
Natural gas — Energy determination

Gaz naturel — Détermination de l'énergie

STANDARDSISO.COM : Click to view the full PDF of ISO 15112:2018



STANDARDSISO.COM : Click to view the full PDF of ISO 15112:2018



COPYRIGHT PROTECTED DOCUMENT

© ISO 2018

All rights reserved. Unless otherwise specified, or required in the context of its implementation, no part of this publication may be reproduced or utilized otherwise in any form or by any means, electronic or mechanical, including photocopying, or posting on the internet or an intranet, without prior written permission. Permission can be requested from either ISO at the address below or ISO's member body in the country of the requester.

ISO copyright office
CP 401 • Ch. de Blandonnet 8
CH-1214 Vernier, Geneva
Phone: +41 22 749 01 11
Fax: +41 22 749 09 47
Email: copyright@iso.org
Website: www.iso.org

Published in Switzerland

Contents

Page

Foreword	v
Introduction	vi
1 Scope	1
2 Normative references	1
3 Terms and definitions	1
4 Symbols and units	6
5 General Principles	7
6 Gas measurement	8
6.1 General	8
6.2 Volume measurement	9
6.3 Calorific value measurement	9
6.3.1 Measurement techniques and sampling	9
6.3.2 Direct measurement — Calorimetry	10
6.3.3 Inferential measurement	10
6.3.4 Correlation techniques	10
6.3.5 Pressure and temperature	10
6.3.6 Gas quality tracking	10
6.4 Volume conversion	10
6.4.1 General	10
6.4.2 Density	11
6.4.3 Compression factor	11
6.5 Calibration	11
6.6 Data storage and transmission	11
7 Energy determination	12
7.1 Interfaces	12
7.2 Methods of energy determination	14
7.2.1 Direct determination of energy	14
7.2.2 Indirect determination of energy	14
8 Strategy and procedures	16
8.1 General	16
8.2 Strategies for energy determination	17
8.2.1 Strategies for single interfaces	18
8.3 Plausibility checks	22
9 Assignment methods	23
9.1 Fixed assignment	23
9.1.1 Fixed assignment of a measured calorific value	23
9.1.2 Fixed assignment of a declared calorific value	24
9.2 Variable assignment	25
9.2.1 Input at two or more different stations with zero floating point	25
9.2.2 Input at two or more different stations with comingled gas flows	26
9.3 Determination of the representative calorific value	27
9.3.1 Arithmetically averaged calorific value	27
9.3.2 Quantity-weighted average calorific value	27
9.3.3 Gas quality tracking	27
10 Calculation of energy quantities	30
10.1 General formulae for energy	30
10.2 Calculation of averaged values — Calculation from average calorific values and cumulative volumes	31
10.2.1 Arithmetic average of the calorific value	31
10.2.2 Quantity-weighted average of the calorific value	32

10.3	Volume and volume-to-mass conversions.....	32
10.4	Energy determination on the basis of declared calorific values	32
11	Accuracy on calculated energy.....	32
11.1	Accuracy.....	32
11.2	Calculation of uncertainty	33
11.3	Bias.....	34
12	Quality control and quality assurance.....	35
12.1	General.....	35
12.2	Check of the course of the measuring data.....	35
12.3	Traceability	36
12.4	Substitute values	36
Annex A	(informative) Main instruments and energy-determination techniques.....	38
Annex B	(informative) Different possible patterns in the change of the calorific value.....	42
Annex C	(informative) Volume conversion and volume-to-mass conversion.....	45
Annex D	(informative) Incremental energy determination	46
Annex E	(informative) Practical examples for volume conversion and energy quantity calculation.....	48
Annex F	(informative) Practical examples for averaging the calorific value due to different delivery situations.....	52
Annex G	(informative) Ways of determining substitute values.....	57
Annex H	(informative) Plausibility check graphical example.....	59
Annex I	(informative) Uncorrected data, bias correction and final result graphical example.....	60
Annex J	(informative) Single-reservoir calorific value determination.....	62
Annex K	(informative)	64
Bibliography		70

Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

The procedures used to develop this document and those intended for its further maintenance are described in the ISO/IEC Directives, Part 1. In particular the different approval criteria needed for the different types of ISO documents should be noted. This document was drafted in accordance with the editorial rules of the ISO/IEC Directives, Part 2 (see www.iso.org/directives).

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. ISO shall not be held responsible for identifying any or all such patent rights. Details of any patent rights identified during the development of the document will be in the Introduction and/or on the ISO list of patent declarations received (see www.iso.org/patents).

Any trade name used in this document is information given for the convenience of users and does not constitute an endorsement.

For an explanation on the voluntary nature of standards, the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the World Trade Organization (WTO) principles in the Technical Barriers to Trade (TBT) see the following URL: www.iso.org/iso/foreword.html.

This document was prepared by Technical Committee ISO/TC 193, *Natural gas*.

This third edition cancels and replaces the second edition (ISO 15112:2011), which has been technically revised.

The main changes compared to the previous edition are as follows:

- [Figures 7](#) and [8](#) have been redrafted;
- [Clause 9](#) has been updated;
- [Annex K](#) has been added.

Any feedback or questions on this document should be directed to the user's national standards body. A complete listing of these bodies can be found at www.iso.org/members.html.

Introduction

Since the early 1800s, it has been general practice for manufactured gas and, subsequently, natural gas to be bought and sold on a volumetric basis. Much time and effort has therefore been devoted to developing the means of flow measurement.

Because of the increasing value of energy and variations in gas quality, billing on the basis of thermal energy has now become essential between contracting partners and the need to determine calorific value by measurement or calculation has led to a number of techniques. However, the manner in which calorific value data are applied to flow volume data to produce the energy content of a given volume of natural gas has been far from a standardized procedure.

Energy determination is frequently a necessary factor wherever and whenever natural gas is metered, from production and processing operations through to end-user consumption. This document has been developed to cover aspects related to production/transmission and distribution/end user. It provides guidance to users of how energy units for billing purposes are derived, based on either measurement or calculation or both, to increase confidence in results for contracting partners.

Other standards relating to natural gas, flow measurement, calorific value measurement, calculation procedures and data handling with regard to gas production, transmission and distribution involving purchase, sales or commodity transfer of natural gas can be relevant to this document.

This document contains eleven informative annexes.

STANDARDSISO.COM : Click to view the full PDF of ISO 15112:2018

Natural gas — Energy determination

1 Scope

This document provides the means for energy determination of natural gas by measurement or by calculation, and describes the related techniques and measures that are necessary to take. The calculation of thermal energy is based on the separate measurement of the quantity, either by mass or by volume, of gas transferred and its measured or calculated calorific value. The general means of calculating uncertainties are also given.

Only systems currently in use are described.

NOTE Use of such systems in commercial or official trade can require the approval of national authorization agencies, and compliance with legal regulations is required.

This document applies to any gas-measuring station from domestic to very large high-pressure transmission.

New techniques are not excluded, provided their proven performance is equivalent to, or better than, that of those techniques referred to in this document.

Gas-measuring systems are not the subject of this document.

2 Normative references

The following documents are referred to in the text in such a way that some or all of their content constitutes requirements of this document. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO 6976, *Natural gas — Calculation of calorific values, density, relative density and Wobbe index from composition*

3 Terms and definitions

For the purposes of this document, the following terms and definitions apply.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

- ISO Online browsing platform: available at <https://www.iso.org/obp>
- IEC Electropedia: available at <https://www.electropedia.org/>

3.1

accuracy of measurement

closeness of the agreement between the result of a measurement and a true value of the measurand

[SOURCE: ISO/Guide 98-3:2008, definition B.2.14]

3.2

adjustment

<of a measuring instrument> of bringing a measuring instrument into a state of performance suitable for its use

Note 1 to entry: Adjustment may be automatic, semi-automatic or manual.

3.3

assignment method

<energy determination> method to derive a calorific value to be applied to the gas passing specified interfaces having only volume measurements

3.4

availability

probability, at any time, that the measuring system, or a measuring instrument forming part of the measuring system, is functioning according to specifications

[SOURCE: EN 1776:1998]

3.5

bias

systematic difference between the true energy and the actual energy determined of the gas passing a gas-measuring station

3.6

calibration

set of operations that establish, under specified conditions, the relationship between values of quantities indicated by a measuring instrument or measuring system, or values represented by a material measure or a reference material, and the corresponding values obtained using working standards

[SOURCE: ISO 14532:2014, definition 2.5.1.1, modified — Definition has been slightly changed and Notes to entry have been removed.]

3.7

superior calorific value

energy released as heat by the complete combustion in air of a specified quantity of gas, in such a way that the pressure, p_1 , at which the reaction takes place remains constant, and all the products of combustion are returned to the same specified temperature, T_1 , as that of the reactants, all of these products being in the gaseous state except for water formed by combustion, which is condensed to the liquid state at T_1

[SOURCE: ISO 14532:2014, definition 2.6.4.1, modified — Definition has been slightly reworded and Notes to entry have been removed.]

3.8

inferior calorific value

energy released as heat by the complete combustion in air of a specified quantity of gas, in such a way that the pressure, p_1 , at which the reaction takes place remains constant, and all the products of combustion are returned to the same specified temperature, T_1 , as that of the reactants, all of these products being in the gaseous state

[SOURCE: ISO 14532:2014, definition 2.6.4.2, modified — Definition has been slightly reworded and Notes to entry have been removed.]

3.9

calorific value station

installation comprising the equipment necessary for the determination of the calorific value of the natural gas in the pipeline

3.10

adjusted calorific value

calorific value measured at a measuring station compensated for the time taken for the gas to travel to the respective volume-measuring station

3.11

corrected calorific value

result of correcting a measurement to compensate for systematic error

3.12**declared calorific value**

calorific value that is notified in advance of its application to interfaces for the purpose of energy determination

3.13**representative calorific value**

calorific value which is accepted to sufficiently approximate the actual calorific value at an interface

3.14**charging area**

set of interfaces where the same method of energy determination is used

3.15**conversion**

determination of the volume under reference conditions from the volume under operating conditions

3.16**correction**

value added algebraically to the uncorrected result of a measurement to compensate for systematic error

Note 1 to entry: The correction is equal to the negative of the estimated systematic error.

Note 2 to entry: Since the systematic error cannot be known perfectly, the correction cannot be complete, see [Annex I](#).

3.17**correction factor**

numerical factor by which the uncorrected result of a measurement is multiplied to compensate for a systematic-error object

Note 1 to entry: Since the systematic error cannot be known perfectly, the correction cannot be complete, see [Annex I](#).

3.18**determination**

set of operations that are carried out on an object in order to provide qualitative or quantitative information about this object

Note 1 to entry: In this document, the term “determination” is only used quantitatively.

3.19**direct measurement**

measurement of a property from quantities which, in principle, define the property

Note 1 to entry: For example, the determination of the calorific value of a gas using the thermoelectric measurement of the energy released in the form of heat during the combustion of a known amount of gas.

[SOURCE: ISO 14532:2014, definition 2.2.1.2, modified — The word “that” has been replaced by “which” in the definition.]

3.20**energy**

product of gas quantity (mass or volume) and calorific value under given conditions

Note 1 to entry: The energy may be called energy amount.

Note 2 to entry: Energy is usually expressed in units of megajoules.

3.21

energy determination

quantitative determination of the amount of energy of a quantity of gas based either on measurement or calculation using measured values

3.22

energy flow rate

energy of gas passing through a cross-section divided by time

Note 1 to entry: Energy flow rate is usually expressed in units of megajoules per second.

3.23

fixed assignment

application without modification of the calorific value measured at one specific calorific-value-measuring station, or the calorific value declared in advance, to the gas passing one, or more, interfaces

3.24

gas transporter

company that conveys gas from one place to another through pipelines

3.25

gas quality tracking

determination of gas quality properties (e. g. the calorific value) at the exit points of a gas grid based on flow calculation; the calculation requires topology data, gas quality data at entry points, volume data at entry and exit points and grid pressures as input information

3.26

interface

place on a pipe used for the transportation or supply of gas at which there is a change of ownership or physical custody of gas

Note 1 to entry: Generally, an interface has an associated measuring station.

3.27

local distribution company

LDC

company that delivers gas to industrial, commercial and/or residential customers

3.28

measuring station

installation comprising all the equipment, including the inlet and outlet pipework as far as the isolating valves and structure within which the equipment is housed, used for gas measurement in custody transfer

[SOURCE: EN 1776:1998]

3.29

measuring system

complete set of measuring instruments and auxiliary equipment assembled to carry out specified measurements

[SOURCE: ISO/IEC Guide 99:2007, definition 3.2, modified — Definition has been slightly reworded.]

3.30

measuring instrument

device intended to be used for making measurements, alone or in conjunction with one or more supplementary devices

[SOURCE: ISO/IEC Guide 99:2007, definition 3.1, modified — Definition has been slightly reworded.]

3.31**plausibility**

property of a value to be within reasonable limits

3.32**producer**

company that extracts raw natural gas from reservoirs which, after processing and (fiscal) measurement, is supplied as dry natural gas to the transportation system

3.33**regional distributor**

company that conveys gas to local distribution companies and/or industrial, commercial or residential customers

3.34**residential customer**

person whose occupied premises are supplied with gas, wholly or in part, such gas not being used for any business purpose, commercial or industrial

3.35**systematic error**

mean that would result from an infinitive number of measurements of the same measurand carried out under repeatability conditions minus a true value of the measurand

3.36**traceability**

property of the result of a measurement or the value of a standard whereby it can be related to stated references, usually national or International Standards, through an unbroken chain of comparisons all having stated uncertainties

Note 1 to entry: This chain of comparisons is called a traceability chain.

3.37**uncertainty**

parameter, associated with the result of a measurement, that characterizes the dispersion of the values that could reasonably be attributed to the measurand

3.38**variable assignment**

application of a calorific value for an assignment procedure based on measurement(s) at calorific value station(s) to the gas passing one, or more, interfaces

Note 1 to entry: That applied calorific value may take into account the time taken for the gas to travel from the calorific value station to the respective volume-measuring stations and other factors, to derive an average calorific value for a network, a state reconstruction of the variation of calorific values through a network, etc.

3.39**zero floating point**

position in a grid conveying gas where there is a boundary with different gas qualities on either side

3.40**non-plausible data**

measurement data that are obviously wrong taking into account the measurement situation at a measuring station and the gas flow situation

3.41**grid node**

connection of two or more pipes in a gas grid, grid nodes typically exist at interfaces (entry/exit) or at points where the pipe geometry changes

3.42

standard load profile

SLP

standard load profile (SLP) is a model to predict the expected hourly or daily energy consumption of customers where the reading is taken only periodically (e.g. once per year)

4 Symbols and units

Symbol	Meaning	SI unit	Customer unit
E	energy	MJ	kWh
e	energy flow rate	MJ/s	kWh/h
H	calorific value	MJ/m ³ ; MJ/kg	kWh/m ³

NOTE 1 Where the calorific value is in megajoules per cubic metre and the gas volume is in cubic metres, or where the calorific value is in megajoules per kilogram and the gas mass is in kilograms, then the calculated energy is in megajoules.

Where the calorific value is in kilowatt-hours per cubic metre and the gas volume is in cubic metres, or where the calorific value is in kilowatt-hours per kilogram and the gas mass is in kilograms, then the calculated energy is in kilowatt-hours.

To convert the number of megajoules to the number of kilowatt-hours, divide the number by 3,6.

M	mass	kg	t
p	pressure (absolute)	Pa, kPa	bar, mbar
Q	quantity of gas	m ³ , kg	ft ³

NOTE 2 When the quantity is given in cubic metres, it is necessary that it should be qualified by temperature and pressure.

sq_v	volume flow rate	m ³ /h, m ³ /s	
q_m	mass flow rate	kg/s, kg/h	
T	temperature (absolute)	K	
t	time	s, h, d	s, h, d
V	volume (gas)	m ³	
Z	compression factor		
ρ	density	kg/m ³	
ϑ	temperature	°C	°F

Subscripts

i	inferior calorific value
j	number of time intervals

n	normal reference conditions (273,15 K; 101,325 kPa)
r	ISO-recommended standard reference conditions (288,15 K; 101,325 kPa)
s	superior calorific value

5 General Principles

The quantity of energy, E , contained in a given quantity of gas, Q , is given by the multiplication of the calorific value, H , by the respective quantity of gas.

Energy may be either measured directly (see [Figure 1](#)) or calculated from the quantity and the calorific value of the gas (see [Figure 2](#)).

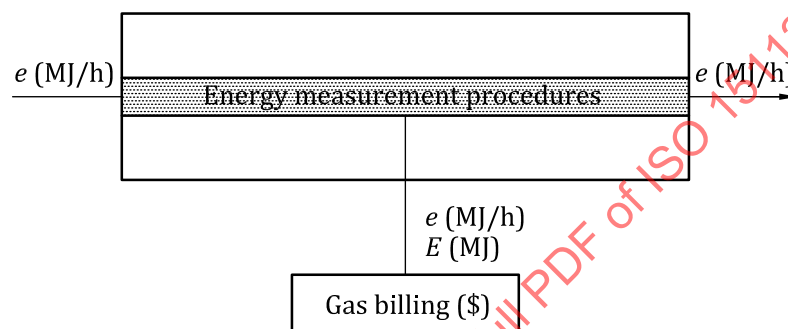


Figure 1 — Energy-measurement scheme

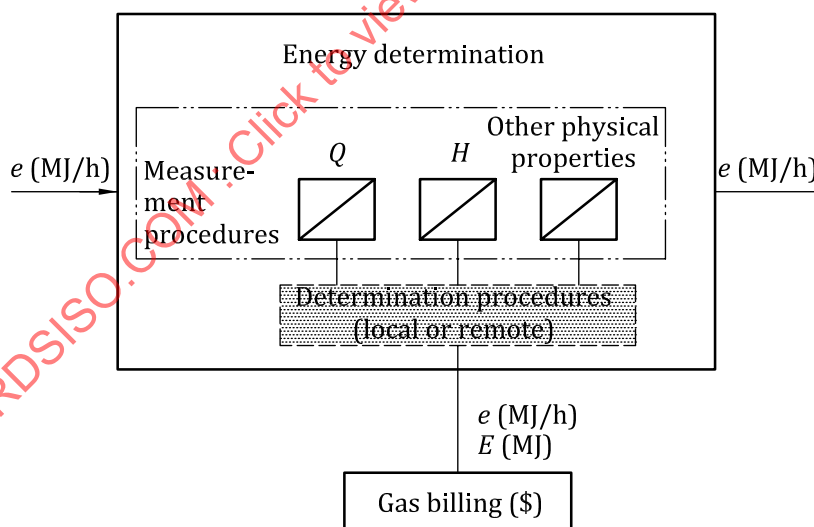


Figure 2 — Energy-determination scheme

Generally, the quantity of gas is expressed as a volume and the calorific value is on a volumetric basis. In order to achieve accurate determinations of energy, it is necessary that both the gas volume and calorific value be under the same reference conditions. The determination of energy is based either on the accumulation over time of calculation results from consecutive sets of calorific values and the concurrent flow rate values, or on the multiplication of the total volume and the representative (assigned) calorific value for that period.

Especially in situations of varying calorific values and when flow rates are determined at a place different from that of the (representative) calorific value, the effect on the accuracy caused by the

difference in time between the determination of the flow rate and the calorific value shall be considered (see [Clause 11](#)).

The gas volume may either be measured and reported as the volume under the ISO-recommended standard reference conditions or be measured under some other conditions and converted to an equivalent volume under the ISO-recommended standard reference conditions, using an appropriate method of volume conversion. The method of volume conversion used at a specific gas-volume-measuring station may require gas quality data determined at other places. For the purpose of this document, the ISO-recommended standard reference conditions of 288,15 K and 101,325 kPa, as defined in ISO 13443, should be used.

NOTE For the gas supply, other conditions can be used, corresponding to national standards or laws. Methods for conversion between different conditions for dry natural gases are given in ISO 13443.

The calorific value may be measured at the gas-measuring station or at some other representative point and assigned to the gas-measuring station. It is also possible for the quantity of gas and the calorific value to be expressed on a mass basis.

This general principle of energy determination is extended in [Clause 10](#) to those cases when the quantity of gas is expressed on either a volumetric or a mass basis.

To achieve the calculation of the quantity of energy of the gas passing a gas-measuring station over a period of time, the methods of energy determination in [Clauses 7](#) to [10](#) are used. Such methods involve an integration over the time period; that integration may be

- of the energy flow, or
- of the gas flow rate over time to obtain the quantity of gas, which is then multiplied by the representative calorific value.

The method of integration may depend on contractual agreements or national legislation.

The general principles of energy determination in [Clauses 7](#) to [10](#) are independent of the method with which the integrations are carried out. The method of integration influences the uncertainty of the determined energy; these effects are considered in [Clause 11](#).

6 Gas measurement

6.1 General

The types of measuring devices and methods used in real measuring stations depend among other things on

- the respective national requirements,
- the flow rate,
- the commercial value of the gas,
- the gas quality variations,
- the need for redundancy, and
- the instrument specification.

Only proven methods and measuring devices/products used at the respective interfaces should be used. An overview of the techniques and procedures currently used in different countries is shown in [Annex A](#).

Methods used for flow and calorific value measurement shall be in accordance with standards, contractual agreements and/or national legislation, as appropriate. If no national legislation exists, the OIML recommendation R140 should be applied.

Action should be taken to identify and reconcile systematic effects. For example, use of different national standards, regulations and/or operating procedures can introduce systematic differences; contract partners should determine the appropriate means to overcome these differences.

The quality of the measurement results, in general, depends on the following factors:

- operating conditions;
- maintenance frequency and quality;
- calibration standards;
- sampling and clean-up;
- changes in gas composition;
- ageing of measurement devices.

A high accuracy can be achieved if the requirements fixed by the manufacturers and by officials are met and all operating procedures for operating, calibration and maintenance are strictly observed.

6.2 Volume measurement

The volume flow-metering system of a natural-gas-measuring station consists of one or more meter runs. Generally, the meters measure the gas volume flow under actual operating conditions. Standards for orifice meters (ISO 5167-1) and turbine meters (ISO 9951) exist.

The selection of a flow-metering system for a specific application depends, as a minimum, on the following:

- conditions of flow;
- flow-measuring range;
- operating conditions, especially operating pressure;
- acceptable pressure loss;
- required accuracy.

For natural-gas volume flow measurement, the instruments mostly used at the interfaces 1 to 6 (see [7.1](#)) are shown in [Annex A](#).

6.3 Calorific value measurement

6.3.1 Measurement techniques and sampling

A calorific-value measuring system consists of a sampling system and a measurement device taken from one of the following groups:

- a) direct measurement (e.g. by combustion calorimeters);
- b) inferential measurement [e.g. by a gas chromatograph (GC)];
- c) correlation techniques;
- d) gas quality tracking using measured entities.

To achieve a high accuracy of calorific value measurement, representative sampling is required. Guidelines are given in ISO 10715.

Depending on the measuring system, the operating procedures, the fluctuation of composition of the gas, and/or the quantity of gas delivered, one of the following sampling techniques can be used:

- continuous direct sampling;
- periodical spot sampling;
- incremental sampling.

Samples are taken for either online analysis or offline analysis.

6.3.2 Direct measurement — Calorimetry

With direct measurement, natural gas at a constant flow rate is burned in an excess of air and the energy released is transferred to a heat-exchange medium resulting in an increase in its temperature. The calorific value of the gas is directly related to the temperature increase.

Calorimetry is used for interfaces 1 to 3 and 5. ISO 15971 gives details of the measurement of combustion properties.

6.3.3 Inferential measurement

With inferential measurement, the calorific value shall be calculated from the gas composition in accordance with ISO 6976.

The most widely used analytical technique is gas chromatography. Procedures for the determination of the composition with defined uncertainty by gas chromatography are given in ISO 6974 (all parts). GC measurement is used at interfaces 1 to 3 and 5.

6.3.4 Correlation techniques

Correlation techniques make use of the relationships between one or more physical properties and the calorific value of the gas. Also, the principle of stoichiometric combustion can be used.

6.3.5 Pressure and temperature

Pressure and temperature measurements can be necessary for the conversion of the gas volume under operating conditions to a volume under standard reference or normal conditions. Details are given in ISO 15970.

6.3.6 Gas quality tracking

On the basis of input data (grid topology, gas quality, gas quantity, pressures), flow conditions throughout the grid shall be determined using a flow mechanics calculation. On this basis, an individual calorific value can be assigned to each interface.

6.4 Volume conversion

6.4.1 General

Conversion of a volume of natural gas measured under operating conditions to a volume under reference or base conditions is based either on a gas pressure, temperature and compression factor (pTZ -conversion) or on gas densities under operating and base conditions (density conversion).

For details, see [Annex C](#), [E.1](#) and [E.2](#), ISO 12213 (all parts) and EN 12405-1.

6.4.2 Density

The density under reference conditions (sometimes referred to as normal, standard or base density) can be required for conversion of volume data. Density under operating conditions may be measured for mass-flow determination and volume conversion.

Details are given in ISO 15970.

6.4.3 Compression factor

For gas volume conversion, the compression factor is

- calculated from the gas composition using a molar analysis (see [E.2](#) and ISO 12213-2),
- calculated using physical properties and some constituents (see [E.1](#) and ISO 12213-3), or
- measured by a Z-meter.

Details are given in ISO 15970.

The compression factor under reference conditions may also be calculated according to ISO 6976. Depending on the quantity of gas delivered and variations in pressure, temperature and gas composition at the specific metering point, the compression factor either may be set constant or shall be calculated from time to time.

The user of this document shall take account of the gas composition, especially with respect to the molar relationships of the higher hydrocarbons to each other and at high pressure. Depending on the gas composition and the pressure, methods for calculating the Z-factor on the basis of ISO 12213-2, rather than on the basis of ISO 12213-3, should be considered to avoid systematic errors.

6.5 Calibration

Quality of calibrations has a significant impact on the trueness of a measurement result. The frequency of the calibrations shall be determined according to the stability of the measurement devices. Calibrations should be traceable to appropriate standards and reference materials.

A representative calibration should be performed under conditions close to those at which the meter operates. For calorific measurement devices, calibration gases that are close to the expected calorific value or composition of the gas to be measured (see, for example, ISO 15971) should be used.

If, upon verification of any measuring instrument used for energy-determination purposes, an agreed deviation between the instrument reading and the corresponding value realized by a standard is exceeded, a calibration of the measuring instrument shall be carried out in order either

- to make adjustments to the instrument that establishes the smallest possible difference between the measured value and the value given by the standard, or
- to derive a correction that is applied to the measured value for subsequent periods to produce the correct value.

The actual process of adjustment or correction may be either manual or automatic, depending on the type of instrument.

If, at the calibration of the calorific-value-measuring device, a difference between measured and certified values occurs, for subsequent periods a correction of the measured values or adjustment shall be performed.

6.6 Data storage and transmission

All relevant data for determining energy shall be securely stored. The length of storage time and place of storage shall take into account national regulations and/or contractual conditions.

The data incorporate

- information contributing to and/or consisting of the amount of energy supplied and, where available,
- information on the data validity or the functioning of the metering station (hardware and software).

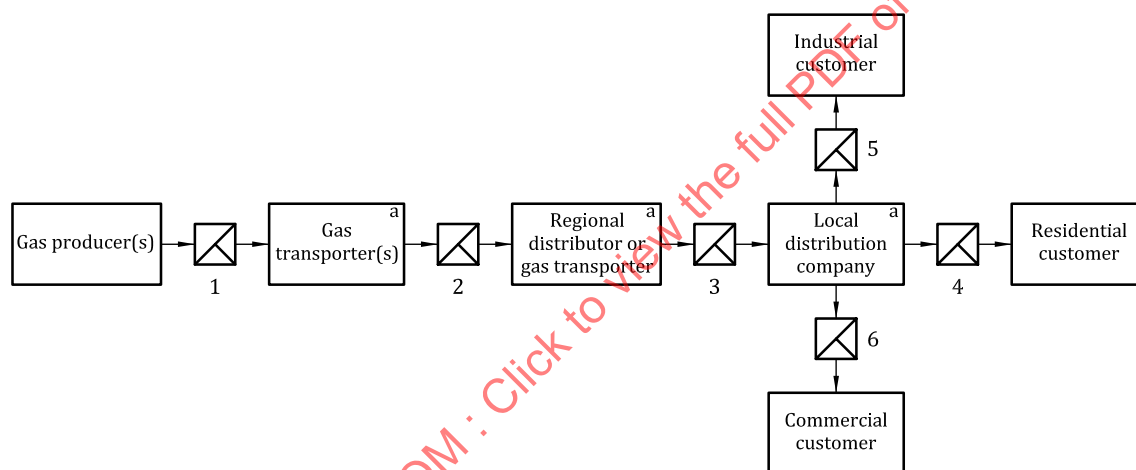
For data transfer, safe procedures shall be taken to ensure the integrity of the data.

7 Energy determination

7.1 Interfaces

Natural gas custody transfer between contract parties is, in general, performed from the producer(s) or gas storages to the end user via intermediate stages involving some or all of the following:

- gas transporter(s);
- regional distributors;
- local distribution company(ies).

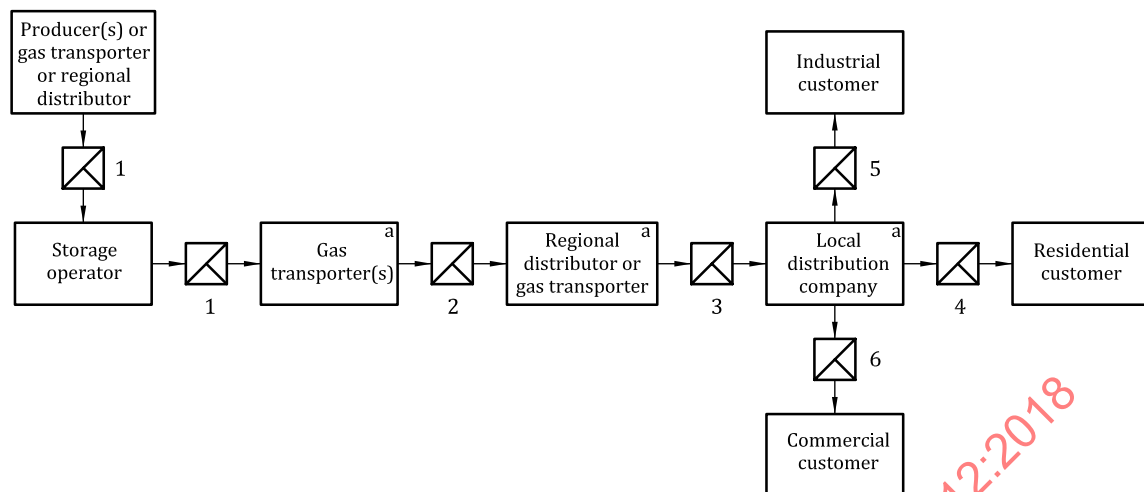


Key

1 to 6 interface

^a If this entity exists.

Figure 3 — Possible interfaces for energy determination from producer(s) to end users



Key

1 to 6 interface

a If this entity exists.

Figure 4 — Possible interfaces for energy determination from producer(s) to end users including gas storages

The boxes numbered 1 to 6 in Figures 3 and 4 represent the different interfaces within a delivery chain; they may consist physically of a real measuring station or may be regarded only as virtual interfaces without any measurement to define the point of delivery or redelivery within contractual situations. Energy determination in a delivery chain between contract parties is performed at interfaces 1 to 6 (see Figures 3 and 4), often also named points of delivery and/or points of redelivery. Figure 3 shows the delivery chain from the producer to the end user; Figure 4 includes, additionally, a storage operator, who usually stores gas for producers, gas transporters or regional distributors for future take-off. The kinds of interfaces may differ within the different countries. They may be used as gas-billing interfaces, if they are actual measuring stations.

Three different models of different delivery situations are given as examples.

- a) The gas transporter supplies gas directly to an industrial customer.

For energy determination at interface 5, the gas volume is measured at interface 2 or 5; because there is no regional distributor/storage company or local distribution company (LDC) involved, the calorific value measured at interface 2 can be used if nearly constant gas quality (see Figure B.1) can be expected.

- b) The gas producer supplies gas directly to an industrial customer.

The pipeline system is used by several gas transporters and regional distributors for transportation; LDCs are not involved. On its way to the industrial customer, no gas quality changes occur. For energy determination at interface 5, the gas volume is measured at interface 5 and the calorific value at, for example, interfaces 5, 3 or 2.

- c) The LDC supplies gas to the end user, commercial and industrial customer.

The LDC is supplied by a regional distributor, or gas transporter or storage company. For energy determination, volume measurement is performed at interfaces 4 to 6. Due to different gas qualities (see Figure B.3), the regional distributor operates a state reconstruction programme for calorific-value determination at interface 3; that calorific value is taken by the LDC for energy determination at interfaces 4 to 6.

The method of energy determination depends on a number of important factors; they shall be taken into account for the suitable energy-determination strategy to support the user of this document to perform a correct energy determination. They include

- grid topology,
- flow directions,
- take-off structure or consumption profile,
- course of calorific value,
- technical equipment,
- contractual requirements, and
- national regulations.

It is the main goal of the methods given in [7.2](#)

- a) to support a satisfactory energy balancing within the transportation grid, and
 - b) to provide a justified energy determination at interfaces,
- taking into account economic aspects.

7.2 Methods of energy determination

7.2.1 Direct determination of energy

For direct measurement (see [Figure 5](#)), individual physical parameters (e.g. Q , H) are not measured. The energy flow and energy quantity are calibrated and shown at the measuring point. At the time of preparation of this document, direct energy-measurement instruments have entered the marketplace, but they are not yet proven technology for custody transfer. No International Standards exist at the moment.

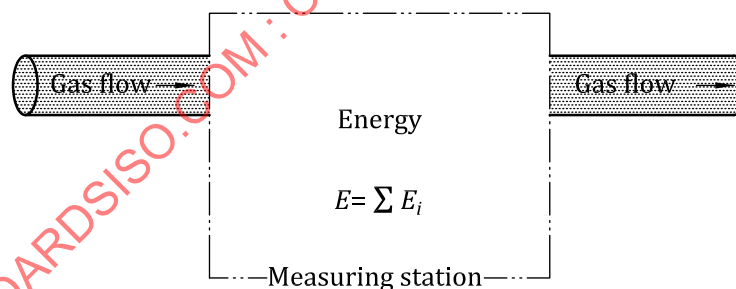


Figure 5 — Direct determination of energy

7.2.2 Indirect determination of energy

For indirect energy determination, the energy is determined on the basis of previously measured or calculated values for volume/mass, calorific value and other entities.

7.2.2.1 Measurement of volume or mass and calorific value at the same station

For indirect determination of energy, the volume or mass, calorific value and additional physical entities, such as CO_2 , density, etc., of the gas are measured separately in a measuring station (see [Figure 6](#)); the measurement devices are individually calibrated. The volume flow rate and energy quantity are typically displayed at the measuring point. For large gas quantities Q_1 and Q_2 , for example at border crossings, it can be necessary to determine the calorific values H_{S1} and H_{S2} by means of two calorific-value measurement devices at each station (see [Figure 15](#)).

Another method is to collect the calorific value and volume data in the measuring station and to transmit the data to a different central energy-determination station where energy flow and energy quantity are determined.

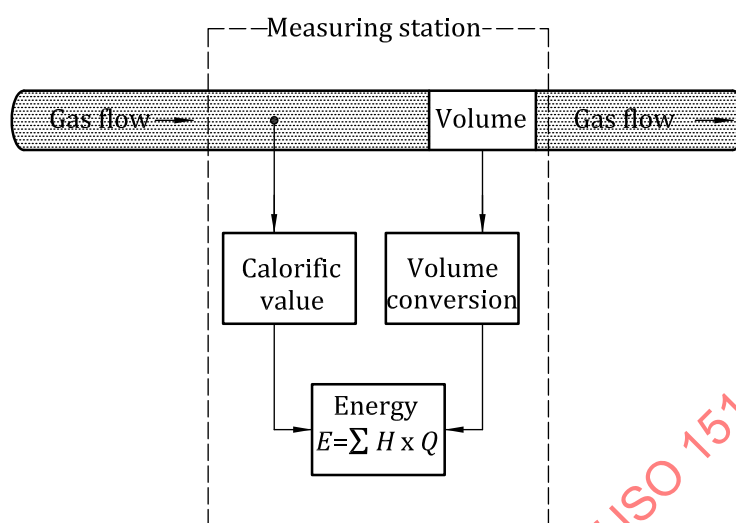


Figure 6 — Local online calorific value measurement

Due to the local gas-quality situation and for economic reasons, it is sometimes of use to take samples of gas (time- or flow-controlled) within the measuring station and to determine the calorific value at a different place (see Figure 7).

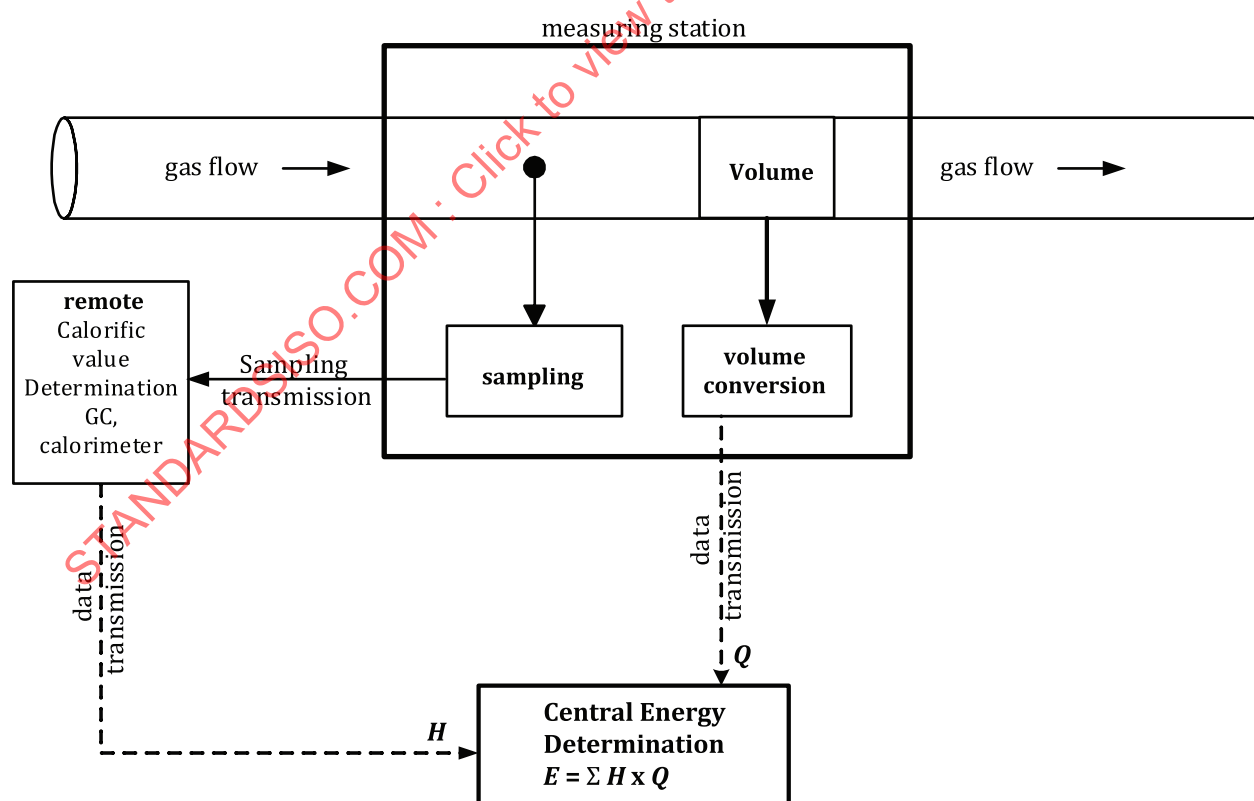


Figure 7 — Local offline calorific value determination

7.2.2.2 Measurement of volume or mass and calorific value at different stations

Whereas the gas volume is measured at every delivery point between contract parties, it can be too expensive to operate a calorific-value measurement device there, too. Thus, the most common method (especially in extensive supply systems) is to assign a representative calorific value (see [Clause 9](#)) to the volume. The calorific values assigned to those interfaces (volume-measuring points) are values measured elsewhere or a value formed from several representative measured values (see [Figure 8](#)). These values are the basis for energy determination. The kind of assignment is determined by the location of the input/output stations in the grid and the conditions of gas flow (see [Clause 9](#)).

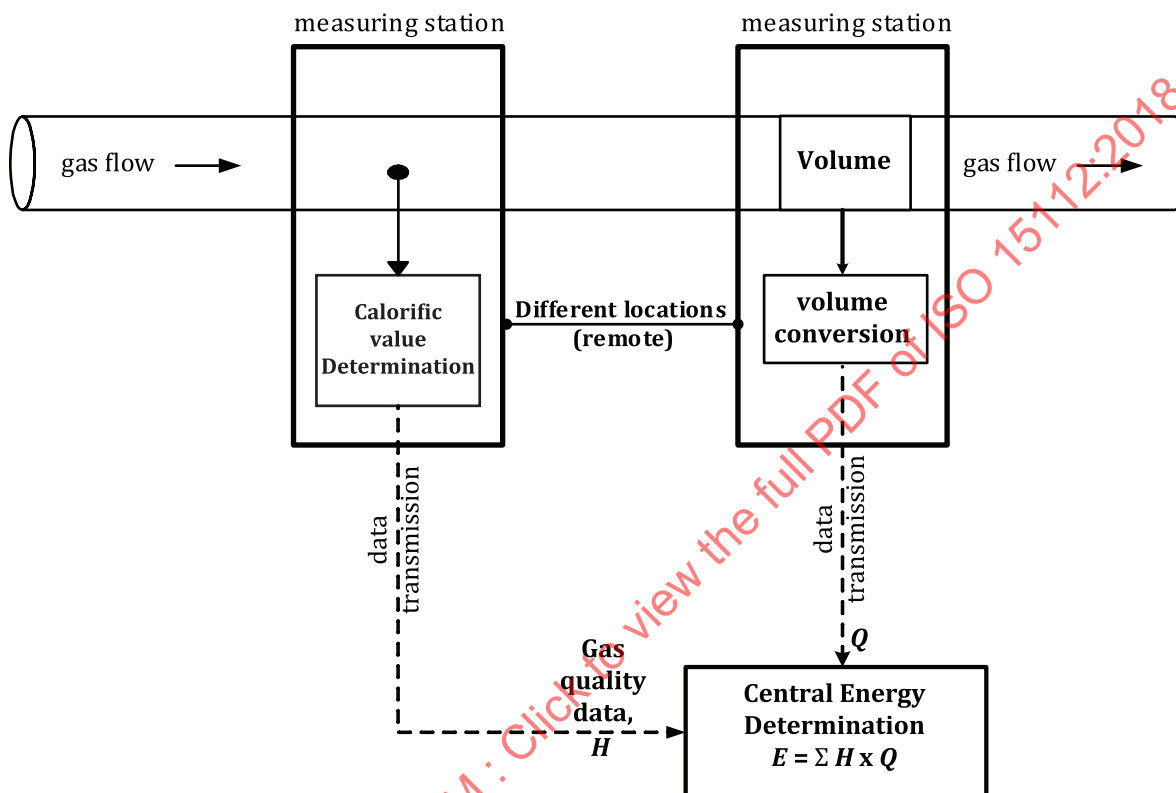


Figure 8 — Remote calorific value determination (example)

8 Strategy and procedures

8.1 General

“Design” in the context of this document encompasses the requirements of what information is necessary and how it should be obtained to fulfil the needs of the energy-determination strategy, taking into account the expected course of data.

Energy determination starts with an assessment of a reasonable energy-determination strategy, followed by a plausibility check of the measured data. The next steps are the assignment of the representative calorific value and the combination of the data (calculation procedures). Finally, a quality-control procedure is performed.

An energy-determination scheme, including “start” and “end” points, is shown in [Figure 9](#).

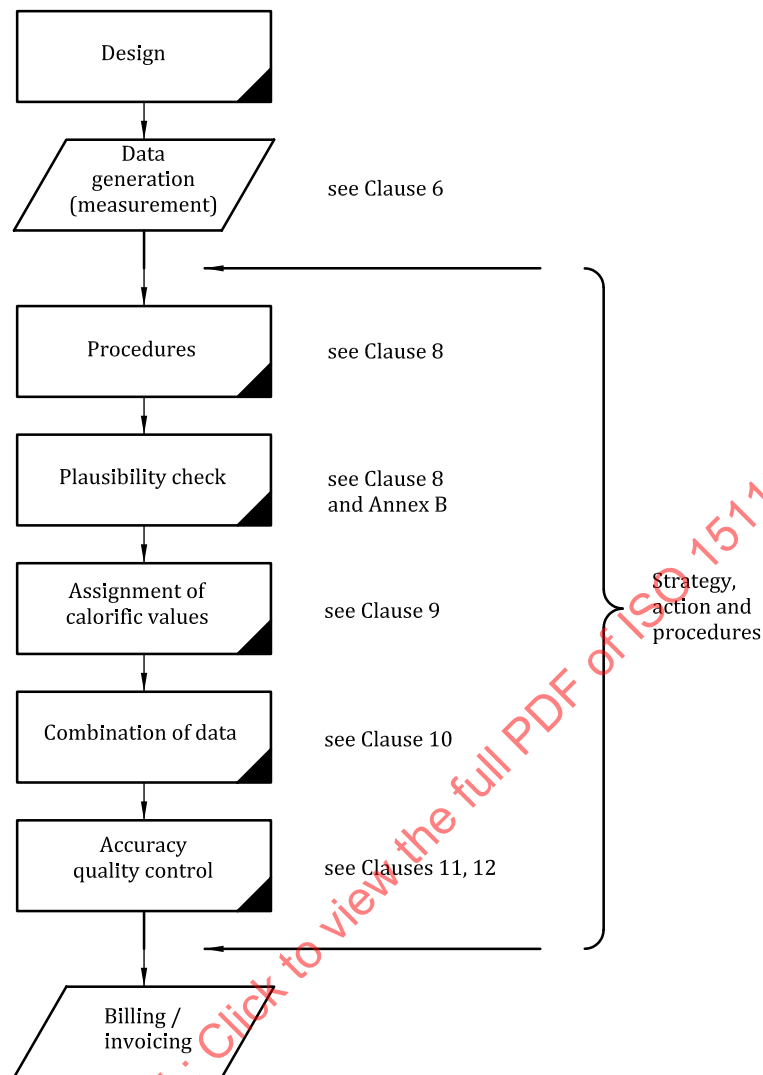


Figure 9 — Strategy for an indirect energy determination

Principally, elaborated strategy and the applicable methods and procedures for energy determination shall be applied without changes. They may be changed only if

- it can be assured that the accuracy of the results will be better or as a minimum not worse, or
- the methods or procedures are no longer applicable, due to the changed gas flow and/or gas quality conditions.

Additionally, due to significant changes with respect to the economics, it may be necessary to change the applicable methods/procedures.

Gas measurement systems are not the subject of this document. However, data generated by new systems or modified devices can cause a better or worse quality of the data; this can lead to modifications of energy determination.

8.2 Strategies for energy determination

For an energy-determination period, it is necessary that a reasonable energy-determination strategy take account of

- the course of the calorific value (calorific value changes) with respect to the relevant supply situation, and

— the correctness of the measuring data (i.e. the raw data)

for a specified interface.

Taking into account the changes in gas quality at an interface is a key factor for the justification and accuracy of the energy determination, i.e. the traceable determination of the calorific value for the relevant interface.

This is important because even a non-corresponding energy-determination strategy at the interfaces can be disadvantageous for one partner and shall be avoided. This can happen, for example, by using a non-matching, i.e. non-representative, calorific value or other non-representative physical data (i.e. density, CO₂) at those interfaces where only the gas volume is measured, or by taking raw data without any critical check.

Generally, changes in gas quality depend on the gas supply situation at that interface, i.e. it is necessary to prove

- a) whether gas from only one origin has passed the subsequent interfaces (see [Figures 12, 13 and 15](#)), or
- b) whether gases from more than one origin have passed the subsequent interfaces (see [Figures 14 and 15](#)).

Supply situations according to [Figures 14 and 15](#) very often result in reasonable changes in gas quality due to the delivery and take-off situations.

Thus, the changes of the daily or monthly averaged calorific values over one billing period (one month, respectively, one year) shall be carefully monitored to decide whether an energy determination period for billing purposes shall be divided into smaller periods.

Changes in gas quality within one day do not usually result in a further subdivision into hourly values. For examples, see [Annex B](#).

8.2.1 Strategies for single interfaces

8.2.1.1 General

In general, large quantities of gas passing interfaces 1 to 3 (see [Figures 3 and 4](#)) are subject to measuring and gas charging. For interfaces 1 to 3, the calorific value for gas charging is determined by local measurement (see [Figures 6 and 7](#)), remote calorific value measurement (see [Figure 8](#)) or appropriate assignment methods or quality tracking (see [9.3.3](#)). It is up to the contract partners and local authorities to agree upon the matching method.

At those points on the gas grid where a direct measurement of the calorific value and other important gas quality entities for gas charging cannot be performed due to technical and/or economic reasons, a determination of these entities as the calorific value, gas density etc. by an indirect way can sometimes be performed (see [9.2, 9.3.1, 9.3.2 or 9.3.3](#)).

8.2.1.2 Interface 1

For the calculation of the volume under reference conditions, a volume conversion is either performed by

- p , T and Z , or
- ρ_n and ρ .

At that interface, the calorific value shall be measured online (see [6.3](#)).

In the case of gas production from a single reservoir where the gas composition may be expected not to change over time, online measurement for the determination of the calorific value may not be necessary. In this case, the calorific value may also be calculated from the gas composition obtained from offline

analysis of periodic samples. For an example, see [Annex J](#). In all other cases, calorific measurement according to the rules that apply for interface 1 shall be followed.

The development over time of separate calorific values relative to the initial calorific value is assessed statistically. If the values exceed an agreed limit (for example 0,5 %, see [Figure J.1](#)) over time, the method shall be changed from offline measurement to online measurement.

The application of this method for calorific-value determination should take into account the fact that the delivery composition of the gas is strongly dependent on the type of gas treatment installation.

For the determination of the compression factor, Z , see 6.4.4.

8.2.1.3 Interface 2

The calculation of the volume under reference conditions is performed as described in [8.2.1.2](#).

The determination of the calorific value for energy-determination purposes is performed by measurement (see [6.3](#)) or assignment (see [Clause 9](#)).

8.2.1.4 Interface 3

The determination of the values for energy determination is performed as described in [8.2.1.3](#). The user of this document should take into account that methods and/or facilities at interface 3 to determine the calorific value, especially for the subsequent interfaces, can differ from country to country.

8.2.1.5 Interface 4

At interface 4 of the gas grid, a measurement of the calorific value and other important gas quality entities can usually not be performed for technical and/or economical reasons. For this interface, assignment methods (see [Clause 9](#)) are necessary.

Before the gas passes the measuring device, upstream a stable pressure of the gas shall be ensured by means of pressure control. The local distribution company (LDC) determines procedures to set the relevant temperature and pressure that can be used for energy-determination purposes, taking into account the ambient pressure. Due to the low pressure, the compression factor is not calculated and set to "1".

For this interface, either a declared or assigned calorific value can be applied, using the upstream calorific value of interface 3.

If only reasonably small changes in gas quality can be expected (see [Figure B.1](#)), it is advisable to use only a declared calorific value for annual energy-determination periods. A declared calorific value is set by the LDC at a fixed value, taking into account the calorific values of the previous 12 months (see [Figure B.1](#)). During the energy-determination period, the LDC regularly checks the calorific values at (upstream) interface 3 from where the gas is feeding interfaces 4.

If the difference between the declared calorific value and the calorific value determined upstream at interface 3 is larger than the permitted difference (for example 1 %), for example when the calorific value has changed significantly (see [Figures B.2](#) and [B.3](#)), these calorific values shall be assigned to the energy-determination periods. For example, in [Figure B.2](#), the calorific value H_{S1} shall be assigned for energy-determination purposes for the time periods t_1 and t_2 , and the calorific value H_{S2} for the time period t_3 .

To ensure a reasonably accurate energy-determination process for those customers, a distinction shall be made between

- a gas grid separated from other gas qualities, and
- a gas grid (open grid) not separated from other gas qualities.

8.2.1.5.1 Gas grid separated from other gas qualities

If gases with different calorific values are kept separately in different gas grids, no mixing of these gases can occur and the calorific value measured or determined at interface 3 may be taken as the basis for energy determination at interfaces 4, as follows.

The average of the calorific value can be calculated either arithmetically or on an hourly volume- or quantity-weighted basis in the manner described in [10.2](#).

- a) At first, at the end of every day or another interval within the energy determination period, the energy quantity of the gas passing interface 3 is calculated by averaging the measured/determined individual calorific values [see [Formula \(6\)](#)] and multiplying that averaged calorific value by the gas volume or quantity that had passed interface 3 during the same interval [see [Formula \(7\)](#)] in the following manner.
 - For this purpose, the individual calorific values within 1 h are averaged using [Formula \(6\)](#). (“hourly-based averaged calorific values”).
 - These hourly-based averaged calorific values are used to calculate a daily averaged calorific value using [Formula \(6\)](#) or, for weighted averaging, [Formula \(8\)](#).
 - Finally, that daily averaged calorific value is multiplied by the gas volume or quantity that had passed interface 3 during the same day.
- b) Secondly, at the end of the energy-determination period, the energy quantities of all intervals will be summed up and divided by the sum of all gas volumes/quantities of all intervals of that energy-determination period [see [Formula \(8\)](#)].

The resulting averaged calorific value can be applied for the calculation of the energy of any interface 4 on the grid. For further details, see [9.1](#). Practical examples are given in [Annex F](#).

8.2.1.5.2 Gas grid not separated from other gas qualities

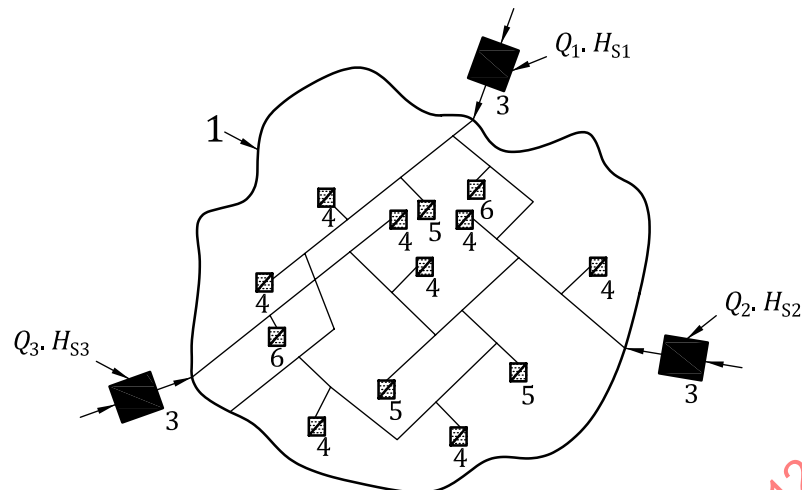
If end users at interface 4 are supplied via several interfaces 3, the calorific values measured/determined at all such interfaces 3 are relevant.

If the calorific values change over time at these interfaces, it is usually impossible to perform a calorific value measurement at each interface 4 due to economic reasons, even if, according to local conditions (i.e. complicated topology, low pressure), the supply situation is unclear. Mostly due to relatively small gas quantities at that interface, technical measures, such as sampling, calorific-value measurement, state reconstruction, etc., cannot usually be justified as a means of increasing the accuracy.

If different gas qualities are fed into a gas grid at several interfaces 3, the calorific values for the interfaces 4 are calculated by arithmetically averaged calorific values (see [9.3.1](#)) or by the following procedure.

- a) At first, the energy quantity for each interface 3 is calculated for the energy-determination period as described in [8.2.1.5.1](#).
- b) The energy quantities of all interfaces 3 are then summed up and divided by the sum of all gas volumes or quantities of all interfaces 3 of that period to give the weighted-average grid calorific value for that period.
- c) If none of the weighted-average calorific values of any interface 3 deviates by more than a permitted value from the weighted-average grid calorific value, then the weighted-average grid calorific value may be applied for the calculation of the energy of every interface 4.

A practical example is given in [Figure 10](#).

**Key**

1 charging area

Figure 10 — Determination of a weighted-average calorific value (example)

EXAMPLE A charging area, where the energy determination is being performed, is supplied at the entry-point interfaces 3 by the gas quantities Q_1 , Q_2 and Q_3 with the corresponding calorific values H_{S1} , H_{S2} and H_{S3} . These calorific values have been either measured at the entry points or determined upstream at interfaces 1 or 2 and assigned to these interfaces 3 as described in [Clause 9](#). Then the averaged calorific value H for energy-determination purposes in the charging area is calculated as described at the above-mentioned points [8.2.1.5.1 a\)](#) and [b\)](#).

For interfaces 5 and 6, this averaged calorific value can also be taken for energy-determination purposes or measured individually at these interfaces. In the latter case, the energy-determination procedure for all other interfaces in the charging area shall take account of the energy quantities determined individually via local calorific value measurement at interfaces 5 and/or 6.

If the deviation exceeds the permitted limit, the national authorities shall be informed about the measure of deviation and the procedure applied. The information is not necessary if, in extraordinarily rare cases, short-term (maximum one week) deviations occur, for example caused by measures to ensure the supply of gas.

If a reasonably accurate energy-determination process can be assured (see [8.2.1.5](#)), arithmetic averaging of the calorific values may be considered. If the difference of the weighted averaged calorific value at interface 3 and the weighted averaged grid calorific value in the charging area is higher or lower than permitted due to strongly changing gas qualities at interfaces 3, the following additional measures inside the charging area can be taken. These measures are necessary to determine the representative calorific value for single parts of the pipeline system as representative as possible, for example, by the use of sampling techniques that take into account the quantities Q_1 , Q_2 and Q_3 and the measured or calculated take-off structures of subsequent interfaces 4 to 6. In these cases, different average calorific values can be calculated for the various parts of the energy-determination system using the calorific values at the entry-point interfaces and the calorific values determined by sampling methods. For further details, see [9.2](#).

8.2.1.6 Interface 5

For industrial customers, i.e. interface 5, the relevant calorific value is determined either upstream by the gas transporter or by the local distribution company. Due to the large quantities and for economical reasons, the relevant calorific value for charging purposes is very often also determined at this interface. According to the gas quantities and pressures, energy determination at interface 5 is performed as it is at interfaces 1 to 4 (see [8.2.1.5.2](#)).

8.2.1.7 Interface 6

The requirements for this interface are similar to those for interface 4. For details, see [8.2.1.5.2](#).

8.3 Plausibility checks

The first step for energy determination is the critical check of the plausibility of the measured, transmitted or recorded data. Non-plausible data can be caused for example by

- malfunction of a measuring device,
- external impacts, such as electromagnetic fields on communication lines,
- breakdown of recording devices, etc.

Other reasons for non-plausible data shall be carefully checked.

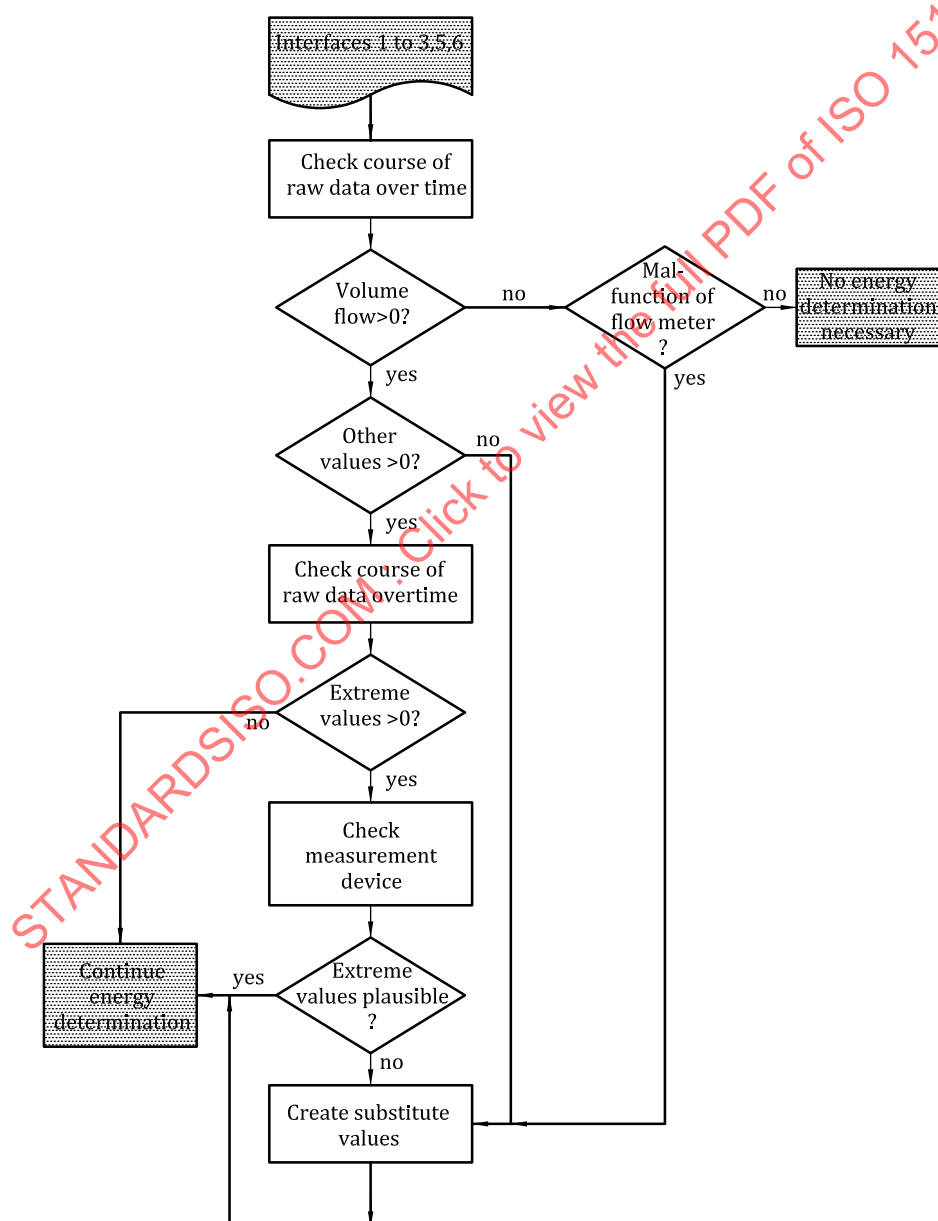


Figure 11 — Data plausibility check

The functional diagram in [Figure 11](#) shows how to perform a plausibility check in a formal way. If the volume flow equals “0”, it should be checked whether or not it is true. If there was a flow and the data are “0”, a problem has obviously occurred, such as a malfunction of the meter or associated devices such as signal transmitters, electronic devices, data storage devices or others. Obviously false data shall not be used. For false or missing data, suitable substitute values (see [12.4](#)) shall be determined.

The box “other values” means entities as pressure, p , temperature, T , density or others. “Extreme values” means values shown to be at the end of the scale.

A practical example is given in [Annex H](#).

9 Assignment methods

9.1 Fixed assignment

A fixed assignment of a calorific value within a charging area for energy-determination periods can generally be performed in simple, separated grids if the following conditions are met.

- The direction of the gas flow between the calorific value and the volume-measuring points is constant.
- The variation of the gas quality and the transit time of the gas between calorific value and quantity-measuring points are reasonably small (see [10.4](#) and [Figure B.1](#)) in that energy-determination period. [Clause 11](#) can be used to check if the required accuracy of the energy determination can be met.

Generally, the methods described in [9.1.1](#) and [9.1.2](#) are possible.

9.1.1 Fixed assignment of a measured calorific value

The calorific value is measured at a calorific-value-measuring station. The data show that the variation of the gas quality is very small (see [Figure B.1](#)). Thus, it is justified to assign the mean of the previously measured calorific value as a fixed value to all subsequent interfaces. The assignment from a single calorific-value-determination station can be demonstrated as follows.

[Figure 12](#) illustrates a single source of gas, whose calorific value, H_s , is determined at the entry point to the pipeline that is operated by a gas transporter/regional distributor and that feeds a number of interfaces at the pipeline. The calorific value assigned to all the interfaces is that determined at the entry point and is not modified to reflect the different times of transit of the gas to different interfaces.

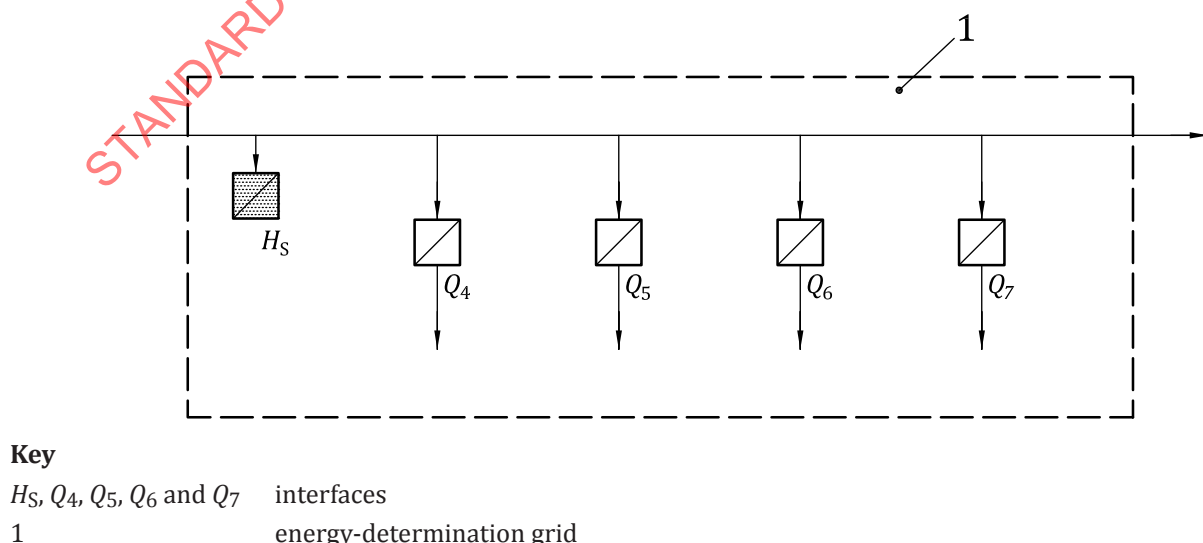
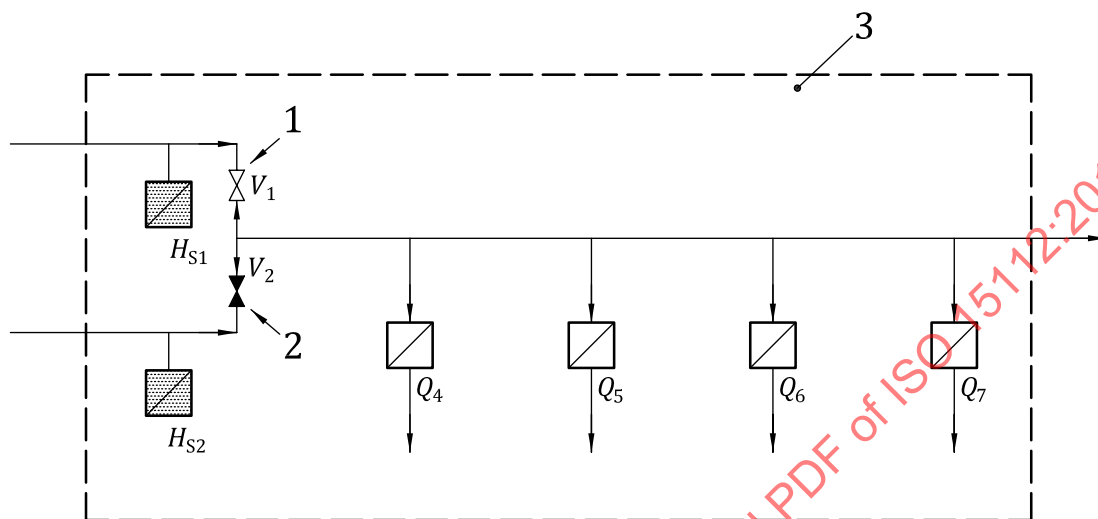


Figure 12 — Example of a fixed assignment for one gas quality — Unidirectional gas flow

The assignment from two or more calorific-value-determination stations can be demonstrated as follows.

Figure 13 illustrates a system in which a gas transporter has the possibility of two different gases entering a pipeline. The two calorific values, H_{S1} and H_{S2} , one for each of the two gases, are determined at a point upstream of the entry (entry point) to the pipeline and there is no calorific-value measurement downstream of the entry point.



Key

H_{S1} , H_{S2} , Q_4 , Q_5 , Q_6 and Q_7 interfaces

1 valve 1

2 valve 2

3 energy-determination grid

Figure 13 — Example of a fixed assignment for two measured gas qualities — Unidirectional gas flow

The gas transporter decides to use a fixed assignment from one or the other of the calorific-value-determination points on the basis that

- a constant supply can be assured at all times from one source,
- the gas supply to the pipeline with the different gas qualities H_{S1} and H_{S2} , resulting in a gas mixture with a calorific value different from H_{S1} and H_{S2} , never happens (both valves are never open at the same time),
- the supply periods with each of the different gas qualities are recorded, and
- for assignment purposes, the calorific value, either H_{S1} or H_{S2} , is used for subsequent interfaces corresponding to the same supply period.

9.1.2 Fixed assignment of a declared calorific value

The calorific value is assumed to be reasonably constant over time within an energy-determination period. The calorific value is measured for checking purposes at a calorific-value-measuring station.

The data acquired confirm that the variation of the gas quality is very small (see [Figure B.1](#)). Thus, it is justified to declare a calorific value and assign that value to all subsequent interfaces.

EXAMPLE A local distribution company (LDC) has a gas grid supplying various customers, as domestic, commercial and small industrial consumers. There are a couple of interface entry points to a gas grid that are supplied with gas from a single pipeline. The calorific value of the gas flowing through the pipeline shows only small variations, except at times of peak demand during the winter when the calorific value can rise by up to 1 % in relation to the average calorific value.

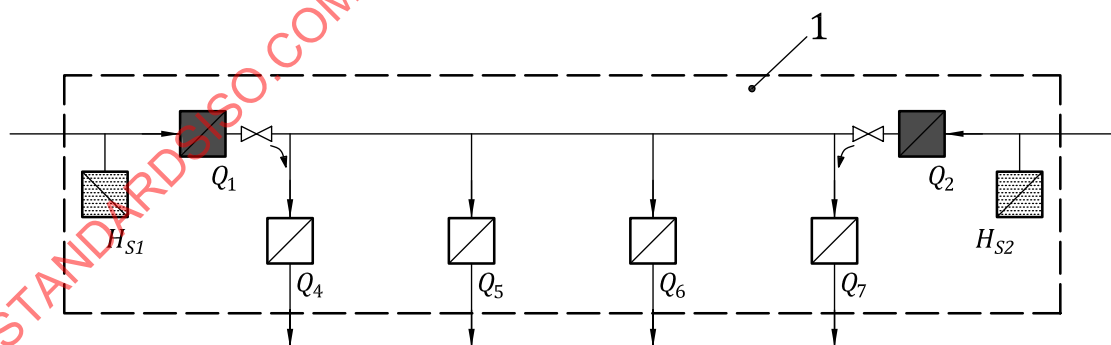
The LDC decides to use a fixed assignment of a declared calorific value for all interfaces on its gas grid; the calorific value is declared for a period of time on the basis that

- the determined calorific value of the gas supplied to consumers is, on average, equal to, or higher than, the declared calorific value (to the nearest 0,1 MJ/m³),
- the average calorific value of the gas supplied to consumers is calculated by averaging, for each day in the declared period, the lowest calorific value of gas,
- the calorific values of all gases entering the grid are determined on a daily basis, and
- if, for any period, the determined calorific value falls below the declared figure, then the LDC will revise the declared value for the following period such that, over the two periods, the determined value is equal to, or higher than, the average declared value.

9.2 Variable assignment

Especially in open gas grids, gas qualities at the interfaces can vary significantly with time (see [Figure B.3](#)). In this case, the requirements for a fixed assignment are no longer applicable and it is necessary to adapt the assignment/calculation method that is suitable for conditions that change with time. The choice of a suitable assignment method changes according to changes in the gas quantities at the input stations, as well as changes in the take-off structures at subsequent interfaces. Thus, careful procedures for variable assignment of the calorific values shall be undertaken. There are two different situations, which shall be generally distinguished; they are described in [9.2.1](#) and [9.2.2](#).

9.2.1 Input at two or more different stations with zero floating point



Key

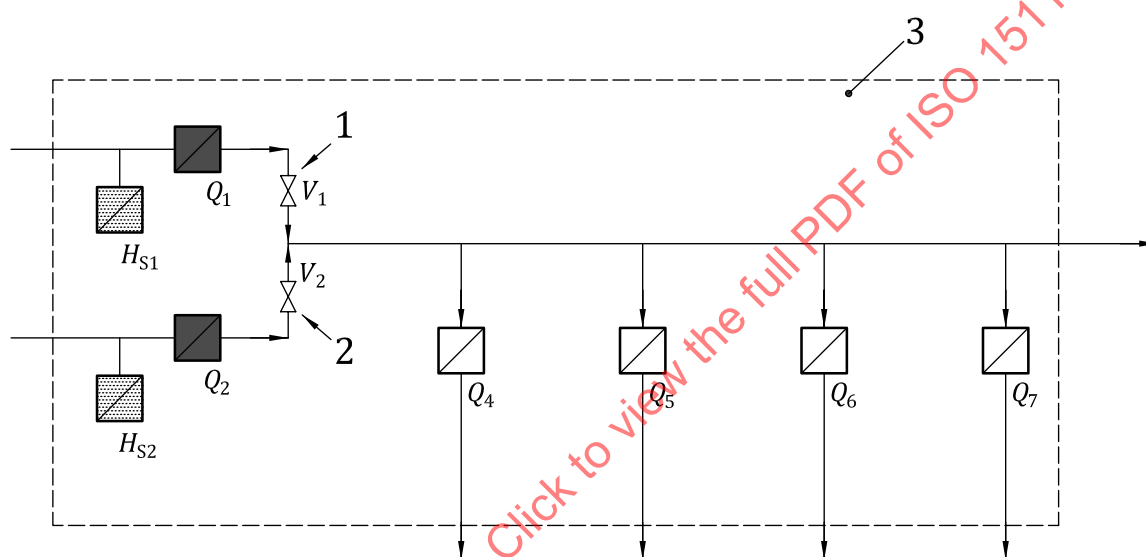
H_{S1} , H_{S2} , Q_1 , Q_2 , Q_4 , Q_5 , Q_6 and Q_7 interfaces
1 energy-determination grid

Figure 14 — Variable assignment — Example for two measured gas qualities and bidirectional gas flows

EXAMPLE For an energy-determination period (see Figure B.3), as shown in Figure 14, different gas quantities and qualities pass the interfaces 1 and 2 (input stations). The defined zero floating point can be located between two interfaces (for example, between two neighbouring interfaces or between an input station and a neighbouring interface). According to the take-off structures at the interfaces 4 to 7, gas with the calorific value H_{S1} can supply interfaces 4 and 5, whereas gas with the calorific value H_{S2} can supply interface 7. A mixture of gas from interfaces 1 and 2 can pass interface 6. Thus, the calorific value H_{S1} can be assigned to interfaces 4 and 5 and the calorific value H_{S2} can be assigned to interface 7. For interface 6, the representative calorific value can be either measured at that interface or determined taking into account the partial gas quantities from Q_1 at interface 1 and Q_2 at interface 2 and the applicable calorific values H_{S1} and H_{S2} using flow or arithmetic weighted averaging (see 10.2.2). The zero flow in the main pipeline can be situated at or between the interfaces 4 to 7.

For the period during which the defined zero floating point has a fixed position within the grid, calorific values can be assigned to each interface on the basis of the gas flows from the input stations to the respective interfaces.

9.2.2 Input at two or more different stations with comingled gas flows



Key

- H_{S1} , H_{S2} , Q_1 , Q_2 , Q_4 , Q_5 , Q_6 and Q_7 interfaces
 1 valve 1
 2 valve 2
 3 energy-determination grid

Figure 15 — Variable assignment — Example for two measured gas qualities and unidirectional gas flow

EXAMPLE During an energy-determination period, the quantity Q_1 at interface 1 with the calorific value H_{S1} and the quantity Q_2 at interface 2 with the calorific value H_{S2} are measured. The two calorific values are always different from each other and can also change during the energy-determination period. Due to this situation, for the calorific value assignment to interfaces 4 to 7, a pattern of the resulting calorific value for each interface 4 to 7 similar to that in Figure B.3, for example, can result at interface 4.

If the calorific-value-measuring stations H_{S1} and H_{S2} are far away from the interfaces Q_4 to Q_7 , the run time of the gas from these measuring stations H_{S1} and H_{S2} to the interfaces Q_4 to Q_7 shall additionally be taken into account; this may be hours or days. A calculation is performed on the basis of the quantities Q_4 to Q_7 , the respective pressures and the pipeline sectional area including the line pack.

A quantity-weighted averaged calorific value shall be calculated at the mixing points behind valve 1 and valve 2 for the gas quantities Q_4 to Q_7 supplied to the respective interfaces 4 to 7 for the energy-determination period, taking into account the run times from the calorific-value-measuring stations for H_{S1} and H_{S2} to the mixing point.

9.3 Determination of the representative calorific value

The accuracy of the determined representative calorific value depends on the completeness and accuracy of the data and the topology of the gas grid. For determining the representative calorific value at the mixing point, the calorific value is calculated using the gas quantities and qualities. To get the representative calorific value, the run time from the input stations to the mixing point, as well as the run time to the subsequent interfaces, may be taken into account. For the determination of the representative calorific value for the energy-determination period at each interface, in all cases except fixed assignment, the methods given in [9.3.1](#), [9.3.2](#) and [9.3.3](#) may be used.

9.3.1 Arithmetically averaged calorific value

For an entry-point interface, an arithmetically averaged calorific value is calculated for the energy-determination period by dividing the sum of periodic single calorific-value measurements at the entry point by the number of measurements of the calorific value (see [10.2.1](#)).

9.3.2 Quantity-weighted average calorific value

For an entry-point interface, a quantity-weighted average calorific value is calculated at this interface for the subsequent interfaces (see [10.2.2](#)) for the energy-determination period.

9.3.3 Gas quality tracking

If gases with different calorific values are fed to a grid, mixing and transition zones will be formed which are characterized by frequent changes in the gas quality during gas transport and gas distribution. If gas is billed on the basis of an average calorific value, this may lead to not acceptable uncertainties. According to [8.2.1.1](#), second paragraph, it may not be justified due to economic reasons to measure the calorific value at all interfaces (exit points) of the gas grid using calibrated instruments or to adjust the calorific value by conditioning but to apply gas quality tracking.

In this context, this chapter describes the procedure for gas quality tracking to determine calorific values at all interfaces (exit points), which can then be used for charging end customers.

Before its application, a gas quality tracking system is subject to a validation process. This includes providing evidence that the system complies with the tolerance limits for calorific value.

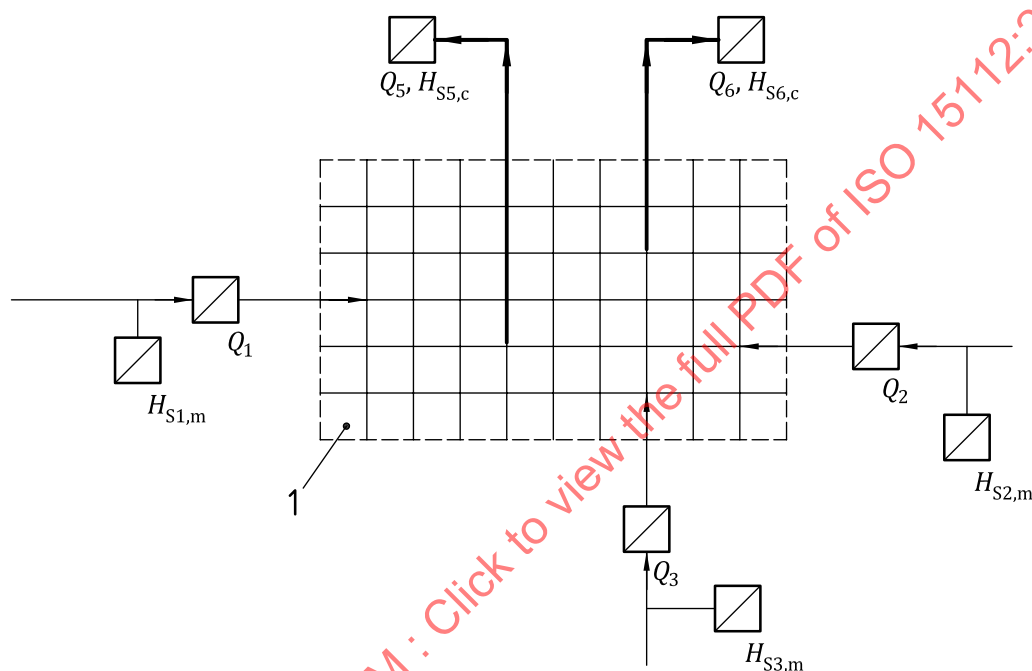
9.3.3.1 Description

The application of a gas quality tracking system requires detailed information on the grid topology (including pipeline length, pipe diameter and pipe roughness), calorific values or further gas quality properties at the interfaces (entry points) and gas quantities at all entry and exit points, for example according to [Figure 14](#) and [Figure 15](#). Further data such as measured grid pressures may also be used in the calculation to enforce the determination of the volume flow through the grid. Calorific values at the entry points may be determined either by calibrated measurement instruments (such as process gas chromatographs) or by an upstream gas quality tracking system. Measured calorific values should be provided as hourly averaged values. Calorific values determined by an upstream gas quality tracking system may be provided as hourly averaged or daily averaged values. Volume data should be provided on an hourly basis. Gas quantities shall be measured by volume meters at the interfaces (entry points). At the exit points, gas quantities may also be determined by so called standard load profiles, see example in [K.2](#). This applies especially to regional gas distribution grids where volume measurement devices are frequently not installed at the corresponding interfaces (e.g. at the exit points to downstream gas grids).

On the basis of these input data, flow conditions throughout the grid can be determined using a flow calculation. This includes the determination of flow speeds and the temporal and geographical distribution of the gas quality within the entire gas grid. On this basis, an individual calculated calorific value may be assigned to each interface (exit point). With respect to the smallest time unit of the input data (hourly based values), the calculation should be performed at least at hourly intervals. The representative gas quality properties for each exit point — e.g. for the monthly calorific value — shall be determined as the volume-weighted average on the basis of hourly values.

If downstream gas distribution grids — especially with respect to interfaces 2 and 3 — are connected to the grid where gas quality tracking is applied, a review shall be carried out to determine to what extent the interfaces (entry points) are hydraulically connected (meshed). In the event of meshing of this type, the calorific value differences of the gas shall be reviewed to determine whether an averaged calorific value can be applied to the subsequent interface 2 or 3 (downstream gas grid).

The time unit required for the gas quality properties determined (e.g. the calorific value) depends on the defined charging period. Normally, the gas quality properties are determined as volume-weighted monthly averages. In certain cases, it may also be necessary to state the results as daily averaged or hourly averaged values, for example for customers with time-based gas charging tariffs or if the gas quality properties are to be used as input variables for a downstream gas quality tracking system. In this context, it should be noted that the verification of compliance with tolerances as described below refers to the underlying time unit in each case.



Key

Q_1, Q_2, Q_3, Q_5 and Q_6 interfaces
1 grid model

Figure 16 — Example of a reconstruction-based gas-quality tracking scheme

9.3.3.2 Validation

The purpose of validation is to verify that the method used complies with the tolerance limits for calorific value and, if applicable, for other gas quality properties.

It should be noted that the accuracy of a gas quality tracking system strongly depends on the quality of the input properties. In particular gas volume data have a significant influence on the calculated result. In general, a gas quality tracking system applied to grids where all interfaces (entry and exit points) are measured with calibrated meters (as it is usually the case in transmission grids) is expected to reach a higher accuracy as if exit gas volumes at exit points were determined e.g. to a large extent based on standard load profiles (as it may be the case in distribution grids). Therefore it is essential to apply the validation procedure as described below to evaluate the uncertainty and correctness of the system.

9.3.3.3 Master validation for commissioning

Before a gas quality tracking system is used for gas charging, a fundamental validation of the system shall be carried out. Validation may be performed as uncertainty calculation in combination with comparison measurements with suitable gas quality measuring instruments. A possible procedure to determine the uncertainty is described in informative [K.4](#).

Measurement uncertainty may be verified by comparison measurements using suitable gas quality measurement instruments (e.g. process gas chromatographs). Both remote mobile and locally fixed measurement instruments may be used. As an alternative, gas sampling techniques (see [6.3.1](#)) may be used. The suitable locations for measurement shall be selected in accordance with the expected flow situation. Measurements shall be performed preferably at exit points where gas quality fluctuations and mixtures of different gas qualities are to be expected. These are those exit points where higher uncertainties may be expected.

Measurement uncertainty shall be deemed to have been verified if deviations in the gas quality properties over the measurement period are within the tolerances.

If an uncertainty calculation is not possible, more complex validation procedures using measurements are required.

9.3.3.4 Regular validation during operation

Measurement uncertainty shall also be verified at regular intervals during operation.

The uncertainty calculation described in [K.4](#) should be repeated as often as necessary, especially in the event of changes in the grid topology or the mode of operation of the grid. Regular monthly uncertainty calculations are recommended provided that the effort is reasonable.

Comparison measurements with suitable gas quality measuring instruments shall also be repeated at regular intervals. The frequency of the comparison measurements depends on several factors including the grid complexity and the measurement uncertainties calculated. Following approval and commissioning, it is recommended that measurements should be carried out at least once per year. When sufficient operating experience has been obtained, the measurement interval may be extended.

9.3.3.5 Software and data processing

General requirements are given in [12.1](#). The software applied for gas quality tracking shall ensure that any accidental, intentional or unintentional influence on the results is avoided. This may for example be achieved by protecting all calculation routines (computing kernel) by check sums. The reproducibility of the calculation results shall also be ensured.

The gas quality tracking software shall meet the following requirements:

- The software version used (computing kernel) including parameters which may be changed shall be clearly identified via a checksum.
- The computing kernel shall not be changed after the approval procedure undertaken by the manufacturer/operator. Any such changes shall be subject to approval.
- In data output on the screen and in results reports, (approved) data relevant for gas charging purposes shall be clearly identified.
- Changes in the grid topology shall be taken into consideration by the software with an hourly resolution.

All input and result data shall be archived for a period of at least one year. Action shall be taken to ensure that any data for charging purposes already generated may be reproduced at any time by repeated calculation.

9.3.3.6 Documentation

The following documentation shall be provided:

- Overall plan of gas grid indicating entry and exit points as well as relevant topology elements (valves, controllers, etc.).
- List of all gas quality and gas quantity measuring instruments relevant for the system.
- Operating manual issued by the supplier of the gas quality tracking software used.
- Test report on fundamental validation and regular validation.
- Procedure instruction of the operator for the operation of the system.
- Quality assurance procedure for gas quality measurement instruments used for validation according to [9.3.3.2](#).

10 Calculation of energy quantities

10.1 General formulae for energy

In accordance with [Figure 2](#), energy determination of a gas flow is based on entities changing in time:

Current flow $q = q(t)$

Current calorific value $H = H(t)$

For the determination of the energy flow, $e(t)$, the basic differential formula is given in [Formula \(1\)](#):

$$e(t) = H(t) \cdot q(t) \quad (1)$$

The quantity of energy, $E(t_j)$ flowing within a period of time t_0 to t_j (for example within an energy-determination period; see [Figures B.1](#) to [B.3](#)) is calculated by integration of [Formula \(1\)](#) from time t_0 to time t_j and giving $E(t_j)$ as given in [Formula \(2\)](#):

$$E(t_j) = \int_{t_0}^{t_j} e(t) dt = \int_{t_0}^{t_j} H(t) \cdot q(t) dt \quad (2)$$

NOTE For example, for a gas flow measured over 24 h or one day $E(t_j)$ becomes $E(24)$ that corresponds the quantity of energy of that period.

The shortest period of time for energy determination for billing purposes is 1 h or multiples thereof (i.e. days, weeks, months, one year).

For the energy determination during 1 h, the following two procedures are possible (see also [7.2.2.1](#) and [7.2.2.2](#)):

- multiplication of the calculated volume under reference conditions with the averaged calculated calorific value of the same hour;
- *in situ* energy calculation in the volume-conversion device using the actual measured entities for the calculation of energy based on the calculation of Q_r and H_r , followed by summing these single energy quantities over 1 h.

The relevant hourly values can subsequently be added to derive daily, monthly or annual quantities.

Calculation formulae:

A small time interval, Δt , is set such that the calorific value, $H(t)$, may be assumed to be a constant entity, H_{const} . In practice, H_{const} matches the result of measuring devices that periodically determine and yield the last measured value between measuring cycles.

To simplify the process, the period t_0 to t_j is subdivided into j time intervals in order to meet the assumption, as given in [Formula \(3\)](#):

$$E(t_j) = H_{\text{const},1} \times \int_{t_0}^{t_1} q(t)dt + H_{\text{const},2} \times \int_{t_1}^{t_2} q(t)dt + \dots + H_{\text{const},j} \times \int_{t_{j-1}}^{t_j} q(t)dt \quad (3)$$

The integrals in [Formula \(3\)](#) correspond to the transported quantities, Q , of gas in the respective time intervals, Δt ; they can be calculated by integration of the actual gas flow over the time, t , as given in [Formula \(4\)](#):

$$Q(t) = \int q(t)dt \quad (4)$$

In practice, however, measuring systems yield the quantity, Q , directly as the result of a measurement.

Thus, [Formula \(4\)](#) can be rewritten as [Formula \(5\)](#):

$$E(t_j) = \sum_{m=1}^j E_m = (H_{\text{const},1} \times Q_1) + (H_{\text{const},2} \times Q_2) + \dots + (H_{\text{const},j} \times Q_j) = \sum_{m=1}^j (H_{\text{const},m} \times Q_m) \quad (5)$$

[Formula \(5\)](#) can be used for any energy-determination period, i.e. from 1 h to days or months.

The monthly average for calculating thermal energy can, for example, be computed from daily values.

For the practical application of incremental energy determination, see [Annex D](#).

10.2 Calculation of averaged values — Calculation from average calorific values and cumulative volumes

When the calorific value is constant during the time period t_0 to t_n (see [Figure B.1](#)), no special calculation is required; if the calorific value changes during this period of time, the procedures for calculation of the appropriate calorific value described in [10.2.1](#) and [10.2.2](#) are used (see [Figures B.2](#) and [B.3](#)).

10.2.1 Arithmetic average of the calorific value

In practical use, the calorific value is often measured at a representative point of the pipeline grid and allocated to volume-measuring stations located at other points. Thus, the arithmetic average of the calorific value, H_{arith} , is derived from n single measurements, as given in [Formula \(6\)](#):

$$H_{\text{arith}} = \sum_{m=1}^j H_m / j \quad (6)$$

[Formula \(5\)](#) can be simplified, if the single factors, $H_{\text{const},m}$, are similar to the arithmetic average value, H_{arith} , as given in [Formula \(7\)](#):

$$E = H_{\text{arith}} \times Q \quad (7)$$

10.2.2 Quantity-weighted average of the calorific value

If the energy of the gas quantity transported from time t_0 to t_n is set in relationship to the gas quantity, $Q(t_n)$, that was transported in the same time period, the so-called “quantity-weighted average of the calorific value” is given by [Formula \(8\)](#), taking into account [Formula \(5\)](#):

$$H(t_j) = E(t_j) / Q(t_j) = \sum_{m=1}^j H_{\text{const},m} \times Q_m / \sum_{m=1}^j Q_m \quad (8)$$

Each of the j single calorific values, $H_{\text{const},m}$, is weighted by the respective quantity Q_m .

For practical examples, see [Annex F](#).

10.3 Volume and volume-to-mass conversions

Where gas-quality data are required for conversion purposes at the interfaces, it is necessary that those data shall relate to the gas flowing to those interfaces, for example by having been measured at the appropriate calorific-value-measuring station. If the calorific value is expressed in volume units and the quantity of gas is measured in mass units, then the calorific value shall be converted from volume units to mass units.

See [Annex C](#) for a general description. For practical examples based on physical properties, see [E.1](#); for practical examples based on the gas composition, see [E.2](#).

10.4 Energy determination on the basis of declared calorific values

To enable a practical, easy energy determination mainly at interface 4, but also applicable to interfaces 5 and 6 (see [Figures 3](#) and [4](#)), a calorific value may be declared within an energy-determination period for a charging area with a set of different interfaces. Prior to the energy-determination procedure, the declared calorific value shall be determined by means of the weighted-average calorific value. For example, if the energy-determination period is one year, the calorific value is determined for each month as an arithmetic or a weighted mean. At the beginning of the month in which energy determination is performed, the weighted annual mean is calculated using the last 12 months' means and taking into account the monthly take-off quantities for the relevant customer/customers.

For longer energy-determination periods, that month in which energy determination is performed shall not be taken into account for calculating the mean value.

A declared calorific value is set by the LDC in the case of an annual energy-determination procedure. The difference between the declared calorific value and the actual mean should be assessed annually. If the difference is more than 1 %, the actual mean should be taken. It is important that there be, over a long time period, no significant bias between the declared and calculated averaged calorific values.

A practical example is given in [Annex B](#).

11 Accuracy on calculated energy

11.1 Accuracy

11.1.1 The accuracy on the quantity of energy determined as passing a particular interface is a combination of

- uncertainties, and
- biases.

11.1.2 Uncertainties can have two sources:

- uncertainty of the measurements of the gas quantity and calorific values made for the determination of energy;
- variability of the parameter being measured (of particular importance when the numerical value of that parameter used in the determination of energy is obtained by some process from measured values).

Uncertainties can be quantified but not eliminated.

11.1.3 Bias results from systematic differences between the actual quantity and calorific value used in the calculation of energy at a gas-measuring station and the true values for those parameters.

A possible bias in the determined energy has a number of sources, such as

- an error in a calibration factor that affects subsequent measurements,
- use of fixed factors, for example in the conversion of measured gas volumes to the corresponding volume under reference conditions, or
- a calorific value attributed to an interface having only gas-volume measurement that may be unrepresentative of the gas passing that interface.

General information on the calculation of uncertainty and identification of bias on the determined quantity of energy is given in [11.2](#) and [11.3](#). A graphical example to show the principal difference between uncorrected measurements, biases and the final result of a measurement (for example a corrected calorific value) by means of a graph, based on ISO/IEC Guide 98-3 (GUM), is given in [Annex I](#).

11.2 Calculation of uncertainty

The relative uncertainty, $u(E)$, of the energy is calculated from [Formula \(9\)](#), which is derived from the general formula for the determination of energy, [Formula \(10\)](#):

$$u(E) = \left[u^2(H) + u^2(Q) \right]^{1/2} \quad (9)$$

where

$u(H)$ is the relative uncertainty of the calorific value;

$u(Q)$ is the relative uncertainty of the quantity of gas.

When calculating uncertainties in energy, it is necessary to take into account the uncertainties of all known influencing factors [see ISO/IEC Guide 98-3 (GUM)].

The total uncertainty of the calculated energy is also affected by the manner in which integration is carried out with respect to the period of time over which the energy calculation is made. If at a gas-measuring station both volume and calorific value are measured and over very short intervals of time the energy is calculated and each of these individual energy sums for the total period are added together, then the integration has a relatively small effect on the total uncertainty. At the other extreme, where the total gas volume delivered over a period of months is multiplied by an average calorific value for that period to obtain the energy for the period, then the effect of the integration on the total uncertainty can be significant, particularly when the rate of gas usage and the actual calorific value vary over the time period. When applying a calorific-value assignment method, the effect of the time delay on the uncertainty shall be taken into account. The practical calculations of uncertainty [see ISO 12213 (all

parts) for the calculation of the uncertainty of Z] depend upon the way in which the quantity of energy passing an interface is calculated (see [Clause 8](#)).

NOTE The uncertainties of the flow rate, p , T and Z , can be determined as given in the flow-measurement standards and ISO 12213 (all parts). As a first approximation, the relative uncertainty for an individual energy calculation can be regarded as equal to the relative uncertainty of the energy calculated over longer periods of time (even over a billing period) obtained by integrating the small portions of energy. This approximation is only applicable when

- j) the relative uncertainty of the measured calorific value is constant over the range of measured calorific values, and
- k) the relative uncertainty of the measured quantity is constant over the measuring range of the flow meter. In practice, this assumption is valid only over a part of a flow meter's range. For example, for some orifice plate metering systems, the uncertainty is essentially constant over 30 % to 100 % of $Q_{\max}$. In some cases, it can be acceptable to use the maximum relative uncertainty for the range of flow rates that are experienced by the flow meter.

The energy, E , is calculated from [Formula \(10\)](#), the general form of [Formula \(7\)](#):

$$E = H \times Q \quad (10)$$

where H is the average or assigned calorific value for the period.

The uncertainty $u(E)$ is calculated from [Formula \(9\)](#), where $u^2(H)$ is the uncertainty of the average or assigned calorific value.

The uncertainty of the average or assigned calorific value contains three elements:

- uncertainty of measurement;
- uncertainty due to variation of the calorific value of the gas over the averaging period;
- time delay.

11.3 Bias

11.3.1 Wherever possible, a bias (see [Annex I](#)) should be identified and eliminated.

Where both the calorific value and volume and/or mass are measured at an interface, bias in the determined energy can arise from potential sources, such as

- calibration errors, or
- use of fixed factors instead of online data, for example, in the conversion of volume from operating to recommended standard reference conditions.

Identification of the source of bias should be achieved through a comparison of data (see [9.2.2](#)), for example a determination of the composition and calorific value of a certified test gas to verify the properties of a calibration gas. In some instances, it can be possible to compare the outputs from measuring systems that are in series, for example, two different measuring systems at the same interface operated by different upstream and downstream measuring partners.

The potential for bias in the determined energy is increased when calorific values are assigned to an interface and/or where simple meters are in use. It can be possible to identify circumstances in which biases are introduced but the options to quantify and eliminate a bias can be limited to procedural changes.

11.3.2 Bias can result from the manner in which the gas volume measured by the meter is converted to an equivalent volume expressed under standard reference conditions. The possibility of bias is high for the relatively simple meters used at interface 4 for domestic consumers, such as

- where the conversion of the measured gas volume at operating temperature to the equivalent volume at standard temperature involves an assumed, fixed value for the operating temperature of the meter; in such cases where the operating temperature is higher than the assumed temperature, the gas volume calculated at the standard temperature is erroneously high, and vice versa, or
- where the conversion of the measured gas volume at operating pressure to the equivalent volume at standard pressure involves the use of a fixed average pressure, for example derived from an average altitude to compensate for atmospheric-pressure variation with altitude; in such cases where the altitude of the interface is higher than the fixed altitude, the gas volume calculated at the standard pressure is too high and vice versa.

11.3.3 Bias can also result from the fact that the calorific value assigned to the interface is not sufficiently representative of the gas passing the interface caused by, for example:

- use of a time-averaged calorific value at an interface where the flow rate of gas varies significantly, possibly even falling to zero, with time;
- use of a flow-weighted average calorific value for a network in which the majority of interfaces are systematically supplied from sources having calorific values different to the flow-weighted average value;
- use of a declared calorific value which, for reasons of regulation, is required to be lower than the calorific value of the gas passing any interface in a network.

12 Quality control and quality assurance

12.1 General

The energy-determination system shall be monitored to ensure its proper functioning and to keep it at its intended level of accuracy and integrity, e.g. by redundant systems. This covers all maintenance, verification and calibration activities pertaining to the system during its lifetime and can be accomplished by maintaining a record of the system performance. Data handling and data transmission shall be performed using safe and secure procedures.

If matching data for a comparison over time are given, the data shall be assessed to recognize possible bias and to evaluate it. This step can usually be performed only if a suitably large quantity of data has been collected, for example after one year. Therefore, the data generated in accordance with [Clause 10](#) are directly used for billing purposes.

Quality control should be embedded in the existing (maintenance) organization of the user.

12.2 Check of the course of the measuring data

To avoid permanent disadvantages of partners causing cross-subsidies of other customers, bias should be avoided at all interfaces. The detection of biases can be performed by comparison on the basis of statistical methods using graphical tools and/or calculation tools, for example, the CUSUM method (see ISO 7870-4).

When using statistical methods on the basis of calculation tools for the detection of bias, the user should take into account that the application of these methods is usually reasonable only for interfaces with large quantities of gas, where extended data acquisition and data analysis, including historical data, can be submitted.

Generally, the following possibilities for performing a data comparison are available:

- calibration standard $\Leftrightarrow$ calibration standard;
- calibration means $\Leftrightarrow$ measuring device;
- measured values $\Leftrightarrow$ calculated values;
- measuring device $\Leftrightarrow$ measuring device;
- input station $\Leftrightarrow$ sum of output stations.

In general, the detection of a bias can reasonably be accomplished only when taking into account a suitably long time period.

For calorimeters, such an interval may be within two calibration periods, for example one month. Usually a comparison of volume-measuring stations is reasonable only after one year, taking into account random summer/winter effects. Where biases (systematic errors) are identified, action shall be taken to quantify and eliminate the bias; in the meantime, suitable substitute values or correction factors shall be used.

12.3 Traceability

Given the physical implications of custody transfer of natural gas, accuracy of measurement of the delivered energy is of fundamental importance. Inaccuracy in the sense of bias or systematic error favours one party at the expense of the other. Random errors have a neutral effect.

Most measurement instruments operate on a comparative technique, and the measurement accuracy is fundamentally influenced by the accuracy of the calibration standard used. Hence the interest in reference materials and in traceability. Traceability is the property of a result of a measurement whereby it can be related, with stated uncertainty, to stated references, usually national or International Standards, through an unbroken chain of comparisons.

12.4 Substitute values

Energy determination shall be ensured at all times as long as energy is transmitted via an interface, even if the measuring instruments have partly or completely ceased to function.

If a measuring instrument has failed, substitute values are required until the proper function of the measuring instruments is restored. They shall be obtained on the basis of the most plausible available values and may be derived from measuring instruments, auxiliary equipment and/or computational models for quantitative estimation.

Substitute values and their origin shall be made known and explained to the contracting parties directly affected by the malfunctioning instrument. They require, in each case, the approval of the parties directly affected (generally the contracting parties). Substitute values shall be clearly distinguished from the other values.

The contracting parties directly concerned shall agree on the general procedures to obtain substitute values within a reasonable amount of time before energy transmission. Such procedures may involve

- upstream and downstream measuring instruments,
- linear regression,
- interpolation of the last plausible value before malfunction to the first plausible value after malfunction,
- hourly, daily, weekly, monthly or annual comparisons with equivalent past periods,
- comparison of output or plant efficiency in the case of industrial customers,

- comparison based on samples,
- comparison of flow conditions in multi-stream measuring systems with constant flow resistance,
- calculations according to physical laws of flow, etc.

Examples of such procedures are given in [Annex G](#).

STANDARDSISO.COM : Click to view the full PDF of ISO 15112:2018

Annex A (informative)

Main instruments and energy-determination techniques

Table A.1

Country	Interface(s) at which the following volume measurement techniques take place							Backup of volume metering
	Diaphragm	Rotary displacement	Vortex	Turbine	Orifice	Ultrasonic	Other	
Belgium	—	3, 5, 6	—	1, 2, 3, 5, 6	—	—	—	—
China	4, 6	4, 5, 6	3, 5, 6	2, 3, 5, 6	1, 2, 3, 5, 6	1, 2, 3, 5	—	—
Germany	4, 6	3, 4, 5, 6	2, 3	1, 2, 3, 5, 6	1, 2	2	—	1, 2, 3, 5
France	4, 6	3, 5, 6	—	1, 2, 3, 5, 6	1, 2	—	—	1, 2
United Kingdom	4, 6	5	—	1, 5	1	4, 5, 6	—	1, 5
Italy	4, 6	3, 5	—	3, 5	1, 3, 5	—	—	1, 3, 5
Netherlands	4, 6	3, 5, 6	—	1, 2, 3, 5, 6	1	1, 2	—	1, 2, 3, 5
Austria	—	—	—	1, 2	1, 2	—	—	1, 2
Russian Federation	1, 3, 5, 6	—	—	1, 3, 5	1, 2, 3, 5	—	—	3, 4
Hungary	4, 5, 6	3, 5, 6	—	3, 5, 6	1, 3, 5	—	—	1, 3
USA	4, 5, 6	1, 2, 3, 4, 5, 6	3, 5	1, 2, 3, 5, 6	1, 2, 3, 5	2, 3, 5	5 ^a	1, 2, 3, 5

^a Venture nozzle and mass.

Table A.2

Country	Interface(s) at which the following activities take place											
	Measurement of		Data storage		Measurement of density at		Compres- sibility factor calcula- tion	Volume con- version	Calori- fic mea- sure- ment	Calori- fic value correc- tion	Substi- tute values at errors	Direct energy mea- sure- ment
Volume	Tempe- rature pres- sure	Inside station	Outside station	Operat- ing condi- tions	Normal condi- tions							
Belgium	1, 2, 3, 4,5,6	2, 3	—	—	—	—	3 ^b	1, 2, 3	2	—	2	—
China	1, 2, 3, 4, 5, 6	1, 2, 3, 5, 6	1, 2	—	—	—	1, 2, 3, 5, 6	—	—	—	—	—
Germany	1, 2, 3, 4, 5, 6	1, 2, 3, 5, 6	1	—	1, 2, 5	1, 2, 5, (1, 2) ^a	1	1	1, 2, 5	1	1	—
France	1, 2, 3, 4, 5, 6	1, 2, 3, 5, 6	1	—	(1, 2) ^a	2 (1, 2) ^a	1	1	1, 2	—	1	—
United Kingdom	1, 4, 5, 6	1, 2, 5, 6	1	1	1 (1, 2, 5) ^a	1 ^a	1	1	1, 2, 5	1	1	—
Italy	1, 3, 4, 5, 6	1, 3, 5	1	—	1 ^a	3, 5 (1, 3, 5) ^a	1	1	1, 3, 5	—	1	—
Netherlands	1, 2, 3, 4, 5, 6	1, 2, 3, 5, 6	1, 2, 5, 6	2	1 (1, 2, 5) ^a	1, 2 (1, 2, 3, 5) ^a	1, 2, 4, 5, 6	1, 2, 3, 4, 5, 6	1, 2, 3, 5	5, 6	1	—
^a Calculated density value.												
^b Plus Z-meter.												

^a Calculated density value.

^b Plus Z-meter.

Table A.2 (continued)

Country	Interface(s) at which the following activities take place										Substitute values at errors	Direct energy measurement
	Measurement of Volume	Temperature pressure	Data storage Inside station	Outside station	Measurement of density at Operating conditions	Normal conditions	Compressibility factor calculation	Volume conversion	Calorific measurement	Calorific value correction		
Russian Federation	1, 2, 3, 4, 5, 6	1, 2, 3, 4, 5, 6	1, 2, 3, 4, 5, 6	—	(1, 2, 3, 5) ^a	1, 2, 3, 4, 5, 6 (1, 2, 3, 5) ^a	1, 2, 3, 4, 5, 6	1, 2, 3, 4, 5, 6	—	—	1, 2, 3, 4, 5, 6	—
Hungary	1, 3, 4, 5, 6	1, 3, 5, 6	1	1	(1, 3, 5) ^a	(1, 3, 5) ^a	1	—	3, 5	—	1	—
USA	1, 2, 3, 4, 5, 6	1, 2, 3, 4, 5, 6	1, 2, 3, 5, 6	1, 2, 3, 5, 6	1 to 6, (1 to 6) ^a	1, 2, 3, 5, 6 (1, 2, 3, 5, 6) ^a	1, 2, 3, 5, 6	1, 2, 3, 5, 6	1, 2, 3, 5	1, 2, 3, 5	1, 2, 3, 5, 6	1, 2, 3, 5
^a Calculated density value.												
^b Plus Z-meter.												

Table A.3

Country	Interface(s) at which the following activities take place							Backup of calorific value measurement
	Density at Operating conditions	Normal conditions	Temperature, pressure measurement	Calorific value measurement by GC	Wet calorimetry	Dry calorimetry	Other	
Belgium	—	(2, 3, 5, 6) ^a	2, 3, 5, 6	(2, 3, 5, 6) ^c	—	—	—	2
China	(3, 5) ^a	(3, 5) ^a	3, 5	(3, 5) ^d	—	—	—	—
Germany	(1, 2, 5) ^b	(1, 2, 5) ^b ; (1, 2) ^a	1, 2, 3, 5, 6	(1, 2, 5) ^c ; 2 ^d	1, 2, 5	—	—	1, 2
France	(1, 2) ^a	(2, 5) ^b ; (1, 2) ^a	1, 2, 3, 5, 6	(1, 2) ^c	1	—	—	1, 2
United Kingdom	1 ^b ; (1, 2, 5) ^a	1 ^a	1, 2, 5, 6	(1, 2, 5) ^c	—	—	—	1, 2, 5
Italy	1 ^a	(3, 5) ^b ; (1, 3, 5) ^a	1, 3, 5	(1, 5) ^c ; (1, 3, 5) ^d	—	—	—	—
Netherlands	(1, 5) ^b ; (1, 2, 5) ^a	(1, 2, 5) ^b ; (1, 2, 3, 5) ^a	1, 2, 3, 5, 6	(1, 2, 5) ^c ; 1 ^d	1	—	—	1, 2, 5
Russian Federation	(1, 2, 3, 5) ^a	(1, 2, 3, 5) ^{a,b}	1, 2, 3, 5, 6	—	—	5	1, 2, 3, 5	—
Hungary	(1, 3, 5) ^a	(1, 3, 5) ^a	1, 3, 5, 6	(1, 5) ^c	—	—	—	—
USA	(1, 2, 3, 5, 6) ^{a,b}	(1, 2, 3, 5, 6) ^{a,b}	1, 2, 3, 4, 5, 6	(1, 2, 3, 5) ^{c,d}	1, 2, 3, 5	1, 2, 3, 5	—	1, 2, 3, 5
^a Calculated value.								
^b Measured value.								
^c Online.								
^d Offline.								

Table A.4

Country	Interface(s) at which the following activities take place					
	Compressibility factor	Volume conversion	Data storage	Generation of substitute values	Correction of metered calculated values by means of correction factor	Direct energy measurement
Belgium	1 ^{b,c}	1 ^d	1 ^{d,e}	1 ^{f,g}	—	—
China	1, 5 ^a	1, 5 ^d	1 ^d	—	—	—
Germany	1 ^b	1 ^d	1 ^d	1 ^{f,g}	1	—
France	1 ^b	1 ^d	1 ^d	1 ^{f,g}	—	—
United Kingdom	1 ^b , 5 ^a	1, 5 ^d ; (4, 5, 6) ^e	1 ^{d,e}	1 ^g	1	—
Italy	1 ^a	1 ^d	1 ^d	1 ^{f,g}	—	—
Netherlands	(1, 3, 4, 6) ^a ; (2, 4, 6) ^b	(1, 2, 3, 4, 5, 6) ^d ; 2 ^e	(1, 2, 5, 6) ^d ; 2 ^e	1 ^g	5, 6	—
Russian Federation	(1, 2, 3, 4) ^a	(1, 2, 3, 5) ^{d,e}	(1, 2, 3, 5) ^d	(1, 2, 3, 5) ^{f,g}	—	1, 5
Hungary	(1, 3, 5) ^a ; (1, 3, 5) ^b	(1, 3, 5) ^{d,e}	(1, 3, 5) ^{d,e}	(1, 3, 5) ^{f,g}	—	—
USA	(1, 2, 3, 5, 6) ^a	(1, 2, 3, 5, 6) ^{d,e}	(1, 2, 3, 5, 6) ^{d,e}	(1, 2, 3, 5, 6) ^f ; (1, 2, 3, 5) ^g	1, 2, 3, 5	1, 2, 3, 5
^a Volume conversion by the AGA 8 method; see Reference [35]. ^b Volume conversion by the S-GERG88 method; see Reference [36]. ^c Z-meter. ^d Inside station. ^e Outside station. ^f Local. ^g Remote.						

Table A.5

Country	Interface(s) at which the following activities take place						
	Energy determination on the basis of locally metered values	Energy values on the basis of calculated calorific values	Energy values, calculated on state reconstruction or mathematical models	Minimum time periods for energy determination for charging			
				Hourly	Daily	Monthly	Other
Belgium	2, 3, 5, 6	4	4	2, 3, 5, 6	3, 5, 6	5	(4, 6) ^c
China	3, 5	—	—	5	—	—	4 ^c , 5 ^a
Germany	1, 2, 3, 5	2, 3, 4, 5, 6	3, 5	1, 2, 3, 5, 6	1, 2, 3	3, 4, 5, 6	(4, 6) ^c
France	1, 2, 3, 5	3, 4, 5, 6	3, 5	1, 2, 5	1, 2, 3, 5	3, 4, 6	4 ^a
United Kingdom	1, 2, 5, 6	1, 4, 5, 6	—	—	1, 4, 5	5, 6	4 ^b
Italy	1, 3, 5	3, 4, 5, 6	—	—	1, 4, 5	3, 4, 5, 6	—
Netherlands	1, 2, 3, 5	4, 5, 6	—	1, 2, 3, 5	—	6	(4, 6) ^c
^a Weekly. ^b Quarterly. ^c Annually.							

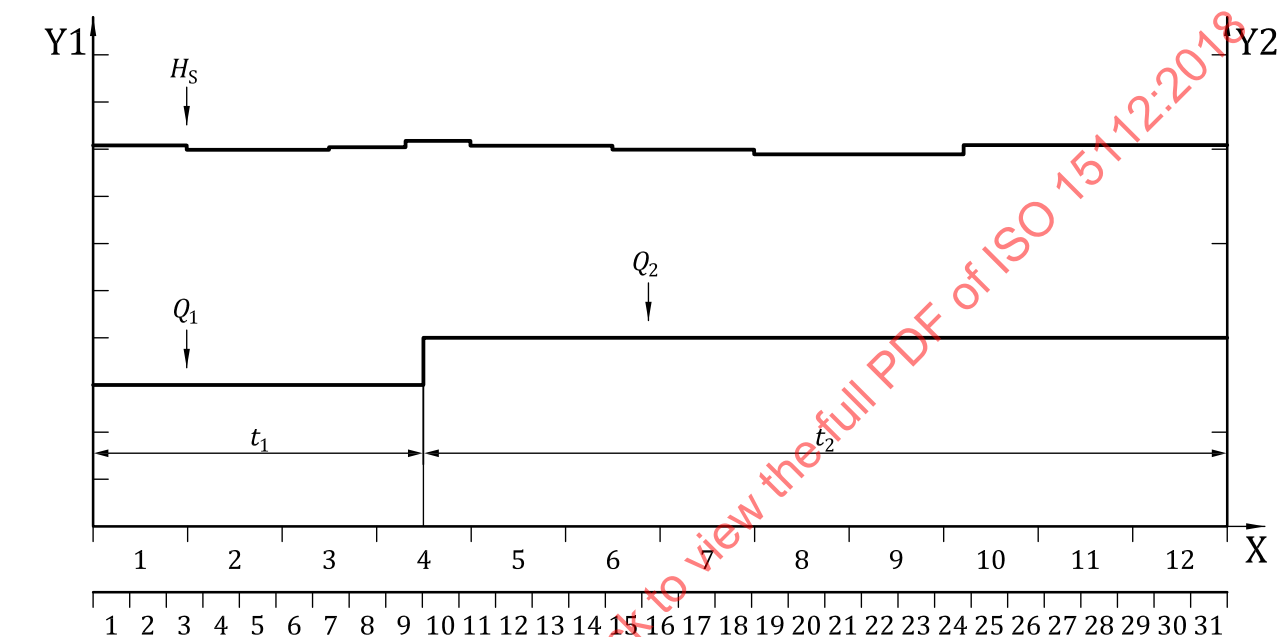
Table A.5 (continued)

Country	Interface(s) at which the following activities take place						
	Energy determination on the basis of locally metered values	Energy values on the basis of calculated calorific values	Energy values, calculated on state reconstruction or mathematical models	Minimum time periods for energy determination for charging			
				Hourly	Daily	Monthly	Other
Russian Federation	1, 2, 3, 4, 5, 6	—	1, 2, 3, 4, 5, 6	—	—	1, 2, 3, 4, 5, 6	—
Hungary	1, 3, 5	3, 4, 5, 6	—	—	1	3, 4, 5, 6	4 ^c , 5 ^a
USA	1, 2, 3, 5	1, 2, 3, 5	4, 5, 6	1, 2, 3	1, 2, 3, 5	4, 6	—
^a Weekly. ^b Quarterly. ^c Annually.							

Annex B (informative)

Different possible patterns in the change of the calorific value

Figures B.1 to B.3 show three different examples of possible patterns in the change of the calorific value for an energy-determination period.



Key

X months (1 = January, 12 = December) or days (1st to 31st of each month)

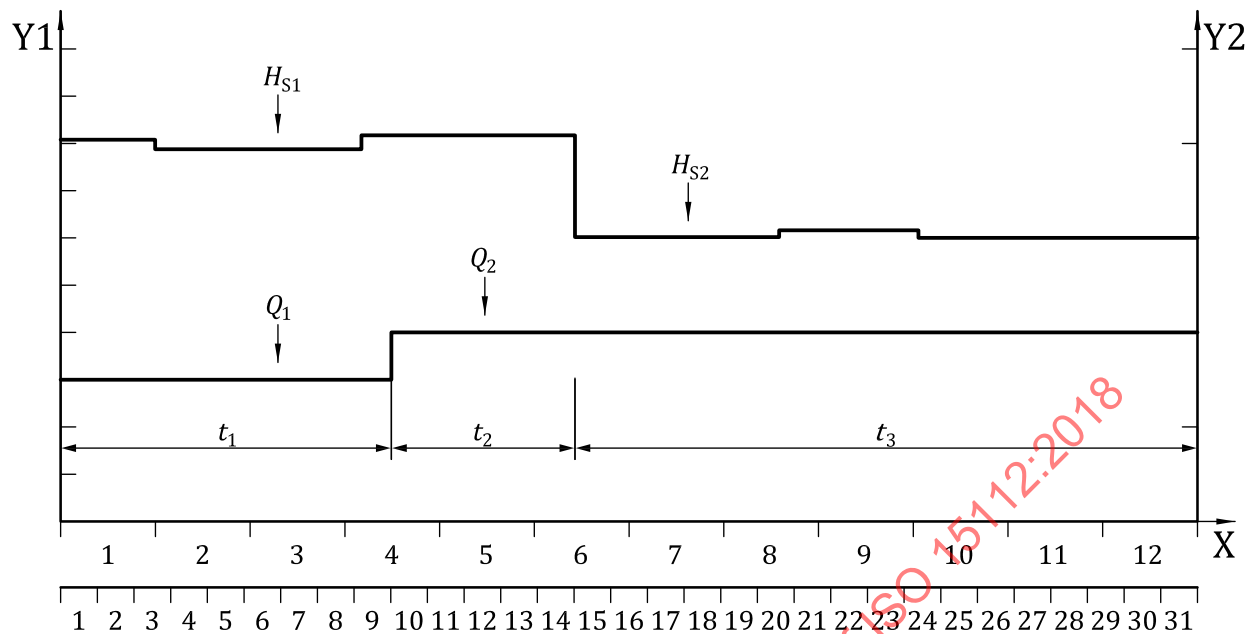
Y1 H_s , expressed in megajoules per cubic metre

Y2 q_v , expressed in cubic metres per day or cubic metres per month

Figure B.1 — Annual or monthly energy-determination period — Only normal changes in gas quality

In Figure B.1, the calorific value in a energy-determination period, for example one month, is nearly constant. Thus, it is justified to calculate an averaged calorific value for the complete month (see Clause 10 and especially 10.4 as a justified method). For energy determination, it is important to take into account that for the period from the 1st to the 10th, the gas flow rate was smaller than for the period from the 10th to 31st. Therefore, an energy quantity, E_1 , can be calculated for the time period, t_1 , and the energy quantity, E_2 , for the time period, t_2 .

If the time scale is one year, it is justified to calculate an annual averaged calorific value due to the shape of the calorific value curve. For such a year, an energy quantity, E_1 , can be calculated for January to April, and for May to December an energy quantity, E_2 .

**Key**

X months (1 = January, 12 = December) or days (1st to 31st of each month)

Y1 H_s , expressed in megajoules per cubic metre

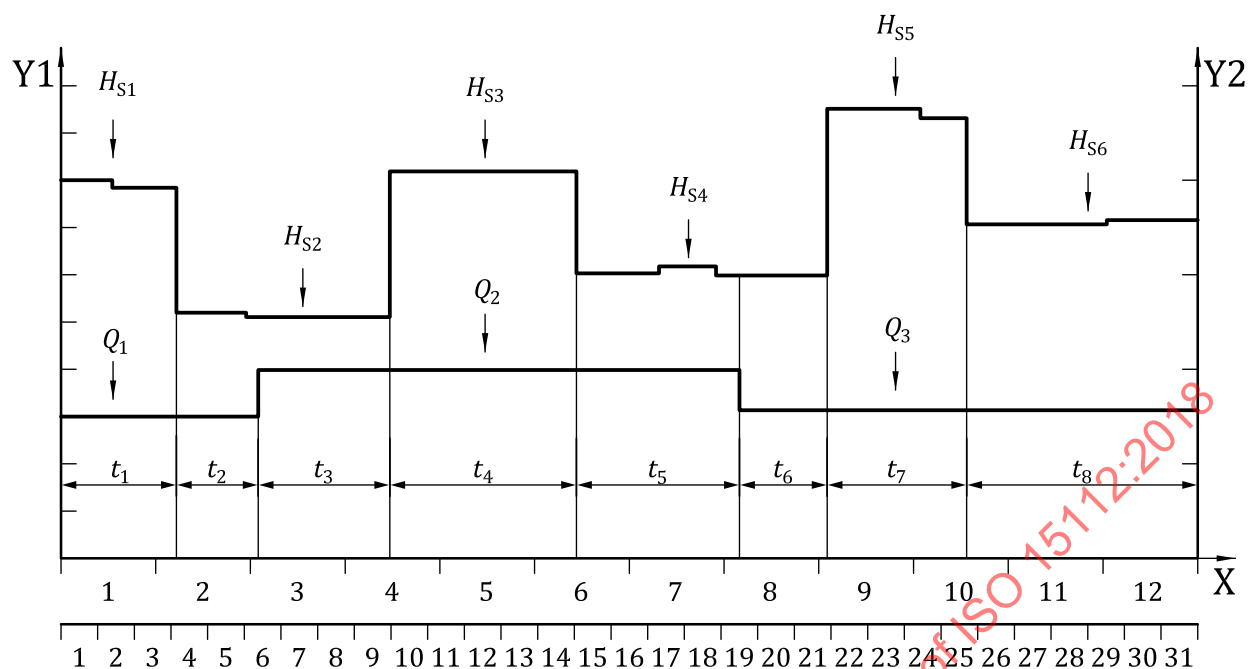
Y2 q_v , expressed in cubic metres per day or cubic metres per month

Figure B.2 — Annual or monthly energy-determination period — Two different calorific values

In [Figure B.2](#), for example, the calorific value within the energy-determination period is nearly constant up to the 15th; the calorific value decreases then to a significantly lower level. For the energy determination, either the month shall be separated into two periods from the 1st to the 15th and from the 16th to the 31st, or an averaged calorific value shall be performed (see [Clause 9](#) and [Annex F](#)). For large quantities and, therefore, for economic reasons, a separation into two periods shall be performed.

In this case, for energy-determination purposes, the energy quantities, E_1 , E_2 and E_3 , for the respective time periods, t_1 , t_2 and t_3 , shall be determined separately to compensate for the different quantities from the 1st to the 10th day and the change in calorific value at the 15th day.

If the time scale is one year, the patterns in the change of the calorific value justify separating the year into the determination periods, t_1 , t_2 and t_3 . Thus, the energy quantity, E_1 , can be calculated for the period from January to April; E_2 , for the period from May to June; and E_3 , for the period from July to December.



Key

X months (1 = January, 12 = December) or days (1st to 31st of each month)

Y1 H_s , expressed in megajoules per cubic metre

Y2 q_v , expressed in cubic metres per day or cubic metres per month

Figure B.3 — Annual or monthly energy-determination periods — Several different calorific values

In [Figure B.3](#), the calorific value within an energy-determination period, for example one month, changes several times; thus, either the month shall be separated into a couple of periods or an averaged calorific value shall be calculated (see [Clause 9](#) and [Annex F](#)). For large quantities and, therefore, for economic reasons, a subdivision shall be made to take the calorific values H_{s1} to H_{s6} into account individually.

In this case, the energy quantities, E_1 to E_8 , associated with the time periods t_1 to t_8 , respectively, are calculated separately, using the different quantities, Q_1 , Q_2 and Q_3 , and the different calorific values associated with each time period. The total energy quantity is obtained by summing the E_1 to E_8 values [see [Formula \(5\)](#)].

If the time scale is one year, the patterns in the change of the calorific value justify separating the year into different energy-determination periods with H_{s1} , H_{s2} , H_{s3} , etc. and to determine for each period an averaged calorific value for energy determination.

Annex C (informative)

Volume conversion and volume-to-mass conversion

As the common International Standards for flow (ISO 5167-1 and ISO 9951) generally provide the flow rate in mass per second or volume (under operating conditions) per second, in the latter case a conversion to volume under reference conditions is necessary.

Starting from the continuity in mass, EN 1776:1998, Annex C, derives formulae for volume conversion and determination of mass from volume and density under operating conditions. These are given, using symbols conforming with the ISO directives, as [Formulae \(C.1\)](#) and [\(C.2\)](#) for calculating the converted volume, V_r , under reference conditions, expressed in cubic metres, and [Formula \(C.3\)](#) for calculating the mass, M , expressed in kilograms.

[Formulae \(C.1\)](#) to [\(C.3\)](#) can be used for calculations relevant to this document.

$$V_r = V \cdot \frac{p \cdot T_r \cdot Z_r}{p_r \cdot T \cdot Z} \quad (\text{C.1})$$

$$V_r = V \cdot \frac{\rho}{\rho_r} \quad (\text{C.2})$$

$$M = V \cdot \frac{p \cdot M_m}{T \cdot Z \cdot R} \quad (\text{C.3})$$

where

- p is the pressure under operating conditions, expressed in kilopascals (bar);
- p_r is the pressure under ISO-recommended reference conditions, expressed in kilopascals (bar);
- T is the operating temperature, expressed in Kelvin;
- T_r is the ISO-recommended reference temperature, expressed in Kelvin;
- V is the volume under operating conditions, expressed in cubic metres;
- Z is the compression factor under operating conditions;
- Z_r is the compression factor under ISO-recommended reference conditions;

NOTE For compression factor, see 6.4.4.

- M_m is the molar mass, expressed in kilograms per mole;
- R is the universal gas constant, equal to 8,314 510 kJ/(kmol × K);
- ρ is the density under operating conditions, expressed in kilograms per cubic metre;
- ρ_r is the density under the ISO-recommended reference conditions, expressed in kilograms per cubic metre.

Annex D
(informative)

Incremental energy determination

In this method, measurements of the calorific value are made at short intervals of time and are multiplied by the quantity of gas registered by the meter in the interval between successive measurements to obtain the quantity of energy for that interval. The time intervals are typically a few minutes, the only requirement being that the calorific value of the gas should remain essentially constant over the chosen time interval.

In many cases, the time interval is equal to the cycle time of the GC used to determine the calorific value. The individual quantities of energy for all the time intervals in the billing period are then added together to give the energy. The method is illustrated in [Figure D.1](#).

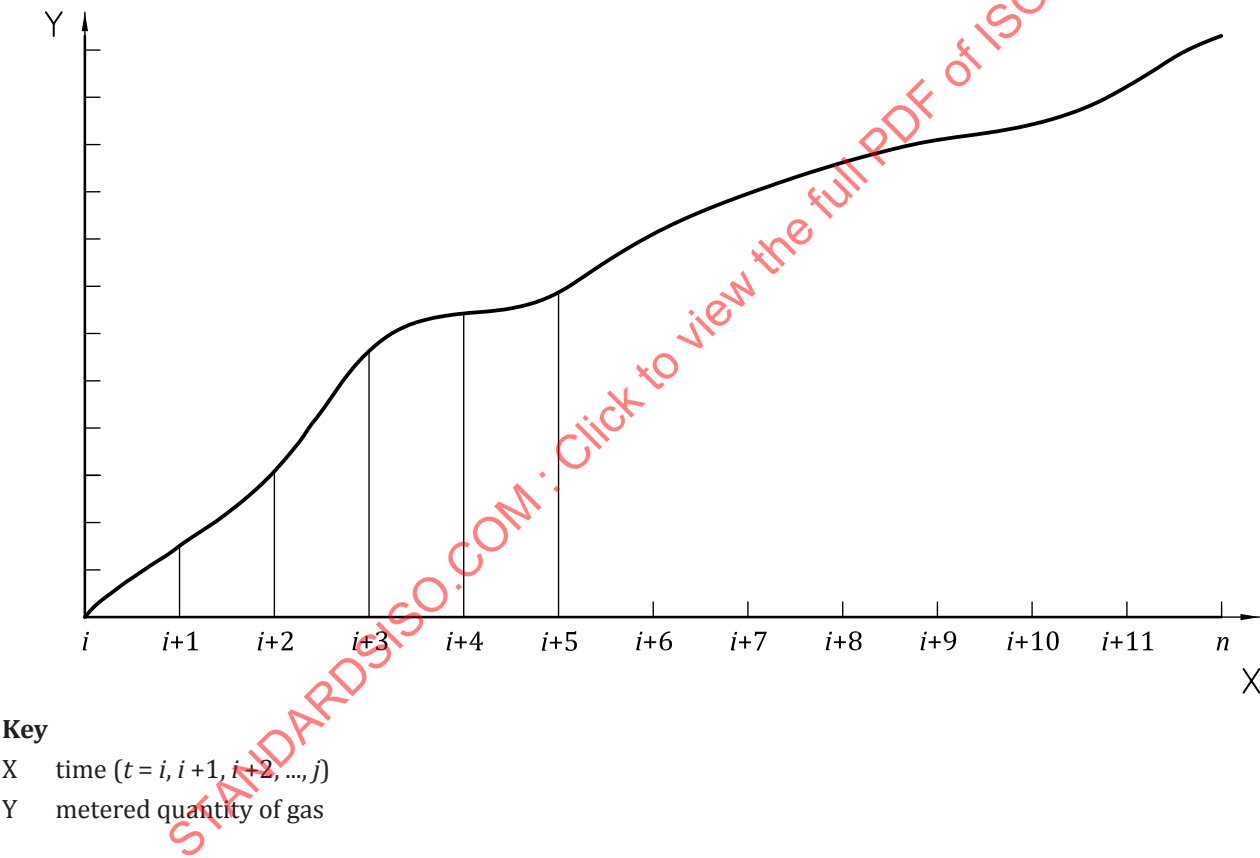


Figure D.1 — Incremental energy determination

At time i , the quantity of gas registered on the meter is $Q_{t=i}$ and the calorific value of gas is measured as $H_{t=i}$.

At time $i + 1$, the quantity of gas registered on the meter is $Q_{t=i+1}$ and the calorific value is measured as $H_{t=i+1}$.

Then the quantity of energy for the interval from $t = i$ to $t = i + 1$ is given in [Formula \(D.1\)](#):

$$E_{t=i \text{ to } t=i+1} = (Q_{t=i+1} - Q_{t=i}) \cdot H_{t=i} \quad (\text{D.1})$$

Then, for any period from time i to time j , the total energy, E , is obtained by adding together all the discrete portions of energy, as given in [Formula \(D.2\)](#):

$$E = (E_{t=i \text{ to } t=i+1} + E_{t=i+1 \text{ to } t=i+2} + \dots + E_{t=j-1 \text{ to } t=j}) \quad (\text{D.2})$$

Practically, this method is implemented using a flow computer to record the quantity of gas registered by the flow meter and the calorific value measurements being input into the flow computer.

Generally, this method is found where both quantity and quality measurements are carried out at the interface. However, state-of-the-art information systems can be used to provide online, fixed-assignment calorific values that can be used in this method.

Annex E (informative)

Practical examples for volume conversion and energy quantity calculation

E.1 Calculations using ISO 12213-3

E.1.1 General formulae

The calculation of the volume, V_n , expressed in cubic metres under normal reference conditions, from the volume under operating conditions is given by [Formula \(E.1\)](#):

$$V_n = V \cdot z \quad (\text{E.1})$$

where

V is the volume under operating conditions, expressed in cubic metres;

z is the z-factor, calculated as given in [Formula \(E.2\)](#):

$$z = \frac{T_n}{T} \cdot \frac{p_{\text{amb}} + p_g - p_{\text{H}_2\text{O}}}{p_n} \cdot \frac{Z_n}{Z} \quad (\text{E.2})$$

where

T_n is the temperature under normal reference conditions, expressed in Kelvin;

T is the operating temperature, expressed in Kelvin;

p_{amb} is the average of the ambient air pressure at the meter, expressed in kilopascals (bar);

p_g is the operating pressure (gauge), expressed in kilopascals (bar);

$p_{\text{H}_2\text{O}}$ is the partial pressure of water in natural gas, expressed in kilopascals (bar);

p_n is the pressure under normal reference conditions, expressed in kilopascals (bar);

Z is the compression factor under operating conditions;

Z_n is the compression factor under normal reference conditions.

Z_n/Z can be calculated from H_S , p_n , and the concentrations of CO_2 , N_2 and H_2 , for example using ISO 12213-3 (S-GERG88; see Reference [36]).

E.1.2 Example calculation

The calculation for the converted volume, V_n , is carried out as demonstrated using the following parameters, as given in ISO 12213-3 (S-GERG88; see Reference [36]):

- average of the ambient air pressure at the meter, p_{amb} 99,66 kPa (0,996 6 bar);
- operating pressure (gauge), p_g 700 kPa (7,0 bar);
- operating temperature, T 288,15 K;
- calorific value, $H_{S,n}$ 11,901 kWh/m³;
- density, ρ_n 0,822 7 kg/m³;
- concentration of CO₂, C_{CO_2} 1,12 mol %;
- concentration of N₂, C_{N_2} 0,80 mol %;
- concentration of H₂, C_{H_2} 0 mol %;
- partial pressure of water in natural gas, p_{H_2O} <0,1 kPa (<0,001 bar);

NOTE 1 p_{H_2O} can be expressed by the product of φ (relative humidity of the gas) and p_{sat} (partial pressure of steam in saturated gases); in dry natural gases p_{sat} is usually $p_{sat} \leq 0,1$ kPa (0,001 bar). Thus, for dry natural gases the expression ($p_{H_2O} = \varphi \cdot p_{sat}$) can usually be set to zero.

- Z_n/Z ratio 1,017 52.

NOTE 2 This value is calculated using the first eight values above in ISO 12213-3 (S-GERG 88; see Reference [36]).

The substitution of the measured values into [Formula \(E.2\)](#) results in the following:

$$z = \frac{273,15 \text{ K}}{288,15 \text{ K}} \cdot \frac{99,66 \text{ kPa} + 700 \text{ kPa} - 0 \text{ kPa}}{101,325 \text{ kPa}} \cdot 1,017 52$$

$$z = 0,947 94 \cdot 7,892 03 \cdot 1,017 52$$

$$z = 7,612 24$$

At a measurement station, a quantity, Q [$V = 1\,000 \text{ m}^3$; $T = 288,15 \text{ K}$; $p_g = 700 \text{ kPa}$ (7,0 bar); $p_{amb} = 99,6 \text{ kPa}$ (0,996 6 bar)], has been measured. Performing the volume conversion to normal conditions using [Formula \(E.1\)](#) yields

$$V_n = 7\,612,24 \text{ m}^3$$

The energy quantity, E , is calculated in accordance with [Formula \(10\)](#) as given below.

$$E = 7\,612,24 \text{ m}^3 \cdot 11,901 \text{ kWh/m}^3$$

$$E = 90\,593,27 \text{ kWh} = 326\,135,77 \text{ MJ}$$

E.2 Calculations using ISO 12213-2

E.2.1 General formulae

The same general formulae and principles as given in [E.1.1](#) hold, except that Z_n/Z can be calculated from the gas analysis, using ISO 12213-2.

E.2.2 Example calculation

The calculations for the energy, E , are carried out as demonstrated using the following parameters, measured at one of the interfaces.

— average of the ambient air pressure at the meter, p_{amb}	99,66 kPa (0,996 6 bar);
— operating pressure (gauge), p_g	5 000 kPa (50,0 bar);
— operating temperature, T	283,15 K;
— concentration of CO ₂ , C_{CO_2}	2,22 mol %;
— concentration of N ₂ , C_{N_2}	0,77 mol %;
— concentration of O ₂ , CO ₂	0,01 mol %;
— concentration of CH ₄ , C_{CH_4}	87,62 mol %;
— concentration of C ₂ H ₆ , $C_{\text{C}_2\text{H}_6}$	8,75 mol %;
— concentration of C ₃ H ₈ , $C_{\text{C}_3\text{H}_8}$	0,53 mol %;
— concentration of <i>i</i> -C ₄ H ₁₀ , $C_{i\text{-C}_4\text{H}_{10}}$	0,03 mol %;
— concentration of <i>n</i> -C ₄ H ₁₀ , $C_{n\text{-C}_4\text{H}_{10}}$	0,04 mol %;
— concentration of <i>i</i> -C ₅ H ₁₂ , $C_{i\text{-C}_5\text{H}_{12}}$	0,01 mol %;
— concentration of <i>n</i> -C ₅ H ₁₂ , $C_{n\text{-C}_5\text{H}_{12}}$	0,01 mol %;
— concentration of C ₆ H ₁₄₊ , $C_{\text{C}_6\text{H}_{14+}}$	0,01 mol %;
— calorific value (calculated from analysis), $H_{\text{S},n}$	11,581 kWh/m ³ ;
— density (calculated from analysis), ρ_n	0,813 3 kg/m ³ ;
— partial pressure of water in natural gas, $p_{\text{H}_2\text{O}}$	<0,1 kPa (<0,001 bar);

NOTE 1 $p_{\text{H}_2\text{O}}$ can be expressed by the product of φ (relative humidity of the gas) and p_{sat} (partial pressure of steam in saturated gases); in dry natural gases, p_{sat} is usually $p_{\text{sat}} \leq 0,1$ kPa (0,001 bar). Thus, for dry natural gases the expression ($p_{\text{H}_2\text{O}} = \varphi \cdot p_{\text{sat}}$) can usually be set to zero.

— Z_n/Z ratio:	1,152 073 7.
------------------	--------------

NOTE 2 This value is calculated using the concentrations CO₂ to C₆H₁₄₊ in ISO 12213-2 {AGA:8-92DC, [Formula \(8\)](#); see Reference [35]}.

The substitution of the measured values into [Formula \(E.2\)](#) results in the following:

$$z = 0,964\,68 \cdot 50,329\,73 \cdot 1,152\,073\,7$$

$$z = 55,935\,58$$

At a measurement station, a quantity, Q [$V = 10\,000\text{ m}^3$; $T = 283,15\text{ K}$; $p_g = 5\,000\text{ kPa}$ (50,0 bar); $p_{\text{amb}} = 99,66\text{ kPa}$ (0,996 6 bar)], has been measured. The volume is converted to V_n under normal conditions using [Formula \(E.1\)](#):

$$V_n = 559\,355,8\text{ m}^3$$

The energy quantity, E , is calculated in accordance with [Formula \(10\)](#):

$$E = 559\,355,8\text{ m}^3 \cdot 41,6916\text{ MJ/m}^3$$

$$E = 23\,320\,438,27\text{ MJ} = 6\,477\,899,52\text{ kWh}$$

STANDARDSISO.COM : Click to view the full PDF of ISO 15112:2018

Annex F
(informative)

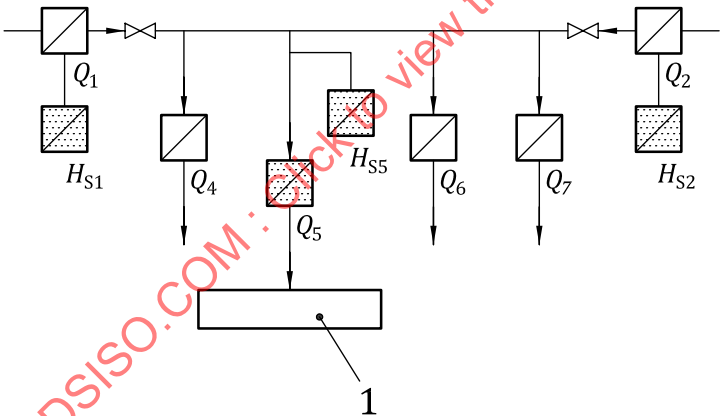
Practical examples for averaging the calorific value due to
different delivery situations

The following examples demonstrate the calculation of the delivered energy for interface 5 (an industrial customer) on the basis of the following:

- arithmetic-average calorific value: see [Figure F.1](#) and [Table F.1](#);
- weighted-average calorific value/fixed assignment: see [Figure F.2](#) and [Tables F.2](#) and [F.3](#);
- weighted-average calorific value/variable assignment: see [Figure F.3](#) and [Tables F.4](#), [F.5](#) and [F.6](#).

NOTE Calorific value correction by means of a correction procedure (see [6.5](#), [12.2](#) and [Annex I](#)) is in use for interfaces 1 to 3 and 5.

[Table F.1](#) shows the calculation of the energy from separate measured calorific values, $H_{S5,n}$, under normal reference conditions and separate gas quantities, $V_{Q5,n}$, under normal reference conditions for an industrial customer at interface 5. Entry-point interfaces 1 and 2 (see [Figure F.1](#)) supply a series of interfaces 4 to 7. H_{S5} , Q_5 , p_5 , T_5 , density and CO₂ concentration are measured at interface 5.



Key
 Q_1 , Q_4 , Q_5 , Q_6 and Q_7 interfaces
1 industrial customer

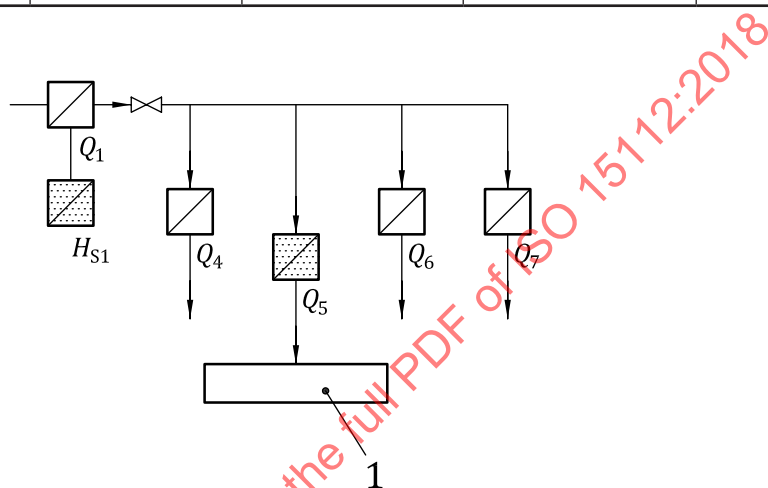
Figure F.1 — System with entry-point interfaces 1 and 2, which subsequently supply a series of interfaces 4 to 7

Table F.1 — Calculation of the energy using separate calorific value measurements at interface 5

Hour	Calorific value $H_{S5,n}$		Gas quantity $V_{Q5,n}$ m ³	Energy E	
	MJ/m ³	kWh/m ³		MJ	kWh
1	39,89	11,08	5 100	203 429	56 508
2	39,82	11,06	4 950	197 089	54 747
3	39,82	11,06	4 880	194 303	53 973

Table F.1 (continued)

Hour	Calorific value $H_{S5,n}$		Gas quantity $V_{Q5,n}$ m ³	Energy E	
	MJ/m ³	kWh/m ³		MJ	kWh
...	...	...	...	...	...
...	...	...	...	...	...
...	...	...	...	...	...
744	39,64	11,01	5 000	217 998	60 555
sum	—	—	3 868 800	153 343 757	42 595 488

**Key** Q_1 , Q_4 , Q_5 , Q_6 and Q_7 interfaces

1 industrial customer

Figure F.2 — System with one entry point at interface 1, which subsequently supplies a series of interfaces 4 to 7**Table F.2 — Data for calculating the quantity-weighted average calorific value used for fixed assignment at interface 5 (industrial customer)**

Hour	Calorific value ^a $H_{S1,n}$		Gas quantity $V_{Q1,n}$ m ³	Energy E	
	MJ/m ³	kWh/m ³		MJ	kWh
1	39,89	11,08	101 000	4 028 688	1 119 080
2	39,82	11,06	105 000	4 180 680	1 161 300
3	39,82	11,06	107 000	4 260 312	1 183 420
...	...	...	...	...	...
...	...	...	...	...	...
744	39,64	11,01	98 000	3 884 328	1 078 980
sum	—	—	72 912 785	2 895 220 864,8	804 228 018

^a The monthly quantity-weighted average calorific value, $H_{S1,n}$, is calculated from hourly data at the entry point as follows:

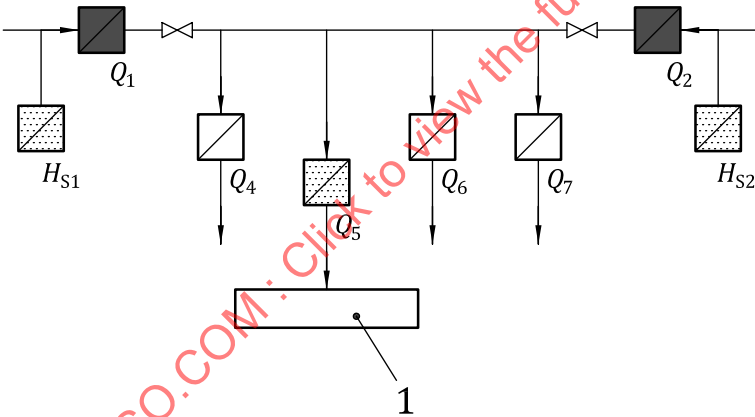
$$\begin{aligned}
 H_{S1,n} &= E/V_{Q1,n} = 2\,895\,220\,844,8 \text{ MJ} / 72\,912\,785 \text{ m}^3 \\
 &= 39,71 \text{ MJ/m}^3 \\
 &= 11,03 \text{ kWh/m}^3
 \end{aligned}$$

Table F.3 — Calculation of the energy at interface 5, using a quantity-weighted average calorific value for fixed assignment

Hour	Quantity-weighted average calorific value		Gas quantity $V_{Q5,n}$ m ³	Monthly energy	
	MJ/m ³	kWh/m ³		MJ	kWh
1	—	—	5 100	—	—
2	—	—	4 950	—	—
3	—	—	4 880	—	—
...	—	—	...	—	—
...	—	—	...	—	—
744	—	—	5 500	—	—
sum	—	—	3 809 280	—	—
—	39,71	11,03	—	151 258 888,8	42 016 358

Figure F.3 shows a gas-transportation system with two entry points, interfaces 1 and 2, which supply a number of interfaces 4 to 7 in a bidirectional gas flow. The change in the gas quality is assumed to be as in Figure B.2.

An industrial customer is supplied at interface 5, where Q_5 , p_5 and T_5 are measured. The applicable calorific value, $H_{S1,n}$, (see Table F.6) for energy determination is assigned on the basis of averaged $H_{S1,n}$ at interface 1 (see Table F.4) and averaged $H_{S2,n}$ at interface 2 (see Table F.5).



Key
 Q_1 , Q_4 , Q_5 , Q_6 and Q_7 interfaces
1 industrial customer

Figure F.3 — System with two entry points, interfaces 1 and 2, which supply interfaces 4 to 7 from two directions

Table F.4 — Data for calculating the monthly quantity-weighted average calorific value at interface 1

Day	Calorific value ^a $H_{S1,n}$		Gas quantity $V_{Q1,n}$ m ³	Energy E	
	MJ/m ³	kWh/m ³		MJ	kWh
1	39,82	11,08	1 150 251	45 881 208	12 744 781
2	39,82	11,06	1 200 500	47 799 108	13 277 530
...	...	...	...	...	...
31	39,64	11,01	1 080 500	42 826 698	11 896 305
sum	—	—	37 747 354	1 497 513 027,6	415 975 841

^a The monthly quantity-weighted average calorific value, $H_{S1,n}$, is calculated from daily measured data at interface 1 as follows:

$$H_{S1,n} = E/V_{Q1,n} = 1\,497\,513\,027,6 \text{ MJ} / 37\,747\,354 \text{ m}^3$$

$$= 39,672 \text{ MJ/m}^3$$

$$= 11,02 \text{ kWh/m}^3$$
Table F.5 — Data for calculating the monthly quantity-weighted average calorific value at interface 2

Day	Calorific value ^a $H_{S2,n}$		Gas quantity $V_{Q2,n}$ m ³	Energy E	
	MJ/m ³	kWh/m ³		MJ	kWh
1	38,88	10,80	600 500	23 347 440	6 485 400
2	39,528	10,98	580 540	22 947 584	6 374 329
...	...	...	...	...	...
31	39,564	10,99	520 000	20 573 280	5 714 800
sum	—	—	17 577 413	692 660 160	192 405 600

^a The monthly quantity-weighted average calorific value, $H_{S2,n}$, is calculated from daily measured data at interface 2 as follows:

$$H_{S2,n} = E/V_{Q2,n} = 192\,405\,600 \text{ kWh} / 17\,577\,413 \text{ m}^3$$

$$= 10,95 \text{ kWh/m}^3$$

$$= 39,420 \text{ MJ/m}^3$$
Table F.6 — Calculation of the energy for interface 5 using variable assignment of a monthly quantity-weighted average calorific value

Day	Calorific value ^a $H_{S1+S2,n}$		Gas quantity $V_{Q5,n}$ m ³	Monthly energy E	
	MJ/m ³	kWh/m ³		MJ	kWh
1	—	—	5 100	—	—
2	—	—	4 950	—	—
...	—	—	...	—	—

^a The calculation is valid on the condition that the difference between H_{S1} and H_{S2} is less than $\pm 2\%$. Then, the quantity-weighted monthly average calorific value, $H_{S1+S2,n}$, for interfaces 4 to 7 is calculated as the sum of the total energies at interfaces 1 and 2 divided by the sum of the total volumes at interfaces 1 and 2.

$$H_{S1+S2,n} = \frac{(415\,975\,841 + 192\,405\,600) \text{ kWh}}{(37\,747\,354 + 17\,577\,413) \text{ m}^3}$$

$$= 11,00 \text{ kWh/m}^3$$

$$= 39,600 \text{ MJ/m}^3$$

Table F.6 (continued)

Day	Calorific value ^a $H_{S1+S2,n}$		Gas quantity $V_{Q5,n}$ m ³	Monthly energy E	
	MJ/m ³	kWh/m ³		MJ	kWh
31	—	—	5 000	—	—
sum	—	—	3 809 280	—	—
—	39,600	11,00	—	150 847 488	41 902 080
^a The calculation is valid on the condition that the difference between H_{S1} and H_{S2} is less than $\pm 2\%$. Then, the quantity-weighted monthly average calorific value, $H_{S1+S2,c}$, for interfaces 4 to 7 is calculated as the sum of the total energies at interfaces 1 and 2 divided by the sum of the total volumes at interfaces 1 and 2.					
$H_{S1+S2,n} = \frac{(415\,975\,841 + 192\,405\,600) \text{ kWh}}{(37\,747\,354 + 17\,577\,413) \text{ m}^3}$ $= 11,00 \text{ kWh/m}^3$ $= 39,600 \text{ MJ/m}^3$					