
Sampling airborne radioactive materials from the stacks and ducts of nuclear facilities

*Échantillonnage de substances radioactives en suspension dans l'air
dans les émissaires de rejet et les conduits des installations nucléaires*

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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 document 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).

ISO draws attention to the possibility that the implementation of this document may involve the use of (a) patent(s). ISO takes no position concerning the evidence, validity or applicability of any claimed patent rights in respect thereof. As of the date of publication of this document, ISO had not received notice of (a) patent(s) which may be required to implement this document. However, implementers are cautioned that this may not represent the latest information, which may be obtained from the patent database available at www.iso.org/patents. ISO shall not be held responsible for identifying any or all such patent rights.

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

For an explanation of 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 www.iso.org/iso/foreword.html.

This document was prepared by Technical Committee ISO/TC 85, *Nuclear energy, nuclear technologies and radiological protection*, Subcommittee SC 2, *Radiological protection*.

This fourth edition cancels and replaces the third edition (ISO 2889:2021), of which it constitutes a minor revision.

The main changes are:

- clarification of the circumstances where numerical modelling may be used to perform or assist with meeting the qualifications for sample extraction locations;
- clarification of passages allowing the use of alternate aerosol particle sizes for the purpose of testing to meet various performance criteria described in this document;
- changes for the discussion of standard uncertainty with regard to setting action levels ([Annex I](#)).

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

This document focuses on monitoring the activity concentrations and activity releases of radioactive substances in air in stacks and ducts. Other situations for monitoring the activity concentrations and activity releases of radioactive substances in air (environmental or workplace monitoring) are being addressed in subsequent standards. This document provides performance-based criteria for the use of air-sampling equipment, including probes, transport lines, sample collectors, sample monitoring instruments and gas flow measuring methods. This document also provides information covering sampling programme objectives, quality assurance, development of air monitoring control action levels, system optimization and system performance verification.

ISO 2889 was first published in 1975 as a guide to sampling airborne radioactive materials in the ducts, stacks, and working environments of installations where work with radioactive materials is conducted. Since then, an improved technical basis has been developed for each of the major sampling specialities. The focus of this document is on the sampling of airborne radioactive materials in ducts and stacks.

The goal of achieving an unbiased, representative sample is best accomplished where samples are extracted from airstreams in which potential airborne contaminants are well mixed in the airstream. This document sets forth performance criteria and recommendations to assist in obtaining valid measurements of the concentration of airborne radioactive materials in ducts or stacks.

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Sampling airborne radioactive materials from the stacks and ducts of nuclear facilities

1 Scope

This document sets forth performance-based criteria and recommendations for the design and use of systems for sampling of airborne radioactive materials in the effluent air from the ducts and stacks of nuclear facilities.

The requirements and recommendations of this document are aimed at sampling that is conducted for regulatory compliance and system control. If existing air-sampling systems are not designed to the performance requirements and recommendations of this document, an evaluation of the performance of the system is advised. If deficiencies are discovered, a determination of whether or not a retrofit is needed and practicable is recommended.

It can be impossible to meet the requirements of this document in all conditions with a sampling system designed for normal operations only. Under off-normal conditions, the criteria or recommendations of this document still apply. However, for accident conditions, special accident air sampling systems or measurements can be used.

This document does not address outdoor air sampling, radon measurements, or the surveillance of airborne radioactive substances in the workplace of nuclear facilities.

NOTE Reference [1] addresses the instrumentation that is frequently used in nuclear air monitoring. Reference [5] addresses air sampling in the workplace of nuclear facilities. References [6] and [7] describe the performance characteristics of air monitors.

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 10780, *Stationary source emissions — Measurement of velocity and volume flowrate of gas streams in ducts*

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

abatement equipment

apparatus used to reduce contaminant concentration in the airflow exhausted through a stack or duct

3.2

absorbent

material that takes up a constituent through the action of diffusion, allowing the constituent to penetrate into the structure of the absorbent (if a solid) or dissolve in it (if a liquid)

Note 1 to entry: When a chemical reaction takes place during absorption, the process is called chemisorption.

3.3

accident conditions

any unintended event, including operating errors, equipment failures and other mishaps, the consequences or potential consequences of which are not negligible from the point of view of protection and safety

3.4

accuracy

closeness of agreement between a measured quantity and the true quantity of the measurand

3.5

action level

threshold concentration of an effluent contaminant at which it is necessary to perform an appropriate action

3.6

adsorbent

material, generally a solid, that retains a substance contacting it through short-range molecular forces that bind the adsorbed material at the surface of the material

3.7

aerodynamic diameter

D_a
for a particle of arbitrary shape and density, the diameter of a sphere with density 1 000 kg/m³ that has the same sedimentation velocity in quiescent air as the arbitrary particle

3.8

aerosol

dispersion of solid or liquid particles in air or other gas

Note 1 to entry: An aerosol is not only the *aerosol particles* (3.9).

3.9

aerosol particle

solid or liquid particle constituents of an *aerosol* (3.8)

3.10

analyser

device that provides for near real-time data on radiological characteristics of the gas (air) flow in a sampling system or duct

Note 1 to entry: An analyser usually evaluates the concentration of radionuclides in a sampled air stream. However, some analysers are mounted directly in or outside a stack or duct.

3.11

aspiration ratio

ratio of particle mass or number concentration in the nozzle inlet to the concentration in the free stream

3.12

bend

gradual change in direction of a sample transport line

Note 1 to entry: The radius of curvature of a bend should be at least three times the inside diameter of the tubing.

3.13**bulk stream**

air flow in a stack or duct, as opposed to the sample flow rate

3.14**burial**

imbedding of a particle into a filter medium or the masking of a particle by subsequent deposits of particulate matter

3.15**calibration**

operation that, under specified conditions, initially establishes a relation between the quantity values with measurement uncertainties provided by measurement standards and corresponding indications with associated measurement uncertainties and then uses this information to establish a relation for obtaining a measurement result from an indication

3.16**coefficient of variation**

C_v

quantity that is the ratio of the standard deviation of a variable to the mean value of that variable

Note 1 to entry: It is usually expressed as a percentage.

3.17**collector**

component of a sampling system that is used to retain radionuclides for analysis

EXAMPLE A filter that is used to remove from a sample stream *aerosol particles* (3.9) that carry alpha-emitting transuranic radionuclides or other radionuclides.

3.18**conditioning system**

apparatus that can be used to purposefully, in a controlled manner, change the *aerosol particle* (3.9) concentration, gas composition, *particle-size distribution* (3.52), temperature or pressure in a *sample stream* (3.68)

3.19**continuous air monitor****CAM**

near-real-time sampler and associated detector that provide data on radionuclides [e.g. concentration of alpha-emitting *aerosol particle* (3.9)] in a *sample stream* (3.68)

3.20**continuous monitoring**

continuous near-real-time measurements of one or more sampling characteristics

3.21**continuous sampling**

either uninterrupted sampling or sequential collection of samples obtained automatically at intervals short enough to yield results that are representative for the entire sampling period

Note 1 to entry: The sample may be analysed in near-real-time (i.e. equivalent to monitoring) or it may be analysed post-sample-collection in a remote laboratory.

3.22**curvature ratio**

ratio of bend radius to the tube diameter

3.23

depositional loss

loss of constituents of the sample on the internal walls of a sampling system

Note 1 to entry: See also [3.84](#).

3.24

decision threshold

value of the estimator of the measurand, which, when exceeded by the result of an actual measurement using a given measurement procedure of a measurand quantifying a physical effect, is used to decide that the physical effect is present

Note 1 to entry: The decision threshold is defined such that in cases where the measurement result exceeds the decision threshold, the probability that the true value of the measurand is zero is less or equal to a chosen probability for a wrong decision, α .

[SOURCE: ISO 11929-1:2019, 3.12 modified – definition identical, but Note 1 to entry changed and Note 2 to entry not included here.]

3.25

detection limit

smallest true value of the measurand which ensures a specified probability of being detectable by the measurement procedure

Note 1 to entry: With the *decision threshold* ([3.24](#)), the detection limit is the smallest true value of the measurand for which the probability of wrongly deciding that the true value of the measurand is zero is equal to a specified value, when, in fact, the true value of the measurand is not zero.

[SOURCE: ISO 11929-1: 2019, 3.13 modified – definition identical, but last sentence of Note 1 to entry not included here as well as Note 2 to entry.]

3.26

droplet

liquid *aerosol particle* ([3.9](#))

3.27

effective dose

sum of the products of the dose absorbed by an organ or a tissue and the factors relative to the radiation and to the organs or tissues that are irradiated

3.28

effluent

waste stream flowing away from a process, plant, or facility to the environment

Note 1 to entry: This document applies to the effluent air that is discharged to the atmosphere through stacks and ducts.

3.29

emission

contaminants that are discharged into the environment

3.30

emit

discharge contaminants into the environment

3.31

extractive sampling

diverting a part of the airflow from a stack or duct for the purpose of the collection of a sample of the air

Note 1 to entry: See [3.69](#) and [3.72](#).

3.32**flow rate**

rate at which a mass or volume of gas (air) crosses an imaginary cross-sectional area in either a sampling system tube or a stack or duct

Note 1 to entry: The rate at which the volume crosses the imaginary area is called the volumetric flow rate and the rate at which the mass crosses the imaginary area is called either the mass flow rate or the volumetric flow rate at standard conditions.

3.33**geometric mean (of a variable)**

x_g
value for N observations of a random variable x_i given by

$$\ln x_g = \frac{1}{N} \sum_{i=1}^N \ln x_i$$

3.34**geometric standard deviation**

s_g
the geometric standard deviation for N observations of a random variable, x_i , calculated from

$$\ln^2 s_g = \frac{1}{N-1} \sum_{i=1}^N (\ln x_i - \ln x_g)^2$$

where x_g is the geometric mean of the random variable

3.35**high-efficiency particulate air filter****HEPA filter**

high-efficiency filter used for removing *aerosol particles* (3.9) from an air stream

Note 1 to entry: A HEPA filter usually collects aerosol particles at the most penetrating particle size (between 0,1 µm and 0,3 µm diameter) with a high efficiency and is designed to collect greater fractions of aerosol particles with diameters either larger or smaller. The minimum efficiency of a HEPA filter is not defined in this document.

3.36**hydraulic diameter**

type of equivalent duct diameter for ducts that do not have a round cross-section

Note 1 to entry: Generally, it is four times the cross-sectional area divided by the perimeter.

3.37**impaction**

process by which *aerosol particles* (3.9) are removed from an air stream by striking an object in the air stream

Note 1 to entry: Curvature of air streamlines, principally on the front side of the object, causes particles with sufficient inertia to strike the object while the airflow passes around it.

3.38**in-line system**

system where the detector assembly is adjacent to, or immersed in, the *effluent* (3.28) stream or stream in the duct or stack

3.39**interception**

process by which *aerosol particles* (3.9) are removed from an air stream by an object in the flow, where the trajectory of the particle's centre of gravity misses the object but the body of the particle strikes the object

3.40

isokinetic

condition that prevails when the velocity of air at the inlet plane of a nozzle is equal to the velocity of undisturbed air in a stack or duct at the point where the nozzle inlet is located

Note 1 to entry: Anisokinetic is the antonym of isokinetic. Sub-isokinetic refers to the condition where the nozzle inlet velocity is less than the free-stream velocity. Super-isokinetic refers to the condition where the nozzle inlet velocity is greater than the free-stream velocity.

3.41

laminar flow

flow regime in stacks or ducts associated with Reynolds numbers less than about 2 200

Note 1 to entry: This regime is not usually encountered in effluent air flows. Mixing in laminar flow results from molecular diffusion, which is a much slower process than mixing in turbulent flow.

3.42

membrane filter

filter medium consisting of thin, organic-based films having a range of selectable porosities and controlled composition

Note 1 to entry: Thin, porous metallic filters are sometimes also called membrane filters.

3.43

mixing element

device placed in a stack or duct that is used to augment the mixing of the contaminant mass with the fluid

3.44

monitoring

continual measurement of a quantity (e.g. activity concentration) of the airborne radioactive constituent or the gross content of radioactive material, at a frequency that permits an evaluation of the value of that quantity in near-real-time, or at intervals that comply with regulatory requirements

3.45

monodisperse aerosol

aerosol (3.8) comprised of (solid or liquid) particles that are all of approximately the same size

Note 1 to entry: In general, the geometric standard deviation of the particle-size distribution of a monodisperse aerosol is less than or equal to 1,1.

3.46

nozzle

device used to extract a sample from an *effluent* (3.28) stream and transfer the sample to a transport line or collection device

Note 1 to entry: Within the nozzle, there is a transition zone where the sample stream adjusts to the conditions in the transport line.

3.47

nozzle exit plane

imaginary plane across the cross-section of a transport system that divides the nozzle region from the transport line

Note 1 to entry: The nozzle is frequently a separate component and the nozzle exit plane is clearly defined as the downstream end of that component. If there is no separate component, the nozzle exit is the end of the transition zone of the nozzle flow.

3.48**nozzle inlet**

imaginary cross-sectional inlet plane of a nozzle where the flow first enters the transport system

Note 1 to entry: In the special case of a shrouded nozzle, the inlet is referenced to the inner nozzle rather than the shroud.

3.49**number size distribution**

representation of the number of particles associated with intervals of particle size, over the full size range encountered in a sample

Note 1 to entry: For samples consisting of *aerosol particles* (3.9), it is a representation of the relative number of particles (measured number of particles in a size interval divided by the total number of particles in the sample) associated with intervals of aerodynamic diameter.

3.50**off-normal condition**

condition that is unplanned and which presents a gap with normal conditions

EXAMPLE Accidents and equipment failure.

3.51**particle**

aggregate of molecules, forming a solid or liquid, ranging in size from a few molecular diameters to several millimetres

3.52**particle-size distribution**

distribution of *particle* (3.51) size as a function of mass or activity rather than number

3.53**penetration**

ratio of the concentration at the outlet of the sampling system, transport lines included, to that in the duct or at the stack

3.54**polydisperse aerosol**

aerosol (3.8) comprised of particles with a range of sizes

Note 1 to entry: In general, the geometric standard deviation of the particle-size distribution of a polydisperse aerosol is greater than 1.1.

3.55**potential emission**

radionuclides that can be released to the environment from a facility in the absence of control equipment

3.56**precision**

closeness of agreement between indications obtained by replicate measurements on the same or similar objects under specified conditions.

Note 1 to entry: A value of precision is obtained by repetitive testing of a homogenous sample under specified conditions. The precision of a method is expressed quantitatively as either the standard deviation computed from the results of a series of controlled determinations or as the coefficient of variation of the measurements.

3.57**probe**

tubing or apparatus inserted into a stack or duct through which a sample of the stream is withdrawn

Note 1 to entry: A probe usually refers to one or more nozzles and part of the transport line.

3.58

profile

distribution of air velocity, of gas concentration or of *particle* (3.51) concentration over the cross-sectional area of the stack or duct

3.59

quality assurance

QA

planned and systematic actions necessary to provide confidence that a system or component performs satisfactorily in service and that the results are both correct and traceable

3.60

radionuclide

unstable isotope of an element that decays or converts spontaneously into another isotope or different energy state, emitting radiation

3.61

record sample

sample that is collected for reporting purposes

Note 1 to entry: Record samples are often analysed off-line.

3.62

reference method

apparatus and instructions for providing results against which other approaches may be compared

Note 1 to entry: The application of a reference method is assumed to define correct results.

3.63

representative sample

sample with the same quality and characteristics for the material of interest as that of its source at the time of sampling

3.64

response time

time required after a step variation in the measured quantity for the output signal variation to reach a given percentage for the first time, usually 90 %, of its final value

3.65

sample

portion of an air stream of interest or one or more separated constituents from a portion of an air stream

3.66

sample-extraction location

location in a stack or duct that coincides with the *sample* (3.65) *nozzle inlet* (3.48)

Note 1 to entry: By extension from the nozzle inlet, the entire plane that is perpendicular to the longitudinal axis of a stack or duct.

3.67

sampler

device that collects or analyses constituents of the air *sample* (3.65)

3.68

sample stream

air that flows through a sampling system

3.69

sampling

process of removing a *sample* (3.65) from the free air and transporting it to a *collector* (3.17) or an *analyser* (3.10) (monitor)

3.70**sampling environment**

conditions of the air flow and gas within a stack or duct that can influence the sampling process

Note 1 to entry: Factors to take into account include pressure, temperature and molecular composition of the gas.

3.71**sampling plane**

cross-sectional area where the *sample* (3.65) is extracted from the air flow

3.72**sampling system**

system consisting of a nozzle, an inlet, a transport line, a flow *conditioning system* (3.18) and a *collector* (3.17) or monitor

Note 1 to entry: A flow conditioning system may be used to change concentration, temperature, humidity, or other characteristics. Depending upon the application, a flow conditioner might not be used in the sampling system.

3.73**sedimentation velocity**

terminal (maximum) velocity an *aerosol particle* (3.9) attains in quiescent fluid (air) as a result of the gravitational force

3.74**sensitivity**

change in indication of a mechanical, nuclear, optical or electronic instrument as affected by changes in the variable quantity being sensed by the instrument

Note 1 to entry: This is the slope of a calibration curve of an instrument, where a calibration curve shows output values of an instrument as a function of input values.

3.75**shroud**

aerodynamic decelerator placed around and extending beyond a sampling nozzle to reduce sampling biases

3.76**standard conditions**

temperature of 298 K (25 °C) and a pressure of 101,325 kPa

Note 1 to entry: Used to convert air densities to a common basis. Other temperature and pressure conditions may be used but should be applied consistently.

3.77**transmission ratio**

ratio of the *aerosol particle* (3.9) concentration at the nozzle outlet to that in the free stream

Note 1 to entry: It is stated whether a mass or activity basis is used.

3.78**transport line**

part of a transport system between the *nozzle exit plane* (3.47) and the entrance plane of a *collector* (3.17) or *analyser* (3.10)

3.79**transport system**

all components of a *sampling system* (3.72), excluding the *collector* (3.17) or *analyser* (3.10)

3.80

turbulent flow

flow regime characterized by bulk mixing of fluid properties

Note 1 to entry: For example, in a tube, the flow is turbulent if the Reynolds number is greater than about 3 000 and laminar if the Reynolds number is below about 2 200. There is little mixing in the laminar flow regime.

3.81

uncertainty

parameter characterizing the dispersion of the value of a measurand, based on the true value of a quantity

Note 1 to entry: The uncertainty is typically stated at a given statistical level of confidence (e.g. 95 %).

3.82

uncertainty analysis

procedure for estimating the overall impact on the *accuracy* (3.4) or *precision* (3.56) of a dependent variable as a result of the estimated uncertainties of the independent variables

3.83

vapour

gaseous form of materials that are liquids or solids at room temperature, as distinguished from non-condensable gases

Note 1 to entry: Vapours are gases but carry the connotation of having been released or volatilized from liquids or solids.

3.84

velocity profile

distribution of the velocity values at a given cross-section in a stack or duct

3.85

volatile

having a high *vapour* (3.83) pressure, which allows significant quantities of material to become gaseous at the prevailing temperature

Note 1 to entry: In this document, the stack or duct temperature is generally considered as the reference.

3.86

wall loss

loss of *sample* (3.65) constituents to the internal walls of a *sampling system* (3.72)

Note 1 to entry: Quantitatively, it is the equivalent concentration lost to the walls of a nozzle, transport line, conditioning system, or transport system divided by the concentration at the inlet plane of the nozzle, transport line, or transport system.

4 Symbols

For the purposes of this document, the following symbols apply.

A	cross-sectional area of a stack or duct, in m^2
$\bar{A}$	average radionuclide stack or duct emission rate over the period of integration, in $\text{Bq}\cdot\text{s}^{-1}$
A_i	cross-sectional area of the i^{th} element, in m^2
C	Cunningham's slip correction for aerosol particles, dimensionless
$C_{\text{cal,pt}}$	Pitot calibration factor, dimensionless

$C_{v,pt}$	velocity-averaging correction factor for a Pitot tube, dimensionless
$C_{v,ta}$	velocity-averaging correction factor for a single point thermal anemometer, dimensionless
$C_{v,af}$	velocity-averaging correction factor for a line average velocity taken with an acoustic flow meter, dimensionless
c_A	effluent activity concentration, in $Bq \cdot m^{-3}$
c_e	activity or mass or aerosol particle concentration at the exit plane of a transport system, in $Bq \cdot m^{-3}$ or $kg \cdot m^{-3}$ or m^{-3} , respectively
$c_{e,j}$	activity or mass or aerosol particle concentration at the exit plane of a component j , in $Bq \cdot m^{-3}$ or $kg \cdot m^{-3}$ or m^{-3} , respectively
$c_{i,j}$	activity or mass or aerosol particle concentration at the inlet plane of a transport system component j , in $Bq \cdot m^{-3}$ or $kg \cdot m^{-3}$ or m^{-3} , respectively
c_∞	activity or mass or aerosol particle concentration in the undisturbed free stream at the nozzle location, in $Bq \cdot m^{-3}$ or $kg \cdot m^{-3}$ or m^{-3} , respectively
c^*	decision threshold of the activity concentration, in $Bq \cdot m^{-3}$
$c^\#$	detection limit of the activity concentration, in $Bq \cdot m^{-3}$
D_a	aerodynamic particle diameter, in μm
De	Dean number of a flow bend, $De = Re/f_{cu}^{1/2}$, dimensionless
d_t	inside diameter of a transport system component (e.g. tube), in m
F_k	fluctuation constant, dimensionless
f_c	ratio of the activity concentration in the sample volume to the effluent activity concentration in the free stream, dimensionless
f_{cu}	curvature ratio ($f_{cu} = r_{cu} \cdot d_t^{-1}$), dimensionless
L	length of a section of tubing, in m
l_w	wall losses of aerosol particles in transport system components, dimensionless
M	mean molar mass of a gas, in $kg \cdot mol^{-1}$
M_p	mixing of radioactive contaminant in the total effluent gas volume, determined as the ratio of the concentration in the sample volume to the concentration in the free stream, dimensionless
N	number of points or observations
n	number of components in a transport system, dimensionless
P	overall penetration of a sample through a transport system, dimensionless
P_{bend}	penetration of a sample through a bend in a tube transport system, dimensionless
P_j	penetration of a sample through the j^{th} component of a transport system, dimensionless
P_s	penetration through a straight tube, dimensionless
p_a	pressure in the stack or duct, in Pa

p_{std}	standard pressure, equal to 101,325 kPa
q_a	volumetric flow rate at actual temperature and pressure conditions, in $\text{m}^3 \cdot \text{s}^{-1}$
q_{std}	volumetric flow rate at standard conditions, in $\text{m}^3 \cdot \text{s}^{-1}$
R	individual gas constant for a particular gas, equal to $R_u \cdot M^{-1}$, in $\text{J} \cdot \text{kg}^{-1} \cdot \text{K}^{-1}$
R_u	universal gas constant, equal to $8,314 \text{ J} \cdot \text{kg}^{-1} \cdot \text{K}^{-1}$
Ra	surface roughness, in μm
Re	Reynolds number of flow in a tube, equal to $\rho \cdot U_m \cdot d_t / \mu$, dimensionless
r	resuspension rate, in s^{-1}
r_{cu}	radius of curvature of a pipe bend, in m
r_n	net count rate (gross minus background) of the sample, in s^{-1}
S	signal
St	Stokes number, equal to $(C \cdot \rho_w \cdot D_a^2 \cdot U_m) / (9 \mu \cdot d_t)$, dimensionless
s	standard deviation
T	temperature, in K
T_a	temperature in stack or duct, in K
T_{std}	standard temperature, equal to 298 K (25 °C)
t	time, in s
t_s	time period over which sampling is performed, in s
u	standard uncertainty
$u(\tilde{A})$	standard uncertainty of the activity emission rate of a radionuclide, in $\text{Bq} \cdot \text{s}^{-1}$
$u(\text{Cal})$	uncertainty associated with determining the calibration factor, dimensionless
$u(F_k)$	standard uncertainty of the fluctuation constant, which is set at 1 for a meter whose readings do not fluctuate, dimensionless
$u(r_n)$	standard uncertainty of the net count rate, in s^{-1}
$u(t_s)$	standard uncertainty associated with the measurement of the sampling time, dimensionless
$u(V)$	total uncertainty in the volume of air, dimensionless
$u(V_T)$	standard uncertainty of the total volume sampled, in m^3
$u(w)$	standard uncertainty of the calibration factor, in s^{-1}
u_C	uncertainty associated with determining the calibration factor, i.e. correcting the indicated flow, dimensionless
$u_r(A)$	relative standard uncertainty of the cross-sectional area, dimensionless
$u_r(\tilde{A})$	relative standard uncertainty of the activity emission rate of a radionuclide, dimensionless

$u_{r,k=2}(\bar{A})$	relative standard uncertainty of the activity emission rate of a radionuclide representing the coverage interval at 95 % ($k = 2$), dimensionless
$u_r(C_{\text{cal,pt}})$	relative standard uncertainty of the Pitot calibration factor, dimensionless
$u_r(f_c)$	relative standard uncertainty of the ratio of the activity concentration in the sample volume to the effluent activity concentration in the free stream, dimensionless
$u_r(P)$	relative standard uncertainty of the overall penetration in a transport system, dimensionless
$u_r(p)$	relative standard uncertainty of the pressure, dimensionless
$u_r(V_a)$	relative standard uncertainty of the volume at actual conditions, dimensionless
$u_r(w)$	relative standard uncertainty of the calibration factor, dimensionless
$u_r(\varepsilon_f)$	relative standard uncertainty of the collection efficiency of the collection medium, dimensionless
$u_r(\varepsilon_d)$	relative standard uncertainty of the detection efficiency, dimensionless
$u_r(\rho)$	relative standard uncertainty of the gas density, dimensionless
u_s	uncertainty in reading the flow meter scale, dimensionless
V_a	volume of effluent that produced the sample at stream temperature, pressure, and gas composition, in m^3
V_i	velocity at the midpoint of the i^{th} element, in $\text{m}\cdot\text{s}^{-1}$
V_{std}	equivalent velocity at standard conditions, in $\text{m}\cdot\text{s}^{-1}$
V_T	total volume of gas (air) sampled, in m^3
v	velocity, in $\text{m}\cdot\text{s}^{-1}$
v_{af}	line-average velocity obtained from an acoustic flow meter, in $\text{m}\cdot\text{s}^{-1}$
v_{pt}	velocity at actual conditions determined from use of the reference Pitot tube, in $\text{m}\cdot\text{s}^{-1}$
v_d	deposition velocity due to Brownian diffusion or turbulent inertial deposition on the wall of a transport tube, in $\text{m}\cdot\text{s}^{-1}$
v_e	effective deposition velocity of contaminant at the wall of a transport tube, in $\text{m}\cdot\text{s}^{-1}$
v_g	sedimentation velocity of an aerosol particle, in $\text{m}\cdot\text{s}^{-1}$
v_{ge}	cross-stream component of gravitational settling velocity, in $\text{m}\cdot\text{s}^{-1}$
v_m	mean velocity of gas (air) in a flow tube, in $\text{m}\cdot\text{s}^{-1}$
$v_{\text{std,ta}}$	velocity obtained from a single-point thermal anemometer at standard conditions, in $\text{m}\cdot\text{s}^{-1}$
$v_{\text{std,ta},i}$	velocity measured with a properly calibrated thermal anemometer at the centroid of the i^{th} element of area, in $\text{m}\cdot\text{s}^{-1}$
w	calibration factor, dimensionless
α	angular coordinate of a tube cross-section, in degrees or radians

ε_a	aspiration efficiency of a sampling nozzle, dimensionless
ε_d	detection efficiency, in $\text{Bq}^{-1}\cdot\text{s}^{-1}$
ε_f	collection efficiency, dimensionless
ε_{Ts}	transport efficiency, dimensionless
θ	flow angle, in degrees or rad
λ	decay constant, in s^{-1}
μ	dynamic viscosity of a gas, in $\text{Pa}\cdot\text{s}$
ρ	gas density in the stack or duct, $\text{kg}\cdot\text{m}^{-3}$
ρ_a	gas density in the stack or duct at actual conditions, in $\text{kg}\cdot\text{m}^{-3}$
ρ_{std}	density of air at standard conditions, equal to $1\,184\,\text{kg}\cdot\text{m}^{-3}$
ρ_w	density of water at $4\,^{\circ}\text{C}$, equal to $1\,000\,\text{kg}\cdot\text{m}^{-3}$
σ	standard deviation
τ	transmission ratio of a nozzle, dimensionless
τ_L	transmission through the transport line, dimensionless
τ_p	transmission through the nozzle, dimensionless
ϕ	angle of inclination of a tube axis relative to horizontal, in degrees or rad

5 Factors impacting the sampling program

This document focuses on the mechanics of obtaining a sample of airborne radioactive constituents in facility ducts and stacks. However, there are important factors that impact the design of sampling systems, in particular the following:

- the purpose of sampling;
- the type of conditions (normal or off-normal conditions);
- the characteristics of the air stream and radioactive constituents;
- the desired measurement sensitivity;
- the concentrations or total emissions which trigger remedial action (action levels).

The impact of these factors on the sampling system should be assessed. Informative guidance concerning the first three of these factors is given in [Annexes G](#) and [L](#). Information relevant to the last two is given in [Annex I](#).

For off-normal conditions, the performance of the sampling system can be affected by the modification of several parameters (temperature, flow rate in ducts or stacks, type of airborne particles). Thus, acceptance criteria introduced in this document for normal conditions should be considered as recommendations for off-normal conditions.

6 Sample extraction locations

6.1 General

The sample extraction location shall provide the possibility to extract a representative sample.

A representative sample is best extracted from a location where the radioactive materials of interest are well mixed within the free stream. The term “well-mixed” includes the several criteria that are given in 6.3. In a well-mixed stream the sampling probe may contain a single nozzle. In circumstances where the well-mixed criteria are not achieved, a multi-nozzle probe may be necessary to get a representative sample. The design and operation of sampling probes are described in 7.3 and Annex M. Tools to determine if the criteria for well-mixed sample extraction locations are described in Annex F. These tools include

- physical testing on the actual ventilation stack,
- physical tests with scale models or geometrically similar stacks, and
- numerical methods validated against physical test data.

6.2 General requirements for sample extraction locations

The stack or duct geometry and the airflow within should be fully understood. The sample extraction probe should be located where the potential radioactive constituents are well mixed with the airflow. To meet the recommendations of the following paragraphs the accommodation for effluent sampling is addressed early in the design of the ventilation system for discharging the airborne effluent.

Usually, for a stack, the sample extraction location should be situated between the discharge of the fan(s) and the discharge to the atmosphere, with the provision that the location should not be so close to the final exit that wind effects can significantly influence the velocity profile at the sampling location. Typically, in a well-mixed airflow, successful sample probe locations are in the range of 5 to 10 hydraulic diameters downstream of a flow disturbance and 3 or more hydraulic diameters upstream of a flow disturbance. There can be instances where greater distances are needed.

Particular attention should be given to the geometry of flow-entry conditions. Any addition of a small secondary air stream close to the wall of the stack or duct should be avoided. Bends, fans, duct junctions and similar disturbances promote mixing, but can also produce distortions in the velocity and the contaminant concentration profile and angularity in the airflow in the first 2 to 3 hydraulic diameters downstream. Therefore sampling locations too close to such disturbances should be avoided, even at the cost of longer sampling lines.

NOTE 1 Features that enhance mixing do so by creating large-scale turbulence. One or more 90° turns, converging airstreams, mixing boxes, perimeter rings and commercial static mixers all enhance mixing. On the other hand, turning vanes and flow straighteners have the opposite effect. The generic tests of References [47], [74], [77] and [96] provide tests of features that promote mixing. These are also summarized in Chapter 15 of Reference [76]. Previously tested configurations can be used and scaled.

NOTE 2 If the user is interested in adopting a previously tested configurations, there are several documented tests in published literature and technical reports: for example References [15], [17], [43], [44] and [89], (or other publicly available test reports such as can be found on OSTI.gov). There are the constraints in F.1.2.5 for adopting the results of previously tested stack or duct configurations.

6.3 provides criteria which describe a satisfactorily well-mixed location in the stack or duct system. These criteria shall be met generally in the centre two-thirds of the cross-sectional area of the duct. Consequently, sampling probes should be located within the centre half of the cross section of the duct where the mixing is optimal. This enables the use of a single nozzle sampling probe, simplifying the probe design and improved sample transport. Modern shrouded nozzles or automatic velocity matching nozzles are accommodating of changes in free stream air velocity.

There are other considerations in locating the sample extraction probe and associated equipment. The location should be readily and safely accessible, it should not present a problem for sampler servicing

and maintenance activities and it should accommodate analysis or collection equipment that does not compromise the quality of the sample. High radiation fields under post-accident conditions can present a problem with respect to worker safety at the sample-extraction location. High ambient temperatures or humidity can also be a problem in some cases. Either of these situations can dictate the requirement for transport lines longer than normally required to accommodate installation of the sample collection and analysis equipment.

Following a careful evaluation (see 7.8), one or more of the following steps should be taken in circumstances where these criteria cannot be satisfied in effluent systems designed and constructed prior to the publication of this document.

- a) Select another location for the sampling probe.
- b) Install features that promote mixing.
- c) Perform an in situ test demonstrating that a representative sample is being collected.

6.3 Criteria for the homogeneity of the air stream at sampling locations

6.3.1 General

The values of the properties that signify a well-mixed location for sample extraction can be characterized by certain criteria which are given in 6.3.2 to 6.3.6 and in Annex F. Achieving the criteria can be demonstrated using a program of physical tests and computation modelling. Measurements of these parameters are made at the in the sampling plane. Physical test results on constructed systems are unambiguous because all hidden construction details and their effects are accounted for. Physical test results from scale models or computational modelling are more ambiguous without accounting for damper construction, damper settings, turning vanes, fan construction, uneven velocity profiles at the discharge of fans and the many hidden construction details.

6.3.2 Angular or cyclonic flow

The presence of a swirl can adversely affect the mixing of particles in the airflow and degrade the performance of sampling nozzles. The mean flow angle between the flow axis and nozzle axis should not exceed 20°.

6.3.3 Air velocity profile

Air velocities are measured at the grid of points described in ISO 10780. The coefficient of variation, C_V , of the measurements is calculated. The C_V should be less than or equal to 20 % across an area that includes at least the centre two-thirds of the area of the stack or duct.

6.3.4 Gas concentration profile

A tracer gas (e.g. alcohol, sulfur hexafluoride, nitrous oxide, helium) is introduced into the airflow upstream of the sampling location. The tracer concentration is measured at the same grid of measurement points used for the velocity uniformity determination. The C_V of concentration at the measurement points is calculated. The C_V of the tracer gas concentration should be ≤ 20 % across at least the centre two-thirds of the cross-sectional area of the stack or duct. Also, at no measurement point should the concentration of the tracer gas differ by more than 30 % from the mean value for all of the points.

NOTE Reference [112] provides a comparison of the use of sulfur hexafluoride and nitrous oxide tracers.

6.3.5 Particle concentration profile

Suitable test aerosol particles are introduced into the airflow upstream of the sampling location. The tracer concentration is measured using the same grid of measurement points as used for the velocity

uniformity determination. The C_V of concentration at the measurement points is calculated. The C_V should be ≤ 20 % across at least the centre two-thirds of the cross-sectional area of the stack or duct.

Test aerosol particles with a D_a of about 10 μm are recommended. This kind of test particle should be used because of the requirement for test aerosol particles whose aerodynamic behaviour clearly exhibits the inertial effects that can adversely influence mixing. They can be relatively easily generated in either monodisperse (single particle size) or polydisperse forms and released into stack or duct flow. Advice on test aerosol injection points is given in [Annex F](#). Favourable results with this particle size result in even better results with aerosol particles of smaller size. See also [Annexes F](#) and [G](#) for further discussion of aerosol particles in different types of facilities.

In cases where additional data about the relevant size distribution (e.g. activity size distribution) under normal, off-normal and anticipated accident conditions are available, the test aerosol particle size may be selected accordingly.

6.3.6 Summary of recommendations for locations to extract samples from a well-mixed air stream

The recommended characteristics for locations from which to extract samples from a well-mixed air stream are summarized in [Table 1](#).

Table 1 — Summary of recommendations for a sampling location

Characteristic	Methodology	Recommendations
Measurement to determine if flow in a duct is cyclonic	ISO 10780	The average resultant angle should be less than 20°.
Velocity profile	Selection of points across a section based on the guidance in ISO 10780 for the centre 2/3 of the area of the stack or duct. Additional points or area may be added to adequately cover the region.	C_V should not exceed 20 % over the centre region of the stack or duct that encompasses at least 2/3 of the stack or duct cross-sectional area.
Tracer gas concentration profiles	Selection of points across a section based on the guidance in ISO 10780 for the centre 2/3 of the area of the stack or duct. Additional points or area may be added to adequately cover the region.	C_V should not exceed 20 % over the centre region of the stack or duct that encompasses at least 2/3 of the stack or duct cross-sectional area.
Maximum tracer gas concentration deviations	Selection of points across a section based on the guidance in ISO 10780 for the entire cross-sectional area. Additional points or area may be added to adequately cover the region.	At no point on the measurement grid should the tracer gas concentration differ from the mean value by more than 30 %.
Aerosol particle concentration profile	Selection of points across a section based on the guidance in ISO 10780. Additional points or area may be added to adequately cover the region.	C_V should not exceed 20 % over the centre region of the stack or duct that encompasses at least 2/3 of the stack or duct cross-sectional area.

7 Sampling system design

7.1 General

The penetration of aerosol particles, gases and vapours of concern from the free stream to the collector or analyser shall be determined.

Performance criteria introduced in this document for monitoring of effluents may be considered as recommendations for sampling systems designed for control monitoring only.

The performance of the sampling system for aerosol particles shall be considered sufficient for normal conditions and for most off-normal conditions if a test under normal conditions with near-monodisperse particles having a D_a of 10 μm yields a penetration value above 50 %. In cases where additional data about the relevant size distribution (e.g. activity size distribution) are available, the test aerosol particle size may be selected accordingly.

The discussion in 7.2 to 7.9 does not cover all possible situations and may be adjusted for particular situations where testing is neither appropriate nor practicable and for local regulations. A risk-based graded approach may be used in the design of systems for sampling radionuclides in stacks and ducts of the nuclear industry. However, formulating such an approach is beyond the scope of this document.

7.2 Volumetric flow measurement

7.2.1 General

Accurate measurements of airflow in both the air sampling system and in the stack or duct being sampled should be provided because they directly impact the accuracy of emissions estimates. Errors are introduced into the calculation of emissions if the emission and sample flow rate units are not based on the same gas density. This becomes significant where airflow is at either elevated or depressed temperature or pressure, for example if the facility is at an elevation of more than 300 m above sea level or if the sample flow meter is on the vacuum side of the air mover. Local regulations may specify the gas density conditions to use for reporting emissions.

In calculating the amount of effluent air, the user should either adjust for the density differences in the air or use measurements based on a standard density. Typical conditions for standard density flow measurements are a pressure of 101,325 kPa and a temperature of 298 K (25 °C). Flow measurements at these conditions are represented by the symbol Q_{std} . The use of so-called mass flow meters in both the emission and sampler airflows, calibrated, can eliminate the need to make density adjustments.

7.2.2 Emission stream flow measurement

The airflow of sampled emission streams should be continuously measured if the flow rate is anticipated to vary by more than 20 % per year (if historical data are available, the 20 % value can be approximated by the standard deviation of the measurements.) Factors such as fan maintenance, the opening of doors and the variations in the number of fans should be taken into account in determining if continuous flow rate is implemented.

If continuous measurement of flow rate is not implemented, then measurements of flow rate should be performed at least annually in accordance with ISO 10780. This standard method, as modified in [Annex A](#), is denoted hereafter as the reference method.

For stacks and ducts where continuous sampling is necessary, the flow measurement and recording system should be capable of determining the mass or volumetric flow rate of the effluent stream with an accuracy that is within 10 % of that measured with the reference method.

Any continuous flow measurement device should be subjected to minimum annual accuracy audits. If the sensor of the continuous flow measurement device is based on electronic or acoustical principles, periodic checks of the instrument zero and span (or linearity) should be made.

7.2.3 Sample air flow rate and volume measurement

The relative accuracy of the sample flow rate measurement and recording system should be within 10 % of a traceable flow standard. The sample flow sensor should be placed in the sampling system so that it does not cause losses of aerosol particles or reactive radioactive gases. As a consequence, the flow sensor is generally located downstream of the sample collector or analyser. Therefore, the gas density at the flow meter differs from the gas density in the stack or duct.

The sample flow rate should be displayed. If a mass flow meter is not used, pressure and temperature instrumentation should be added to enable calculation of the gas density at the sensor.

If the sampling flow rate does not vary by more than 20 % over the sampling period, as a minimum it should be recorded at the start and the end of a sampling period. For such a case, the total volume sampled, V_T , may be calculated using [Formula \(1\)](#):

$$V_T = \frac{q_1 + q_2}{2} \cdot t_s \quad (1)$$

where

- V_T is the total volume sampled, in m^3 ;
- q_1 is the volumetric flow rate indicated by the flow meter at the start of the sampling period, in $\text{m}^3 \cdot \text{s}^{-1}$;
- q_2 is the volumetric flow rate at the end of the sampling period, in $\text{m}^3 \cdot \text{s}^{-1}$;
- t_s is the sampling period, in s.

Continuous flow measurement should be used if the flow rate can vary by more than 20 % during the sampling period. When continuous flow measurement is employed, the flow rate should be recorded at intervals not exceeding 10 min. The total volume of sampled air is based on integration of flow over the entire sampling period. If the time interval between recordings is Δt , expressed in seconds, and the flow rate during the interval (either the true average in the interval, the average of the initial and final values in the interval, or the value at the interval midpoint) is q_i , expressed in cubic metres per second, the total volume of air sampled, V_T , expressed in cubic metres, is calculated using [Formula \(2\)](#):

$$V_T = \sum_{i=1}^N q_i \cdot \Delta t \quad (2)$$

where

- V_T is the total volume of air sampled, in m^3 ;
- q_i is the volumetric flow rate during the i^{th} interval, in $\text{m}^3 \cdot \text{s}^{-1}$;
- N is the number of intervals, dimensionless;
- Δt is the time interval between recordings, in s.

Other integration schemes may be used if the numerically induced errors do not exceed those implicit in the Formula above. The total sample volume, V_T , is based on the flow rate indicated by the flow meter.

If a controller is used to maintain a constant flow rate, the controller should maintain the flow rate within 15 % over conditions that correspond to an initial pressure drop across the collector (usually a filter) or analyser to a value that is twice the initial pressure drop.

If the emission flow rate can vary more than 20 % over a sampling interval, automatic control of the sample flow rate should be used and the sample flow rate should be varied in proportion to the flow rate through the stack or duct. The ratio between sample flow rate and effluent flow rate should be maintained within 20 % of the sample fraction at normal operating conditions.

An exception may be made for that part of the air monitoring system containing a real-time contamination sensor if its operation depends on a constant internal flow rate.

The flow controller should be tested at conditions similar to the operating conditions.

7.2.4 Leak checks

A leak in the sampling system or around the sample collector can cause the indicated sample flow rate to be in error and could cause improper functioning of the sample collector. A sampling system should be inspected for leaks at the time of installation and at any time when either significant maintenance is performed or during a system inspection. The inspection or test methodology should be practical for the installation and should be documented.

Leakage under flowing conditions into a sample collector, measured with apparatus such as that discussed in NOTE 1 or NOTE 2, below, should not exceed 1 % of the operational flow rate when the pressure level across the filter or collector is as recommended in NOTE 1.

NOTE 1 Visual inspection and the observation of foreign materials on samples can identify large leaks. If the sampling system is strategically equipped with full-bore ball valves, then parts of the system can be isolated for vacuum or pressure decay measurements or by observing or measuring flow through a blocked system. An explanation of the latter approach is to block flow through the nozzle(s) (or at a transport line connection close to the sampling probe), then apply vacuum to the transport line and measure the leakage rate. For example, a mass flow meter could be attached downstream of the collector or monitor and the vacuum source connected to downstream side of the mass flow meter. The pressure level in the tubing between the collector or monitor and vacuum source would be adjusted to the nominal value encountered during sampling (typically about 4 kPa for sampling systems that involve use of collection filters) by bleeding air into the line downstream of the flowmeter.

NOTE 2 Reference [56] illustrates the use of a chamber and tracer gas method for leak testing of filter holders and continuous monitors under flowing conditions.

NOTE 3 The recommendations of this subclause do not apply to collectors internal to continuous air monitoring instruments.

7.3 Nozzle design and operation for extracting aerosol particles

7.3.1 General

The information in 7.3.2 to 7.3.7 is applicable to sampling from stacks and ducts that have the potential to emit aerosol particles. Extracting the air sample with a properly designed nozzle from a location where the potential contaminants are well mixed in the airflow provides a representative sample during normal conditions and an adequate sample during accident conditions. Background information on the design parameters of sampling nozzles is given in Annex M. Performance recommendations for nozzles are given in 7.3.2 to 7.3.7.

7.3.2 Nozzle performance

M.1 provides a discussion of the relationships of the nozzle performance factors. Performance characteristics of sampling nozzles are tested with liquid aerosol particles with a recommended D_a of about 10 μm . This kind of test particle should be used because of the requirement for test aerosol particles whose aerodynamic behaviour clearly exhibits the inertial effects that can adversely influence mixing. They can be relatively easily generated in either monodisperse (single particle size) or polydisperse forms and released into stack or duct flow. Comparable behaviour should be demonstrated to provide similar transmission values, because liquid particles adhere to walls, while solid particles can rebound or be re-entrained from a surface.

The test aerosol D_a of about 10 μm is recommended because the results should be conservative in most situations as discussed in Annex G. Because radioactivity bearing particles commonly change size as a facility ages or events occur, favourable test results with this particle size means even better performance with reduced uncertainty for smaller aerosol particles size. In cases where additional data about the relevant size distribution (e.g. activity size distribution) under normal, off-normal and anticipated accident conditions are available, the test aerosol particle size may be selected accordingly.

The presence of a nozzle should not disturb the aerosol particle concentration in the stack or duct. Accordingly, the frontal area of a nozzle, including the shrouds if so equipped, should not be excessive

(e.g. not greater than 15 % of the stack or duct cross-sectional area) and the inlet diameter should not be too small.

7.3.3 Application and performance considerations

7.3.3.1 General

The factors in 7.3.3.2 to 7.3.3.5 should be considered in the selection and use of a sampling nozzle.

7.3.3.2 Location

Sampling should take place at a location where both the aerosol particle concentration and fluid momentum (velocity) are well mixed and thus meet the performance criteria of 6.3.

7.3.3.3 Orientation

For aerosol particle sampling, the nozzle axis should be aligned parallel to the temporal mean flow direction.

7.3.3.4 Transmission and aspiration ratios

A sampling nozzle shall have a transmission ratio within the range of 0,80 to 1,30 over the anticipated range of normal and abnormal conditions for an aerosol with a particle size of $10\ \mu\text{m}$ D_a . Also, the aspiration ratio of a sampling nozzle shall be within the range of 0,80 to 1,50 for the anticipated range of operating conditions. The transmission and aspiration ratios of the selected nozzle design should be traceable to experimental verification of performance for conditions that include the nominal sampling flow rate and range of anticipated sampling flow rates, the nominal free stream velocity and the range of anticipated free stream velocity, and a particle size, D_a , of about $10\ \mu\text{m}$. In cases where additional data about the relevant size distribution (e.g. activity size distribution) are available, the test aerosol particle size may be selected accordingly. If actual testing is used, the means for determining the transmission and wall-loss ratios should be documented. If reference to previous testing is employed, the equivalency of the selected design and the design that was tested should be documented.

7.3.3.5 Nozzle configuration

The leading edge of the nozzle inlet should have a sharp edge and the external cone angle should not exceed 30° . Other configurations may be used if experimental data show either equivalent or superior performance to the sharp-edged nozzle. If the sampling nozzle is shrouded, the shroud should not have a sharp leading edge. For sharp-edged nozzles, the leading edge of the nozzle should be inspected for damage following installation and subsequent to any maintenance procedures in which the nozzle could be damaged. Studies (e.g. References [47], [82] and [83]) have shown that single-nozzle, shrouded probes have reduced wall losses for particles having an aerodynamic diameter, D_a , greater than $5\ \mu\text{m}$, in high-velocity streams (velocity greater than $10\ \text{m}\cdot\text{s}^{-1}$) when compared to isokinetic nozzles.

7.3.4 Sampling probes with multiple-inlet nozzles

The previous edition of this document recommended selecting a sampling location where both fluid momentum and contaminants are well mixed with the airflow. The concept of “well mixed” was loosely defined and the deployment of multiple nozzles in circular ducts larger than 201 mm in diameter, or in rectangular ducts with cross-sectional areas greater than $0,093\ \text{m}^2$, was recommended to ensure the possibility of extracting a representative sample. However, it is now recommended that in place of multiple-point sampling, single-point representative sampling should be used, with the requirement that potential contaminants be “well mixed” at the sample-extraction location, as specified in the performance criteria of 6.3.

If the effluent stream cannot be adequately mixed and sampled at a single point, probes with multiple inlets have been a solution in the past. Now, in this case, an in-place demonstration of the system providing a representative sample is recommended.

Now it is required that a multi-nozzle probe be capable of the same performance as a single-nozzle probe. A multi-nozzle probe shall, then, in principle, consist of a number of acceptable nozzles. The total sample flow is then increased. Nozzles with an internal diameter smaller than about 10 mm are usually not an option because they have a lower flow and higher losses. It is necessary, in fact, that a nozzle on a multi-nozzle probe perform as well as, or better than, a single nozzle because of particle losses in additional bends and joints.

Multi-nozzle probes may be used where they can be demonstrated, with in-place testing, to provide a representative sample. Some designs of multi-nozzle samplers are discussed in [Annex M](#).

7.3.5 Materials of construction

Nozzles should be constructed of materials that do not react with either the aerosol particles or the vapour constituents of the gas stream. The average surface roughness of the internal regions of nozzle that contact the sampled stream should not exceed 0,8 μm . The average surface roughness of the external region of the sampling nozzle from the inlet plane to a distance of two nozzle inlet diameters from the inlet plane should not exceed 1,6 μm . A shroud should have an average surface roughness that does not exceed 3,2 μm .

7.3.6 Maintenance

The sampling nozzle should be checked periodically for alignment, presence of deposits of foreign materials and other factors that can degrade the performance of the sampling system. If there are background aerosol particles that can produce deposits, a cleaning schedule should be established that does not allow the occlusion of over 5 % of the inlet area of a nozzle. For nozzles that are used to sample air, filtered by a high efficiency particulate air filter (HEPA), the nozzle should be cleaned if there are visible deposits of material on either the internal or external regions of the nozzle, although visible deposits would not be expected if the filters were performing satisfactorily.

7.3.7 New concepts

When new approaches are developed for design and operation of nozzles, such designs may be used in ducts and stacks if it can be demonstrated experimentally that the designs meet or exceed the performance specifications given in [7.3](#). The test conditions should include experiments to determine the wall losses and aerosol particle transmission under the following conditions:

- particle sizes, D_a , of about 3 μm , 10 μm and 20 μm at the nominal free-stream velocity and nominal flow rate;
- maximum and minimum operational or anticipated free-stream velocities for a particle size, D_a , of 10 μm at the nominal sampling flow rate;
- maximum and minimum anticipated sampling flow rates for a particle size, D_a , of 10 μm at the nominal free-stream velocity.

7.4 Sample transport for particles

7.4.1 General

The transport of aerosol particles from a sampling nozzle to a collector or analyser should take place in such a manner that changes in concentration and size distribution of airborne radioactive materials are minimized within the constraints of current technology.

7.4.2 Depositional losses

In general, there are some losses of aerosol particles in transport lines due to particle deposition, and any design entails compromises. The design parameters should be carefully chosen to optimize the utility of the overall system. [Annex B](#) provides guidance on assessing particle penetration.

The deposition of particles inside the transport lines from the extraction point to the filter should be determined experimentally using test aerosol particles or by the use of documented computer codes or documented and referenced hand calculations.

The performance of the sampling system is considered sufficient under normal, off-normal and anticipated accidental conditions, if a test with near monodisperse particles of 8 µm to 12 µm (D_a) yields a penetration value above 50 %.

A D_a of 10 µm is mentioned when no information on aerosol size distribution is available. In cases where additional data about the relevant size distribution (e.g. activity size distribution) are available, the test aerosol particle size can be selected accordingly.

Particles with a D_a smaller than 10 µm have a higher penetration due to lower deposition rates. Particles with a D_a larger than 10 µm can be expected to have smaller penetrations. On the other hand, for dry particles, the penetration can increase with particle size due to resuspension. Therefore, penetration measurements with test aerosol particles having a D_a of 10 µm can be considered to address the penetration minimum.

In cases where additional data about the relevant size distribution (e.g. activity size distribution) are available, the test aerosol particle size may be selected accordingly.

Documented computer codes or documented and referenced hand calculations, based on verified experimental data, may also be used to assist in the design of the sampling line and selection of test aerosol particles. The computations may be used to extrapolate measured penetration values to other particle sizes, and may be used also in the case of a modification of the flow rate or changes in sampling-line geometry. [Annex B](#) describes such calculation methods.

The straight sections of transport tubes, particularly horizontal tubing sections, should be kept as short as possible, and the number of bends should be minimized within the geometrical constraints of the application. There should be no inward-facing steps at the tubing connections that cause more than a 1 % reduction in tube diameter. The tubing ends should be free of burrs and crimping. Bends should have a curvature ratio of at least 3. Flattening, which is defined as the ratio of the minimum tube diameter to the original tube diameter, should not be less than 0,85. The user should note that special fabrication techniques can be needed to meet these specifications.

7.4.3 Corrosion

The internal walls of the transport system should be constructed of materials that are minimally reactive to inadvertently deposited aerosol particles or to reactive vapour compounds that can be present in the sample. The materials of construction for external walls and seals between sampling system components should also be compatible with the environment to which they are exposed. Materials recommended for the nuclear industry are stainless steel for general applications and polytetrafluoroethylene for radioiodine.

7.4.4 Electrostatic effects and flexible tubes

If plastic is used in aerosol-particle transport systems, internal electric fields can cause particle losses (see Reference [23]). In particular, plastic tubing that has been flexed can show abnormally high wall deposits as seen in Reference [70]. A transport system should be constructed of materials, such as metals or conductive plastics, that do not maintain internal electrostatic fields. In many applications, it is useful or convenient to employ flexible, non-metallic tubing to connect a sampler or analyser to a transport line, particularly if it is necessary to isolate an analyser from mechanical vibrations in the sample transport line. The inside diameter of the plastic line should not be smaller than the inside diameter of the components with which it is connected and the bend curvature ratio should not be less than three, nor may the curvature of a bend cause more than a 15 % change in the inside diameter of the tube. If non-conductive, flexible tubing is used, the line length should be kept as short as practicable and should not exceed 0,5 m.

Of the flexible tubes that can be categorized as non-conductors, neoprene and natural rubber are recommended to minimize electrostatic deposition of particulate matter (see Reference [23]). If radioiodine is present in the effluent stream, the materials suggested in [Annex C](#) should be used.

7.4.5 Smoothness of internal surfaces

To minimize aerosol-particle depositional losses and to facilitate decontamination, the internal surfaces of transport lines should be as close to hydraulically smooth as practical. Drawn tubing or other types of tubes with Ra/d_t less than approximately 5×10^{-5} are acceptable, where Ra represents the height of surface roughness of the internal tube walls and d_t is the tube diameter. This criterion requires an average surface roughness of approximately 1,6 μm or less for tube sizes that are in the order of 25 mm in diameter.

7.4.6 Condensation

Sample transport lines, collectors and analysers should be designed to avoid condensation of vapour. Condensation takes place when the temperature of air in the sample transport line is less than the saturation temperature of the vapour of interest. It can be necessary to thermally insulate, and in some cases heat, the sample transport line to prevent condensation. For situations in which heating of the sampling line can result in unacceptably high temperatures at a collector or analyser, a dilution system should be considered. However, care should be exercised to ensure that the dilution process does not produce condensation at the mixing location. Experimental or numerical analyses should be performed to demonstrate the effectiveness of any design provisions that are intended to minimize or preclude the formation of condensation in sample transport systems.

7.4.7 Cleaning transport lines

An additional consideration at some facilities is the necessity to clean transport lines. For applications in which the sampled air is HEPA-filtered, cleaning might not be necessary within the expected lifetime of the installation. However, for applications where background aerosol particles or contamination are present in normal and/or off-normal conditions, it can be necessary to periodically remove deposits from the internal walls of the transport system. If, after inspection, there is an indication of deposits inside the nozzle inlet, the transport line should be inspected and, if deposits are visible inside the transport line, the line should be cleaned or replaced. For systems that sample non-HEPA-filtered air containing background aerosol particles, regular inspections are recommended. If an estimate can be made of the rate of deposition of all aerosol particles on the internal walls of the system, the system should be cleaned when the mean mass of deposited material exceeds $1 \text{ g}\cdot\text{m}^{-2}$. Measuring the mass of material deposited is possible only in systems designed to allow it. Alternatively, a cleaning schedule can be set up based on performance of the transport system. The interval between cleanings should be such that accumulations of wall losses cause a reduction of no more than 10 % in the overall penetration of aerosol particles with a D_a of 10 μm through the sampling system.

In cases where additional data about the relevant size distribution (e.g. activity size distribution) are available, the test aerosol particle size may be selected accordingly.

In addition, if there are indications of re-entrainment of deposits from the walls of the sampling system or if there has been sampling of easily re-entrainable aerosol particles (e.g. flakes), either of which can cause anomalous radiological data to be gathered, the system should be cleaned. Decontamination and waste-production should be taken into account for any cleaning procedure.

7.5 Gas and vapour sample extraction and transport

Much of the discussion in [7.4](#) applies generally for sampling particles and gases. However, consideration should be given to extracting and transporting vapour and gases to determine where special system design can be required.

When non-reactive gases and vapour are the only species being sampled, the sampling recommendations are considerably simpler than those for aerosol particles. The recommendations for minimizing particle

line-loss are irrelevant. Deposition in long transport lines and condensation due to temperature changes in the line should be avoided. If the flow can contain only gaseous contaminants, the nozzle design is not critical, but the sampling should take place at a location where the flow is well mixed and meets the criteria of 6.3. The nozzle design can be simply an open-ended or perforated tube. The extraction and transport requirements that apply include extracting the sample from a well-mixed location and avoiding water and vapour condensation in the transport and collection system (except where condensation is used as the collection method).

When non-reactive gases, vapour and particles are being simultaneously sampled, the particle sample extraction and transport requirements should apply, which also ensures adequate delivery of the gas and vapour sample. The remaining consideration, then, is the selection of suitable collection devices. The gas or vapour-collection device should be located downstream of a particle filter to eliminate potential radionuclide interferences by particulate matter. The minimum penetration for vapour or gas samples from the free stream to the collector or analyser should be 50 %.

When working with reactive gases and vapour, particular attention should be paid to the sampling-system construction materials and to avoiding condensation. The construction material should have minimum reactivity with the gas. Consideration should be given to the advantages of providing a separate sampling system for the gases whenever the construction materials that are for the transport of the particle and gas samples are incompatible. In situations where even a low level of reactivity cannot be avoided, the length of the transport line should be kept to a minimum. The penetration of the gas or vapour through the complete extraction and transport system should be documented. The minimum penetration for vapour or gas samples from the free stream to the collector or analyser should be optimized, with the objective of achieving 50 %. If long transport lines are unavoidable, consideration should be given to the effect of transport and detection delay time caused by deposition, chemical transformation and subsequent resuspension. Consideration should also be given to how significant the effect the delay has on the timeliness, interpretation and usefulness of the resulting data. Although rapid changes in the emission can become attenuated over a long time interval relative to the change in emission, the data can still be useful and quantitative when interpreted in that light. [Annexes C, H and K](#) provide guidance on the sampling of radioiodine, tritium and carbon-14, respectively. [Annex B](#) provides guidance on verifying the transport of sampled constituents.

7.6 Collection of particle samples

7.6.1 General

Depending on the purpose of the sample, a wide range of techniques is available for monitoring or collecting airborne particles. Particles can be collected on filters for retrospective determination of total mass, radionuclide activity, or chemical form, in cyclones or cascade impactors for determination of particle size distribution, on electron microscope substrates for determination of particle morphology or they can be observed using light-scattering or time-of-flight techniques for measuring the number, concentration and size. Near-real-time devices, such as alpha continuous air monitors (CAMs), typically collect particles on a filter or impaction substrate and monitor the accumulation of radioactive substance with time. Critical issues for selection and operation of particle collection devices are as follows:

- appropriate presentation of the sample for real-time analyses or preservation of the sample for retrospective analyses;
- adequate flow rates and detection efficiencies to meet sensitivity requirements;
- minimal in-leakage within the collector;
- minimal particle loss within the collection zone.

7.6.2 Filter media

Selection of a particle-collection filter should be based on careful consideration of collection efficiency for the typical particle size in the duct, the area of the filter, the pore size, the filter's resistance to air

flow, the background radioactive material of the filter, filter fragility, cost, self-absorption within the filter and chemical solubility. If the performance characteristics of the front and back surfaces of the filter are not within 5 % of each other for the intended purpose of the sample, there should be a clear means of identifying the appropriate surface for particle collection. The filter should be strong enough to maintain its integrity at the required sample flow rates and during handling activities.

When filter media are used, a backup support that produces a negligible pressure drop should be used behind the filter to prevent filter distortion or deterioration. The filter holder should provide adequate structural support while not damaging the filter, should prevent sampled air from bypassing the filter, should facilitate changing the filter and should facilitate decontamination. If gaskets are used to seal the filter to the backing plate, the gasket should be in contact with the filter along the entire circumference to ensure a good fit. The gasket should be periodically inspected to detect degradation and eliminate build-up of dust or filter material, which can result in sampled air bypassing the filter.

To reduce the uncertainty associated with collection efficiency, filters that are used for sampling airborne radioactive particles should have a minimum efficiency of 95 %. Efficiency values should be applicable to the conditions of use, for example, the collection efficiency depends on the face velocity and humidity (see Reference [23]).

If published or manufacturer's data on filter collection efficiency are not available for the particle sizes of interest, then the efficiency should be determined by the user. This can be done by placing a highly efficient membrane or glass fibre filter behind the filter of interest and then comparing the mass penetrating to the backup filter to the total mass collected on both filters (see Reference [48]). If a filter with an efficiency lower than 95 % is required to meet the overall sampling objectives, then a correction for efficiency should be made. Because filter efficiency is a function of air flow rate, care should be taken to maintain a sample flow rate that is adequate to achieve the desired collection efficiency.

If penetration of radioactive material into the collection media or self-absorption of radiation by the material collected can reduce the count rate by more than 5 %, a correction factor should be used. A dual filter method can also be used to measure the absorption efficiency in the filter medium (see Reference [48]). Evaluation of self-absorption in the material collected may require separate radiochemical analyses.

[Annex D](#) illustrates the type of information that is useful in selecting an appropriate filter for sampling airborne radioactive particles. This includes physical and performance characteristics of a number of typical coarse-fibre, fine-fibre and membrane-type filters.

7.7 Collection of gas and vapour samples

7.7.1 General

Airborne radioactive volatile materials and noble gases (e.g. krypton) are frequently present in nuclear facility effluents. Their sampling and collection require techniques and methods that are different from those used in particle sampling. This topic may be divided into two general methods of sampling:

- a) sampling with retention of specific constituents of the air stream, and
- b) sampling without constituent separation. [Annexes C, H, K](#) and [N](#) provide further guidance specific to radioiodine, tritium, carbon-14 and ruthenium-106, respectively.

7.7.2 Sampling with retention of specific constituents

Sampling with removal and collection of specific constituents requires a detailed knowledge of the chemical and physical properties of the radioactive material of interest, including possible interfering materials, such as particulate matter and accompanying non-radioactive gases (e.g. acids and organic chemicals). The many possible combinations of the properties of the constituents being measured and the accompanying airborne materials require careful study to select the optimum collector. Gases and vapour components can be soluble in water, can be highly reactive with certain solutions, can dissolve in specific non-aqueous solvents, or can be retained on specific solid adsorbents or other specifically

prepared media. In general, continuous or extended samples are taken when separation and removal of a constituent is required. Sampling rates should be established to ensure adequate sensitivity for the selected radioassay method and should be compatible with the collector performance characteristics. Avoiding sample breakthrough should also be considered when choosing the sampling rate and duration. The principal collection methods include solid absorbents (such as carbon, zeolites, silica gel and metal beds), condensation, gas absorption and catalytic or chemical reaction (see Reference [20]).

The retention of certain sampled gaseous or vaporous constituents are specifically addressed in the annexes. [Annex H](#) discusses Tritium, [Annex K](#) discusses ^{14}C and [Annex N](#) covers ^{106}Ru .

7.7.3 Sampling without constituent separation

In some instances, a sample of air containing gaseous radioactive constituents can be desired for measurement of the activity concentration of airborne materials and its trend. Examples are noble gas isotopes, tritium and activated gases near a reactor. Volume collection and flow-through detectors are the two principal methods for total gas sampling or monitoring.

Because the constituent radioactive materials of interest might not be concentrated with a particular flow-through or volumetric collection device, insufficient sensitivity of detection can limit or preclude their use. It is necessary to evaluate each situation individually to determine the feasibility of the gross sample measurement.

Volume collection methods include the following:

- using an evacuated container that can be valved open to the stream of interest, then sealed and returned to a laboratory for measurement of all the activity or the activity of individual constituents;
- passing the stream through the sample vessel until the vessel is completely purged, then closing the inlet and outlet valves;
- pumping the sample stream into deflated bags (of a non-adsorbing material) for subsequent compression and analysis;
- compressing the sample stream into a vessel for real-time or subsequent analysis.

A flow-through sample vessel may also be an ion chamber whose ion current reflects the activity concentration of material in the gas. Care should be taken to keep the gas well above its dew point in the sampling system and ionization chamber. A gradual build-up of contamination in the chamber usually occurs, which increases the observed activity rates.

Flow-through chamber samplers can be similarly monitored by gamma-ray scintillation crystal counters or other detector held adjacent to or inserted in a well in the chamber wall. An increased background from contamination can be expected in these samplers, and the chamber should be periodically decontaminated to avoid errors from this source. Prior filtration of the air stream can assist in keeping the chamber clean when gaseous constituents alone are being measured. A flow-through sampling system, which is frequently used at power reactors for accident monitoring, involves placement of a high- or wide-range detector mounted directly inside or outside the effluent stack or duct.

7.8 Evaluation and upgrading of existing systems

If an existing air sampling system was not designed to the performance requirements and recommendations of this document, an evaluation of the performance of the system, including the location of the probe, is recommended. If deficiencies are discovered, an evaluation study should be performed to determine if a retrofit is recommended and possible. Arriving at a suitable solution requires optimizing among competing factors. Guidance on the process of optimization for radiological protection has been described in Reference [10]. Evaluation of existing systems should be undertaken using proven techniques.

7.9 Summary of performance criteria and recommendations

Throughout this document, performance criteria for various elements have been included in the discussion of each element. For convenience, they are summarized in [Table 2](#). These criteria cover aspects of system design and operation. The approach followed in this document is to give performance criteria and recommendations according to guidance values when no information is available on the type of stream being measured. When information is available on the stream being measured (e.g. aerosol particle size distribution), then these specific values should be used.

A facility sets action levels for a particular radionuclide in an aerial discharge in response to the authorization discharge levels for that facility set by the regulator of that facility. Action levels can be either a control mechanism used by the facility itself, or levels that, if reached, necessitate official notification to the regulator.

Performance criteria for the sampling and measurement of a specific radionuclide are determined by the facility, in consultation with the regulator and [Table 2](#), in order to provide accurate monitoring of the specific radionuclide.

Table 2 — Summary of performance criteria and recommendations

Performance criteria and recommendations ^a	Reference
The performance of the sampling system for aerosol particles shall be considered sufficient for normal, off-normal and anticipated accidental conditions, if a test under normal conditions with monodisperse particles with a D_a of 10 μm yields a penetration value $\geq 50\%$.	Clause 7 and 7.4.2
The penetration of the gases and vapours of concern from the free stream to the collector or analyser should be considered sufficient if it is $\geq 50\%$.	Clause 7 and 7.5
The nozzle should have a transmission ratio $\geq 80\%$ and $\leq 130\%$ for particles with a D_a of 10 μm .	7.3.2
Recommendations for a suitable sampling location are as follows: a) coefficients of variation over the central 2/3 by area of the cross-section within 20 % for with a D_a of 10 μm , gaseous tracer, aerosol particle tracer and gas velocity; b) the tracer gas concentration should not vary from the mean by $>30\%$ at any point on the measurement grid (see 6.3.4). c) Flow angle $\leq 20^\circ$ relative to the long axis of the stack or duct and the nozzle inlet	6.3
Effluent flow rate continuous measurement recommended if flow variation is greater than $\pm 20\%$ in a year.	7.2.2
Effluent and sample flow rate should be measured within $\pm 10\%$.	7.2.2 and 7.2.3
Continuous sample flow rate measurement and control are recommended if the stack or duct flow varies by more than $\pm 20\%$ during a sample interval. Flow control should be within $\pm 15\%$.	7.2.3
NOTE 1 A D_a of 10 μm is mentioned when no information on aerosol size distribution is available. In cases where additional data about the relevant size distribution (e.g. activity size distribution) are available, the test aerosol particle size can be selected accordingly.	
NOTE 2 This table is based on presence of aerosol particles in the stack or duct. If no particles are emitted, criteria can be based on gas characteristics only.	
^a The criteria given in this table are recommendations based on user experience in actual testing. These recommendations for individual uncertainties and biases result in a satisfactory overall uncertainty.	

8 Quality assurance and quality control

Every facility that conducts sampling and monitoring of effluent radioactive substances should have a quality assurance (QA) programme. The purposes of a QA programme are to provide assurance to facility management, regulatory agencies and the public of the validity of the data from the sampling and monitoring of released radioactive substances, and to identify any deficiencies in the sampling

equipment and procedures in order to take corrective action. The tools used to accomplish these objectives include documentation, calibration, maintenance and inspection. As a minimum, the QA programme should address the quality aspects of the sampling of effluent radioactive substances in the following areas:

- a) organization:
 - organizational responsibilities;
 - administrative controls;
 - reporting and notification system;
 - documentation;
 - personnel qualifications;
- b) design of the sampling system:
 - source terms;
 - selection of extraction locations;
 - selection of sampling and monitoring devices;
 - selection of collection procedures;
- c) operating procedures:
 - sample extraction procedures;
 - sample collection procedures;
 - system operation procedures;
 - calibration procedures;
 - data analysis;
 - maintenance and check procedures;
 - maintenance procedures;
 - check and test procedures;
 - status;
 - disposition of non-conformant items and conditions;
 - corrective action programme.

Additional information relating to inspections and calibrations particularly relevant to air sampling systems is presented in [Annex J](#).

Annex A (informative)

Techniques for measurement of flow rate through a stack or duct

A.1 General

The volumetric flow rate, q_a , through a stack or duct is defined as given in [Formula \(A.1\)](#):

$$q_a = \int_A v \cdot dA \quad (\text{A.1})$$

where

q_a is the volumetric flow rate through a stack or duct, in $\text{m}^3 \cdot \text{s}^{-1}$;

v is the velocity at any location across a stack or duct, in $\text{m} \cdot \text{s}^{-1}$;

A is the cross-sectional area of the duct, in m^2 .

A method for determining q_a involves measuring the velocity at a finite number of points in a duct, where each point is chosen as the centroid of an area element. The relationship defining q_a is as given in [Formula \(A.2\)](#):

$$q_a = \sum_{i=1}^N v_i \cdot \Delta A_i \quad (\text{A.2})$$

where

q_a is the volumetric flow rate through a stack or duct, in $\text{m}^3 \cdot \text{s}^{-1}$;

v_i is the velocity at the centroid of the i^{th} area element, in $\text{m} \cdot \text{s}^{-1}$;

A_i is the cross-sectional area of the i^{th} element, in m^2 .

The cross-section of the stack or duct is divided into N elements. In practice, all of the N elements have equal areas. The approach embodied in ISO 10780 serves as the reference method for this document. The requirements for the absence of cyclonic flow given in ISO 10780:1994, Annex C, are also included in the reference method.

The flow rate, q_a , is associated with the air density, ρ , that exists in the stack or duct. The density is calculated from the ideal gas equation for dry air as given in [Formula \(A.3\)](#):

$$\rho_a = \frac{p_a}{R \cdot T_a} \quad (\text{A.3})$$

where

ρ_a is the gas density in the stack or duct at actual conditions, in $\text{kg} \cdot \text{m}^{-3}$;

p_a is the pressure in the stack or duct, in Pa;

R is the individual gas constant, here for air, equal to $287 \text{ J} \cdot \text{kg}^{-1} \cdot \text{K}^{-1}$;

T_a is the temperature in the stack or duct, in K.

The volumetric flow rate at standard conditions, q_{std} , is the parameter that is being calculated for reporting and analysis purposes, and it is related to the actual volumetric flow rate, q_a , as given in [Formula \(A.4\)](#):

$$\rho_a \cdot q_a = \rho_{\text{std}} \cdot q_{\text{std}} \quad (\text{A.4})$$

where

- ρ_a is the gas density in the stack or duct at actual conditions, in $\text{kg}\cdot\text{m}^{-3}$;
- q_a is the volumetric flow rate through a stack or duct at actual conditions, in $\text{m}^3\cdot\text{s}^{-1}$;
- ρ_{std} is the gas density in the stack or duct at standard conditions, in $\text{kg}\cdot\text{m}^{-3}$;
- q_{std} is the volumetric flow rate through a stack or duct at standard conditions, in $\text{m}^3\cdot\text{s}^{-1}$.

The flow rate at standard conditions from [Formula \(A.4\)](#) can then be expressed as given in [Formula \(A.5\)](#):

$$q_{\text{std}} = q_a \cdot \frac{T_{\text{std}}}{T_a} \cdot \frac{p_a}{p_{\text{std}}} \quad (\text{A.5})$$

where

- q_{std} is the volumetric flow rate through a stack or duct at standard conditions, in $\text{m}^3\cdot\text{s}^{-1}$;
- q_a is the volumetric flow rate through a stack or duct at actual conditions, in $\text{m}^3\cdot\text{s}^{-1}$;
- T_a is the temperature in the stack or duct, in K;
- T_{std} is the standard temperature, equal to 298 K;
- p_a is the pressure in the stack or duct, in Pa;
- p_{std} is the standard pressure, equal to 101,325 kPa.

In practice, the value of q_a is determined from velocity measurements at traverse points as specified in ISO 10780, with the value calculated from [Formula \(A.2\)](#). The temperature and pressure in the stack or duct are measured in accordance with the requirements of ISO 10780.

A.2 Special considerations for use of ISO 10780 in sampling stacks and ducts of nuclear facilities

A.2.1 General

The reference method for determining air flow rate through a stack or duct, as given in ISO 10780, is developed for flow rate determinations in non-nuclear stacks and ducts. It is necessary to take into consideration several differences between sampling from non-nuclear stacks and ducts and their nuclear counterparts.

A.2.2 Pitot tubes

An S-type Pitot tube was recommended in ISO 10780 to reduce the risk of dust plugging the ports of the Pitot tube when measurements are made in dusty environments. Because dust loading during velocity mapping in a stack or duct in the nuclear industry is, however, usually not a concern, Prandtl-type Pitot static tubes (Type L in ISO 10780) should be considered as the reference apparatus for sampling under the requirements of this document. However, when dust loading is of concern, an S-type Pitot tube may be considered.

A.2.3 Mean molar mass of the stack or duct gas

In the industrial applications for which ISO 10780 was designed, the gas being tested often contains products of combustion or an elevated water-vapour content resulting from drying operations. In contrast, the gas in most stacks and ducts of the nuclear industry is ventilation air. In the latter case, it is usually unnecessary to determine the mean molar mass, M , in stacks and ducts. However, if it can be anticipated that there can be more than 10 % water vapour in the stack or duct, or if there are other gases that can change the mean molar mass by more than 4 % from the value for dry air (20,896 kg·mol⁻¹), it is necessary to determine the mean molar mass of the gas. In this case, the resulting value is used to calculate the gas constant, R , as given in [Formula \(A.6\)](#):

$$R = \frac{R_u}{M} \quad (\text{A.6})$$

where

R is the individual gas constant, in J·kg⁻¹·K⁻¹;

R_u is the universal gas constant, equal to 8,314 J·mol⁻¹·K⁻¹;

M is the molar mass of the gas, in kg·mol⁻¹.

A.2.4 Thermal anemometers

If the mean molar mass and water content of the stack or duct gas are in accordance with the values stated, the gas can be treated as air. If the dust loading in the stack or duct is such that any deposits on a thermal anemometer probe do not change the calibration of a thermal anemometer by more than 3 % during the course of the velocity measurements and if there is no condensation of water vapour or other vapour on the sensor during flow measurements, a thermal anemometer can be used instead of a Pitot tube.

When a thermal anemometer is used for velocity mapping in a stack or duct, the flow rate based on standard conditions, q_{std} , is determined according to [Formula \(A.7\)](#):

$$q_{\text{std}} = \sum_{i=1}^N v_{\text{std,ta},i} \cdot \Delta A_i \quad (\text{A.7})$$

where

q_{std} is the volumetric flow rate through a stack or duct at standard conditions, in m³·s⁻¹;

$v_{\text{std,ta},i}$ is the velocity measured with a properly calibrated thermal anemometer at the centroid of the i^{th} element of area, in m·s⁻¹;

A_i is the cross-sectional area of the i^{th} element, in m².

A.3 Conversion of data from single point or single line measurements to total flow rate

A.3.1 General

If continuous single-point velocity measurements with a Pitot tube or a thermal anemometer or line-integral measurements from an acoustic flow meter are used to infer the total flow rate through a stack or duct as a function of time, the resulting data should include a correction factor accounting for the shape of the velocity profile. The correction factor is determined by comparing the flow rate obtained with the single-point technique with those of the reference method defined in ISO 10780.

A.3.2 Pitot tube

The velocity-averaging correction factor for a Pitot tube, $C_{v,pt}$, is defined in [Formula \(A.8\)](#):

$$C_{v,pt} = \frac{q_a}{v_{pt} \cdot A} \quad (A.8)$$

where

- $C_{v,pt}$ is the velocity-averaging correction factor for a Pitot tube, dimensionless;
- q_a is the volumetric flow rate through a stack or duct at actual conditions calculated according to [Formula \(A.2\)](#), in $\text{m}^3 \cdot \text{s}^{-1}$;
- v_{pt} is the velocity obtained from a single-point Pitot tube, in $\text{m} \cdot \text{s}^{-1}$;
- A is the cross-sectional area of the duct, in m^2 .

Usually, the single point is located near the centre of the duct. Multiple measurement points can be used, in which case the value of v is the average of the values from the multiple points.

The velocity, v , is continuously monitored during the period that the reference method testing is performed to establish the velocity-averaging correction factor $C_{v,pt}$. If the value of v changes by more than 5 % during the course of testing, the data should be rejected and the test repeated. If the stack or duct is subject to long-term flow rate variations that exceed 25 %, additional tests should be carried out at the highest flow rate (if it exceeds the base condition by more than 25 %) and the lowest operational flow rate (if it deviates from the base condition by more than 25 %) to establish values of the velocity-averaging correction factor at those conditions. A single value of the flow correction factor may be used if the range of flow-correction-factor values is within 7 % of the base condition. If the values of the correction factor at the extreme flow conditions are greater than 7 % of the base condition, then a relationship may be needed between the velocity correction factor and flow rate.

At least two replicate tests should be performed to establish a value of the correction factor, $C_{v,pt}$. During routine use, the flow rate, q_a , is determined from readings of the single-point Pitot tube as given in [Formula \(A.9\)](#):

$$q_a = C_{v,pt} \cdot v_{pt} \cdot A \quad (A.9)$$

A.3.3 Thermal anemometer

A thermal anemometer located at a single point in a flow field provides a reading that is related to the total flow rate at standard conditions, q_{std} , through the relationship given in [Formula \(A.10\)](#):

$$q_{std} = C_{v,ta} \cdot v_{std,ta} \cdot A \quad (A.10)$$

where

- q_{std} is the volumetric flow rate through a stack or duct at standard conditions, in $\text{m}^3 \cdot \text{s}^{-1}$;
- $C_{v,ta}$ is the velocity-averaging correction factor for a single point thermal anemometer, dimensionless;
- $v_{std,ta}$ is the velocity obtained from a single-point thermal anemometer at standard conditions, in $\text{m} \cdot \text{s}^{-1}$;
- A is the cross-sectional area of the duct, in m^2 .

The numerical value of the velocity-averaging correction factor, $C_{v,ta}$, is determined by comparing the readings from a thermal anemometer operated at a single point with simultaneous data from the reference method flow rate test. It is necessary to correct the data from the reference method test to

standard conditions through use of [Formula \(A.5\)](#). The requirements for carrying out the tests are the same as those for the Pitot tube correction factor as given in [A.3.2](#).

Continuous measurements of the effluent flow rate at standard conditions can be obtained by using a rake of parallel thermal anemometers, with the individual anemometer elements placed on an ISO 10780 based grid of measurement points. Provided that the electronic signals are processed properly, the output reading of such a system is the flow rate according to [Formula \(A.10\)](#).

A.3.4 Acoustic flow meter

The reading provided by an acoustic flow meter is a distance-weighted average velocity across a line between a sending transducer and a receiving transducer. As such, the velocity reading is not directly related to flow rate, even along the line, because it is necessary to base the flow rate on an area-weighted average velocity. To obtain the flow rate, q_a , through a stack or duct from acoustic flow meter readings also requires applying a velocity-averaging correction factor, $C_{v,af}$, as shown in [Formula \(A.11\)](#):

$$q_{std} = C_{v,af} \cdot v_{std,af} \cdot A \quad (A.11)$$

where

- q_{std} is the volumetric flow rate through a stack or duct at standard conditions, in $\text{m}^3 \cdot \text{s}^{-1}$;
- $C_{v,af}$ is the velocity-averaging correction factor for an acoustic flow meter, dimensionless;
- $v_{std,af}$ is the line-average velocity obtained from an acoustic flow meter at standard conditions, in $\text{m} \cdot \text{s}^{-1}$;
- A is the cross-sectional area of the duct, in m^2 .

The approach and requirements for determining $C_{v,af}$ are the same as those for the single-point Pitot tube method.

Annex B (informative)

Modelling of particle losses in transport systems

B.1 General

Aerosol particles can be deposited on internal surfaces of transport systems as a result of mechanisms that cause particles to move transversely to air flow streamlines. This includes the phenomena of gravitational settling, inertial impaction, turbulent inertial deposition and Brownian diffusion. For most transport systems, the Brownian diffusion mechanism is of significance only for aerosol particles with sizes smaller than approximately 0,3 µm, whereas the other mechanisms are of importance for particles larger than this size. Turbulent deposition is of consequence for flows with Reynolds numbers >2 200, where the Reynolds number, Re , is as given in [Formula \(B.1\)](#):

$$Re = \frac{\rho \cdot v_m \cdot d_t}{\mu} \quad (B.1)$$

where

- Re is the Reynolds number of flow in a tube, dimensionless;
- ρ is the gas density in the stack or duct, in kg·m⁻³;
- v_m is the mean velocity of the gas in a flow tube, in m·s⁻¹;
- d_t is the tube diameter, in m;
- μ is the dynamic viscosity of a gas, in Pa·s.

For a tube of circular cross-section, the Reynolds number can also be expressed as given in [Formula \(B.2\)](#):

$$Re = \frac{4 \cdot \rho \cdot q_a}{\mu \cdot d_t \cdot \pi} \quad (B.2)$$

where

- Re is the Reynolds number of flow in a tube, dimensionless;
- ρ is the gas density in the stack or duct, in kg·m⁻³;
- q_a is the volumetric flow rate through a stack or duct, in m³·s⁻¹;
- d_t is the tube diameter, in m;
- μ is the dynamic viscosity of a gas, in Pa·s.

where q_a is the volumetric flow rate through the tube and is equal to the product of the mean velocity and the cross-sectional area.

The combination of flow rate and tube diameter of most aerosol-particle sampling systems result in turbulent flow.

Empirical or semi-empirical models for predicting the effects of the various depositional mechanisms exist for most components of a sampling system. For nozzles, the losses are controlled by inertial forces including those associated with flow turbulence, the Saffman force and, occasionally, by gravitational settling. For vertical tubes, it can be assumed that the depositional losses are controlled by turbulent, inertial deposition, and Brownian diffusion. For horizontal tubes, the losses are caused by gravitational settling, turbulent inertial deposition, and Brownian diffusion. In bends, the losses are due to the effects of inertial impaction. The predictive models typically assume that the velocity and concentration profiles are uniform at the entrance section of the component of interest. It is expected that this assumption is not fulfilled in many sampling system components because the flow disturbance created by an upstream component can affect the depositional characteristics in the succeeding component. However, in experimental studies with a composite transport system (nozzle, horizontal tube, inclined tube, vertical tube and bends), References [80] and [111] showed that the use of a model based on a sequential combination of components with assumed undisturbed inlet conditions compared well with experimental data.

B.2 Aerosol particle penetration through transport system components

B.2.1 General

The penetration, P_j , of aerosol particles through the j^{th} component of a transport system is defined in [Formula \(B.3\)](#):

$$P_j = \frac{c_{ej}}{c_{ij}} \quad (\text{B.3})$$

where

- P_j is the penetration of aerosol particles through the j^{th} component of a transport system, dimensionless;
- c_{ej} is the activity or mass or aerosol particle concentration at the exit plane of a component j , in $\text{Bq}\cdot\text{m}^{-3}$ or $\text{kg}\cdot\text{m}^{-3}$ or m^{-3} ;
- c_{ij} is the activity or mass or aerosol particle concentration at the inlet plane of the j^{th} component, in $\text{Bq}\cdot\text{m}^{-3}$ or $\text{kg}\cdot\text{m}^{-3}$ or m^{-3} .

If there are n components in a sampling system, it is assumed the overall penetration, P , can be calculated as if each component were independent, see [Formula \(B.4\)](#):

$$P = \prod_{j=1}^n P_j \quad (\text{B.4})$$

where

- P is the overall penetration of aerosol particles through a transport system, dimensionless;
- P_j is the penetration of aerosol particles through the j^{th} component of a transport system, dimensionless;
- n is the number of components in a transport system, dimensionless.

Estimates of particle losses in sampling systems should be calculated analytically or determined experimentally. Analytical calculations may be performed, for example, using the methods in this annex and in References [18], [62] or [75]. Numerical computations may be performed with one of

several available qualified computer codes¹⁾. As an example, the software Deposition Calculator¹⁾ uses several semi-analytical models to determine the depositional losses in sample lines due to gravitation settling, diffusion and turbulent inertial deposition. The models for each type of depositional loss are further refined based on laminar, turbulent and transitional flow regimes. The laminar gravitation settling model is based on Reference [49]. For the case when the flow is turbulent the model developed in References [93] and [94] is used. The turbulent eddy model, which only applies to turbulent flow, is based on work presented in References [62], [64] and [68]. Different types of transport system components are also addressed, such as certain types of inlet nozzles, straight tubes, bends, splitters and fittings that either enlarge or reduce the flow tube diameter. Losses associated with contractions in fittings are discussed in Reference [82]. Similar capabilities are implemented in other standards²⁾ and codes such as Astec/Sophaeros²⁾[59][28] and PAPAV[106][107][108].

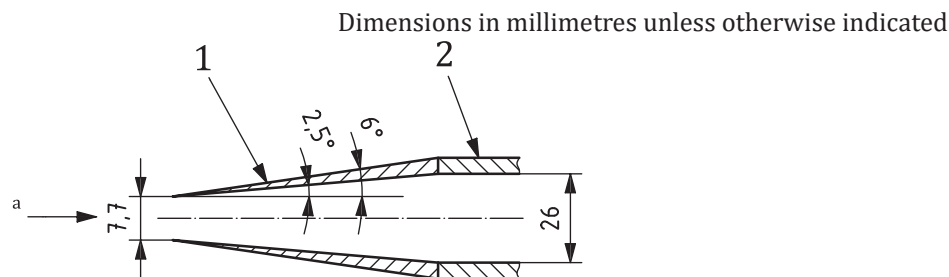
B.2.2 Wall losses in nozzles

At the date of publication of this document, there was no general model for predicting wall losses in nozzles. Reference [36] determined an experimental correlation of wall losses as a function of design and operational conditions for isokinetic nozzles of a Willeke-Okazaki configuration^[109]. These nozzles are similar in design to those in the first edition of this document, except that there is no bend. The model of Reference [36] is based on experiments with particle sizes of only 10 µm and 20 µm, so it cannot be used as a general predictive tool for all nozzle applications. Nevertheless, it provides the basis for estimating internal wall losses in the straight region of a nozzles upstream of the bends.

The software Deposition Calculator includes the models of References [36] and [105] for use with unshrouded isokinetic nozzles. The model from Reference [36] can overestimate the losses in the more modern designs of isokinetic nozzles, such as is illustrated in Figure B.1 from Reference [21]. A shrouded nozzle from Reference [79] is basically a nozzle fitted with a flow decelerator (see Figure B.2). It has lower wall losses than an unshrouded nozzle and it is less susceptible to off-design sampling conditions (e.g. off-angle flow direction, flow turbulence, changes in sampling flow rate or changes in the free-stream velocity) than an unshrouded nozzle (see Reference [22]). If the Deposition Calculator model is applied to a system fitted with a shrouded nozzle, the code calculates aerosol particle transmission based on the model of Reference [45].

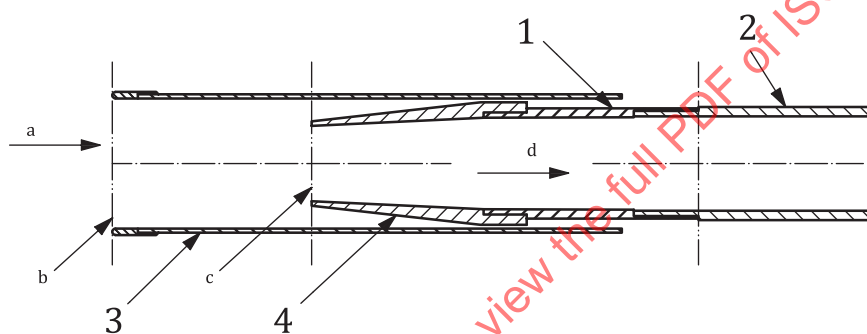
1) Contact address for Deposition Calculator:
Blunt Consulting LLC
283 Heritage Rd
Williston, SC 29853
<http://www.bluntconsulting.com>

2) Contact address for Astec/Sophaeros code:
IRSN/PSN-RES/SAG
CE Cadarache
Bât 702
13115 Saint-Paul-lez-Durance CEDEX
France

**Key**

- 1 nozzle
- 2 transport line
- a Flow.

Figure B.1 — Unshrouded nozzle of Reference [21] with wall losses of aerosol particles with a D_a of 10 μm equal to about half of those of a nozzle with a constant internal diameter

**Key**

- 1 nozzle
- 2 transport line
- 3 shroud
- 4 inner nozzle
- a Stack or duct gas flow.
- b Shroud entrance plane.
- c Nozzle entrance plane.
- d Sample flow to collector or monitor.

Figure B.2 — Shrouded nozzle

B.2.3 Straight tubes

The penetration of particles through a straight tube is calculated as given in [Formula \(B.5\)](#):

$$P_s = e^{-\frac{\pi d_t \cdot v_e \cdot L}{q_a}} \quad (\text{B.5})$$

where

P_s is the penetration through a straight tube, dimensionless;

d_t is the tube diameter, in m;

- v_e is the effective deposition velocity, in $\text{m}\cdot\text{s}^{-1}$;
- L is the length of a section of tubing, in m;
- q_a is the actual volumetric flow rate through the tube, in $\text{m}^3\cdot\text{s}^{-1}$.

The effective depositional velocity is the magnitude of the vector sum of the gravitational settling terminal velocity, which is always directed downward, and the turbulent inertial deposition and Brownian diffusion velocities, which are directed radially outward in a tube. A basic assumption when using this model is that the aerosol particles are well mixed across any cross-section of the tube.

The effective deposition velocity for an inclined tube, [Figure B.3](#), was modelled (see Reference [14]) as given in [Formula \(B.6\)](#):

$$v_e = \frac{1}{2\pi} \int_0^{2\pi} (v_d - v_{ge} \cdot \sin \alpha) d\alpha \quad (\text{B.6})$$

where

- v_e is the effective deposition velocity for an inclined tube, in $\text{m}\cdot\text{s}^{-1}$;
- v_d is the deposition velocity due to the combined effects of thermal (Brownian) diffusion and turbulent inertial deposition, in $\text{m}\cdot\text{s}^{-1}$;
- v_{ge} is the cross-stream component of gravitational settling velocity, in $\text{m}\cdot\text{s}^{-1}$.

NOTE [Formula \(B.6\)](#) is subject to the constraint that $(v_d - v_{ge} \cdot \sin \alpha)$ needs to be greater than zero. If not, it is set equal to zero. The constraint is necessary because otherwise the prediction would be equivalent to aerosol particles being transported from the environment through the top (relative to the horizontal plane) of a tube.

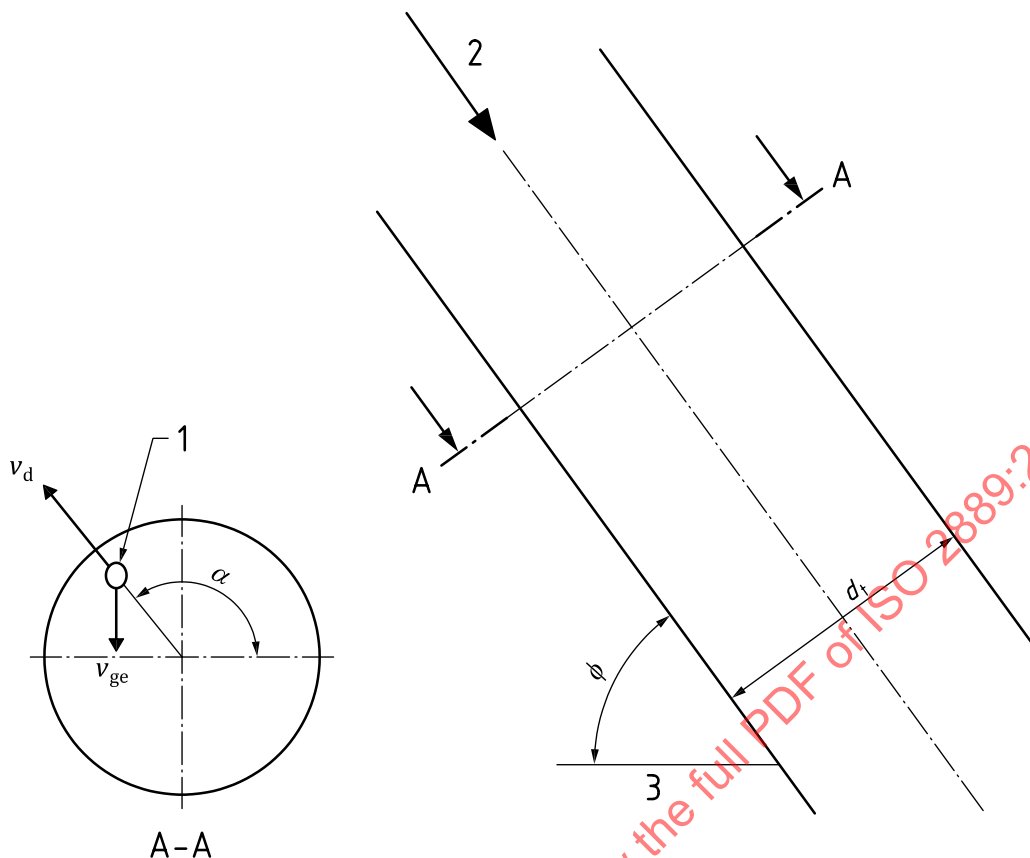
For a tube whose axis is inclined at an angle of ϕ relative to the horizontal plane, v_{ge} is as given in [Formula \(B.7\)](#):

$$v_{ge} = v_g \cos \phi \quad (\text{B.7})$$

where

- v_{ge} is the cross-stream component of gravitational settling velocity, in $\text{m}\cdot\text{s}^{-1}$;
- v_g is the sedimentation velocity of an aerosol particle, in $\text{m}\cdot\text{s}^{-1}$;
- ϕ is the inclination angle to horizontal, in rad or degrees.

If the effects of both gravitational settling velocity in the cross-stream direction and the turbulent inertial deposition velocity are of consequence, there is a tube diameter that optimizes aerosol particle penetration, because for a fixed flow rate, tube sizes smaller than the optimal value have increased turbulent depositional losses and tube sizes larger than the optimum have enhanced gravitational depositional losses.

**Key**

- 1 particle
- 2 flow
- 3 horizontal plane

Figure B.3 — Geometric model to illustrate parameters used to model particle deposition in a straight tube

A particle-deposition velocity in tubes can be correlated with a particle-relaxation time. Several semi-analytical models that lead to this correlation have been proposed. However, at the date of publication of this document, none can predict deposition of particles in the inertial size regime ($D_a > 1 \mu\text{m}$) from basic principles.

The software Deposition Calculator uses several semi-analytical models to determine depositional losses in sample line due to gravitational settling, diffusional deposition and turbulent inertial deposition. The models used for each type of depositional loss are further refined based on laminar, turbulent and transitional flow regimes. The laminar gravitation-settling model is based on Reference [49]. For the case when the flow is turbulent the model developed in References [93] and [94] is used. The turbulent eddy model, which only applies to turbulent flow, is based on work presented in References [62], [64] and [68].

B.2.4 Bends

Particle losses in bends are principally due to the effects of particle inertia where the air flow follows a curved path and the particles tend to go straight. Neglecting Brownian diffusion, turbulent inertial deposition and gravitational effects, analyses show that for two-dimensional channels, the penetration is a function of the Stokes number, St , as given in [Formula \(B.8\)](#):

$$St = \frac{C \cdot \rho_w \cdot D_a^2 \cdot v_m}{9 \cdot \mu \cdot d_t} \quad (B.8)$$

where

St is the stokes number, dimensionless;

C is the Cunningham's slip correction for aerosol particles, [\[37\]](#) dimensionless;

ρ_w is the density of water at 4 °C, equal to 1 000 kg·m⁻³;

D_a is the aerodynamic particle diameter, in µm.

The analysis for particle deposition in bends of circular cross-section is complicated by the fact that the flow is three dimensional. A secondary flow is established as the air passes through the bend, where the secondary flow consists of a set of counter-rotating vortices in which air goes from the inside of the bend to the outside of the bend along the tube cross-sectional radius. [Formulae \(B.9\)](#) and [\(B.10\)](#) hold:

$$De = \frac{Re}{\sqrt{f_{cu}}} \quad (B.9)$$

$$f_{cu} = \frac{r_{cu}}{d_t} \quad (B.10)$$

where

De is the Dean number of a flow bend, dimensionless;

Re is the Reynolds number of flow in a tube, dimensionless;

f_{cu} is the curvature ratio, dimensionless;

r_{cu} is the radius of the curvature of a pipe bend, in m.

Reference [\[86\]](#) noted that for turbulent flow, when the curvature of the bend, f_{cu} , is between 2,5 and 15, this quantity has a small effect on particle trajectories and the deposition depends mostly on St and Re . Experimental data for 90° bends with turbulent flow Reynolds numbers of 6 000 and 10 000 show that the penetration is correlated only with St as given in [Formula \(B.11\)](#):

$$P_{bend} = 10^{-0,963St} \quad (B.11)$$

It should be assumed that the uncertainty in this model increases for Reynolds numbers outside of the test range. For laminar flow [\[102\]](#), numerically modelled particle deposition in 90° bends and their results include the effects of secondary flow. They carried out calculations over a range of curvature ratios and Dean numbers and provided empirical correlations of the results, which show that the efficiency depends on the Stokes number, the curvature ratio and the Dean number.

In the software Deposition Calculator, a turbulent model of Reference [\[86\]](#) is used for $Re > 4\,000$ and $St > 0,1$ while a curve fit of the Reference [\[86\]](#) data for $Re = 1\,000$ is used for all other conditions. This method is presented graphically as [Figure B.4](#).

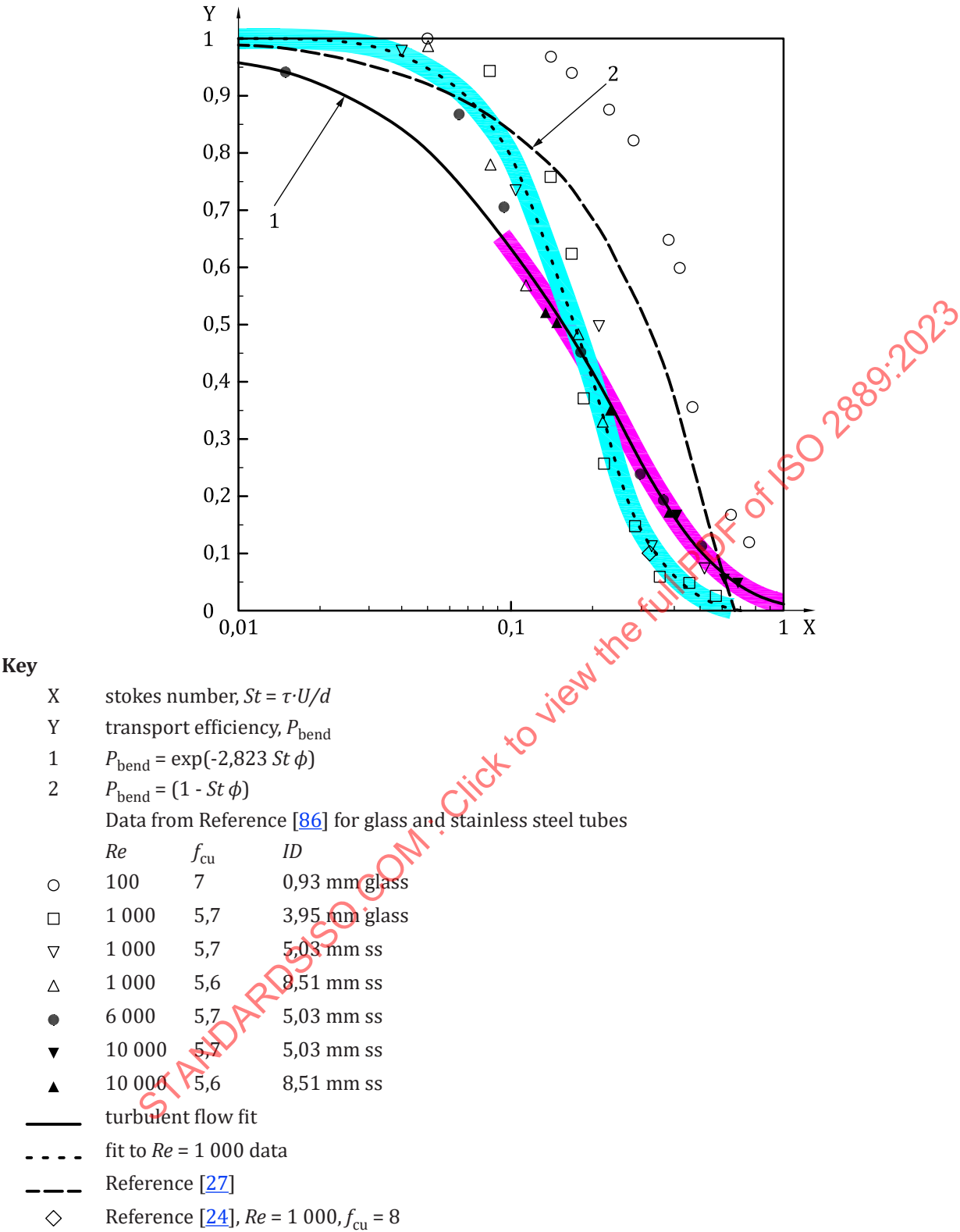


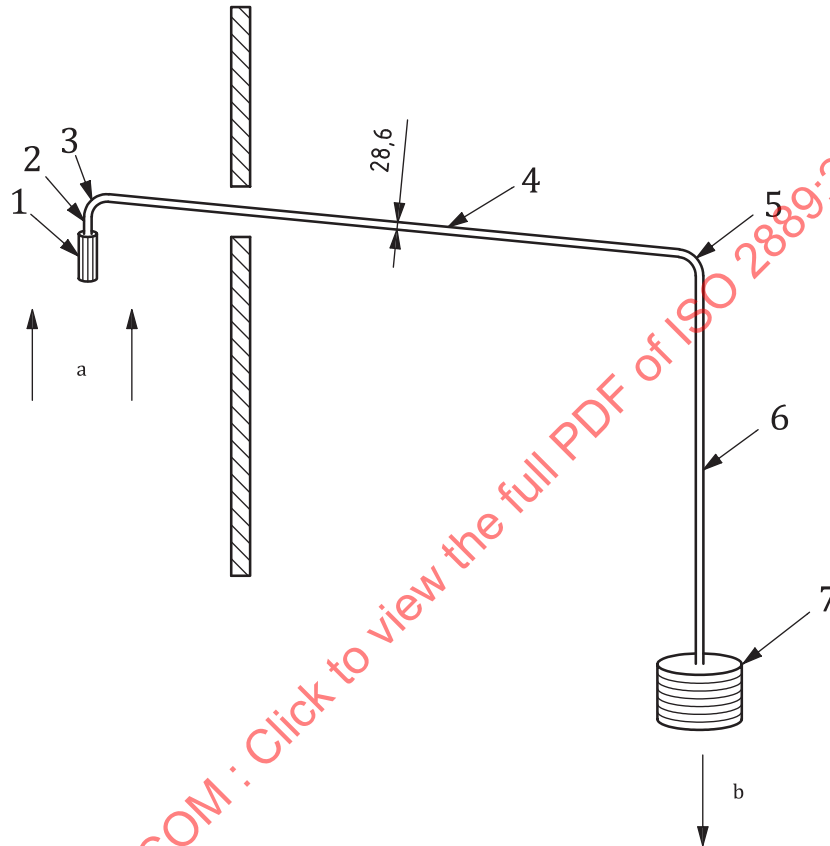
Figure B.4 — Deposition calculator bend model data

B.3 Calculation of sample losses in a transport system

As an illustration of the calculation of particle losses in a transport system, assume the geometrical configuration shown in Figure B.5 for a system that is designed to sample aerosol particles with a

D_a of $10\ \mu\text{m}$ at a flow rate of $56,6\ \text{l}\cdot\text{min}^{-1}$ from a free stream that has a velocity of $10\ \text{m}\cdot\text{s}^{-1}$. Assume a shrouded nozzle in which the velocity inside of a $52,5\ \text{mm}$ diameter shroud is $3\ \text{m}\cdot\text{s}^{-1}$ and the internal nozzle samples isokinetically from the $3\ \text{m}\cdot\text{s}^{-1}$ stream ($18,2\ \text{mm}$ diameter inner probe). Furthermore, it is assumed that the sampling tube has a $28,6\ \text{mm}$ inside diameter ($31,8\ \text{mm}$ outside diameter with a $1,6\ \text{mm}$ thick wall). This leads to the input and output values for the Deposition Calculator code shown in [Table B.1](#). The overall penetration through the system is $74,3\ \%$, with the losses predominantly occurring in the $2\ \text{m}$ long horizontal tube and the bends.

Dimensions in millimetres



Key

- 1 to 6 see [Table B.1](#)
- 7 collector or monitor
- a Flow in stack or duct.
- b Flow to vacuum source.

NOTE Free-stream velocity, equal to $10\ \text{m}\cdot\text{s}^{-1}$, is reduced to $3\ \text{m}\cdot\text{s}^{-1}$ in the shroud.

Figure B.5 — Layout of example aerosol particle transport system

Table B.1 — Example of using the software Deposition Calculator to predict aerosol particle penetration through a six-component transport system

Element		Penetration through the element ^a %
Number	Description	
1	Shrouded probe (RF-2-111): probe angle with free stream, 0°	95,08 (through nozzle)
2	Tube, 0,2 m long, 90° from horizontal	100,0
3	Bend, 90°	99,70
4	Tube, 2 m long, 0° from horizontal	77,78
5	Bend, 90°	99,70
6	Tube, 2 m long, 90° from horizontal	100,0
		Overall penetration: 73,51
^a Conditions: — flow rate: 56,6 l·min⁻¹ — tube diameter: 28,6 mm — particle density: 1 g·cm⁻³ — particle size: monodisperse, D_a equal to 10 µm — free-stream velocity: 10 m·s⁻¹		

Annex C (informative)

Special considerations for the extraction, transport and sampling of radioiodine

C.1 General

Obtaining samples of airborne radioiodine is complicated because radioiodine is present in air effluents in several forms, specifically as particulate matter, as elemental iodine (I_2), as hypoiodous acid (HOI) and in organic form, mainly as methyl iodide (CH_3I). The existence of the HOI form is not universally accepted, but it is the postulated identity of an otherwise indeterminate form with a deposition velocity lower than that of elemental iodine and that can penetrate a cadmium iodide bed but is collected by an iodophenol bed in a species sampler.

These chemical forms of radioiodine, particularly the elemental form, may be expected to deposit initially in ducts and in sampling lines and then subsequently be resuspended and emitted as the same or another form (see Reference [25]). The organic form is the least likely to be deposited and only a small fraction of it is collected by some of the solid adsorbents that are used to limit radioiodine emissions (see Reference [60]). However, it cannot be ruled out that during off-normal events the major form of radioiodine is elemental. Therefore, evaluations of sample transmission under off-normal conditions should assume that form. Reference [42] summarizes the most recent studies on radioiodine sampling and transport and many of the following considerations are based on that summary.

C.2 Sample extraction and transport

The considerations for the extraction of gases and vapour set forth in 7.5 are applicable to radioiodine. In view of the likelihood that at least some of the radioiodine in an air effluent is attached to particulates, all of the considerations applicable to aerosol particles as set forth in the main body of this document and its annexes also apply to the extraction and transport of radioiodine.

Laboratory studies have shown that, in the extraction and transport of radioiodine, materials that come in contact with the radionuclide can interact with it (e.g. copper, PVC, Buna-N). These materials should be avoided. Reference [55] indicated that the preferred materials are PTFE, polyethylene, aluminium, carbon steel and stainless steel.

Condensation of the iodine and water vapour in transport lines should be avoided by heat tracing the lines to at least 323 K (50 °C) and by avoiding abrupt temperature transitions.

Formula (C.1) from Reference [42] relates the penetration at equilibrium, P , the dimensionless ratio of the outlet concentration, c_e , to the inlet concentration, c_i , of radioiodine in transport lines to the deposition velocity and parameters of the sampling system:

$$P = e^{-4 \frac{v_d \cdot L}{v_m \cdot d_t}} \quad (C.1)$$

where

P is the overall penetration in a transport system, dimensionless;

v_d is the deposition velocity due to the combined effects of thermal (Brownian) diffusion and turbulent inertial deposition, in $m \cdot s^{-1}$;

L is the length of a section of tubing, in m;

d_t is the tube diameter, in m;

v_m is the mean velocity over the cross-section of the transport system, in $\text{m}\cdot\text{s}^{-1}$.

From [Formula \(C.1\)](#), it is evident that the penetration of radioiodine vapour is optimized by minimizing the length of the transport line, L , and using the largest diameter, d_t , and the highest flow velocity, v_m , subject to external constraints (e.g. particle transport, space availability, or collector capacity).

[Formula \(C.1\)](#) does not take into account radioactive decay or resuspension. The model in Reference [\[103\]](#) gives an equilibrium relationship that includes these effects, see [Formula \(C.2\)](#):

$$P = \frac{1}{1 + v_d \left(\frac{\pi d_t L}{q_a} \right) \left(\frac{\lambda}{\lambda + r} \right)} \quad (\text{C.2})$$

where

P is the overall penetration in a transport system, dimensionless;

v_d is the deposition velocity due to the combined effects of thermal (Brownian) diffusion and turbulent inertial deposition, in $\text{m}\cdot\text{s}^{-1}$;

L is the length of a section of tubing, in m;

d_t is the tube diameter, in m;

q_a is the volumetric flow rate at actual temperature and pressure conditions, in $\text{m}^3\cdot\text{s}^{-1}$;

r is the resuspension rate, in s^{-1} ;

λ is the decay constant, in s^{-1} .

[Formula \(C.2\)](#) suggests that the greater the resuspension rate, r , the larger the penetration. This is also illustrated in [Figure C.1](#), where penetration is shown as a function of time and resuspension rate. Reference [\[42\]](#), however, indicates that the resuspension rate of deposited radioiodine decreases for each radioiodine species as a function of time. The resuspension rates are also dependent on the amount of iodine initially deposited, some of which seems to remain firmly deposited. The latter has not been directly observed by laboratory studies but can be estimated on the basis of an activity balance following prolonged observation. Finally, Reference [\[42\]](#) states there is no satisfactory experimental verification of predicted penetration factors under either equilibrium or transient conditions. However, the following generalizations can be made from the limited available data, mostly those from Reference [\[103\]](#).

A summary of sampling systems for reactors, [\[42\]](#) indicates that a typical sampling system consists of a 15 mm diameter stainless steel transport line that is about 52 m in length with a flow rate of about $57 \text{ l}\cdot\text{min}^{-1}$. Simulated sample transport lines with a range of similar designs were tested by Unrein et al., [\[104\]](#) who measured short-term (on the order of 2 h) penetration factors of 0,62 for injected ^{131}I (as I_2) through 19 mm diameter by 48 m long tubes. Reference [\[42\]](#) predicted the equilibrium penetration factor to be about 0,75, with an approximate time of two weeks to reach equilibrium.

With the caveat that the penetration factor was not measured in the tests of Reference [\[103\]](#) until 2 h after beginning the iodine injection (lower penetration factors might have been found for measurements earlier in the test sequence), these short-term results provide a conservative estimate of penetration factors following a stepwise increase of radioiodine concentrations in conventional transport lines. The equilibrium value provides a conservative estimate for longer-term sampling of normal concentrations.

Much smaller penetration factors were found for a few tests of systems with long transport lines of 6 mm diameter tubing when operated at flow rates of less than $1,7 \text{ l}\cdot\text{min}^{-1}$. Some accident-air sampling systems used this design to reduce the potential dose at the sample collector by using a low sample flow

rate and a small-diameter transport line. Tests simulating such systems (see Reference [103]) show very poor penetration of the radioiodine to the collector. Consequently, many of these systems were redesigned to collect a low-flow-rate subsample from a high-flow, large-diameter transport line to take advantage of its favourable sample penetration from the stack. An example of the effect of flow rate on the penetration of I_2 in a small diameter transport line is shown in Figure C.2.

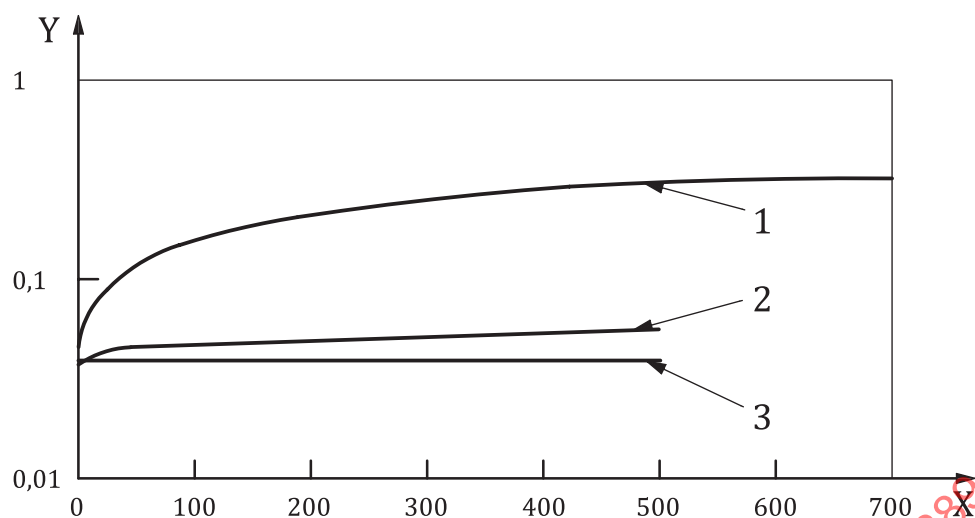
The foregoing applies primarily to the transport of elemental radioiodine and, to a lesser extent, to the hypiodous iodide form. It can be assumed that radioiodine in organic forms, with their much lower deposition velocities is transported with higher efficiencies, so that the assessment from above turns out even more conservative. As summarized in Reference [42], the fraction of organic radioiodine during normal operations appears to be quite variable from facility to facility, and the data do not provide a basis from which the organic fraction in the facility effluent during accident conditions can be estimated. At the start of the accident conditions the predominant form will be elemental or particulate.

An example for a code modelling iodine losses in transport systems is Astec/Sophaeros³⁾. It includes models for losses due to physical phenomena (condensation, sorption), chemical reactions as well as iodine particle losses. Models and examples are described in Reference [12].

C.3 Collection media for radioiodine

While carbon is an efficient collector of I_2 , it is much less efficient for iodine isotopes in organic compounds which have a low deposition velocity. Reference [61] has indicated that the removal mechanism of elemental iodine on carbon adsorbents is primarily by physical absorption, while the removal of hydrogen iodide (HI) is by physical absorption, chemical reaction and isotopic exchange. The removal of organic iodides such as CH_3I is by isotopic exchange. To improve the latter process, carbon filter media treated with potassium iodide (KI) or triethylenediamine (TEDA) should be utilized. Packages with different types of adsorbents in series are available for the collection of radioiodine with separation by chemical form.

3) Contact address for Astec/Sophaeros code:
 IRSN/PSN-RES/SAG
 CE Cadarache
 Bât 702
 13115 Saint-Paul-lez-Durance CEDEX
 France



Key

X time, in h

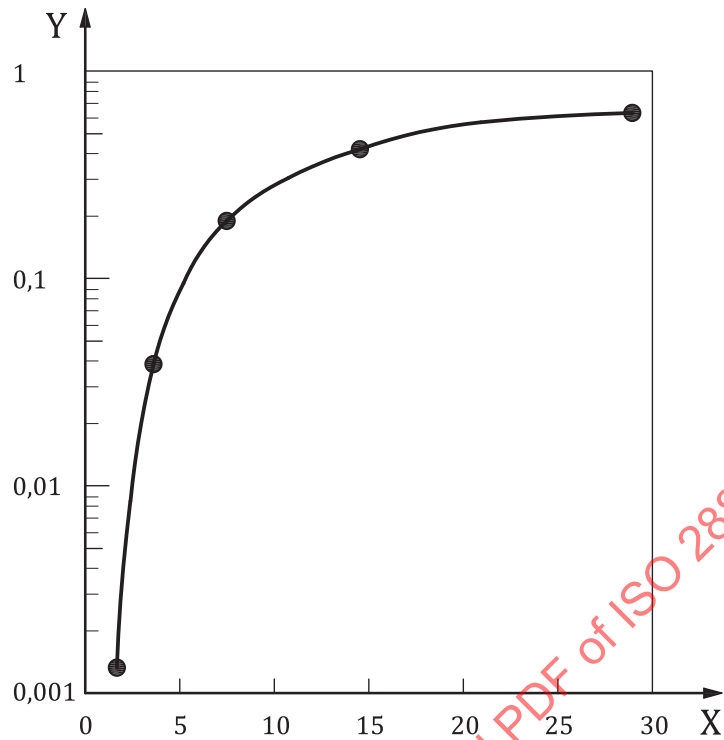
Y ratio of outlet to inlet concentration

1 curve for a resuspension rate of $1,5 \times 10^{-6}$

2 curve for a resuspension rate of $1,5 \times 10^{-7}$

3 curve for a resuspension rate of $1,5 \times 10^{-8}$

Figure C.1 — Predicted penetration of radioiodine as a function of time for various resuspension rates

**Key**

X flow rate, expressed in l·min⁻¹

Y ratio of outlet to inlet concentration

NOTE Tube size: 6,4 mm inside diameter by 43 m long.

Figure C.2 — Predicted initial radioiodine penetration through a stainless steel transport line

Annex D (informative)

Optimizing the selection of filters for sampling airborne radioactive particles

Filters are porous structures with controlled external dimensions, such as thickness and cross-sectional area normal to the flow. Filtration is the most widely used technique for collection of aerosol particles because of its low cost and simplicity. Filters capture particles by a combination of physical processes, which include direct interception, inertial deposition, Brownian diffusion, electrical attraction and gravitational sedimentation. As shown in [Figure D.1](#) (from Reference [63]), filters typically have a minimum collection efficiency for particles that are approximately 0,1 µm to 0,5 µm in diameter. Above this size, filtration efficiency increases due to inertial impaction and below this size, efficiency increases due to Brownian diffusion.

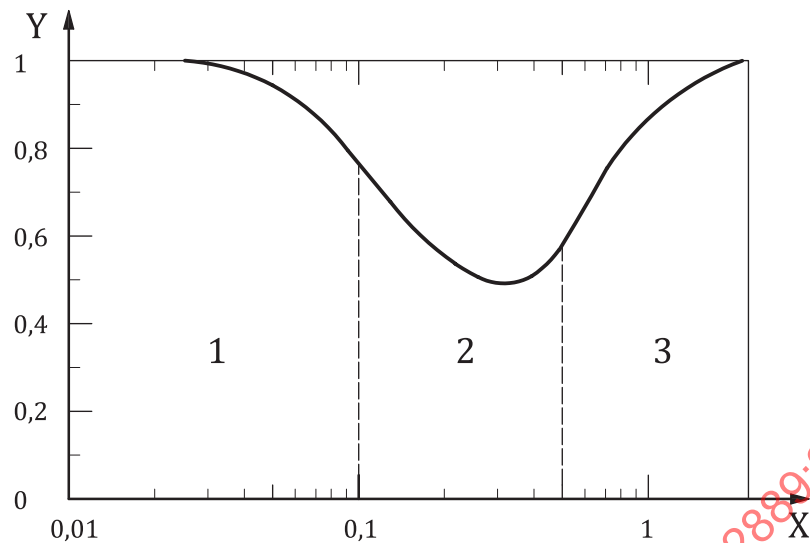
A common misconception is that filters act as sieves and that there is a direct relationship between the pore size of a filter and the minimum particle size that can be collected. In reality, because collection occurs by a complex combination of mechanisms, filters with nominal pore sizes larger than 1 µm can be very efficient collectors of sub-micrometer particles. As demonstrated by Reference [66] membrane filters show no serious degradation of collection efficiency until the pore diameters exceed 5 µm. Filters with a 5 µm pore size are often preferred because they result in lower pressure drops than smaller pore-size filters yet retain high efficiency values.

Many filter media are available for use in the collection of aerosol particles (see for example References [69] and [67]). Materials include cellulose, glass, quartz and plastic fibres. Sintered structures of metals or mineral particles are used for high-temperature filtration. Users are cautioned to be selective in their choice of filter media.

If a sample is being separated from the filter for a particular analytical method, the user should select a filter medium that can be easily dissolved by a method that does not attack the particles of interest. In other cases, it is imperative that the sample be collected on the surface of the filter rather than imbedded in the filter. Reference [50] has demonstrated that absorption of alpha radiation emitted from airborne particles collected on glass-fibre filters does not constitute a major source of error in estimating concentrations of airborne alpha-emitting radionuclides, but the excellent resolution in alpha spectroscopy requires the use of membrane-type filters that are front-surface collectors.

Decisions on changing the filter media should include considerations such as the potential loss of continuity between historical and future sampling results, potential impacts on vacuum-system performance, requirements for analyser retesting, requirements for revision and approval of documentation, retraining requirements for workers and potential impacts on secondary uses of the filter samples, such as periodic chemical analyses for process control. However, some filter media date back many decades and their continued use is not justified simply because of historical precedents.

[Table D.1](#) summarizes the type of information that is useful for selecting an appropriate filter for sampling airborne radioactive particles. This table includes a variety of coarse-fibre, glass-fibre and membrane-type filters, but does not constitute an endorsement of any particular manufacturer or filter type. Conversely, the absence of any particular filter from the example table does not constitute a rejection of that medium. For general sampling applications, information is provided on durability, flow resistance and efficiency. Information related to the collection of radon decay products and the energy resolution for alpha spectroscopy is included for alpha continuous air monitor (CAM) applications. The alpha spectroscopy resolution is based on detection of the 6,0 MeV alpha emission of ^{218}Po (a naturally-occurring decay product of ambient ^{222}Rn , which causes interference in instruments used to detect plutonium or uranium isotopes).

**Key**

- X particle diameter, in μm
- Y efficiency
- 1 diffusion regime
- 2 diffusion and interception regime
- 3 inertial impaction and interception regime

Figure D.1 — Schematic of filter efficiency versus particle size illustrating the different filtration regimes

Filter efficiencies range from >99,999 % at all particle sizes and flow rates for the Millipore type AA, 0,8 μm pore size membrane filter to >50 % for the Whatman 41 cotton cellulose filter at low flow rates and small particle sizes. Typical flow rates range from 4 $\text{l}\cdot\text{min}^{-1}\cdot\text{cm}^{-2}$ at 35 kPa pressure drop for the Millipore Type AA membrane filter to 59 $\text{l}\cdot\text{min}^{-1}\cdot\text{cm}^{-2}$ for the Millipore Fluoropore, 5 μm pore-size polytetrafluoroethylene (PTFE) membrane filter. Resolution for alpha spectroscopy of the ^{218}Po alpha emission at 6,0 MeV ranges from as low as 350 keV (full width at half maximum) for the Fluoropore 3 μm PTFE membrane filter to greater than 1 500 keV for the Whatman 41 cotton-cellulose fibre filter. The poor resolution associated with the Whatman 41 filter makes that filter unsuitable for use in continuous air monitors that are employed to detect plutonium or uranium in the presence of ambient radon decay products. In addition, although the Whatman 41 is easily dissolved for chemical analyses, it has a collection efficiency that decreases dramatically at low flow rates. The Fluoropore filters have a very low pressure drop, good collection efficiency and excellent resolution for alpha spectroscopy, but are not readily dissolved for radiochemical procedures. The selection of the larger pore 5 μm Fluoropore filter over the 3 μm pore option provides a substantial improvement in flow rate with only a modest decrease in sampling efficiency and resolution for alpha spectroscopy. [Table D.2](#) provides other information useful for the selection of sample filters.

As new filter types become available, comparisons such as these can be made by the user to ensure that appropriate filter types are selected for sampling radioactive aerosol particles.

**Table D.1 — Characteristics of filters evaluated for use in sampling radioactive particles
(adapted from References [53] and [72])**

Filter type	Filter composition and durability	Typical flow rate ^a l·min ⁻¹ ·cm ⁻²	FWHM ^b of the ²¹⁸ Po peak keV	Relative radon progeny counts in the Pu ROI ^c	Relative radon progeny collection efficiency ^d	Filter efficiency range ^e %
Millipore type SMWP (5,0 µm pore size), Millipore Corp., Bedford, MA	Mixed esters of cellulose acetate and cellulose nitrate (fragile; electrostatic; both sides identical)	16	670	1	1	98,1 to >99,99
Millipore type AW19 (5,0 µm pore size), Millipore Corp.	Homogeneous, micro-porous polymers of cellulose esters formed around a cellulose web (rugged; both sides identical)	16	470	0,57	0,99 ± 0,01	99,93 to >99,99
Durapore (5,0 µm pore size), Millipore Corp.	Polyvinylidene fluoride (rugged; both sides identical)	14	790	1,55	0,67 ± 0,1	—
Fluoropore (3,0 µm pore size), Millipore Corp.	Polytetrafluoroethylene (PTFE) bonded to polypropylene high-density fibers (rugged; front is membrane; back is fibres; sides barely distinguishable by naked eye)	23	350	0,47	1,04 ± 0,02	98,2 to >99,98
Fluoropore (5,0 µm pore size), Millipore Corp.	Polytetrafluoroethylene (PTFE)	—	—	—	—	—
Versapor 3 000 (3,0 µm pore size, Gelman Sciences, Ann Arbor, MI)	Acrylic copolymer on a nylon fibre support (rugged; both sides identical)	25	590	0,94	0,75 ± 0,02	99,7 to >99,99
Gelman type A/E (~1,0 µm pore size) Gelman Sciences	Borosilicate glass fibre without binder (breakable during handling; both sides identical)	25	≥1 000	1,31	0,92 ± 0,01	99,6 to >99,99

^a Flow rate determined under vacuum at 35 kPa.

^b FWHM is the typical full width at half maximum of the ²¹⁸Po peak obtained with a 2,5 cm diameter filter and a 2,5 cm diameter solid state detector with a 0,5 cm separation distance during sampling of room air at Lovelace Biomedical in Albuquerque, NM.

^c Radon progeny background counts in the Pu ROI for the filter of interest, divided by similar counts obtained simultaneously on a Millipore SMWP filter.

^d Total radon progeny background counts on the filter of interest, divided by similar counts obtained simultaneously on a Millipore SMWP filter. Mean and standard uncertainty for five replicate tests.

^e The range of filter efficiency values given generally corresponds to a particle diameter range of 0,035 µm to 1 µm, a pressure drop of 1 cm to 30 cm Hg, and a face velocity range of 1 cm·s⁻¹ to 100 cm·s⁻¹. Values are from References [69], [51] and [52].

NOTE There are several trade names mentioned in Tables D.1 and D.2. This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of the products named.

Table D.1 (continued)

Filter type	Filter composition and durability	Typical flow rate ^a l·min ⁻¹ ·cm ⁻²	FWHM ^b of the ²¹⁸ Po peak keV	Relative radon progeny counts in the Pu ROI ^c	Relative radon progeny collection efficiency ^d	Filter efficiency range ^e %
Whatman EPM 2 000, Whatman LabSales, Hillsboro, OR	Borosilicate glass microfibre without binder (breakable during handling; both sides identical)	20	≥1 000	1,48	1,00 ± 0,03	—
Whatman 41, Whatman LabSales	Cotton-cellulose filter paper (rugged; currently used primarily for liquid filtration; both sides identical)	25	≥1 500	1,65	0,42 ± 0,01	43 to >99,5
Nuclepore (0,6 µm pore size), VWR Scientific, Pleasanton, CA	Polycarbonate membrane (rugged; thin; very electrostatic; currently used primarily for liquid filtration; collection side recommended by manufacturer is the shiny side)	4	500	0,89	0,85 ± 0,02	53 to >99,5
Millipore type AA (0,8 µm pore size), Millipore Corp.	Mixed esters of cellulose (fragile; electrostatic; collection side is darker)	7	520	0,91	1,05 ± 0,01	99,999 to >99,999

^a Flow rate determined under vacuum at 35 kPa.

^b FWHM is the typical full width at half maximum of the ²¹⁸Po peak obtained with a 2,5 cm diameter filter and a 2,5 cm diameter solid state detector with a 0,5 cm separation distance during sampling of room air at Lovelace Biomedical in Albuquerque, NM.

^c Radon progeny background counts in the Pu ROI for the filter of interest, divided by similar counts obtained simultaneously on a Millipore SMWP filter.

^d Total radon progeny background counts on the filter of interest, divided by similar counts obtained simultaneously on a Millipore SMWP filter. Mean and standard uncertainty for five replicate tests.

^e The range of filter efficiency values given generally corresponds to a particle diameter range of 0,035 µm to 1 µm, a pressure drop of 1 cm to 30 cm Hg, and a face velocity range of 1 cm·s⁻¹ to 100 cm·s⁻¹. Values are from References [69], [51] and [52].

NOTE There are several trade names mentioned in Tables D.1 and D.2. This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of the products named.

Table D.2 — Other characteristics of filters

Filter type	Product	Specification	Binding agent	Pore size/ Fibre diameter	Thickness	Area weight	Collection efficiency	Flow resistance $v_a=0,34 \text{ m/s}$ $\text{Pa}\cdot\text{s}\cdot\text{m}^{-1}$	Temperature max. °C	Burst pressure Wet kPa	Dry kPa	Ash residue %
Fibre filter	GF 6	Borosilicate fibre	Inorganic	—	350	80	99,97	15 000	500	>6	>18	>90
Fibre filter	GF 8	Borosilicate fibre	Inorganic	—	350	75	99	7 000	500	>4	>10	>90
Fibre filter	GF 9	Borosilicate fibre	Inorganic	—	350	70	99,97	7 000	500	>5	>10	>90
Fibre filter	GF 10	Borosilicate fibre	Organic	—	350	70	99,97	6 500	180	>10	>30	>85
Fibre filter	MN 85/90	Borosilicate fibre	Organic (acrylic resin)	0,4 to 1,8	400	90	99,99	6 500	250	—	—	—
Fibre filter	GF/A	Borosilicate fibre	None	—	260	—	—	—	500	2	—	—
Membrane	AE 99	Cellulose nitrate	—	8	—	44	99,95	10 650	125	—	—	—
Membrane	W 41	Cellulose	—	5	220	85	70	5 800	—	1,5	69	0,007
Membrane	SMWP	Cellulose acetate and cellulose nitrate	—	5	—	—	>98	—	—	—	—	—
Membrane	AW 19	Cellulose ester	—	5	—	—	99,9	—	—	—	—	—
Membrane	TE 38	PTFE-laminated supported by polyester-tissue	—	5	270	—	—	8 200	145	—	—	—
Capillary pore filter	—	Polycarbonate	—	0,6	10	—	53 to 99,5	—	—	—	—	—

Annex E (informative)

Evaluating the errors and the uncertainty for the sampling of effluent gases

E.1 General

The error in the determination of a quantity, such as the amount of radioactive substance emitted from a stack or duct during a certain period, can be defined as the difference between its actual value and the measured value. Knowledge of the actual value and the measured value can usually be gained when calibrating or qualifying a measurement procedure or instrument against standards, physical constants or scientific laws. In most situations the upper and lower bounds of uncertainty limit the possible quantifiable errors (see Reference [81]). Generally, the uncertainty consists of random uncertainties (Type A uncertainty) and systematic uncertainties such as, for example, model assumptions (Type B uncertainty, or bias). For a measurement a determination of the limits of uncertainty is needed that properly and completely combines both sources of uncertainty (see References [2] and [8]).

A simple stack-effluent measurement system, such as a filter air sampler (FAS) connected by a transport line to a nozzle, is used to illustrate a typical effluent measurement process. To estimate the amount of radioactive substance emitted from a stack, a sample of the effluent is extracted, transported to a collection medium and collected. It is further necessary to quantify the collected material and relate the quantified amount to the effluent. For the estimation of the effluent concentration, it is necessary to know or estimate the effluent flow rate, the area of the sampling plane, the sample flow rate, the nozzle transmission, the mixing ratio of constituents in the duct, the sample transport penetration, the sample collection efficiency and the analysis efficiency.

The sampling and analysis processes contribute both systematic uncertainties and random uncertainties to the overall uncertainty in the estimated activity. In the case of the extraction-plane parameters and instrument calibrations, both systematic uncertainty and Type A uncertainties are combined into one uncertainty that is assumed to also be relevant during normal operations. For example, during the calibration of the measurement devices, a known and constant input is provided and output or measured response is observed. The systematic uncertainty is determined by the offset of the mean value from the expected standard response and Type A uncertainty by the random variation in the output. In other elements, such as the nozzle inlet or transport line, calibration is not possible but surveys with standard aerosol particles under known, controlled conditions can be conducted to establish bounds related to the expected performance. Similarly, uncertainty estimates can be put on parameters such as the area of the sample-extraction plane, the degree of mixing of contaminants in that plane, and the mean axial velocity of the effluent through that plane. A calibration removes part of the systematic uncertainty but not the combined uncertainty.

The systematic uncertainty that remains embedded in the predetermined parameter or calibrated device elements is not fully known in a particular application. Bounds can be placed on the embedded uncertainties. The random components of the combined uncertainty arise during the calibration. They are estimated by the standard deviations of the variable measurement components, such as the sample flow rate or the activity in the sample. Such standard deviation estimates are derived from the data alone without external reference.

E.2 Uncertainty estimation related to the emission of a radioactive substance

Regulatory limits on radionuclide emissions are generally stated in terms of limits on the resultant dose per year to members of the public. It is therefore necessary that a facility sampling and measurement system generates accurate and reliable estimates of quantities emitted in a given

sampling interval that can then be inserted into models for environmental transport and dose estimation. The average activity emission rate over the period of integration, $\bar{A}$ (becquerel per second), related to a radionuclide that is being transported out of a stack or duct can be represented as given in [Formula \(E.1\)](#):

$$\bar{A} = c_A \cdot v_m \cdot A \quad (\text{E.1})$$

where

- $\bar{A}$ is the activity emission rate of a radionuclide averaged over the period of integration, in $\text{Bq}\cdot\text{s}^{-1}$;
- c_A is the effluent activity concentration of a radionuclide, in $\text{Bq}\cdot\text{m}^{-3}$;
- v_m is the mean axial velocity over the cross-section of the transport system, in $\text{m}\cdot\text{s}^{-1}$;
- A is the cross-sectional area of a stack or duct, in m^2 .

The first factor in [Formula \(E.1\)](#), c_A , the effluent activity concentration, is determined by activity measurement, measured sample volume and constant parameters of the system, as given in [Formula \(E.2\)](#):

$$c_A = \frac{r_n}{V_a \cdot P \cdot f_c \cdot \varepsilon_f \cdot \varepsilon_d} \quad (\text{E.2})$$

where

- c_A is the effluent activity concentration of a radionuclide, in $\text{Bq}\cdot\text{m}^{-3}$;
- r_n is the net count rate of the sample, in s^{-1} ;
- V_a is the volume of effluent that produced the sample at actual conditions, in m^3 ;
- P is the overall penetration in a transport system, dimensionless;
- f_c is the ratio of the activity concentration in the sample volume to the effluent activity concentration in the free stream, dimensionless;
- ε_f is the collection efficiency of the collection medium, dimensionless;
- ε_d is the detection efficiency, dimensionless.

The second factor of [Formula \(E.1\)](#), v_m , is the average effluent flow velocity, typically determined by pre-operational measurements along traverses across the duct. Other methods and devices can be used such that near-real-time data for v_m are obtained.

In ISO 10780, the mean axial velocity of the effluent is computed from the results of multiple velocity determinations, one at each subsection of the cross-sectional area over the sampling plane. This determination is made before and during start-up operations and not during normal operations. The relation between v_m at a particular location and the actual parameters measured with a Pitot tube is as given in [Formula \(E.3\)](#):

$$v_m = v \cdot \cos \theta = C_{\text{cal,pt}} \cdot \cos \theta \cdot \sqrt{\frac{2 \cdot \Delta p}{\rho_a}} \quad (\text{E.3})$$

where

- v_m is the mean axial velocity over the cross-section of the transport system, in $\text{m}\cdot\text{s}^{-1}$;

- $C_{\text{cal,pt}}$ is the Pitot calibration factor, dimensionless;
- Δp is the differential pressure, in Pa;
- ρ_a is the gas density in the stack or duct determined from measured static pressure, molar mass of the gas and temperature, using the equation of state, in $\text{kg}\cdot\text{m}^{-3}$;
- θ is the flow angle, in rad or degrees.

The last factor of [Formula \(E.1\)](#), the flow cross-sectional area, A , is also predetermined from blueprints or from measurements across the stack or duct. That area is divided into equally sized subsections for purposes of measuring an average axial velocity.

The stack or duct emission rate, $\bar{A}$ (becquerel per second), is calculated by combining [Formulae \(E.1\)](#), [\(E.2\)](#) and [\(E.3\)](#) as given in [Formula \(E.4\)](#):

$$\bar{A} = r_n \cdot \frac{A \cdot C_{\text{cal,pt}} \cdot \cos \theta \cdot \sqrt{\frac{2 \cdot \Delta p}{\rho}}}{V_a \cdot P \cdot f_c \cdot \varepsilon_f \cdot \varepsilon_d} = r_n \cdot w \quad (\text{E.4})$$

where

- $\bar{A}$ is the activity emission rate of a radionuclide averaged over the period of integration, in $\text{Bq}\cdot\text{s}^{-1}$;
- r_n is the net count rate, in s^{-1} ;
- A is the cross-sectional area of a stack or duct, in m^2 ;
- $C_{\text{cal,pt}}$ is the Pitot calibration factor, dimensionless;
- Δp is the differential pressure, in Pa;
- ρ is the gas density in the stack or duct determined from measured static pressure, molar mass of the gas and temperature, using the equation of state, in $\text{kg}\cdot\text{m}^{-3}$;
- θ is the flow angle, in rad or degrees;
- V_a is the volume that produced the sample at actual conditions, in m^3 ;
- P is the overall penetration in a transport system, dimensionless;
- f_c is the ratio of the activity concentration in the sample volume to the effluent activity concentration in the free stream, dimensionless;
- ε_f is the collection efficiency of the collection medium, dimensionless;
- ε_d is the detection efficiency, dimensionless;
- w is the calibration factor, dimensionless.

[Formula \(E.4\)](#) is the mathematical model of the emission measurements.

E.3 Quantifying uncertainty

E.3.1 Stack or duct emission measurement uncertainty analysis methods

An emission release rate is a function of a large number of measured parameters as contained in [Formula \(E.4\)](#). A basic analysis of uncertainty can be carried out under the assumption that the

distribution of Type A uncertainties is almost normal, this is valid for most, but not necessarily all, of the related quantities.

E.3.2 Combined uncertainty associated with the measurement process

An uncertainty analysis of the measurement processes that take place either in the pre-operational or operational phases has been developed. Each of the terms in the emission model has been examined and expanded as appropriate. This is a recommended approach that may be more fruitful than attempting to formulate a comprehensive and complex analysis. The total emission rate from the stack or duct is calculated from activity measurements of a continuously taken air sample extracted from a qualified sampling location, and other measurements of parameters in the emission equation. The combined uncertainty can then be estimated by summing over each of the terms of the final uncertainty equation (Reference [81]).

E.3.3 Uncertainty associated with bias

The many components of systematic uncertainty in an emission measurement can be combined into one term following the method given in Reference [4]. These components can be classified as calibration, parameter estimate, and design and test systematic uncertainties for components such as nozzles and transport lines.

The calibration systematic uncertainty represents the overall calibration uncertainty, combining elements of both systematic uncertainty and random uncertainty, that accrue during calibration of instruments or devices. This is typically the residual systematic uncertainty remaining after gross systematic uncertainty is zeroed out (up to an acceptance limit) during calibration. In some cases, it is identified by manufacturers as the accuracy limit of the calibrated scale or readout device.

The parameter systematic uncertainty represents the overall uncertainty (again combining Type A and Type B uncertainties) that is derived from field and laboratory determinations of sample-extraction parameters, such as the degree of particulate mixing, f_c , or mean profile velocity, v_m , that ultimately are applied as a single value to estimations of stack or duct emissions. Other systematic uncertainty contributions related to sampling location effects are treated separately below.

The design and test systematic uncertainty term results from residual uncertainties in the process of design and qualification of sampling nozzles and devices that are engineered from first principles and empirical factors, and then manufactured and tested to confirm that certain performance characteristics, such as nozzle-inlet penetration efficiency, have been met. Such a critical performance characteristic is specified by an acceptable range for a given set of operating conditions, and there is an acceptance limit on deviation between the design performance and the result of confirmation tests with tracer materials.

The combined systematic uncertainty limit serves the same purposes for the systematic uncertainty as does the two-standard-deviation random uncertainty estimate. It can be combined with the 95 % confidence limit estimate of the Type A uncertainty component in calculating the overall uncertainty of measurement. A full uncertainty analysis entails a careful consideration of the many sources of uncertainty and proper combining of each component to generate an overall uncertainty estimate.

The [Table I.1](#) guidance levels for acceptable accuracy and precision of measurement require evaluation of factors that can contribute uncertainty to components of the system. Without detailed analysis, it can be difficult to demonstrate that estimates are reasonably complete and defensible. The discussion in the remainder of this annex is meant to provide a framework for an evaluation of uncertainty in an effluent measurement based on an analytic expression relating radionuclide emission to variable parameters in the measurement, and on residual systematic uncertainty and implicit uncertainty inherent in the methodology of continuous emission monitoring from a single point.

It is worthwhile to reflect here on what interpretation should be given to the “true value” of the measured effluent-radioactive-substance emission rate by the single-point representative sampling method. The fact that flow and mixing parameters are averaged over the profile implies that the intended “true” emission rate is the “conceptual value” at the sampling location, i.e. that the area-averaged emission rate of radioactive substance from the stack or duct based on the axial locations of

the nozzle(s), assuming that the installed instrumentation does not disturb either the concentration or the flow distributions and that the pre-operational measurements of critical parameters, i.e. v_m , P , V_a and f_c , properly reflect sampling under operational conditions. The conceptual value shall be distinguished from the “available value”, which is the emission rate estimated only at the axial location of the nozzle(s), without assumptions about disturbance, mixing or how representative the parameter estimates are. Because it is the well mixed, mean effluent radioactive substance emission rates that are of interest, the effects of the non-uniform distribution of velocity and contaminant mal-distribution shall be added to the list of sources of uncertainties, and uncertainties in the correction factors that account for the non-uniform distribution shall be considered. The estimation of the true emission rate (the conceptual value) depends on parameters measured at other times, under possibly different conditions, and with test aerosol particles and gases rather than the actual radioactive contaminants. So it is evident that the uncertainty in the result of a single-point sample depends on considerations other than explicit operational or pre-operational measurement uncertainties.

E.3.4 Uncertainty associated with conceptual systematic uncertainty

E.3.4.1 General

As noted in [E.1](#), a useful distinction can be made between the uncertainty of the measurement processes and the uncertainty associated with the overall methodology. The measurement uncertainty is attributable to explicit performance aspects of the sampling and analytic hardware. Sampling methodology systematic uncertainty derives from issues related to sample extraction location and related implicit factors that affect how well the sample represents the true emissions from the stack or duct during the period of the sample. These are what Reference [\[81\]](#) terms “pattern factors”, defined as factors that describe variation in velocity and mixing in the profile and an estimate of how the value of the emission rate at the nozzle location compares with the mean value. In some situations, the conceptual systematic uncertainty can be the largest by far. Two of the more significant sorts of conceptual systematic uncertainty derive from assumptions about temporal variation and about the completeness and accuracy of model assumptions.

E.3.4.2 Uncertainty associated with temporal variations

Changes over time in stream conditions following site qualification are assumed not to significantly influence the measurement outcome. Yet, increases or decreases in volumetric sample flow, effluent discharge rate or modifications in the stack or duct use can cause distortions in sample extraction or contaminant mixing. The degree to which a single-point sample is representative can then change. This is why it is particularly important that those factors that are counted on to produce a well-mixed condition at a qualified sample-extraction location be robust with respect to reasonably anticipated changes in stack or duct conditions. Mechanical mixing elements or deflected, colliding flows produce mixing conditions at the selected sample extraction plane that are resistant to change under modifications in facility use or under upset conditions, and so their use is compatible with small values of estimated uncertainty. A judgement is required about assigning the significance of temporal variations and, hence, the magnitude of this uncertainty term.

E.3.4.3 Model systematic uncertainty

There are a number of simplifying model assumptions implicit in the representation of continuous emission monitoring by extractive sampling from a single point. Among these are that the contaminant-transport processes in the sample nozzle and line are well represented by semi-empirical models (see [Annex B](#)), that the measured uniformity of the velocity profile and degree of mixing at the stage of site qualification continue to apply during operations (a pattern assumption), and that the activity-measuring process is well represented by the single-parameter sample collection and radiation-detection efficiencies. To varying degrees, model-based assumptions might not be fully correct for a particular application. Again, a judgment is required to assign an estimate to this uncertainty term. Both the temporal variation uncertainties and model systematic uncertainty can be combined with other fixed uncertainties arising from various measurements in generating an overall systematic uncertainty limit estimate.

The user is advised to become very familiar with the assumptions and limitations of the models used to optimize sample-transport-line design to ensure that special provisions properly take into account the deposition of highly reactive species, particle bounce or resuspension from previously deposited materials. The same applies to models of other aspects of the sampling and measuring process. The use of computational modelling aids should be viewed as a part of a larger process of design and evaluation that should include data from laboratory studies, findings in the peer-reviewed literature and field testing.

E.3.5 Describing the combined uncertainties in emission measurement

The last step of an uncertainty analysis is bringing together the 2σ random uncertainties into a combined estimate of overall uncertainty stated in terms of the emission estimate, $\bar{A}$, the estimated uncertainty limit of the effluent measurement (from References [2], [3] and [8]), as given in [Formulae \(E.5\) to \(E.7\)](#):

$$u^2(\bar{A}) = u^2(r_n) \cdot w^2 + r_n^2 \cdot u^2(w) \quad (\text{E.5})$$

where

$u(\bar{A})$ is the standard uncertainty of the activity emission rate of a radionuclide, in $\text{Bq}\cdot\text{s}^{-1}$;

r_n is the net count rate, in s^{-1} ;

w is the calibration factor, in s^{-1} ;

$u(r_n)$ is the standard uncertainty of the net count rate, in s^{-1} ;

$u(w)$ is the standard uncertainty of the calibration factor, in s^{-1} .

For this document, the calibration factor, w , is interpreted to include many sources of Type A error as shown in [Formula \(E.6\)](#).

$$\begin{aligned} u_r^2(w) &= \frac{u^2(w)}{w^2} = \\ &= u_r^2(A) + u_r^2(C_{\text{cal,pt}}) + u_r^2(p) + u_r^2(\rho) + u_r^2(V_a) + \\ &\quad + u_r^2(P) + u_r^2(f_c) + u_r^2(\varepsilon_f) + u_r^2(\varepsilon_d) \end{aligned} \quad (\text{E.6})$$

where

$u_r(w)$ is the relative standard uncertainty of the calibration factor, dimensionless;

$u_r(A)$ is the relative standard uncertainty of the cross-sectional area, dimensionless;

$u_r(C_{\text{cal,pt}})$ is the relative standard uncertainty of the Pitot calibration factor, dimensionless;

$u_r(p)$ is the relative standard uncertainty of the pressure, dimensionless;

$u_r(\rho)$ is the relative standard uncertainty of the gas density, dimensionless;

$u_r(V_a)$ is the relative standard uncertainty of the volume at actual conditions, dimensionless;

$u_r(P)$ is the relative standard uncertainty of the overall penetration in a transport system, dimensionless;

$u_r(f_c)$ is the relative standard uncertainty of the ratio of the activity concentration in the sample volume to the effluent activity concentration in the free stream, dimensionless;

$u_r(\varepsilon_f)$ is the relative standard uncertainty of the collection efficiency of the collection medium, dimensionless;

$u_r(\varepsilon_d)$ is the relative standard uncertainty of the detection efficiency, dimensionless.

And finally, when a coverage interval is desired, the expanded relative uncertainty is given by [Formula \(E.7\)](#):

$$u_{r,k=2}(\bar{A}) = 2 \cdot u_r(\bar{A}) \quad (\text{E.7})$$

where

$u_{r,k=2}(\bar{A})$ is the relative standard uncertainty of the activity emission rate of a radionuclide representing the coverage interval at 95 % ($k = 2$), dimensionless;

$u_r(\bar{A})$ is the relative standard uncertainty of the activity emission rate of a radionuclide, dimensionless.

E.4 Evaluation of uncertainties

E.4.1 General

According to Reference [19], estimates of the magnitude of most of the uncertainty terms can be described at least to the level of what is attainable at the 95 % confidence level (corresponding to a 2σ interval for random variables). Many of the uncertainties considered above are relatively small and controllable by good practice. Others require more careful consideration.

E.4.2 Uncertainty in sample volume, stack or duct area, and transmission efficiency

Uncertainty in sample volume, V , and stack or duct area, A , are generally small and well understood. Sample volume measurement is readily accomplished and corrected for altitude and temperature. As described in [J.3.3](#), the accuracy of the measurement of the sample flow (and hence sample volume) with a flow meter should be periodically checked with a secondary standard flow meter, and differences maintained to less than 10 % of standard. Expanded uncertainty with a factor of 2σ in the order of 5 % should be achievable.

The cross-sectional area of the effluent flow at the sample-extraction location should be accurately ascertained from engineering drawings of the effluent stack. Systematic uncertainty associated with this determination should be much less than 2 %.

Sample-transport-line penetration in the case of particulate effluent is harder to estimate because the characteristics of the aerosol particles being sampled cannot be fully described in advance. Although the particle-size characteristics of many types of radioactive aerosols have been studied (see [Annex G](#) for a discussion of such studies), many possibilities for high-efficiency particulate air filter (HEPA filter) failure and associated unique aerosol-particle-size distributions exist. Nonetheless, an estimate of the magnitude of uncertainty in this parameter can be made. The estimate may be based on in-place tests with particles of either conservative or realistic size characteristics. The estimate may also be obtained with the aid of suitable particle-penetration models (see [Annex B](#)) and appropriately varied parameters. It is expected that a proposed transport line for a sampling system is designed for optimal performance using a conservative assumption, for example, 10 μm monodisperse aerosol particles. A conservative design establishes the lower limit of penetration because polydisperse aerosol particles with an equal or smaller geometric mean have greater transmission efficiency. So, in the case of the system analysed in [Table B.1](#), the range of possible transmission efficiencies for particles having a D_g smaller than 10 μm exceeds 74,3 %. In cases where additional data about the relevant size distribution (e.g. activity size distribution) are available, the test or design aerosol particle size may be selected accordingly.

For example, assume that there is a polydisperse log-normal distribution of particle sizes with an average activity median aerodynamic diameter (AMAD) of 1,8 μm and $\sigma_g = 2,2$, like that found in a

research and development facility glovebox line by Reference [35]. Further assume that the model predicted transmission efficiency for the fictional sampling system is 94,2 %. For uncertainty analysis purposes, suppose the aerosol particles actually encountered in an accident effluent is more like the fabrication facility average size distribution reported in Reference [35] (AMAD of 4,0 μm , $\sigma_g = 1,7$). Then, the predicted transmission efficiency can be 85,9 %, a relative difference of 8,8 % compared with the 94,2 % estimate. The average predicted deviation using a range of size distribution parameters from four other plutonium handling facilities included in their study was 7 %. A 15 % estimated 95 % confidence level uncertainty relative to predicted performance and assuming a realistic aerosol particle size distribution appears to be attainable.

These estimates are summarized in Table E.1.

E.4.3 Uncertainty in velocity measurement parameters

Uncertainties in the determination of velocity at each equal-area location in a profile are summarized from Reference [19] in Table E.2, assuming that an S-type Pitot tube nozzle is used. Here, as for the parameters in Table E.1, attention to the details of the design and operation of the hardware in use is required.

Table E.1 — Uncertainty in sample volume, stack or duct area and transmission line efficiency

Parameter	Uncertainty %
Sample volume $2u_r(V)/V \times 100 \%$	± 5
Stack or duct area $2u_r(A)/A \times 100 \%$	± 2
Transmission line efficiency $2u_r(\tau_p)/\tau_p \times 100 \%$	± 15

Table E.2 — Velocity measurement parameter errors

Parameter	Uncertainty %
Pitot calibration $2u_r(C_{\text{cal,pt}})/C_{\text{cal,pt}} \times 100 \%$	± 1
Flow angularity $2\tan(\theta)u_\theta \times 100 \%$	± 8
Differential pressure $2u_r(\Delta p)/\Delta p \times 100 \%$	± 14

E.4.4 Uncertainty in measurement parameters

Measurement of activity in a sample can be either an on-line process in a continuous monitor or an off-line process in a laboratory. In all systems, detector efficiency and sample capture efficiency parameters can usually be well defined. Detector efficiency is typically determined by comparison against a transfer standard traceable to the governing national institute of standards and measurements. Uncertainty can be held to a minimum (1 % to 2 %).

Type A uncertainty in the counting process and associated background-interference variability are the largest contributors to this uncertainty. However, bounds can be put on this uncertainty by careful planning.

In most circumstances, a relative 95 % confidence level uncertainty of 5 % to 10 % appears to be attainable by adjustment of sample and background count times. But in practice this might not always be possible due to the unrealistically long count intervals that can result.

E.4.5 Methodological bias

Turning now to the implicit methodological or conceptual uncertainties, there is a large component of engineering judgment required to assign values to these, but bounds can be placed on the estimated uncertainty.

The uncertainty associated with the sample-withdrawal location, for example, can be estimated by the measured coefficient of variations in the mixing of tracer gas and tracer particles that are required as part of qualifying a sample-extraction location for continuous-emission single-point sampling and monitoring. Based on limited studies, it appears that a 95 % confidence limit systematic uncertainty of δ_{RE} equal to 10 % is attainable, and in any event it should be much less than 20 %.

Uncertainty associated with changes in effluent emission conditions over time are difficult to predict. If mixing elements installed in a stack or duct are employed to ensure complete mixing, then an estimate of the 95 % confidence limit on uncertainty due to time varying effects on the order of δ_{TE} equal to 2 % to 4 % appears to be reasonable.

Similarly, model assumption uncertainty, at the 95 % confidence level, on the order of δ_{ME} equal to 2 % to 4 % is feasible as long as proper qualifications of the sampling nozzle, sample transport line, and sample withdrawal location are demonstrated.

E.5 Summary of uncertainty analysis

Estimated uncertainties in [Tables E.1](#) and [E.2](#) (or equivalent from an independent analysis) and the other parameter uncertainty estimates can be substituted into the appropriate equations to obtain a total measurement-process uncertainty estimate.

An example is given as follows. The estimated uncertainty of activity concentration, $u(c_A)$, (5,5 %), sample volume measurement uncertainty, $u(V)$, (5 %) and sampling plane area uncertainty, $u(A)$, (2 %) contribute least to the total. The uncertainties in the sample-transport-line efficiency, $u(\varepsilon)$, (15 %) and emission mean-axial-velocity, $u(v_m)$, (16,2 %) terms contribute the most. The resultant combined uncertainty, as given in [Formula \(E.8\)](#), is on the order of 23 %.

$$u(\bar{A})_{k=2} \pm \sqrt{(0,055)^2 + (0,05)^2 + (0,02)^2 + (0,15)^2 + (0,162)^2} \times 100 = \pm 23 \quad (E.8)$$

This estimate should be understood as an indication of what can be attainable based on the assumptions concerning the measurement procedures carried through the analysis. Differences in the way that mean axial velocity is determined, improvements in reducing uncertainties in volumetric flow measurement and better potential transport-line-loss estimation can be possible in some cases, as can reductions in the uncertainty in certain profile “pattern” parameters, such as any of the statistics describing mixing at the sampling plane.

E.6 Correlated uncertainties

The analysis in this annex to this point has been based on the assumption that the uncertainties are separable. The uncertainties are not separable in all cases. The transport efficiency, ε_{TS} , and the collection efficiency, ε_p , are dependent upon the flow rate. In some cases, the detection efficiency is dependent on the flow rate.

Based on the uncertainty analysis of the emission rate given in [Formula \(E.4\)](#), it may be concluded that the intrinsic uncertainty of the flow-rate can be ignored, but not its extrinsic or correlated uncertainty.

The transport penetration is dependent upon the flow rate. If the transport penetration is high (>90 %) for all particle sizes, flow rate changes of 10 % or less can have little effect on the transport efficiency. When the transport penetration is lower, small changes in the flow rate can greatly affect the transport efficiency for some particles sizes. These changes should be empirically determined or calculated using a code such as Deposition Calculator. A similar relationship for particle collection can be developed, for which the same correlated uncertainty arguments apply.

The detection efficiency can be affected by the flow rate if detection is dependent upon geometry and the geometry is collection-dependent. A simple example is alpha-particle detection on a filter. If the collection of large, high-activity particles takes place primarily near the edge of the filter and, therefore,

the edge of the detector, then the detection efficiency for these particles is diminished. The flow-rate-correlated uncertainty should be determined.

Flow-rate-correlated uncertainties in these cases are further dependent on the particle-size distribution, i.e. the effect of a few large particles can be quite small, while for a significant number of large particles, this effect can dominate.

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Annex F (informative)

Mixing demonstration and sampling system performance verification

F.1 Mixing demonstration methods

F.1.1 General

At least two methods have been used to demonstrate the state of mixing of the potential contaminants with the effluent air stream. They are described in [F.1.2](#) and [F.1.3](#).

F.1.2 Method 1

F.1.2.1 General

Method 1 was developed specifically to assess conformance with [6.3](#).

F.1.2.2 Tracers

To test for contaminant mixing, the tracer should be introduced as far upstream as possible of the sampling probe, yet downstream of feeder ducts, fans and air-pollution-abatement equipment. If it is necessary that a stack or duct be tested for both particles and gases, the same injection location should be used for both tracers. The gaseous tracer should be introduced at five or more locations across the cross-section of the air stream. For a rectangular duct, the injection should be at the centre and near each corner (at or within a distance of 25 % of a hydraulic diameter from a corner). For a round duct, the introduction should be at the centre and near the wall (within 20 % of the diameter from the wall). The aerosol particle tracer may be introduced at only one location, located at the centre of a stack or duct.

The degree of mixing for particles should be tested with particles having a diameter, D_a , between 8 μm and 12 μm , or larger if there can be a significant fraction of the aerosol particle mass or activity associated with sizes larger than a D_a of 10 μm . In cases where additional data about the relevant size distribution (e.g. activity size distribution) are available, the test aerosol particle size may be selected accordingly.

If, in any foreseeable circumstances, only gaseous contaminants can be present, it is not necessary to test for particle tracer uniformity.

Testing to establish the degree of mixing requires sufficient gas or particles to provide an adequate signal at the extraction point. The method of detection and its detection limit are the important considerations in the amount of material introduced. Sufficient material should be introduced to allow detection after dilution in the effluent stream. Examples of methodologies for obtaining data on velocity, tracer gas and aerosol particle profiles are given in References [\[39\]](#), [\[44\]](#), [\[89\]](#) and [\[73\]](#).

Tracer uniformity measurements should be conducted at the location of the sampling probe using the measurement grid developed in [F.1.2.4](#).

F.1.2.3 Measurement conditions

The tests should be conducted while the stack or duct flow rate is approximately the same as the expected normal flow rate. If the stack or duct flow rate is expected to vary more than 25 % from the mean, then the tests should be conducted at the flow rate extremes.

It is essential to establish with confidence that the location chosen for sample extraction is based on demonstrated complete mixing using the above methods and that the location continues to meet the mixing-performance criteria if air flow conditions change relative to those at the time of testing. Historical records of effluent flows may be used to provide evidence of extremes (high or low) of flows that can be encountered in a stack or duct. Calculations of expected flows under accident conditions or very different operating modes may be based on documented engineering judgment. Mixing under flow rates that are considerably different from normal may be substantiated by tests with models or field testing of the stack or duct. Under most conditions, changes in the effluent flow rate does not significantly affect the mixing. In general, if the flow rate increases, acceptable mixing is not jeopardized. However, if the flow rate were reduced to the point where the Reynolds' number becomes much less than 10 000, there can be a major degradation in the mixing effectiveness. This event is generally possible only with a stack or duct having a very small cross-section, such as a tank vent. If this is possible, the flow system should be modified to preclude the onset of near-laminar conditions.

F.1.2.4 Measurement points

It is convenient to use the same measurement points for the flow angle, air velocity and tracer concentration tests and they should be selected in accordance with ISO 10780. It can be necessary to add or adjust measurement points to achieve a suitable grid and because of the proximity of a sampling point to a wall.

F.1.2.5 Transference of qualification test results

It is not always necessary to perform the full qualification test series on all stacks or ducts if a geometrically similar design has already been shown to meet the qualification criteria given in 6.3 and provided that the following apply.

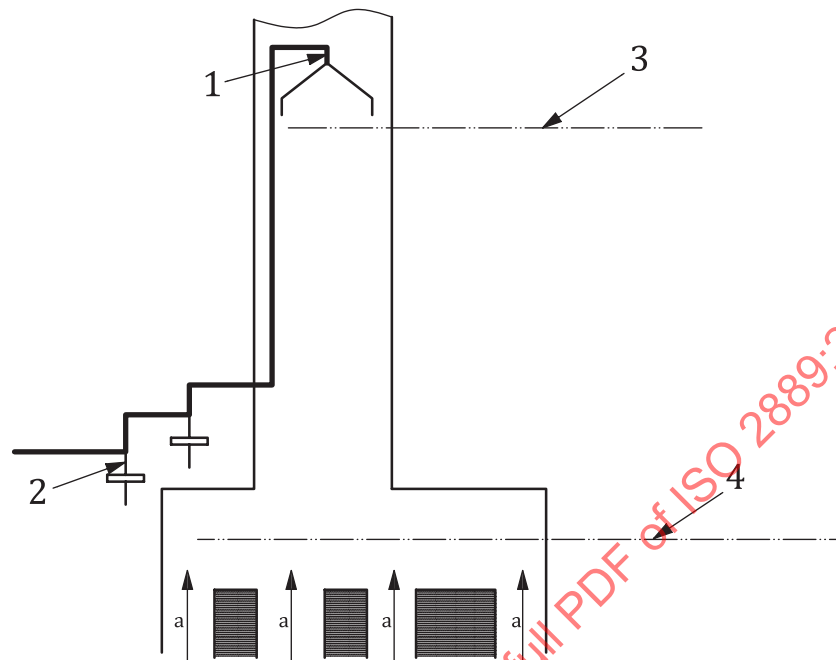
- a) A geometrically similar stack or duct (one with proportional critical dimensions) has been tested and the sampling location has been found to comply with the requirements of 6.3. Critical dimensions are those associated with components of the effluent flow system that can influence the degree of contaminant mixing and the velocity profile. The prior testing may be conducted either on a stack or duct in the field, or it may be conducted on a scale model.
- b) The product of mean velocity [see Formula (A.2)] and hydraulic diameter of the candidate stack or duct is within a factor of six of that of the tested stack or duct, and the hydraulic diameter of the candidate stack or duct is at least 250 mm at the sampling location. The Reynolds numbers based on hydraulic diameter of both the candidate stack or duct and the tested stack or duct are greater than 10 000 [see Formulae (B.1) and (B.2) for examples of expressions that can be used for calculation of Reynolds numbers].
- c) The measured velocity profile in the actual stack or duct should meet the requirements of 6.3.
- d) The difference between the velocity C_V of the two systems is not more than 5 %.
- e) The sampling location in the candidate stack or duct is placed at a location geometrically similar to that in the tested stack or duct.

If these transference requirements are fulfilled, the sampling location in the second stack or duct is considered to be acceptable.

F.1.3 Method 2

Figure F.1 shows how a number of ventilation channels come together in a chamber on top of which stands the stack or duct. Sample extraction for effluent monitoring is done about four diameters up the stack or duct. The test was originally developed to measure particle losses in probes and pipes, but it can also be used for investigating the effects of incomplete mixing of the contaminants in the effluent air.

Figure F.1 shows a two-stage sampler installation. The first stage brings air down to the measurement room at a rate of $60 \text{ l}\cdot\text{s}^{-1}$. There, several samplers withdraw air at $0,5 \text{ l}\cdot\text{s}^{-1}$. The probe in the stack or duct has four to six nozzles.



Key

- 1 sample extraction location
- 2 samplers
- 3 tracer gas detectors
- 4 injection level for tracer gas and particles
- a Air flows from reactor building, turbine building.

Figure F.1 — Illustration of the use of method 2 to determine mixing

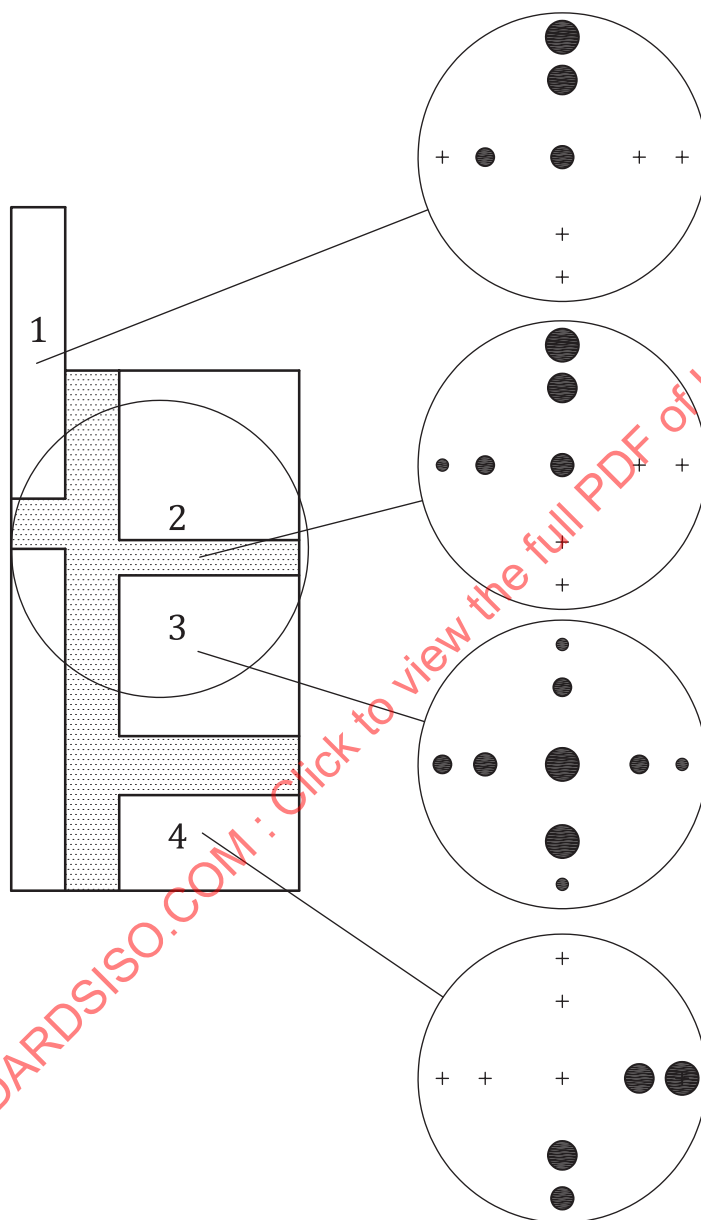
The test is comprised of the injection of known amounts of monodisperse particles in the stack or duct air stream at a suitable point upstream of the sampling installation. To find a suitable injection point, the gas flow is mapped by means of a tracer gas, ethanol, released as a spray at various points in the stack or duct base. The resulting concentration distributions at the sample extraction point are observed by means of an array of gas detectors. An injection point that produces symmetrical distribution over the stack or duct area, and low concentrations at the stack or duct walls, is selected for the particle tests (Figure F.2, key item 3). In this way the losses of tracer particles become small and limited to the flow near the stack or duct wall where the probe doesn't sample.

Monodisperse particles nominally with a D_a of $2 \mu\text{m}$, $4 \mu\text{m}$, $8 \mu\text{m}$, and $16 \mu\text{m}$ tagged with dysprosium are dispersed at the selected injection point. The particle concentrations at the sampler inlets are calculated from the observed gas distribution. The amount calculated as aspirated into the sampler piping is compared with the quantity collected by the ordinary filters of the sampling train to obtain the sampling-line transmission efficiency.

The sample-extraction location is situated about half-way up the stack or duct. This is enough for producing an even velocity field, but it is not enough for mixing. Figure F.2 shows the stack or duct base as seen from above. (The cross-section at the stack or duct base is irregular because several ducts discharge into the base of the stack or duct). Key items 1 to 4 represent the different injection points for the tracer gas. The tracer gas that is injected into the different ventilation channels is not well mixed into the main air stream as it passes the sample-extraction location. The spots in the four circles to the right represent observed concentrations at the sampling level. As is evident from Figure F.2, the air streams from different ventilation systems are not well mixed. The corrective measures outlined in

[Clause 6](#) are advised for this example, and single-point sampling is not appropriate unless the mixing is corrected.

With the data from the tests using the tracer gas and a knowledge of the positions of the individual sampling nozzles of the probe, the response to different concentration patterns can be estimated. It seems that four to six nozzles can be quite satisfactory with geometry as described above. Further details about this test method are reported in Reference [100].



Key

1 to 4 gas injection points in the stack or duct base

Figure F.2 — Stack or duct base as seen from above with diagram of sample results

NOTE The corresponding tracer distributions over the circular stack or duct section at the sampling level are shown to the right. The area of the spot is proportional to the observed concentration. The “+” symbols indicate where no tracer was detected.

F.1.4 Numerical modeling to qualify the sample extraction location

The computational fluid dynamics, CFD, model is useful in visualizing and understanding bulk stream flow behaviour in the final exhaust system to assist in determining whether a more advantageous sampling location is feasible.

Simulation by CFD of particle transport and deposition is generally divided into the following steps: (1) simulation of the flow field, (2) particle tracking, and (3) post-processing to obtain particle-deposition information (Reference [106]). Various models are available including FLUENT and STAR-CD. The model should use a fully developed three-dimensional mesh of the full geometric detail and include the sections from the last disturbance to the exhaust point of the system. The CFD model (including boundary conditions and defined mesh) should incorporate the appropriate aspects of the system (e.g. $Re > 10,000$, $(10 \pm 1) \mu\text{m}$ AD particles, etc.) and run enough simulation iterations to converge on a solution. Output results from CFD are directly influenced by the quality of the model inputs such as the mesh model and running to convergence.

At the selected sampling extraction point, the CFD results should meet the acceptance criteria of 6.3, and the arithmetic difference between the CFD modeled stack or duct velocity C_V and the candidate stack or duct velocity C_V is not more than 5 percentage points. If these requirements are met, the CFD-identified sampling extraction point in the stack or duct then is tested in situ or with a physical scale model. Models are validated against a known model/measurement in order to show the models works properly.

Examples of methodologies for obtaining CFD data are given by References [17], [87], [101] and [104].

F.1.5 Alternative approaches

Other approaches may be used to qualify a location for sampling provided that the accuracy of the methodology is equal to, or exceeds, that based on the criteria given in this annex. For example, in an application dealing with sampling of radioactive gases, it can be possible to inject a tracer gas into the stack or duct at a known mass flow rate. If the mass flow rate of tracer emitted from the stack or duct based on use of a single-point sampler at a candidate location were to show a value within $\pm 20\%$ of the mass flow rate of injected tracer, the sampling location may be considered acceptable. In such a situation, it can be necessary to demonstrate the quality of samples acquired at different flow rates (e.g. average, high and low) of the stack or duct gas.

F.2 When to conduct sampling system performance verification

F.2.1 General

There are instances when a performance verification of a sampling system is advisable. These include the following:

- when the air stream being sampled is not well mixed before a new system becomes operational;
- when an existing system has just come under additional regulatory requirements;
- when the potential to emit contaminants through an existing system has changed significantly;
- when an existing system has had significant changes, for example: changing the stream flow beyond the original design limits; adding a new effluent stream in a manner that destroys the well mixed state at the nozzle location; or changing system operating parameters outside of the design range;
- when the supporting documentation for a newly installed system is deficient.

F.2.2 Approaches to verification

The methods for verifying sample-transmission performance through nozzles and transport systems fall into four categories:

- in-place testing;
- laboratory simulations;
- modelling based on deposition and resuspension rates determined in the laboratory;
- a combination of the above.

[Table 1](#) summarizes the requirements for qualifying sample extraction locations, nozzles and transport lines for particles, gases and vapour. In [7.3.2](#), it is recommended that nozzle performance for particles be tested using liquid aerosol particles. In [7.4.2](#), it is recommended that transport-line performance be assessed either through aerosol particle testing or through calculations with a verified model. In [6.3](#), a general method is provided for use in qualifying the sampling location for particles, gases and vapour using in-place testing. Nozzle and transport-line performance methods for gases and vapour are not specified in this document.

Meeting the performance requirements usually involves a combination of methods. In-place testing can give the most unambiguous result, but it can also be difficult to implement in all situations. The discussion in [F.2.3](#) gives examples of methods in each category.

F.2.3 In-place testing

F.2.3.1 Particle sampling examples

References [\[39\]](#), [\[44\]](#) and [\[89\]](#) provide examples of employing the methods outlined in [Clauses 6](#) and [7](#). Sulfur hexafluoride or nitrous oxide gases and oleic-acid or vacuum pump oil aerosol particles are used as the tracers to qualify the sample extraction location. Oleic-acid aerosol particles are also used to verify the performance of nozzles and transport lines for particles. Examples of other verification methods attempted include the following:

- using 3 µm to 30 µm aluminium and iron powder aerosol particles (see Reference [\[65\]](#));
- using uranium aerosol particles (see Reference [\[92\]](#));
- using fluorescent-dye-tagged dioctylphthalate 20 µm aerosol particles (see Reference [\[98\]](#));
- using sub-micrometre aerosol particles of sodium fluorescein dye (see Reference [\[29\]](#));
- tests performed on several stacks using polystyrene latex microspheres using optical particle counters on samples from the nozzle inlets and the exits of the transport lines (see Reference [\[91\]](#));
- tests conducted on sampling systems at the Waste Isolation Pilot Plant using salt aerosol particles (see Reference [\[83\]](#));
- systems tested using powdered tracer aerosol particles with geometric mean diameters of 1,3 µm and 8,5 µm (see Reference [\[41\]](#)). Temporary sample collectors were arrayed across the stack to characterize the mixing and to determine the average emission rate for comparison against the existing system;
- systems tested using powdered test aerosol particles with geometric mean diameters of 1 µm (TiO₂), 3 µm (SrTiO₃), 8,5 µm (Mo₂C), 200 µm (brass) and 800 µm (WC)^[107]. The test aerosol particles had been injected into the stack as well as into some nozzles. With these tests, the concentration distribution of test aerosol particles over the sampling plane as well as the penetration of these test aerosol particles from the stack and from the nozzle entrance was determined;
- using cascade impactors and optical particle counters to test several systems without employing tracer aerosol particles (see Reference [\[57\]](#)).

These last examples might not necessarily meet the current guidelines of this document, but provide insight into other approaches.

F.2.3.2 Radioiodine sampling examples

Sampler performance was tested^[98] for radioiodine by injecting both depositing and non-depositing forms of iodine into the ventilation stream. Samples were collected both in the stack at the elevation of the sampler nozzles and at the regular sample-collection point. The non-depositing form was ^{131}I -tagged methyl iodide and $^{131}\text{I}_2$ was used as the depositing form.

Stable methyl iodide and elemental iodine were injected into the stack or duct flow upstream of the fan (Reference [29]). Collected iodine samples were analysed using neutron activation.

In Reference [65], ^{131}I injected into a stack or duct, which was sampled at several locations in the cross-section using charcoal traps. These data, together with velocity data and tracer aerosol particle tests, were used to determine the contaminant profiles for the stack or duct.

F.2.4 Laboratory simulation

F.2.4.1 General

Laboratory simulations are more rapid and convenient than in-place tests. It is unlikely that complete systems can be simulated and the effects of surface contaminants in older systems can significantly affect real performance.

F.2.4.2 Aerosol particle examples

Reference [91] described a test of a full-scale sampling system of simple design. The tests were conducted using polystyrene latex microspheres and laser particle counters sampling from the nozzle inlet and from the end of the transport line. Reference [76] shows tests of a simulated sampling system using oleic acid aerosol particles tagged with sodium fluorescein. Wind-tunnel tests comparing the performance of probes using shrouded nozzles to those using tapered inlet isokinetic nozzles are described in Reference [40].

F.2.4.3 Radioiodine examples

Radioiodine line-loss tests to simulate air samplers used at several nuclear generating stations are described in References [103] and [33]. The tests cover the range of air sampler characteristics commonly observed at reactor sites. The sample transport tubes were either 304 or 316 stainless steel as clean as received from the distributor. The results from these and other tests are summarized in Reference [42].

F.2.5 Modeling

F.2.5.1 General

Modeling is often used to address the performance of transport lines. Modeling does not completely address all performance aspects of a sampling system, notably the adequacy of contaminant mixing at the sampling plane. This weakness may be overcome in the future as illustrated in Reference [38] where a three-dimensional fluid mechanics model was used to identify potential flow-measurement locations in the off-gas ductwork of a power station. They also compared the model results with velocity-traverse data.

F.2.5.2 Particle examples

Reference [36] provides an example of the use of the Deposition code and compares the results against tests of a simulated air sampling system. Examples of the use of earlier models are found in References [88], [13] and [94].

[Annex B](#) describes most of the elements that it is necessary to take into account in a particle-loss model. An example calculation is also given.

F.2.5.3 Radioiodine examples

Examples of modelling radioiodine transmission through several sampler-transport lines are found in References [\[42\]](#) and [\[91\]](#). See also [Annex C](#).

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Annex G (informative)

Transuranic aerosol particulate characteristics — Implications for extractive sampling in nuclear facility effluents

G.1 General

The engineering of stack or duct monitoring and sampling systems for nuclear facilities requires close attention to the design and placement of the sample-extraction nozzle and transport line to ensure the most representative sample possible (see Reference [78]). With respect to the physical characteristics of the effluent that it is necessary to sample under normal conditions and particularly under emergency conditions, little has been said other than that the most significant accident in a facility in terms of both an event and its consequences is likely to be fire (see Reference [26]). Fire can cause radioactive sources to release airborne radioactive aerosols, and smoke can plug filtration systems causing the filter to lose its integrity by rupture of the medium or seals. Therefore, while there are no definite answers concerning the aerosol particle characteristics that a sampling nozzle and transport line should be capable of handling, a number of investigations have been made of the expected filtration performance of HEPA filters under standard operating conditions. Also, a number of investigations have been made of the characteristics of aerosol particles present in gloveboxes or generated when containment structures, flammable liquids or mixed radioactive materials are spilled and burned. Because these aerosol particles can be expected to be present in a fire, they provide a first approximation of the character of aerosol particles that it can be necessary to sample in effluent stacks or ducts. The discussion in this annex is meant to place some reasonable bounds on the size of particles that can typically be present in the event containment is lost and to provide some perspective on the type of design and testing considerations that it is necessary to apply to sampling nozzles and transport lines.

G.2 HEPA filtration effects

Nuclear-facility stack or duct emissions are typically controlled by multiple stages of HEPA filters. The HEPA filter is designed to remove particulates from a gas stream with an efficiency of at least 99,97 %. Selective penetration of HEPA filters by sub-micrometre particles (0,1 µm to 0,4 µm) and negligible penetration by other sizes of particles is predicted by filtration theory (see Reference [95]). Therefore, it is sometimes concluded that it is not necessary to design sampling systems for HEPA filtered stack or duct that take into account inertial effects in the sampling-nozzle inlet and transport line. However, this conclusion is invalid.

In Reference [84] studies at the Rocky Flats Plant of particulate emissions in stack or duct effluents, particles were sampled and sized downstream of HEPA filters. A laser particle counter was employed to obtain number/size distribution data over the size range from the sub-micrometre range to over 10 µm (optical diameter). While only a small fractional percentage of particle counts correspond to diameters greater than 5 µm, when these data are converted to a volume distribution (hence reflecting the actual distribution of particle mass, and possibly activity, in the samples), the volume percentages corresponding to particles with diameters greater than 5 µm are quite significant (greater than 30 %). The authors also observed the presence of white fibres in the vent discharge and speculated that fibres and other large particles might have been shed by the HEPA filters.

A different explanation for the presence of the larger particles downstream of HEPA filters comes from studies in Reference [95] of leak phenomena in HEPA filter systems (pinhole leaks, frame seal leaks, etc.). Particle penetration through filter perimeter seals and the filter pack was determined separately and in combination. Penetration is observed only when the challenge aerosol particles are introduced into the system, ruling out the possibility that the observed particles were shed from the filter pack. He found that whereas filtration theory predicts a penetration fraction of 10^{-16} for 0,7 µm particles,

the observed penetration was approximately 10^{-5} . Reference [95] concluded that system leakage phenomena and the size distribution of the challenge aerosol particles can override filtration theory considerations in predicting the size distribution of particles penetrating the HEPA filter systems.

This is consistent with studies of multiple HEPA banks,[35] which data are summarized in Table G.1. In this case, the presence of a small but significant fraction of supra-micrometre-sized particles after the second and third stages is indicated because the geometric standard deviations, s_g , remain large. But the respective activity concentrations are very small due to the reduced challenge and narrowing spectrum of particle size at each successive stage. At the same time, failure of earlier stages can be expected to result in both higher release concentrations and larger quantities of particles in the inertial size range (with a D_a equal to $1\text{ }\mu\text{m}$ and larger). Therefore, the design of extractive sampling systems in HEPA-filtered stacks or ducts should reflect a consideration of the presence of large particles, even under the presumption of normal operating conditions and HEPA filtration.

Table G.1 — HEPA efficiency and particle penetration of Pu aerosol particles

HEPA stage	Particle size AMAD ^a of challenge μm	Mean measured efficiency %	Remaining activity ^b $\text{Bq}\cdot\text{m}^{-3}$
1	0,7 to 2,1, $\sigma_g = 2$ to 3	99,998 76	10^7 to 10^8
2	0,45 to 0,82, $\sigma_g = 1,5$ to 2	99,998 17	10^2 to 10^4
3	0,37 to 0,70, $\sigma_g = 1,3$ to 1,8	99,864 92	1 to 5

^a AMAD is the activity median aerodynamic diameter.

^b Challenge aerosol particle concentration is equal to $10^{12}\text{ Bq}\cdot\text{m}^{-3}$ to $10^{14}\text{ Bq}\cdot\text{m}^{-3}$.

G.3 Transuranic aerosol particulate characteristics under accident conditions

HEPA filter failure under a variety of accident conditions adds another dimension to the concern for being prepared to sample particulate radioactive substances in the larger size ranges. But now it is necessary to ask, what, if a substantial HEPA failure occurs, is the upper particle-size limit that one can expect that it is necessary to sample efficiently in order to properly represent the majority of the activity in the effluent? In other words, can the size distribution of effluent aerosol particles at the sampling plane suddenly shift to one characterized by the occurrence of a significant particle-size mode in a region above D_a equal to $10\text{ }\mu\text{m}$ to $15\text{ }\mu\text{m}$? It is understandably difficult to characterize aerosol particles that can be expected as a result of HEPA failure. As is the case with normal, intact HEPA filter banks, the characteristics of the aerosol particles penetrating a failed HEPA are determined by the characteristics of the challenge aerosol particles. The literature, derived from studies of aerosol particles associated with accidental spills and fires in nuclear facilities and of in situ dust and debris in uranium/plutonium gloveboxes and ducts, provides the best indications of what to expect. The following synopses of a few cases are indicative of what is known and expected.

- In a study of plutonium particle sizes in air samples taken in operational areas at the Rocky Flats Plant many years ago when maintenance operations on gloveboxes resulted in loss of containment (see Reference [58]), it was found that operations such as machining, oxide crushing and fluorination of plutonium produced airborne particles with mass median aerodynamic diameters (MMAD) of $2\text{ }\mu\text{m}$ to $4,5\text{ }\mu\text{m}$ (assumed density of $11,45\text{ g}\cdot\text{cm}^{-3}$). Conditions related to glove failure in a glovebox for burning plutonium metal leading to worker exposure produced larger airborne particles having a MMAD of $13,8\text{ }\mu\text{m}$. The author noted that these data agree very closely with the activity median aerodynamic diameters (AMAD) of particles measured at the AERE radiochemical laboratories in Reference [97].
- In Reference [34] the authors placed sampling nozzles in process lines or gloveboxes under “worst normal” conditions (i.e. when aerosol-particle generation as a result of routine operations was highest) in a study of challenge aerosol particle characteristics and the response of multiple HEPA filters. Facility operations included research and development activities, fabrication, and chemical recovery. Activity concentrations in challenge aerosol particles were in the range of $108\text{ Bq}\cdot\text{m}^{-3}$ to $1\text{ }010\text{ Bq}\cdot\text{m}^{-3}$. Fabrication operations produced fairly large aerosol particles (predominant

AMAD, D_a , equal to 3 μm to 5 μm), while recovery operations consistently produced particles in the sub-micron range (0,1 μm to 1,0 μm). Research and development operations generated particles predominantly in the intermediate range (1 μm to 4 μm). The largest reported size bracket in the log-normally distributed impactor data from all sites was a 10,9 μm bracket (normalized frequency of 4 % by activity), from a research and development facility.

- c) Apart from accidents, processing facilities age and contaminated structural materials can become suspended downstream of HEPA filter systems. The contamination can occur gradually or during process and filter upsets. Reference [71] summarizes the historical measurements of particle-size distributions in a plutonium finishing and reclamation complex downstream of HEPA filter systems. The AMAD of plutonium-bearing particles ranged from 1,3 μm to 20 μm . Ventilation ducts damaged by exposure to acidic fumes can generate significant concentrations of slightly contaminated rust particles.
- d) In case of an accident, the potential contribution of fire-generated aerosol particles should be included. Reference [46] describes studies of burning radioactively contaminated materials, uranium was used as a surrogate for plutonium. Combustion aerosol particles containing uranium from contaminated plastics produced fairly large particles with an MMAD equal to 1 μm to 5 μm . Compounds in glovebox gloves (polychloroprene) produced the largest particles (MMAD of 19,9 μm). Burning cellulose produced particles with an MMAD from less than 1 μm to as large as 10,5 μm . Conversion of these numbers to aerodynamic diameter is uncertain as the density and shape factors are unknown.
- e) Accident conditions can involve leaks or spills of liquid and powder forms of radioactive substances that generate airborne materials. In studies with uranium and other surrogates (see Reference [16]), aerosol particles with a wide range of MMADs between 3 μm and 20 μm were measured. Liquid spills appear to produce the largest particles from splash droplets that start large but get smaller as the liquid evaporates. The particle size distributions resulting from powder spills have more variability due to agglomeration effects in the bulk state.
- f) Agglomeration in accident-generated aerosol particles has been shown to produce larger particles in a polydisperse aerosol of smaller particles. However, it appears that this process does not yield extraordinarily large particle-size modes. Using data from the Oak Ridge National Laboratory nuclear safety pilot-plant experiments with burning sodium in a containment structure, Reference [54] predicted (and confirmed with observation) a relatively stable evolution of the mean aerodynamic diameter. Diameters remained below a D_a of 5 μm for 5 days following release and confinement.
- g) In some facilities, there is the potential for involvement of plutonium metal in fire scenarios. Studies have been made of the release of aerosol particles under reducing and oxidizing environments. Reference [32] found that the activity-median diameters of plutonium aerosol particles generated from plutonium metal pellets and foils were variable, but ranged from a D_a of 4 μm to approximately 10 μm .

G.4 Implications for nozzle design

The available data indicate that the most common mode of particle size for plutonium and uranium aerosol particles under a wide variety of conditions of generation is a D_a of between 1 μm and 5 μm with measurable percentages of particles up to 10 μm , or even 20 μm . The appearance of particles in the inertial size range (with a D_a above 1 μm) can be anticipated, even under routine operating conditions and certainly under a wide range of accident conditions. Inlets of sampling nozzles for particulate emissions should, then, be tested for transmission performance in the range of 3 μm to 15 μm . The shrouded nozzle inlet, for example, is designed so that the transmission of inertial-sized particles through the inlet is between 83 % and 103 % under the flow conditions of intended use. Predicted performance of a design is confirmed with measurements in a wind tunnel using test aerosol particles with a D_a of 10 μm . The design is iterated until there is good agreement between predicted and measured performance, at which point the shrouded nozzle is qualified for use. Unless it is known that a facility stack or duct effluent can contain a sizeable mode of very large-sized particles due to the

nature of the materials being handled, it is not necessary to require performance testing in size ranges beyond 15 µm.

G.5 Implications for other nuclear facilities

Although the discussion in this annex has relied largely on data from plutonium facility experience, the concern for proper sampling and monitoring of the large-particle components of effluents downstream of HEPA filtration in other types of nuclear facilities is equally important. Reference [30] reviews a wide range of literature on particle-size distributions of radioactive aerosol particles measured in workplaces throughout the nuclear industry and government laboratories, it was found that in a total of 52 papers reporting 160 measurements of particle AMAD, the measurements of AMAD in the nuclear power industry and fuel handling facilities follow distributions similar to those in workplaces as a whole, with median values of about 4 µm. The exception seems to be for uranium mills, where the median is about 7 µm. Reference [99] reviewed the characteristics of accident-generated aerosol particles in Swedish power reactors. A large body of literature exists on aerosol particles, vapour, and gases generated in postulated power-reactor accidents. Little is available concerning less consequential off-normal events. The implication is clearly that the challenge aerosol particles presented to HEPA filtration in practically any nuclear facility contain a significant component of particles with a D_a larger than 2 µm to 3 µm and, thus, can be present downstream of the filtration banks where sample extraction takes place.

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Annex H (informative)

Tritium sampling and detection

H.1 Tritium chemistry

Tritium, an isotope of hydrogen, generally behaves in a manner similar to hydrogen. Typically, it is found in two primary forms in the exhaust stream:

- a) in the elemental form as a gas;
- b) in the oxide form as water vapour.

It is of particular interest that tritium in the oxide form has a boiling point slightly above 100 °C. For some regulatory analysis this allows the oxide form to be considered a liquid instead of a gas, which in turn allows the use of a liquid physical form factor instead of that for a gas.

Also, tritium is sometimes found in the exhaust stream as a component of methane or other volatile organics, or as a component of particulate matter. One such example is LiOH, where the hydrogen is replaced with a tritium atom, LiOT. This compound is a solid material at temperatures up to 400 °C.

H.2 Sampling considerations

H.2.1 General

The first step in selecting an appropriate sampling system is to determine the chemical form of tritium in the exhaust stream. If it is present in multiple forms, multiple sampling techniques may be employed. When the oxide form is present, it is necessary to consider carefully the temperature and moisture content of the exhaust stream. If the exhaust stream contains water in droplet form, then the tritium can also be in this form and sampling as though particles were present is recommended. Sampling for a vapour only is appropriate when the oxide is not expected to condense.

H.2.2 Sampler nozzle

The sampler nozzle should be located in the appropriate place depending on the chemical form of tritium. The location and nozzle configuration should conform to practices outlined elsewhere in this document.

H.2.3 Heat tracing

The use of heat tracing on sample lines designed for tritium sampling should be evaluated very carefully. Several of the tritium sample-collection methods rely on either absorption of water vapour into a medium or condensation in a condenser apparatus. If the temperature of the sample is maintained too high, the tritium can desorb from the medium. Not all of the vapour condenses and, therefore, some of the absorbing solution in the bubblers can be lost. All of these conditions lead to biased results.

On the other hand, if the physical state of the tritium in the exhaust is gaseous or vapour and the exhaust stream contains high humidity, then heat tracing can be necessary to avoid condensation of sample in the sample lines and sample chamber. Condensation can cause the sample collector to plug and, in the case of an ionization chamber, the reading can be disrupted because of shorting of the central electrode to ground or of the high-voltage electrode.

H.2.4 Medium location

Except for tritium existing in a particulate form, the tritium sample medium is generally located downstream of a particulate filter. This keeps particles from plugging the sample medium.

H.3 Sample media

H.3.1 General

There are several generally acceptable methods and/or media available for sampling tritium when it is not in the particulate form. When it is in the particulate form, then sampling methods for particulate matter discussed elsewhere in this document should be used.

Although there are many factors that affect the sensitivity of a method, sensitivities on the order of 40 000 Bq·m⁻³ are possible with sampling followed by laboratory analysis. Typical factors, but certainly not all possible factors, that can affect sensitivity are sample flow rate, temperature of sampling, pressure of sampling, analytical method and sample medium.

Often, information on the concentrations of both the oxide and the elemental gas forms of tritium is desired. The uptake of tritium in the oxide form is very efficient, on the order of 99 % in Reference [11]. However, only 0,004 % of the elemental tritium entering the body is converted to the oxide form and adsorbed. Therefore, releases of tritium in the elemental form have a much lower dose than that from the same quantity of tritium oxide. Combinations of the following methods can be used to determine total tritium and oxide levels, with the difference being the elemental tritium in the stream being sampled. Direct measurement is also possible by first removing the tritium oxide, then converting the elemental tritium to an oxide form, followed by additional sampling.

H.3.2 Silica gel

This method of sampling tritium oxide is the simplest to perform. This is a continuous sample collected over a period of days to weeks. It involves placing a canister of silica gel in the sample stream and absorbing the tritium as water vapour on the silica gel. For sampling tritium oxide, a coloured silica gel may be used. The anhydrous silica gel is blue. As it adsorbs moisture, the colour changes, either to a different blue or to another colour. The sample, once collected, is sent to a laboratory where it is heated to desorb the tritiated water.

If tritium in the elemental form is present, then a catalyst, such as palladium, can be installed upstream of the sample chamber. The catalyst converts the elemental tritium to the oxide form, which can be absorbed on the silica gel.

H.3.3 Molecular sieves

This method is identical to the silica gel method, except molecular sieves are used in place of silica gel. This method has two advantages over silica gel.

- a) The media can be better dried initially, resulting in a lower background.
- b) A palladium catalyst, which converts the elemental tritium to the oxide form, can be coated directly on the molecular sieve.

A primary drawback of molecular sieves is the desorption of the tritium. This typically involves heating the medium to 500 °C in an evacuated furnace. Also, molecular sieves have a lower moisture-handling capacity than silica gel, but the medium is a more efficient drier. Therefore, when the moisture content of the sampled exhaust is high, silica gel is probably a better medium. However, when the moisture content is low, molecular sieves can be a better choice.

H.3.4 Bubblers

Although a variety of absorbing materials can be used in bubblers, ethylene glycol or water are most often used. This method provides advantages in the laboratory, in that sample desorption is not required. The primary disadvantage is that the bubblers, which are typically of glass, and the liquid media are difficult to handle when it is necessary to use them in the field or plant environment.

H.3.5 Condensation

In high-moisture exhaust streams, condensation is likely to be the most suitable method, since the other methods are limited by exhaustion of the absorption media. However, this method can be difficult to use. This method is based on the condensation of the tritiated water with a dehumidifier or condenser. The sample is routed through a mechanical cooling system and the condensate is collected in the liquid state. A loss or reduction of cooling capacity of the condenser allow the tritium-containing moisture to escape from the system in the exit gas. To ensure representative sampling, regular equipment maintenance is required.

H.3.6 Catalysts

All of the above methods rely on the tritium being in a vapour form, generally water vapour. When tritium is present as an elemental gas, it is then necessary to convert it, using a catalyst, to the oxide form before it can be sampled. Although any catalyst that can convert elemental hydrogen into the water or oxide form can be used, a palladium catalyst is the most common choice.

When tritium is present as an organic chemical species, it is often necessary to use a combustion catalyst. An example is a platinum on aluminium oxide catalyst in a heated combustion chamber. The tritium in the organic compound is oxidized to HTO and collected using the methods described above.

H.4 On-line detection

H.4.1 Ionization detectors

This is a very simple detector that can detect both elemental and oxide forms of tritium. The sensitivity can be as low as $0,04 \text{ MBq}\cdot\text{m}^{-3}$ depending on chamber volume. The major drawback to this detector is that it is sensitive to any gamma field in the general area and to any other ionization occurring in the chamber. A second chamber is sometimes used to compensate for external gamma fields by exposing the second chamber only to the field and not the exhaust stream. An additional chamber with a silica gel or molecular sieve pre-treatment is sometimes used to discriminate between oxide and elemental tritium.

This type of detector can be used when tritium is present as an organic vapour, such as tritiated methane.

H.4.2 Proportional counters

This type of counter detects tritium by using a rise-time discrimination principle. Since the soft beta of tritium has a short drift time, this detector can discriminate between tritium and other radionuclides, such as noble gases or other gamma emitters. The sensitivity of these instruments is around $0,4 \text{ kBq}\cdot\text{m}^{-3}$.

Annex I (informative)

Action levels

I.1 General

An action level is an effluent contaminant concentration threshold at which it is necessary to perform an appropriate action. The type of action performed depends on the circumstances. The action can entail generation of alarms, diversion of effluent through added effluent treatment or intervention in the process creating the contaminant. There are inevitable consequences of whatever response is taken. Some responses are relatively minor, others are much more significant in terms of cost, damage to equipment and, possibly, even human health and safety. Careful consideration, taking into account all such consequences, should be given to the setting of an action level for an effluent sampling or monitoring system. There can be not only false negative outcomes (i.e. a true release of significance that is not investigated), but also false positive outcomes (i.e. worker responses to alarms, risks associated with rapid shutdown, and costs that are incurred needlessly). Both types should be anticipated but, depending on the hazard potential of a particular effluent and other factors, false positive outcomes can be of more consequence to facility operations and worker safety than false negative ones, due to work stoppage and evacuation of areas. Facility administrators should be cognizant of all reasonably anticipated outcomes.

Action levels involving events with significant releases and potential risks to members of the public are generally set either by the regulator or through discussions between the regulator and the licensee. Releases above these regulatory alarm levels require reporting to the regulator. In addition, the licensee may set his own, lower alarm levels (also called administrative levels or limits). These may be set to avoid reaching the regulatory action levels or they may be set for other reasons, such as costs involved in the releases. For example, tritium releases from heavy water reactors may be based on the economic cost of losses of heavy water. There is usually an internal reporting procedure when administrative levels or limits are exceeded. Administrative levels are usually set somewhat above normal release levels. Alarms based on increasing release rates may be used in control monitoring to warn the operators that conditions have changed and immediate action may be required to avoid exceeding an action level or administrative level.

The process of selection of an appropriate action level requires a consideration of

- a) the physical and chemical characteristics of the contaminant,
- b) the characteristics of the sampling system required to obtain a sample of the contaminant for analysis and counting (e.g. the nozzle design characteristics, the transport line design or sampling location), and
- c) the type, intensity and variability of interference with the measurement.

Each of these three factors can contribute relative uncertainty to the contaminant-concentration estimate and, therefore, affect the level of confidence that can be assigned to the decision. The selection of an appropriate action level is separate from, and precedes, considerations of the required sensitivity of the sampling and measurement systems.

It is useful in the context of discussing action levels to draw distinctions among the following:

- control monitoring: sampling for purposes of providing adequate warning so that an operator can take action to protect workers and the public from excessive exposure (i.e. continuous monitoring with alarm);

- system availability: tracking sampling-system availability and response so the facility operators are alerted if equipment failure takes a system off-line or seriously degrades performance;
- performance sampling: regulatory compliance sampling that yields data of such quality and type that the facility owner can identify and quantify the most significant radionuclides present in the effluent and support demonstration of regulatory compliance by meeting all requirements for sample-extraction location, instrument calibration and maintenance, sample handling and chain of custody.

When determining action levels, consideration should be given to accuracy, precision and relative uncertainty. These terms apply to both the process of sampling and the process of measurement. The concern for accuracy is directed at the elimination of systematic uncertainty in the sampling and measurement processes. Regular calibration of sampling and measuring equipment using accepted procedures and traceable standards is used to establish accuracy.

Statistical measures of the dispersion of results about a measurement-population mean are used to calculate precision. Sampling precision can be determined by replicate samples obtained under the same conditions. Measurement precision is obtained from the statistics of repeated measurements on replicate samples and by detailed analysis and propagation of uncertainties in component measurements. Sampling and measuring precision are combined statistically to obtain an estimate of overall precision.

The concern of precision determinations is the estimation and, where possible, reduction of random uncertainties in the sampling and measuring processes.

The departures of measured values from either the true values (accuracy effects) or from the mean of measured values (precision effects) are measures of relative uncertainty in the sampling and measurement system results. Contributions to relative uncertainty in sampling or measurement are best determined independently and combined by statistical propagation of relative uncertainty. A detailed discussion of sampling and measurement random errors and error determination methods leading to an assessment of overall effluent sampling system relative uncertainty is found in [Annex E](#).

Table I.1 — Guide to standard uncertainty of sampling and measurement

Factor or consideration	Record sampling	Control monitoring
Frequency of		
a) sampling	continuous	continuous
b) measurement	weekly	near real-time
Relative standard uncertainty of sampling system		
a) overall accuracy	±15 %	±20 %
precision	±15 %	±20 %
b) sampling accuracy	±10 %	±10 %
precision	±10 %	±10 %
c) measurement accuracy	±10 %	±18 %
precision	±10 %	±18 %
System availability	>90 %	>95 %

Each facility should set data quality objectives for its particular sampling and measurement systems. Guidance for recommended levels of relative uncertainty as they pertain to accuracy and precision of sample extraction and transport, and of measurement, is given in [Table I.1](#). Most of the recommendations for control monitoring in [Table I.1](#) are not as stringent as the recommendations for record sampling. Near-real-time radiation detection by a CAM, for example, usually cannot yield as accurate a measurement as can be expected of a laboratory counting system because variable systematic uncertainty introduced by the presence of interfering background activity can often be significantly reduced or eliminated in the laboratory. It should be recognized that the accuracy and precision recommendations of [Table I.1](#) are not meant to be absolutes that can be equally appropriate in all cases and conditions. For example, effluents containing highly reactive constituents such as radioiodine can be particularly difficult

to extract and transport without significant sampling bias, leading to estimated sampling accuracy relative uncertainties higher than 10 %. In contrast, measurement accuracy in some systems can be easily held well below 10 %, given the characteristics of the instrumentation and measurement processes and should be so reported.

There can be justifiable reasons for sacrificing some degree of measurement accuracy in control systems to achieve higher instrument reliability, extended range of response, more effective background compensation or other optimization goals. The overall system-relative uncertainty limit values are derived by summing the respective relative-variance estimates of the sampling accuracy or precision components. For example, the recommended limit of 20 % for the overall relative uncertainty in system accuracy for continuous monitors is the square-root of the sum of squares of the sampling accuracy component, equal to 10 %, and the measurement accuracy component, equal to 18 %, of relative uncertainty. Mentioning these components separately calls attention to the fact that if the sampling nozzle and transport line are not properly designed and properly placed, relative uncertainty is created in sampling accuracy, and no amount of attention to measurement accuracy in the system can prevent the system from generating poor, biased data and faulty alarm responses.

I.2 Action levels for control monitoring

The discussion in the previous part of this annex provides a basis for incorporating accuracy and precision considerations in the setting of appropriate action levels. There are no hard-and-fast rules concerning how to set these levels. It should be kept in mind that there is a trade-off between having high confidence in an alarm being triggered only by a “true” contaminant release, and having sufficient alarm sensitivity to the presence of the contaminant, albeit at lower confidence. It is in this context that it is necessary to assess the true costs of false alarms as part of the decision, along with the desire to detect the contaminant at the lowest concentration and shortest integrating time. In some CAM instruments, net counts are converted to activity-concentration estimates by dividing the counts by detector efficiency, volume sampled and time. All of these factors add relative uncertainty to the results. However, for the purposes of an alarm, the largest relative uncertainty is typically contributed by the counting relative uncertainty, as shown in [Annex E](#). If the costs of false alarms in an effluent monitoring system are large, then a decision can be made to set the alarm threshold at a relatively high level and accept the risk of not detecting a lower-level release in a timely fashion. In some contexts, monitoring the trends of the measurement results can be a useful tool to aid in maintaining control below a chosen action level. The user should determine an action level that can be attained, and design and operate the system so the detection limit for that sampling and measurement process is sufficiently below the action level to avoid false alarms.

Control monitoring, using a CAM with an alarm, does not imply that there can be a more relaxed attitude toward achieving a representative sample. In the case of continuous monitoring for particulate radioactive substance in effluent streams, there might not always be a “sufficient” number of small particles in a release to cause an alarm. Nor should it be assumed that, in a poorly designed sampling system, a few large particles can get through to trigger an alarm. Such assumptions are ill-advised and unacceptable. The danger resulting from a lack of attention to the CAM sampling system design and placement is that the component of the sample that is not well represented (possibly the larger size particles, for example) can be the very component that provides the best chance for early warning and, hence, control, worker protection and impact limitation. The inherent limitations of providing radiation detection in the sampler during the sample collection process are due to the large background component in the detected signal. The choice of a relatively insensitive sampling system can lead to alarm thresholds consistent with an acceptable false-alarm rate. However, the system can then be susceptible to excessive false negative responses.

I.3 Action levels for record sampling

A facility or work area that has the potential for radioactive emissions (categories 1 to 3) should carry out record sampling at an appropriate frequency. Record samples are collected continuously in an integrated sample, and then are analysed by subsequent counting (off-line). The levels of activity that can be detected by these means are typically many orders of magnitude smaller than those detectable

by on-line monitors. Additional sensitivity can be achieved by preparing composite samples from several week-long samples into monthly or even quarterly samples. The decision to attempt to achieve a certain detection limit goal requires a balance among costs, time and other factors. It is necessary that the relative uncertainty in the final estimate of quantities, concentration and rates of emission be derived from uncertainties in each of the factors entering into the respective calculation.

Another layer of analysis can be applied after determining the central estimates and their uncertainties, and logging them over time. Here the question is not whether a particular measurement is above the decision level for that contaminant, but whether a given trend in the data is normal or indicative of an off-normal condition, or whether a given datum is an outlier or is actually an elevated concentration. For this purpose, certain statistical tests and trending procedures (e.g. a control chart) are important. A retrospective action level may be set above the trend line in the data (say, at the 3σ level) to help decide whether a datum should be regarded as belonging to the family of normal values for the parameter being measured (e.g. the mean concentration or amount of radioactive substance emitted during the previous month or quarter), or belonging to a new, unidentified situation that requires investigation and intervention. An investigation can determine whether a systematic uncertainty had been introduced in the analytical procedure for radionuclide determination, or can aid in ascertaining if a small leak exists in the filtration system of the facility stacks or ducts being sampled or monitored. The decision on the appropriateness of a given multiplier defining the action level hinges on the estimated costs associated with either being wrong in concluding that an excursion beyond the action level has occurred, and that controls in a facility are breaking down (false positive), or being wrong about thinking that emission controls and analytical procedures are normal, only to discover later that chronic low-level releases have been undetected (false negative). If the achieved detection limit for a sampling system is well below the action level, there is sufficient latitude to eliminate false negatives by trending.

I.4 System sensitivity needed to achieve selected action levels

Once an action level has been determined, another issue that should be addressed is whether the performance of the proposed real-time monitoring system is sufficient. This is not a question of accuracy, but of whether a chosen system is able to provide the needed net response above background. Assume that a real-time monitoring system has been operating in a given airborne effluent long enough to establish the population of, n , blank responses (meaning zero concentration of radioactive material of concern present, but varying levels of background activity) large enough to reliably establish an estimate of the population mean of the blank, μ_B , and standard deviation, σ_B . While in operation, if there is no contaminant present, but there is an interfering background, the number of net activity concentration equal to gross activity concentration minus background activity concentration, would be expected to have a limiting mean of zero, and a standard deviation of $\sigma_0 = \sigma_B \cdot \sqrt{1 + \frac{n-1}{n \cdot (n-3)}} \cdot (n \geq 4)$.

The decision threshold of the activity concentration, c^* , is given in [Formula \(I.1\)](#):

$$c^* = k_{1-\alpha} \cdot \sigma_0 \quad (I.1)$$

When the net activity concentration exceeds the decision threshold, there is a α % chance of making the error of concluding that activity concentration is present when there is truly only a background level present (a false positive). Said differently, an alarm level set at c^* would provide a false alarm rate of $X = \alpha$ %. Usually in the field of radioactivity measurements, we consider $\alpha = 2,5$ % or 5 %. From [Table I.2](#), if we consider $\alpha = 5$ % then $c^* = 1,645 \cdot \sigma_0$. Considering that a real-time monitoring system provides a measurement every minute which means there are daily 1 440 output data and if the alarm is set up at the decision threshold level, 72 false alarms per day ($X = \alpha = 5$ % of 1 440) are expected and this may not be acceptable in terms of human factor.

Table I.2 — Alarm setup parameter K and its associated false alarm rate X

$K = k_{1-\alpha}$	$X = \alpha$ %
1,282	10
1,645	5
1,960	2,5
2,327	1
3,091	10^{-1}
3,29	5×10^{-2}
3,720	10^{-2}
4	$3,2 \times 10^{-3}$
4,075	$2,3 \times 10^{-3}$
4,267	10^{-3}
4,756	10^{-4}
5	$2,9 \times 10^{-5}$
5,203	10^{-5}
6	$9,9 \times 10^{-8}$
Key K alarm setup parameter $k_{1-\alpha}$ Quantile of a standard normal distribution for a probability $1 - \alpha$, dimensionless X false alarm rate, in % α probability of a false positive, in %	

The detection limit, which represents the metrological capability of the measuring system, is the smallest value of the activity concentration detectable with a probability, $1 - \beta$.

The detection limit of the activity concentration, $c^\#$, can be approximated by [Formula \(I.2\)](#):

$$c^\# = 2 \cdot c^* \quad (\text{I.2})$$

considering that $\beta = \alpha$

If $\alpha = \beta = 5\%$ is considered, then $c^\# = 2 \cdot c^* = 3,29 \cdot \sigma_0$. From [Table I.2](#), if the alarm level is set up at the detection limit value, i.e. $K = 3,29$ corresponding to a false alarm rate $X = 5 \times 10^{-2}\%$, and if a daily 1 440 output data is still considered, then about 20 false alarms per day are expected and still may not be acceptable in terms of human factor.

If one false alarm per month ($X = 2,3 \times 10^{-3}\%$) is considered as acceptable then the alarm level can be set up at $4,075 \cdot \sigma_0$ which corresponds to the minimum decision threshold of the activity concentration, c^* , of the system. However, as the multiplier X is increased, the false alarm rate decreased but there is an increased probability that low-level concentrations of contaminants do not trigger an alarm. The upper limit, $c_{\min}^\triangleright$, of the confidence interval of the $c_{\min}$ for a given probability, $1 - \gamma$, expressed as, $c_{\min}^\triangleright = c_{\min} + k_{1-\frac{\gamma}{2}} \cdot u(c_{\min})$, shall be less than or equal to the regulatory activity concentration limit R_L .

It is preferable that the regulatory activity concentration limit, R_L , is much higher than $c_{\min}^\triangleright$ so the alarm level set up, $S_0 = c_{\min}^*$, at the value of c^* can be only an administrative alarm level decided by the licensee for operational reasons.

Figure I.1 summarizes all the metrological requirements previously described about the system sensitivity needed to achieve selected action levels [6]. This subject is discussed further in References [4] [5] and [6].

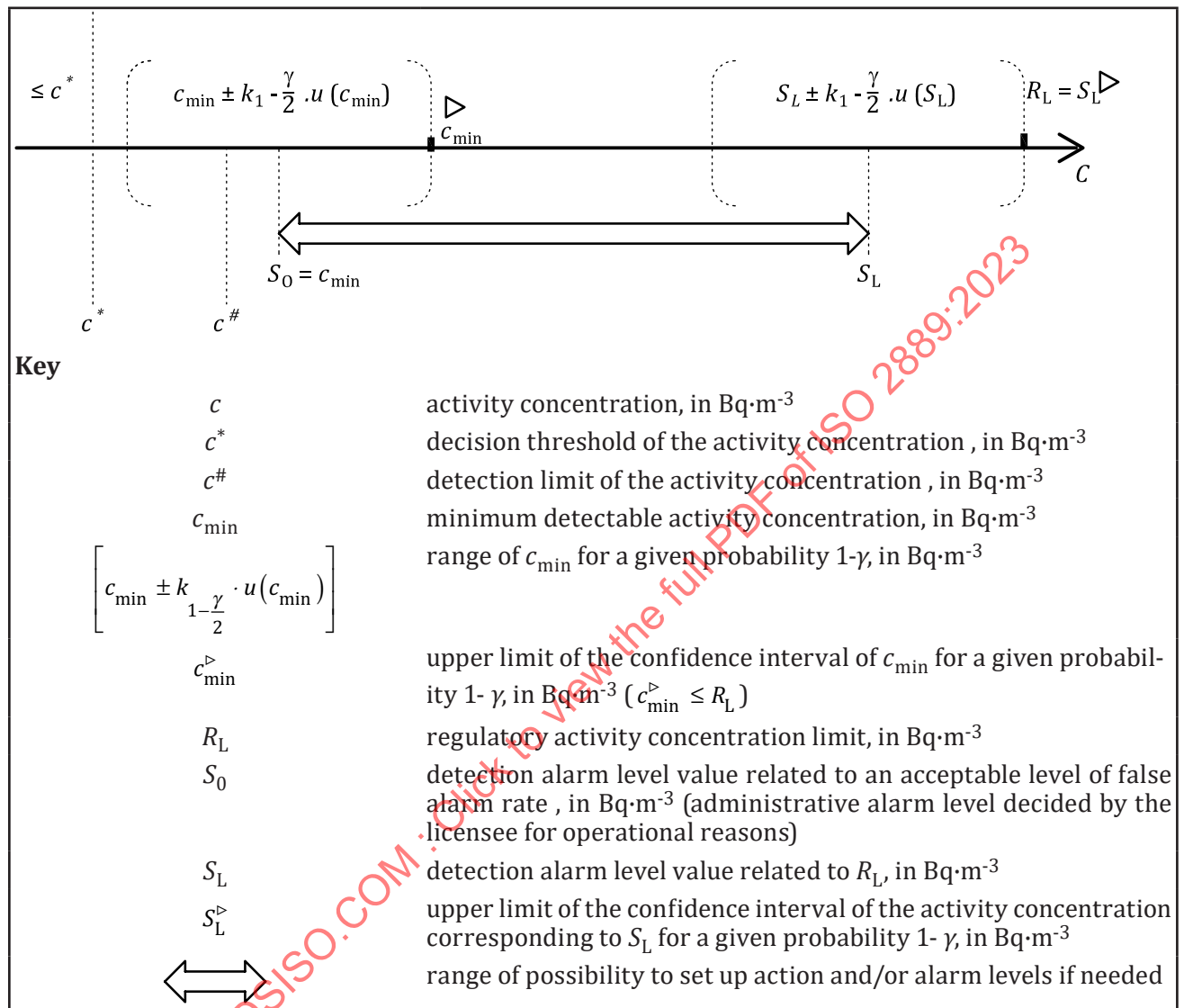


Figure I.1 — Synthesis of the metrological requirements to achieve selected action levels

It should also be noted that the performance of a real-time monitoring system does not only depend on the metrological aspect characterized by the decision threshold, the limit of detection and the $c^\#$ but also on its dynamic capacity characterized by its response time when facing a released activity event and especially in case of puff release of short time. In that situation if the response time is too long the actual value of the activity concentration cannot be measured.

The ideal performance is to have $c^\#$ as low as possible associated with a very short response time, but unfortunately these two criteria are in opposition. It may be then necessary to find a compromise between AMD and the response time in the choice and the settings of a real-time monitoring system in order to take into account both the dynamic characteristics of the released activity events and the respect of the requirements needed to achieve selected action levels.

I.5 System performance and availability alarms

System performance and availability alarms are a separate consideration from action levels based on effluent releases. The designer of the sampling system should consider the requirement for alarms activated by system-component failure that results in the inability to sample properly. Such system-failure alarms should be differentiated from alarms triggered by effluent-release action levels because a very different response is required. Establishing system-failure alarms should be based on a statistical evaluation and consequence analysis, considering acceptable levels of false positives and false negatives as discussed in [I.4](#).

System failure can take two forms. The first is a complete failure. Complete failure can be a system shutdown caused by an interruption of power or by the loss of a vital component. The complete failure should be indicated by an alarm to ensure that action is taken to restore operation. A complete failure due to the loss of a vital component can require a separate alarm for each mode of failure because the different failure modes can require different responses. The importance of the alarm and the priority of response should be determined and entered into the facility alarm-and-response plan.

The second form is a partial failure that compromises the quality of the output, renders the output unusable and causes the system to fall below safety or regulatory requirements. This type of failure should require a system alarm and a graded response because there can be differences in operation that require interpretation of instrumental data before activating an alarm. Two examples of partial failure are a significantly reduced sample flow rate and a significant leak in the sample-transport system. There are many other possibilities for partial failure and it can be impractical to trigger alarms for them all. Consequently, there are guidelines given for maintenance and inspections in [Annex J](#).

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Annex J **(informative)**

Quality assurance

J.1 General

Documentation, maintenance, inspection and calibration are key components of ensuring the quality of air samples.

J.2 Documentation

J.2.1 General

The quality assurance programme should ensure that the air sampling system and its components are characterized and documented.

J.2.2 Source term

This includes changes to the ventilation system or changes to processes that can affect the airborne effluent discharged. The nature of the processes serving each stack or duct should be identified, including information about the identity of the radionuclides as well as their chemical and physical forms. The air-cleaning systems associated with each stack or duct should be identified as well as the probable nature of releases resulting from the possible failure of these systems.

J.2.3 Effluent flow characterization

The results of studies to characterize the flow conditions of the effluents should be documented (e.g. spatial and temporal variations in velocity across the stack or duct, determination of cyclonic flow, estimates of particle size distributions, etc.). The documentation should include or list all procedures employed, times and dates of the measurements, individuals involved, equipment used and any pertinent information regarding facility operations.

J.2.4 Design and construction

Documentation that describes the objectives of each sampling system and lists all radionuclides and their potential physical and chemical forms should be available. If a particular component is present but not sampled, the reasons should be discussed.

The rationale and any supporting evidence for sampling at a particular location along the duct or stack should be documented. Similarly, the rationale for sampling at particular point(s) within (across) the stack or duct should be documented. Documentation that explains the rationale for the design of the sampling system should be available. This includes documentation regarding the choice of the transport system, the material, diameter and configuration of the sampling lines, the choice of filters or absorbers, the selection of flow meters, etc.

Also, there should be a means for allowing verification that the installed sampling equipment is that described in the documentation. This can be accomplished by identification marks on the installed components. An evaluation of particulate losses in the sampling lines should be documented. Other design documents that should be maintained include engineering change-control documents, equipment manuals and vendor-supplied information.

J.3 Maintenance and inspection

J.3.1 General

The requirements for maintenance and inspection depend on the nature of the sampling equipment. Routine maintenance may be performed as described in the manufacturer's equipment manuals. Non-routine maintenance should also be performed on the basis of the results of inspections. The guidance provided here can be used as appropriate, such as in cases where there are no manufacturer recommendations.

Inspection and maintenance activities should be described in procedures. Checklists should be employed as part of the inspection protocols, and, after use, a checklist should become a part of the record of the inspection. The inspection and maintenance records should include the nature of the inspection or maintenance, reasons for the inspection or maintenance, names of the individuals involved, times and dates, identity of the equipment employed and a description of any replacement parts or materials. All deficiencies identified during scheduled and unscheduled inspections should be recorded. Recommended maintenance and inspection guidelines are given below. Regularly scheduled inspections should be performed at least once a year, possibly concurrent with calibrations. Ideally, the same individuals responsible for the calibrations are also responsible for the inspections.

J.3.2 Inspections

Inspections should be performed routinely, quarterly or annually as appropriate and practicable, possibly concurrent with other maintenance. Inspections should include, but are not limited to, the following:

- position and orientation of sampling nozzles or inlets;
- condition of nozzle or inlet openings;
- dust accumulation in the sampling nozzles, inlets and transport lines;
- corrosion, physical damage or dust loading to the transport lines and equipment;
- filter-holder gaskets;
- leakage in the overall sample-transport system;
- tightness of all fittings and connections;
- condition of flow sensors;
- calibration of flow meters (the value of the flow rate determined by the test should not deviate from the nominal value more than 10 %).

J.3.3 Sampling system flow meter inspections

Mass flow meters should be checked at least annually with a secondary or transfer standard, where a transfer standard is typically a calibrated mass flow meter placed in series with the unit being tested. Unscheduled calibrations can be needed if any maintenance to the sampling system has been conducted that can affect the performance of the flow meter. The flow rate at which the mass flow meter is checked should be at a level that is within ± 25 % of the nominal design sampling rate of the system. If the flow rate, q_{std} , of the flow meter being tested differs by more than 10 % from the value indicated by a secondary standard, the flow meter should be removed from service for maintenance and calibration.

Flow through critical flow venturis should be checked at the start of each sampling period by observing the values of Δp_m (differential pressure across the meter) and Δp_f (differential pressure across the filter). If the value Δp_m is less than that needed for critical flow, the vacuum system should be checked to determine the cause. If the value of Δp_f is less than 70 % of that normally observed when the particular filter or collector is used, the critical flow meter should be inspected for blockage, or the sampling system should be checked for other possible problems. The critical flow meter should be removed from

service for cleaning and re-calibration if it is the cause of the erroneous reading. If the value of Δp_f is greater than 130 % of that normally observed, the filter or collector should be inspected for possible problems.

It might not be necessary to check rotameters in the field with secondary standards unless maintenance has been done or changes have been made to the sampling system that can affect its accuracy. A rotameter should be inspected at the start of each sampling interval to ensure that no foreign matter has been deposited on the inside surfaces in the measurement tube. If foreign matter is visible, the rotameter should be removed from service, cleaned and re-calibrated.

J.3.4 Continuous effluent flow measurement apparatus

On an annual basis, response checks should be made of the flow-rate readings from installed equipment through use of a reference Prandtl-type Pitot tube. If a thermal anemometer or Pitot tube is used in the stack or duct, the reference Pitot tube should be placed in the vicinity of the installed device at a point where, based on previous measurements (see [Annex A](#)), the velocity reading is either the same as that of the installed device or a known correction factor can be applied to provide a ratio of the two velocity readings. If the installed sensor is a Pitot tube, the velocities calculated from use of the two tubes should be within ± 10 % (after taking into account any correction factors). If the installed sensor is a thermal anemometer, the velocity, V , determined from use of the reference Pitot tube should be converted to the equivalent velocity, V_{std} , at standard conditions as given in [Formula \(J.1\)](#):

$$v_{std, ta} = v_{a, pt} \cdot \frac{T_{std}}{T_a} \cdot \frac{p_a}{p_{std}} \quad (J.1)$$

where

$v_{std, ta}$	is the velocity obtained from a single-point thermal anemometer at standard conditions, in $m \cdot s^{-1}$;
$v_{a, pt}$	is the velocity at actual conditions determined from use of the reference Pitot tube, in $m \cdot s^{-1}$;
T_a	is the temperature in the stack or duct, in K;
T_{std}	is the standard temperature, equal to 298 K;
p_a	is the pressure in the stack or duct, in Pa;
p_{std}	is the standard pressure, in kPa.

The ratio of the velocity at standard conditions indicated by the installed sensor and the reference sensor should be within ± 10 %.

If the velocity value from either an installed Pitot tube or thermal anemometer is outside of the specified range, the cause of the difference should be determined. It can be necessary to recalibrate the device. Also, if a sensor requires maintenance that can affect the calibration, the device should be recalibrated.

If the flow sensor is a Pitot tube, response checks should be made at least quarterly to verify the functionality of any pressure gauges used in conjunction with the Pitot-tube readout. This check may be a simple test to show that the application of a pressure differential causes an appropriate output of the gauge.

If an acoustic flow meter is used as the installed equipment, at least quarterly performance checks should be made by comparing the average velocity determined with the acoustic flow meter to the velocity at a reference point determined with a Prandtl-type Pitot tube. Based on the reference-method measurements (see [Annex A](#)) taken during calibration of the acoustic flow meter, a ratio can be established between the average velocity and the velocity at the selected reference point. The velocity measured with the acoustic flow meter should agree within ± 10 % of the single-point Pitot-tube measurement when the latter is corrected with the velocity ratio.

J.4 Calibration

J.4.1 General

Measurement and test equipment should be calibrated using standards whose calibration is traceable to the governing national institute of standards and measurements or derived from accepted values of natural physical constants. The principal calibration activities on a sampling system involve the verification of sample flow rate, sampling time and effluent flow rate. The suggested calibration frequency is annually for systems operated under normal or controlled environmental conditions. For systems used under extreme conditions, the calibrations should be conducted more frequently, e.g. every six months.

The methods used in calibrating all equipment and systems should be clearly described in procedures. The results of all calibrations should be recorded. This includes flow-meter and timer calibrations. The records should include the names of the individuals involved, times and dates, and the types and serial numbers of the calibration equipment.

J.4.2 Calibration of sampling system flow meters

The goal of the flow-meter calibration is to help ensure that the uncertainty in the measurement of the total volume of air sampled is $\pm 10\%$. [Annex E](#) describes a number of considerations for uncertainty analysis. All flow meters should be calibrated at least annually against devices that are either based on first principles (bubble meters or proof meters) or that are traceable to the governing national institute of standards and measurements.

The internal sensing region of a flow meter should be inspected before calibration. If there is any indication of surface deposits, the internal components of the flow meter should be cleaned or replaced.

Mass flow meters should be calibrated at conditions corresponding to 40 %, 70 %, 100 %, 130 % and 170 % of the nominal flow rate in terms of standard conditions. Other values may be used. However, technical justification should be documented to show that the use of the selected points provides calibration data of a quality equivalent, or superior, to the recommended points. If the flow rate through the sampling system can, under normal conditions or anticipated or accident conditions, exceed the limits recommended herein for flow calibration, additional calibration points should be used to encompass the possible operating range.

It can be necessary to calibrate critical venturi flow meters only at a single point that corresponds to operating conditions with a sufficient pressure differential across the meter such that the velocity at the throat of the meter is sonic. The temperature at the entrance of the critical flow meter during calibration should be within $\pm 5\text{ }^{\circ}\text{C}$ of the average temperature anticipated at that same location during sampling. The absolute pressure at the entrance of the critical flow meter should be within $\pm 2\%$ of the average absolute pressure anticipated at that location.

Rotameters should be calibrated at flow-rate conditions that correspond to the average anticipated flow rate during sampling, and at 75 % and 125 % of the anticipated sampling flow rate.

J.4.3 Calibration of effluent flow-measurement devices

An effluent flow-measurement system should be calibrated at least annually against the reference method discussed in [Annex A](#). The goal of the calibration is to measure flow rate relative to the reference method that is accurate to within $\pm 10\%$.

J.4.4 Calibration of timing devices

Timing devices should be calibrated at least annually. The uncertainty should not be greater than 1 min per month.

Annex K (informative)

Carbon-14 sampling and detection

K.1 Carbon-14 chemistry

Carbon-14 is a radioactive isotope of carbon with behaviour similar to that of the stable carbon isotopes. The nuclear reactions that produce ^{14}C in reactors with thermal neutrons are $^{17}\text{O}(\text{n},\alpha)^{14}\text{C}$ ($\sigma = 0,24 \text{ b}$), $^{14}\text{N}(\text{n},\text{p})^{14}\text{C}$ ($\sigma = 1,82 \text{ b}$) and $^{13}\text{C}(\text{n},\gamma)^{14}\text{C}$ ($\sigma = 0,0009 \text{ b}$).

NOTE $1 \text{ b} = 10^{-28} \text{ m}^2$.

Carbon-14 is of concern because of its very long half-life (5 730 a), the mobility of carbon in the environment and its ubiquitous presence in biological systems. Carbon-14 has been identified in airborne effluents from nuclear power plants in the forms of particulate ^{14}C , gaseous $^{14}\text{CO}_2$ and non- CO_2 gases. The latter can be ^{14}CO or various organic gases, e.g. $^{14}\text{CH}_4$.

K.2 Sampling considerations

K.2.1 General

When selecting a suitable sampling medium for ^{14}C , it is important to consider the presence of other contaminants (both radioactive and non-radioactive) in the sampling stream in addition to the physical and chemical forms of ^{14}C being collected. For example, if the effluent stream has high humidity, it is necessary to remove the moisture before using a molecular sieve to sample the $^{14}\text{CO}_2$, otherwise the moisture will saturate the molecular sieve. Likewise, it is necessary to remove HTO from the sample stream before collecting ^{14}C to avoid interference in the counting of ^{14}C , unless a chemical step is added before counting to remove the substances that interfere with the ^{14}C signal.

K.2.2 Particulate ^{14}C

The same requirements for sampling other particulate radioactive substance apply to particulate ^{14}C . These include considerations of sampling location, nozzle design and line penetration.

K.2.3 Gaseous ^{14}C

The same requirements for sampling other gaseous radioactive substance apply to gaseous ^{14}C .

K.3 Sampling media

K.3.1 Particulate ^{14}C

The filters installed to collect other particulate radioactive substances should also collect particulate ^{14}C . One very important consideration, though, is the filter material. Because it is normally necessary to combust the sample to separate the ^{14}C from other radionuclides, it is advantageous to use a non-combustible filter material, e.g. glass fibre. Furthermore, the filters sometimes collect only minute amounts of particulate carbon, so that it is necessary to add a carbon carrier to carry out the analysis. Lampblack compressed into a pellet has been found suitable as a carrier when glass-fibre filters are combusted in a tube furnace. Lampblack is generally made from fossil carbon, so it contains no ^{14}C , but this should be verified by combusting blank pellets. The radiochemical yield can be determined by the recovery of the carbon from the lampblack carrier.