	Description	Unit	Convert to	Excel constant	Multiply by	Conversion
	mega cubic meter	Mm3	m3		1.0 E+06	Exact
	liter	L	m3		1.0 E-03	Exact
	cubic inch	in3	m3		16.38706 E+06	Exact
	cubic foot	ft3	m3		28.31685 E -03	Exact
	kilo cubic foot	kft3	m3		28.31685	Exact
	million cubic foot	MMft3	m3		28.31685 E +03	Exact
	barrel (42 US liquid gallons exactly)	bbl	m3		0.158987295	Exact
	US liquid gallon (231 cubic inches exactly)	US.gal	m3		3.785411784 E-03	Exact
Standard Volume	Imperial (U.K.) gallon standard cubic meter	UK.gal	m3		4.54609 E-03	Exact
Standard volume		sm3	2		105:02	E
	kilo standard cubic meter	ksm3	sm3		1.0 E+03	Exact
	mega standard cubic meter	Msm3	sm3		1.0 E+06	Exact
	standard cubic foot	scf	sm3		28.31685 E -03	Exact
	kilo standard cubic foot	kscf	sm3		28.31685	Exact
	million standard cubic foot	MMscf	sm3		28.31685 E +03	Exact
	barrel (standard)	bbl (s)	sm3		0.158987295	Exact
Iormal Volume	normal cubic meter	m3(n)	5115		0.130301233	LAUCE
Normal volume					105.00	
	kilo normal cubic meter	km3(n)	m3(n)		1.0 E+03	Exact
	mega normal cubic meter	Mm3(n)	m3(n)		1.0 E+06	Exact
olume per Volume	volume fraction	vol/vol				
	volume percentage	%vol	vol/vol		1.0 E-02	Exact
/olume per Pulse	cubic meters per pulse	m3/pulse	-			
	cubic centimeters per pulse	cm3/pulse	m3/pulse		1.0 E-06	Exact
			m3/pulse		1.0 E-03	
	litres per pulse	l/pulse	/1			Exact
	cubic inches per pulse	in3/pulse	m3/pulse		1.63871E-05	Exact
	cubic feet per pulse	ft3/pulse	m3/pulse		0.0283168	Exact
/olume per Time	cubic meter per second	m3/s				
	cubic meter per hour	m3/hr	m3/s		1/3600	Exact
	cubic meter per day	m3/d	m3/s		1/86400	Exact
		-			· · ·	
	kilo cubic meter per second	km3/s	m3/s		1.0 E+03	Exact
	kilo cubic meter per hour	km3/hr	m3/s		(1/3600) E+03	Exact
	kilo cubic meter per day	km3/d	m3/s		(1/86400) E+03	Exact
	mega cubic meter per hour	Mm3/hr	m3/s		(1/3600) E+06	Exact
	mega cubic meter per day	Mm3/d	m3/s		(1/86400) E+06	Exact
	cubic feet per hour	ft3/hr	m3/s		0.02831685/3600	Exact
	· · · · · · · · · · · · · · · · · · ·	•			· · ·	
	cubic feet per day	ft3/d	m3/s		0.02831685/86400	Exact
	kilo cubic feet per hour	kft3/hr	m3/s		28.31685/3600	Exact
	kilo cubic feet per day	kft3/d	m3/s		28.31685/86400	Exact
	million cubic feet per hour	Mft3/hr	m3/s		28316.85/3600	Exact
	million cubic feet per day	Mft3/d	m3/s		28316.85/86400	Exact
	barrels per hour	bbl/hr	m3/s		0.158987295/3600	Exact
	•					
Standard Volume Der Time	barrels per day standard cubic meter per second	bbl/d sm3/s	m3/s		0.158987295/86400	Exact
	standard cubic meter per hour	sm3/hr	sm3/s		1/3600	Exact
	standard cubic meter per day	sm3/d	sm3/s		1/86400	Exact
	kilo standard cubic meter per second	ksm3/s	sm3/s		1.0 E+03	Exact
	kilo standard cubic meter per hour	ksm3/hr	sm3/s		(1/3600) E+03	Exact
	kilo standard cubic meter per day	ksm3/d	sm3/s		(1/86400) E+03	Exact
	mega standard cubic meter per hour	Msm3/hr	sm3/s		(1/3600) E+06	Exact
	mega standard cubic meter per day	Msm3/d	sm3/s		(1/86400) E+06	Exact
	standard cubic feet per hour	scf/hr	sm3/s		0.02831685/3600	Exact
	standard cubic feet per day	scf/d	sm3/s		0.02831685/86400	Exact
	kilo standard cubic feet per hour		sm3/s		28.31685/3600	Exact
	•		· · · · · · · · · · · · · · · · · · ·		· · · · · · · · · · · · · · · · · · ·	
	kilo standard cubic feet per day	kscf/d	sm3/s		28.31685/86400	Exact
	million standard cubic feet per hour	MMscf/hr	sm3/s		28316.85/3600	Exact
	million standard cubic feet per day	MMscf/d	sm3/s		28316.85/86400	Exact
	barrels per hour (standard)	bbl/hr	sm3/s		0.158987295/3600	Exact
Normal Volume per	barrels per day (standard) normal cubic meter per second	bbl/d m3(n)/s	sm3/s		0.158987295/86400	Exact
Time						
	normal cubic meter per hour	m3(n)/hr	m3(n)/s		1/3600	Exact
	normal cubic meter per day	m3(n)/d	m3(n)/s		1/86400	Exact
					-,	

Type of unit	Description	Unit	Convert	Excel	Multiply	Conversion
			to	constant	by	
	kilo normal cubic meter per second	km3(n)/s	m3(n)/s		1.0 E+03	Exact
	kilo normal cubic meter per hour	km3(n)/hr	m3(n)/s		(1/3600) E+03	Exact
	kilo normal cubic meter per day	km3(n)/d	m3(n)/s		(1/86400) E+03	Exact
	mega normal cubic meter per hour	Mm3(n)/hr	m3(n)/s		(1/3600) E+06	Exact
	mega normal cubic meter per day	Mm3(n)/d	m3(n)/s		(1/86400) E+06	Exact

Terminology

Term	Description	Same as
Heating Value	Usually the same as Gross Heating Value	
Calorific Value	Usually the same as Superior Calorific Value	
Superior Calorific Value	Heating value when assuming that water formed at the combustion stays in the gaseous	Gross Heating Value
	state. From ISO6976.	
Inferior Calorific Value	Heating value when assuming that water formed at the combustion has totally condensed to the liquid state. From ISO6976.	Net Heating Value
Gross Heating Value	Heating value when assuming that water formed at the combustion stays in the gaseous state Term used in GPA2172.	Superior Calorific Value
Net Heating Value	Heating value when assuming that water formed at the combustion has totally condensed to the liquid state. Term used in GPA2172.	Inferior Calorific Value
Molar Mass Ratio	Ratio of molar mass of gas and molar mass of air at the base conditions	Specific Gravity Ideal Specific Gravity Ideal Relative Density
Relative Density	Ratio of real mass density of gas and real mass density of air at the base conditions	Real Relative Density Real Specific Gravity
Specific Gravity	Ratio of real mass density of gas and real density of air at the base conditions	Molar Mass Ratio Ideal Specific Gravity Ideal Relative Density

Standard composition

The Standard Composition is a standard array of mole fractional values that is used by all functions that require a (partial) compositional analysis.

The following table defines the sequence of the components and also defines which function uses which component.

Component	Used in AGA8 / AGA10	Used in ISO6976	Used in GPA2172	Used in AGA5	Used in GERG 2008	Used in MR113
Methane		\checkmark				
Nitrogen						
Carbon Dioxide						
Ethane						
Propane						
Water			2)			3)
Hydrogen Sulphide						
Hydrogen						
Carbon Monoxide						
Oxygen						
-Butane						
n-Butane						
-Pentane						
n-Pentane						
n-Hexane						
n-Heptane						
n-Octane						
n-Nonane						
r-Decane						
Helium						
Argon						
Neo-Pentane	(1)		(1)		(1)	(1)

1) Depending on function input 'Neo-Pentane mode' the value is added to i-Pentane or n-Pentane or it is neglected.

2) GPA2172 uses the specified water fraction for wet gas calculation only.

3) MR113 requires the input water fraction to be 0. The output water fraction is calculated from absolute humidity

6 Revisions

Revision A

Date March 2017

- Initial, release
- Added IUPAC Ethylene and IAWS-IF97 functions
- Updated and corrected IO functions, moved to separate chapter.
- Updated incorrect page-header in TOC.
- Removed PPRV output from fxPeriodLatch(..) function.
- Added extended ISO6976 function that takes all 55 components of the standard
- Added GERG2004 and GERG2008 functions
- Minor editorial changes
- Added new natural gas viscosity calculation
- Added C11 C21 components to the extended ISO6976 functions
- Added "Quarter circle" and "Conical entrance" orifice calculations
- Added GSSSD method MR 113-03 function
- Raised the pressure limit of API 2004 functions (tables 53, 54, 59, 60) from 150 to 250 bar.
- Removed duplicated functions.
- Added GOST30319 calculations.

Revision B

Date November 2017

- Update to new ABB lay-out
- New document code: CM/FlowX/FR-EN
- Reintroduce revisions chapter

Revision C

Date March 2018

- New calculations are added.
- Various existing functions are corrected.



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