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Dear Member

IEC COMMITTEE DRAFT (CD)

REPLY TO CSC@BSIGROUP.COM BEFORE 10TH OCTOBER 2017

Please find attached:

10/1026/CD - IEC 60480 ED3: Specification for re-use of SF6 and its mixtures in electrical equipment

IEC National Committees have been invited to comment on the above document. As a member of the responsible BSI committee you are asked to give your comments on the document. Please send any comments that you wish to be considered for submission as UK comments to IEC by the above date.

When submitting comments please ensure that they are entered into the [IEC comments template](#). If you have any queries in how to use the template then please do not hesitate to contact the Committee Service Centre.

It should be noted that this is often the final stage for the submission of major technical comment on the standard, as the national comments submitted to IEC determine whether this standard can progress to the next stage, i.e. circulation as a draft international standard (CDV).

Please also bear in mind that acceptance of a draft IEC standard means agreement in principle to it being the basis of a new British Standard, as it is BSI policy to implement all IEC projects as BS IEC standards unless any of the following situations apply:

- UK voted negatively at the FDIS stage.
- There is a current BS which covers the scope of the international standard and the BS continues to be the preferred document at the national level.
- There is an implemented EN standard covering the scope of the international publication.
- The International standard is subsequently agreed for UAP procedure in CLC and publication of the EN is expected within 12 months of the availability of the IEC publication.

If we do not hear from you by the above date we will submit 'no comment' to the IEC.

Yours sincerely,

Committee Service Centre

PROJECT NUMBER: IEC 60480 ED3	
DATE OF CIRCULATION: 2017-08-18	CLOSING DATE FOR COMMENTS: 2017-11-10
SUPERSEDES DOCUMENTS: 10/1021/RR	

IEC TC 10 : FLUIDS FOR ELECTROTECHNICAL APPLICATIONS	
SECRETARIAT: Italy	SECRETARY: Mr Massimo Pompili
OF INTEREST TO THE FOLLOWING COMMITTEES: TC 14, SC 17A, TC 20, SC 36A, TC 38, TC 112	PROPOSED HORIZONTAL STANDARD: <input type="checkbox"/> Other TC/SCs are requested to indicate their interest, if any, in this CD to the secretary.
FUNCTIONS CONCERNED: <input type="checkbox"/> EMC <input checked="" type="checkbox"/> ENVIRONMENT <input checked="" type="checkbox"/> QUALITY ASSURANCE <input type="checkbox"/> SAFETY	

This document is still under study and subject to change. It should not be used for reference purposes.

Recipients of this document are invited to submit, with their comments, notification of any relevant patent rights of which they are aware and to provide supporting documentation.

TITLE: Specification for re-use of SF6 and its mixtures in electrical equipment

NOTE FROM TC/SC OFFICERS:

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INTERNATIONAL ELECTROTECHNICAL COMMISSION

**SPECIFICATION FOR RE-USE OF SF₆
AND ITS MIXTURES IN ELECTRICAL EQUIPMENT**

FOREWORD

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International Standard IEC 60480 has been prepared by IEC technical committee 10: Fluids for electrotechnical applications.

This third edition cancels and replaces the second edition, published in 2004, and constitutes a technical revision.

The main changes with respect to the previous edition are listed below:

- the requirements for the re-use of SF₆ has been confirmed;
- this new edition contains specification for the re-use of SF₆ mixtures, namely SF₆/N₂ and SF₆/CF₄ mixtures.
- there is a new repartition of annexes of the three standards IEC 60376, IEC 60480 and IEC 62271-4. This new edition contains now five annexes, which are:
 - Description of the different methods of analysis (on-site and laboratory)
 - By-products of SF₆ and its mixtures

- Procedure for evaluating the potential effects on health from by-products of SF₆ and its mixtures
- Reclaiming recommendations
- Cryogenic reclaim of SF₆.

The text of this standard is based on the following documents:

FDIS	Report on voting
10/XX/FDIS	10/XX/RVD

Full information on the voting for the approval of this standard can be found in the report on voting indicated in the above table.

This publication has been drafted in accordance with the ISO/IEC Directives, Part 2.

The committee has decided that the contents of this publication will remain unchanged until the maintenance result date indicated on the IEC web site under "<http://webstore.iec.ch>" in the data related to the specific publication. At this date, the publication will be

- reconfirmed;
- withdrawn;
- replaced by a revised edition, or
- amended.

SPECIFICATION FOR REUSE OF SF₆ AND ITS MIXTURES IN ELECTRICAL EQUIPMENT

1. Scope

Sulphur hexafluoride (SF₆), Nitrogen (N₂) and Carbon Tetrafluoride (CF₄), are gases commonly used for electrical equipment. Taking into account environmental concerns, particular attention shall be paid to re-use criteria for SF₆ and its mixtures with N₂ and CF₄ as used in electrical equipment.

This International Standard gives criteria for re-using of SF₆ and its mixtures after recovery and reclaiming from electrical equipment (e.g. for maintenance, at the end-of-life). Procedures for recovering and reclaiming used SF₆ and its mixtures are out of the scope of this Standard and are described in IEC 62271-4.

Description of the different methods of analysis, by-products, procedure for evaluating the potential health effects from by-products, cryogenic reclaiming of SF₆, and reclaiming recommendations are provided in the annexes of this standard.

Storage, transportation and disposal of SF₆ and its mixtures are covered by IEC62271-4 and must be carried out according to local regulations. Procedures to determine SF₆ leakages are described in IEC 60068-2-17.

2. Normative references

The following referenced documents are indispensable for the application 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.

IEC 60050(191): *International Electrotechnical Vocabulary (IEV) – Chapter 191: Dependability and quality of service*

IEC 60050(212): *International Electrotechnical Vocabulary (IEV) – Chapter 212: Insulating solids, liquids, gases*

IEC 60050(441): *International Electrotechnical Vocabulary (IEV) – Chapter 441: Switchgear, controlgear and fuses*

IEC 60050(826): *International Electrotechnical Vocabulary (IEV) – Chapter 826: Electrical installations of buildings*

IEC 60376: *Specification of technical grade sulphur hexafluoride (SF₆) and complementary gases that can be used in its mixtures for use in electrical equipment*

IEC 62271-1: *High-voltage switchgear and controlgear – Part 1: Common specifications*

IEC 62271-4: *High-voltage switchgear and controlgear – Part 4: Handling procedures for sulphur hexafluoride (SF₆) and its mixtures*

IEC 60068-2-17 *Basic environmental testing procedures – Part 2: Tests – Test Q: Sealing*

106 3. Terms and definitions

107 3.1 General

108 For the purposes of this document, the terms and definitions contained in IEC 60050(191)
109 IEC 60050(212), IEC 60050(441) and IEC 60050(826), as well as the following, apply.

110 3.2 Electrical equipment

111 item used for such purposes as generation, conversion, transmission, distribution or utilization of
112 electrical energy, such as electric machines, transformers, switchgear and controlgear, measuring
113 instruments, protective devices, wiring systems, current-using equipment, insulated bushings, surge
114 arresters

115 3.3 Enclosure

116 part providing protection of equipment against certain external influences and, in any direction,
117 against direct contact

118 [IEV 826-03-12]

119 3.4 Maintenance

120 combination of all technical and administrative actions, including supervision actions, intended to
121 retain an item in, or restore it to, a state in which it can perform a required function

122 [IEV 191-07-01]

123 3.5 Container

124 vessel suitable for the containment of gases either in gaseous or liquid phase, according to local
125 and/or international safety and transportation regulations

126 3.6 Used sulphur hexafluoride

127 SF₆ which has been introduced into electrical equipment

128 3.7 reclaiming

129 process of contaminants removal from an insulating liquid or gas

130 [IEV 212-09-05, modified]

131 3.8 Recovery

132 process of transferring gas from electrical equipment to an alternate container

133 3.9 SF₆ mixture

134 gas mixture formed by SF₆ and a complementary gas, typically N₂ or CF₄

135 3.10 Contaminants

136 foreign substances or materials in an insulating liquid or gas which usually has a deleterious effect on
137 one or more properties

138 [SOURCE: IEC 60050-212:2010, 212-17-27, modified]

139 3.11 By-products

140 contaminants which are formed by the degradation of SF₆ and its mixtures by electrical arcs or sparks

141 4. Contaminants and their sources

142 4.1 General

143 SF₆ recovered from electrical equipment in operation contains several kinds of contaminants.
144 Contaminants in recovered SF₆ come both from gas handling and from use.

Table 1 summarizes the main contaminants and their sources, additional information is available in Annex B.

Table 1 – SF₆ contaminants

SF ₆ situation and use	Origin	Possible contaminants
Handling and in service	Leaks and incomplete evacuation Desorption	For pure SF ₆ : Air, Oil, H ₂ O For SF ₆ mixtures: Air, Oil, H ₂ O, N ₂ , CF ₄
Insulating function	Partial discharges: Corona and sparking	Gaseous by-products: HF, SO ₂ , SOF ₂ , SOF ₄ , SO ₂ F ₂
Switching equipment	Switching arc erosion	Gaseous by-products: HF, SO ₂ , SOF ₂ , SOF ₄ , SO ₂ F ₂ , SF ₄ , CF ₄ , WF ₆ Solid by-products: Metal dusts, particles, AlF ₃ , FeF ₃ , WO ₃ , CuF ₂
	Mechanical erosion	Metal dusts, particles
Internal arc	Melting and decomposition of materials	Gaseous by-products: HF, SO ₂ , SOF ₂ , SOF ₄ , SO ₂ F ₂ , SF ₄ , CF ₄ , WF ₆ Solid by-products: Metal dusts, particles, AlF ₃ , FeF ₃ , WO ₃ , CuF ₂

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149 **4.2 Contaminants from handling and use**

150 Filling and recovering gas leads to the additional contamination with air and water (humidity).

151 Moisture desorbs from internal surfaces of the equipment and from polymeric parts. Oil from handling
152 equipment (pumps and compressors) may also be inadvertently introduced.

153 When using gas mixtures, the possibility of cross contamination shall be considered.

154 **4.3 SF₆ by-products in equipment that only have an insulating function**

155 The essential process is the decomposition of SF₆ by partial discharges (corona and sparking). The
156 immediate products are fragments of SF₆, such as SF₅, SF₄ and F, that combine with O₂ and H₂O to
157 form compounds, mainly HF, SO₂, SOF₂, SOF₄ and SO₂F₂. Due to low energy of the partial
158 discharges, the accumulated quantities of these compounds are usually negligible.

159 **4.4 SF₆ by-products in switching equipment**

160 During current interruption, the existence of high temperature arcs leads to the formation of
161 decomposition products of SF₆, vaporized electrode metal, plastics and impurities. In addition,
162 chemical reactions take place between the products formed (see Table 1).

163 The quantity of these by-products depends on the number of operations, the cumulative short circuit
164 current, the design of equipment and the use of adsorbers (solid adsorbents).

165 Switching equipment may also contain particles and metal dust coming from the rubbing of contacts.

166 **4.5 SF₆ by-products from internal arcs**

167 The occurrence of an internal arc is extremely rare. The expected impurities in SF₆ in faulted
168 equipment are similar to those normally found in switching equipment. The difference lies in the

quantity of compounds, which create a potential toxic risk (see Clause 7). In addition, significant vaporization of metallic material occurs and creates additional reaction products such as dust.

4.6 SF₆ mixtures specific by-products

For SF₆ mixtures, the usual SF₆ by-products mentioned in Table 1 and specific mixture by-products, such as Nitrogen Oxide(s) and Nitrogen Fluoride(s) for SF₆/N₂ and Fluorocarbon(s) for SF₆/CF₄, are produced. The quantities depend on the mixture composition, contaminants and energy introduced. For typical SF₆ mixtures, the gas decomposition rates are not expected to exceed those for SF₆.

Within the decomposition products generated in mixtures, SF₆ by-products are generally predominant in terms of quantity and toxicity. Safety procedures related to the presence of the usual SF₆ by-products shall also apply in applications with SF₆ mixtures.

4.7 Requirements for reclaimed SF₆

Table 2 – Requirements for reclaimed SF₆

Substances ^(a)	Concentration
SF ₆	>97% volume
Air and/or CF ₄	< 30000 µl/l (3 % volume)
H ₂ O	< 200 µl/l
Mineral oil	< 10 mg/kg ^(b)
Acidity	< 50 µl/l total or 12 µl/l for (SO ₂ +SOF ₂) or 25 µl/l HF
a) H ₂ S and CO have been considered irrelevant due to lack of valuable data. b) If gas handling equipment (pump, compressor) containing oil is used, it may be necessary to measure the oil content of the SF ₆ . If all equipment in contact with the SF ₆ is oil-free, then it is not necessary to measure oil content.	

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For determination of total acidity, the sum of all acidic compounds is reported as one value and expressed as total acidity or 12 µl/l for (SO₂+SOF₂) or 25 µl/l HF

4.8 Requirements for SF₆ mixtures

Table 3 – Requirements for SF₆/N₂ mixtures

Substances	Concentrations
Complementary gas used with SF ₆	Must be identical to the OEM specifications
SF ₆ percentage	±5 % volume of the rated value ^(a)
Air and CF ₄	<3 % volume ^(a)
H ₂ O	<200 µl/l
Mineral oil	<10 mg/kg (b)
Total acidity	<50 µl/l total or 12 µl/l for (SO ₂ +SOF ₂) or 25 µl/l HF
Storage conditions	
shall comply to J.7 of IEC 62271-4 in order to prevent liquefaction of SF ₆	

- a) or unless otherwise specified by the Original Equipment manufacturer (OEM).
- b) If gas handling equipment (pump, compressor) containing oil is used, it may be necessary to measure the oil content of the SF₆. If all equipment in contact with the SF₆ is oil-free, then it is not necessary to measure oil content.

Table 4 - Requirements for SF₆/CF₄ mixtures

Substances	Concentrations
Complementary gas used with SF ₆	Must be identical to the OEM specifications
SF ₆ percentage	±5 % volume of the rated value (a)
Air and N ₂	<3 % volume (a)
H ₂ O	<200 µl/l
Mineral oil	<10 mg/kg (b)
Total acidity	< 50 µl/l total or 12 µl/l for (SO ₂ +SOF ₂) or 25 µl/l HF
Storage conditions	
shall comply to J.7 of IEC 62271-4 in order to prevent liquefaction of SF ₆	
a) or unless otherwise specified by the Original Equipment manufacturer (OEM).	
b) If gas handling equipment (pump, compressor) containing oil is used, it may be necessary to measure the oil content of the SF ₆ . If all equipment in contact with the SF ₆ is oil-free, then it is not necessary to measure oil content.	

5. Reclaim of SF₆ and SF₆ mixtures

5.1 Feasibility and process

The quality of reclaimed SF₆ shall meet the requirements of this standard.

All occurring contaminants are formed in normal operation and can generally be eliminated on-site. Table 5 lists the type of contamination and the methods recommended for removing the impurities.

Table 5 – Suggested reclaiming operations

Operational contamination	Moisture (water vapour)	Gaseous decomposition products	Solid decomposition products	Air, N ₂ and/or CF ₄	Contaminated with mineral oil
Reclaiming method	Adsorption with molecular sieve	Adsorption with activated aluminium oxide	Retaining with solid filters	SF ₆ gas separation device (*)	Activated charcoal filter
(*) NOTE: For SF ₆ mixtures, these gaseous contaminants could not be removed easily on-site. In each situation, an evaluation of the reclaiming options should be done to determine if the mixture could be reclaimed on-site.					

Various types of adsorbent materials are available to remove contaminants from SF₆ gas (see Table 6).

Table 6 – Typical adsorbents for various SF₆ impurities

Adsorbent	Contaminants removed
-----------	----------------------

Molecular sieve 4A	Water, SO ₂ , SOF ₂ , SF ₄
Molecular sieve 13X	Water, SO ₂ , SOF ₂ , SF ₄ (also adsorbs some SF ₆)
Activated aluminium oxide	Water, SO ₂ , SOF ₂ , SF ₄ , HF
Soda lime (CaO-NaOH)	Water, SO ₂ F ₂ , HF
Activated charcoal	Oil vapour

200

201 Through the corresponding handling with service devices, the gas will be purified and finally verified. If
 202 the salvaged material meets the requirements for an unreserved re-use in electrical installations, the
 203 direct refilling with a filling and evacuating device will be required. This handling covers the normal
 204 operation of electrical equipment.

205 If the testing of the gas shows an unacceptable level of decay compared to the requirements of this
 206 standard, a decision regarding the reclaiming method has to be made depending on the level and type
 207 of contamination. In general, re-purifying the gas on-site with a service device plus a separation
 208 device will be the most favourable way. However, if re-use is not possible, disposal will be necessary.
 209 In this case, the gas must be sent to the SF₆ manufacturer or reclaimer.

210 Figure 1 defines the selection procedure to determine the best use of SF₆ after recovery for potential
 211 treatment.

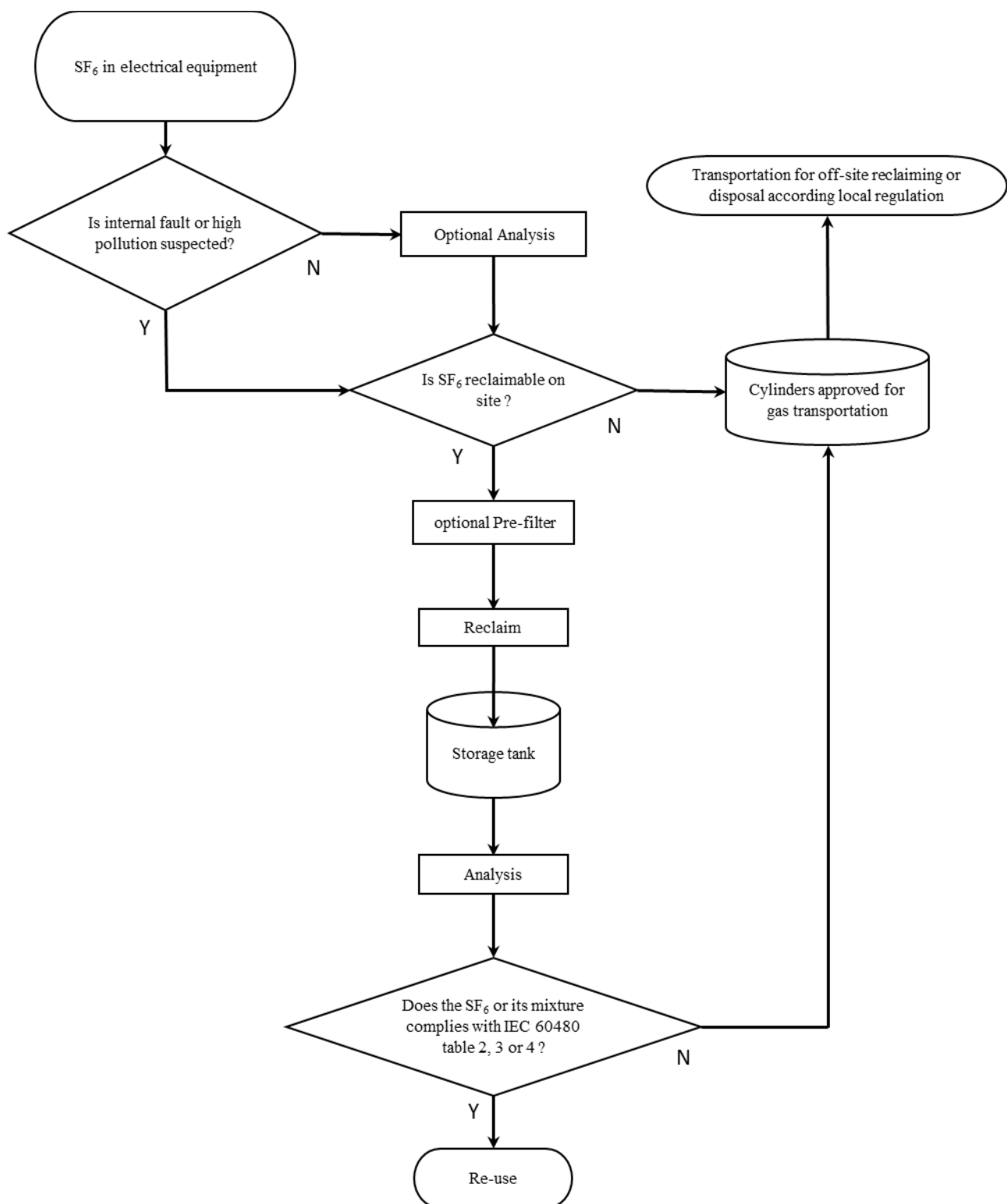


Figure 1 – Decision flow chart for the destination of recovered SF₆

- For contaminants of water or decomposition products, the question of whether the SF₆ is reclaimable on-site depends only on the performance of the filters available. The addition of

external pre-filters may be required to increase the efficiency of the reclaiming process. If the SF₆ is not reclaimable on-site, then it shall be returned to the SF₆ manufacturer or sent to a reclaiming or disposal company.

- The case of contamination with air, N₂ and/or CF₄ shall be considered separately.
 - For non-mixed SF₆, if the concentration of air and/or CF₄ exceeds the maximum acceptable impurity level as given in Table 2, and if the container from which the sample has been taken contains liquid SF₆, then transfer SF₆ from the gas phase into a second container. The transfer should be continued until a sample from the first reservoir satisfies the maximum acceptable level. The contents of the second container cannot be reclaimed on-site. Any container that contains no liquid SF₆, i.e. only the gas phase, requires the analysis of only one sample for air and CF₄ to determine if it is suitable for re-use or cannot be reclaimed on-site;
 - For mixed SF₆, if the concentration of air, N₂ and/or CF₄ exceeds the maximum acceptable impurity level as given in Table 3 or 4, an evaluation of the reclaiming options should be done to determine if the mixture could be reclaimed on-site.

5.2 Detection techniques for checking the quality of the gases

5.2.1 General

The detection techniques shall be used on gas samples. Details of the procedures for obtaining a sample and of the analytical methods are provided in Annex A.

5.2.2 On-site analysis

If on-site analysis systems are used, they shall be equipped with a gas recovery system. The release of SF₆ to the atmosphere shall be avoided and the safety of personnel shall be assured. On-site available analytical methods are shown respectively in Table 7 and Annex A.

In case of internal arc fault suspected or a circuit-breaker which failed to clear, caution should be taken to not damage the measuring instrument.

Table 7 – On-site methods

Substances	Methods available
SF ₆	Portable gas chromatograph with thermal conductivity detector (GC-TCD) Infrared spectroscopy Electrochemical sensor
N ₂ in SF ₆ mixture	Portable gas chromatograph with thermal conductivity detector (GC-TCD) Infrared spectroscopy ¹ Electrochemical sensor ^{1, 2}
CF ₄ in SF ₆ mixture	Portable gas chromatograph with thermal conductivity detector (GC-TCD) Infrared spectroscopy Electrochemical sensor ^{1, 2}
Decomposition products: SO ₂ , SOF ₂ , SO ₂ F ₂ (See Note)	Portable gas chromatograph with thermal conductivity detector (GC-TCD) Infrared spectroscopy Electrochemical sensor
HF	Infrared spectroscopy Electrochemical sensor
Air and CF ₄	Density meter (for the % of SF ₆) Portable GC-TCD Infrared spectroscopy
Water (humidity)	Electronic hygrometer

	Chilled mirror hygrometer Infrared spectroscopy Electrochemical sensor
Oil	Tube for mineral oil
<p>NOTE SO₂F₂ may be present in the gas but cannot be measured with detector tubes.</p> <p>Note 1: Indirect quantification done by the subtraction of the SF₆ content</p> <p>Note 2: Mixture identification and composition must be known before analysis</p> <p>Note 3: Gas chromatography is only used to determine the concentration of N² or CF₄ in SF₆ mixtures</p>	

5.2.3 Laboratory analysis

If no equipment is available on-site, the following recommended techniques shall be used. Laboratory methods are intended to provide a quantitative assessment of the contaminants in a gas sample.

Water is the only contaminant that cannot be determined accurately from a sampling cylinder. The water content of a sample in a cylinder taken from a container is not representative of the water content in the container because water adsorbs on all surfaces. Therefore, the water analysis should always be carried out directly on the container on-site. There is no recommended order of analysis.

Table 8 – Laboratory methods

Impurities	Methods available
SF ₆	Gas chromatograph (Note 2) Infrared absorption
N ₂ in gas mixture	Gas chromatograph (Note 2)
CF ₄ in gas mixture	Gas (Note 2) Infrared absorption
Air: oxygen and nitrogen	Gas chromatograph (Note 2)
CF ₄ as contaminant	Gas chromatograph (Note 2) Infrared absorption
Oil	Infrared absorption Gas chromatograph (Note 2)
Decomposition products (total acidity): SO ₂ , SOF ₂ , SO ₂ F ₂ , SF ₄ , HF ¹	Gas chromatograph (Note 2) Ion chromatography Infrared absorption Wet chemistry
<p>NOTE 1 Ion chromatography, infrared absorption and wet chemistry are the only method available to quantify HF</p> <p>NOTE 2 Gas chromatography can be performed by using adequate detectors for the determination of different species of contaminations or to control the mixing gas concentrations.</p>	

6. Handling, storage and transportation

Refer to IEC 62271-4.

7. Safety and first aid

7.1 General safety rules

7.1.1 General

Before starting any maintenance/service work in electric power equipment, a detailed inspection of the state of the equipment shall be carried out and documented. In addition to the local safety regulations, at least the following general safety rules shall be followed:

- 261 • Switch off and isolate;
- 262 • Secure against re-closing;
- 263 • Verify that equipment is de-energised;
- 264 • Earth and short-circuit the equipment;
- 265 • Cover or fence off nearby live parts.

266 Written documents giving permission to work on the electric power equipment should be agreed and
 267 signed by both the Owner / User of the equipment and the service provider.

268 Table 9 lists the major issues to consider when working with SF₆ electric power equipment.

269 **Table 9 – Measures when working with SF₆ electric power equipment**

Item	Work in the vicinity of equipment (operation of equipment, visual check, room-cleaning)	Filling, recovery, evacuation of gas compartments	Opening of gas compartments, work on open compartments
Material safety data sheet/operational manuals	Not required	Applicable	Applicable
Training	Applicable [NOTE]	Applicable	Applicable
SF ₆ handling equipment	Not required	Applicable	Applicable
Cleaning/ neutralising equipment	Not required	Not required	Applicable
Personal protection equipment	Not required	Not required	Applicable
NOTE General information should be specified according to type of work and installation, according to local safety regulations.			

270

271 A notice stating that open fire, naked flames (e.g. matches), smoking, use of heat engines, heating to
 272 more than 200 °C and welding without special precautions are prohibited and instructions for giving
 273 first-aid (see 7.3) should be displayed while SF₆ is being handled in any location.

274 When a gas compartment is opened after the electric power equipment has been in service, in order
 275 to avoid contact with the fine solid by-products, which may be present, personnel should wear suitable
 276 protective clothing. Particular attention should be given to protecting the eyes and the respiratory
 277 tract. Personnel working in or near to opened gas compartments, which have contained normally
 278 arced or heavily arced SF₆ should:

- 279 • use suitable tools and equipment;
- 280 • wear suitable protective clothing (see Table 10 and 7.1.5);
- 281 • observe high standards of personal hygiene;
- 282 • clean themselves and their equipment using disposable materials, before leaving the work area;
- 283 • remove protective clothing and wash them thoroughly as soon as possible after having left the
 284 work area;
- 285 • ensure that clothing, tools and components that have been in contact with by-products are
 286 securely packed in sealed bags or other sealed containers and are subsequently treated to
 287 neutralise any residues.

7.1.2 Protection of personnel

SF₆ is handled by certified personnel, only. Different training modules may be adopted, depending on the kind of work to be performed on the electric power equipment.

Specific training covering the recovery of used SF₆ from a gas-filled compartment is required for personnel opening or entering the gas compartment. The Operating Instruction Manual of the equipment provided by the Original Equipment Manufacturer should be strictly followed.

Table 10 gives an overview of the potential risks, safety precautions as well as safety equipment and tools required when opening or entering a gas compartment. No substantial difference between indoor and outdoor conditions is given as the operator opens or enters the compartment.

Table 10 – Safety measures when opening or accessing gas compartments

Item	Any compartment which contained normally or heavily arced SF ₆	Any compartment which contained non-arc'd SF ₆
Potential risk	Fumes of cleaning substances O ₂ starvation Remaining used SF ₆ Residual reactive gaseous by-products Solid by-products and adsorber materials	Fumes of cleaning substances O ₂ starvation Remaining used SF ₆ or other gas from production process
Safety precaution	Removal of solid by-products and adsorber materials Ventilation Measurement of O ₂ concentration when entering Wear personal protective equipment Protect solid by-products against hydrolysis	Ventilation Measurement of O ₂ concentration before entering
Safety equipment and tools	Suction ventilator or vacuum cleaner O ₂ concentration measuring device Single use protective overalls, protective footwear, hair cap Acid proof safety gloves Full face mask (preferred) or, at least, breathing protective mask Protective goggles Environmental protection against rain or wind (outdoor only)	Suction ventilator or vacuum cleaner O ₂ concentration measuring device

298

Like any gas but oxygen, a concentration greater than 190 ml/l (19 % by volume) of SF₆ in the air is considered as a potential risk of asphyxiation. This is because it reduces the oxygen concentration down to 160 ml/l (16 % by volume), which is usually considered as the threshold for asphyxiation. As a consequence it is recommended that the oxygen content in the gas compartment be measured prior to accessing it. In addition to that, the oxygen content in the ambient environment may be checked when working in confined spaces.

Eating, drinking and smoking are prohibited when accessing or opening a gas compartment. It is recommended that clothes should be changed and the skin washed as soon as possible after the work to prevent potential danger of irritation or burns.

7.1.3 Handling of contaminated safety equipment and tools

Equipment and tools, which have been in contact with solid by-products or adsorber materials are considered to be contaminated. They are collected afterwards and placed in plastic bags. The plastic bags are sealed with tape and labelled. Disposal is done according to the local regulations.

Reusable equipment and tools should be washed and neutralised in a water/soda solution with 10 % by weight liquid soda or equivalent and then rinsed with clean water. Examples of neutralising solutions are given in Table 11.

Disposal of both the waste water/soda solution and the waste washing water is done according to the local regulations.

Table 11 – Neutralising solutions

Active agent	Formula	Concentration kg/100 l	T1 [NOTE 1] hours	T2 [NOTE 2] hours	Reference
Lime	Ca(OH) ₂	Saturated	Not applicable	24	[9]
Sodium carbonate (washing soda)	Na ₂ CO ₃	1,1 3 10 [NOTE 3] 10-14 [NOTE 3] 3	Not applicable Wash not applicable 1 not applicable	24 not applicable 0,25 48 not applicable	[10] [9] [11] [12] [13]
Sodium bicarbonate	NaHCO ₃	1 [NOTE 4]	Not applicable	not applicable –	[9]
<p>NOTE 1 Reusable safety equipment, tools, gas compartments, and internal parts of compartments which contained normally arced SF₆ should, where practicable, be treated with a neutralising solution for a time period T1. They should then be rinsed with clean water.</p> <p>NOTE 2 Reusable safety equipment, tools, gas compartments, and internal parts of compartments which contained heavily arced SF₆ should, where practicable, be treated with a neutralising solution for a time period T2. They should then be rinsed with clean water.</p> <p>NOTE 3 When using alkaline solutions at such high concentrations, care should be taken to avoid contact with the skin and eyes.</p> <p>NOTE 4 Recommended for washing the skin.</p>					

318

319 7.1.4 Pressurised equipment and tools or measuring devices

320 As with any pressurised gas, a sudden volume expansion results in a local temperature drop and may
321 cause freezing. Suitable insulated gloves (e.g. leather, not latex) and protective goggles should be
322 worn when working on high-pressure piping, valves or connectors, during filling operations.

323 All equipment and tools used during SF₆ handling potentially contain gaseous or liquid SF₆ under high
324 pressure. They should be handled with care.

325 7.1.5 Personal safety and protective equipment

326 The use of personal safety and protective equipment is not related to the presence of the SF₆ electric
327 power equipment itself. Standard safety shoes, helmet, and protective goggles may be required
328 according to local regulations when working in the vicinity of the switchgear.

329 In case of abnormal release of SF₆ due to external fire or internal arc fault, additional safety rules
330 apply for entering a switchgear room, according to 6.

331 Depending on the type of work to be performed on site and according to Table B.6, workers engaged
332 in handling SF₆ are provided with the following personal safety and protective equipment:

- 333 • Protective gloves: suitable acid resistant gloves made of e.g. neoprene, PVC, rubber;
- 334 • Protective goggles: chemical type industrial goggles according to local regulations (e.g. European
335 Standard EN 166);

- Breathing protective mask: to be used in conjunction with protective goggles, the breathing protective mask helps to protect mouth and nose against dust and should be used in case of exposure to minor dusty areas only;
- Full face mask: for short-term inspection and work where ventilation can be provided but where the concentration of by-products may exceed the appropriate maximum level, a face mask with changeable active charcoal filter is used according to local regulations (e.g. European Standards EN 140, EN 141 and EN 143 specify masks, gas filters and particle filters, respectively. Combined filters of type A2/B2/E2/K2/P3 manufactured to these standards are available and are able to provide protection against by-products including particles with a diameter greater than 1 µm);
- Single use protective overall: dust proof protective clothes to wear over normal clothes, shoe covers, hair cap. Pocket-less, hooded, non-permeable (e.g. bonded polypropylene) disposable industrial grade overalls having elastic ankle and wrist grips, overlapping the footwear and gloves;
- Respirator: when entering indoor applications after major leakage or internal arcing when the concentration of O₂ or the amount of by-products are at unsafe levels, a respirator is used according to local regulations (e.g. European Standard EN 136);
- O₂ concentration measurement device for permanent monitoring of the O₂ content in the environment;
- Environmental protection for outdoor work locations: temporary shelter to prevent the ingress of rain and the wind dispersing solid by-products (if any) while the gas compartment is open;
- Vacuum cleaner: a high efficiency dedicated vacuum cleaner, equipped with a filter capable of trapping particles in the range of 1 µm, and a non-metallic open-ended nozzle according to local regulations (e.g. a type H machine in accordance with BS 5415, Supplement No. 1, 1986);
- Suction ventilator: equipment for forced ventilation of enclosed spaces and other inaccessible areas. Such equipment might be portable or permanently installed, depending on the size of the installation.

7.1.6 Facilities and services

Where gas compartments containing used SF₆ have to be recovered and opened, it is desirable that adequate washing facilities for workers be available, and a supply of water for preparing cleaning solutions may be required.

7.2 Additional safety measures in case of abnormal release of SF₆ due to external fire or internal arc fault

General safety recommendations to adopt when working with SF₆ on site are given in 7.1. Clause 7.2 describes additional safety measures in case of abnormal release of SF₆ due to external fire or internal arc fault.

Under these circumstances, personnel trained on modules C1 or C2 (see IEC 62271-4 Annex C) are allowed to enter and clean the switchgear room or to access the electric power equipment. Table 12 gives an overview of the potential risks, safety precautions as well as safety equipment and tools required.

NOTE Fire fighting can enter the switchgear room in the terms described in the local regulations.

376

Table 12 – Additional safety measures

Item	Abnormal release of heavily arced SF ₆	Abnormal release of non-arc'd or normally arced SF ₆
Potential risk	Fumes of cleaning substances O ₂ starvation SF ₆ abnormally released Residual reactive gaseous by-products Solid by-products	Fumes of cleaning substances O ₂ starvation SF ₆ abnormally released
Safety precaution	Removal of solid by-products Ventilation Measurement of O ₂ concentration when entering Wear personal protective equipment	Ventilation Measurement of O ₂ concentration when entering
Safety equipment and tools	Suction ventilator or vacuum cleaner O ₂ concentration measuring device Single use protective clothes, shoe covers, hair cap Acid proof safety gloves Full face mask (preferred) or, at least, breathing protective mask and protective goggles	Suction ventilator or vacuum cleaner O ₂ concentration measuring device

377

378 The same rules apply to areas below the level at which the release occurred, poorly ventilated or
 379 unventilated areas (e.g. cable trenches, inspection pits, drainage systems). Outdoor equipment needs
 380 standard measures for outdoor conditions to be applied (e.g. rain protection, wind protection). Natural
 381 ventilation is normally enough to prevent the potential risk of O₂ starvation.

382 A second person being in continuous visual and audible contact should be present, when entering
 383 areas that may have a low oxygen concentration.

384 7.3 First aid equipment and treatment

385 7.3.1 General

386 First aid equipment includes:

- 387 • Normal industrial first-aid equipment including eyewash equipment containing a saline solution;
- 388 • Means for contacting emergency services;
- 389 • Guidance for medical doctors.

390 The application of the general safety rules (see 7.1) should minimise the likelihood of accidents. In
 391 case of an accident, first aid treatment should be applied as described in the following clauses.

392 7.3.2 Irritation of the skin

393 In case of signs of skin irritation, the personnel are removed from the area. Contaminated clothing is
 394 removed and the affected part washed with cool running water. Professional medical advice should be
 395 sought.

396 NOTE While seeking for professional medical advice, the affected part can be treated with calcium gluconate gel (HF
 397 antidote gel) as a remedy after washing for fluoride acid on skin.

398 7.3.3 Irritation of the eyes

399 In case of signs of eyes irritation, the personnel evacuate the area. Irrigation of the eye or eyes is
 400 carried out immediately and continued until a medical professional advises the patient to stop.

401 **7.3.4 Breathing difficulty**

402 All personnel should immediately evacuate the affected area and move into the fresh air.
403 Contaminated clothing on a person with breathing difficulties should be removed and this person
404 covered with a blanket and kept still and under continuous observation. Emergency medical
405 assistance shall be summoned without delay. If the patient stops breathing, artificial respiration must
406 be immediately commenced by trained and qualified personnel.

407 **8. Environmental aspects**

408 Most created by-products are recombining to SF₆ inside the gas compartment or are transferred to
409 solid by-products, some remain as toxic gaseous by-products but are trapped by molecular sieve. At
410 the end of life all gaseous by-products and molecular sieve are recovered from gas-filled compartment
411 and shall not be released to the environment.

412 Low leakage rates, according to IEC 62271-1, during lifetime are leading to immediate dilution due to
413 natural ventilation and therefore having negligible impact on the environment.

414 Major failures causing gas releases are extremely rare as records from 50 years of experience show.
415 The quantities released in such extreme cases are again very limited by the fact that standard design
416 of products is compartmented, limiting the fault to the place where it originates.

Annex A (informative)

Description of the different methods of analysis (on-site and laboratory)

A.1 Sampling

A sample must be representative of the gas composition in a container. The sample should be taken from the liquid phase. If there is no liquid SF₆ in the container then the sample will be truly representative of the gas.

Before taking a sample from the container, the volume of all the connections from the container to the analytical equipment should be evacuated with a vacuum pump. In the case of a container that is connected to a sampling port, via a long section of narrow piping, the pipe section should be purged with gas from the container. This will ensure that the sample is representative of the SF₆ in the container.

As far as possible, the components in contact with the SF₆ should be constructed of stainless steel or other chemically resistant material to minimize reactions and contamination during the sampling process. The sample to be analysed must not pass through an active filter (molecular sieve, aluminium oxide) that could alter the gas composition. However, a particle filter may be necessary to ensure that the sample is not contaminated with particles.

A.1.1 On-site sampling connection

A direct connection is required for all on-site methods of analysis. The connecting line should be stainless steel tubing, 3 mm to 6 mm (nominal) outer diameter in size, as shorter as possible and no more than 2 m long. The ends of the line should have self-sealing fittings or valves to seal the line from ambient air contamination when not in use.

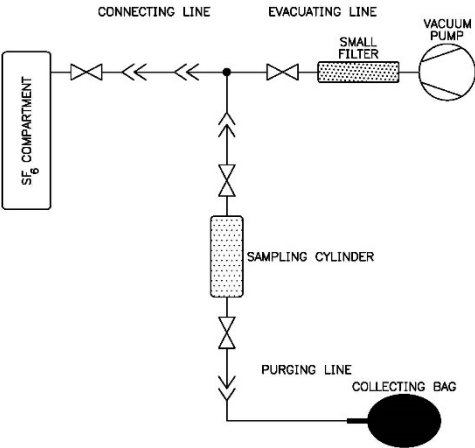
A.1.2 Sample cylinder for laboratory analysis

The cylinder volume shall be in the range of 150 ml to 500 ml (a larger volume, up to 1 000 ml, may be needed for the Fourier transformed infra-red analysis (FTIR)). All materials of the cylinder and valve in contact with the sample shall be stainless steel or chemically resistant materials, such as PTFE (polytetrafluoroethylene). Ideally the sampling material should be treated internal to create an inert surface. The cylinder and connections need to operate at high pressure (5 MPa).

The following procedure is recommended in order to remove any impurity (Figure A.1):

- heat the cylinder to 100 °C and evacuate for 1 h (primary vacuum);
- close the valve and cool the cylinder completely to room temperature before using;
- evacuate the gas;
- keep the cylinder evacuated.

451 **A.1.3 Sampling methods for laboratory analysis**
452 **A.1.3.1 One sampling cylinder method**

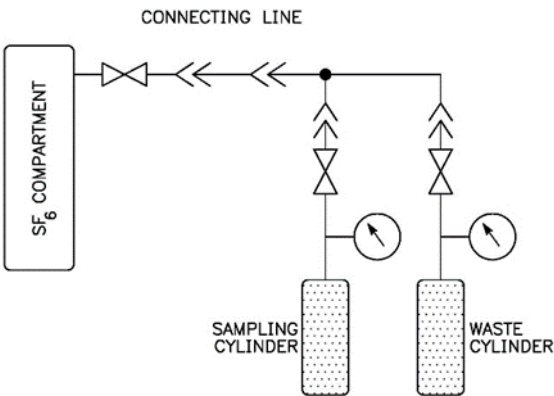


453
454 **Figure A.1. – One sampling cylinder method set-up.**

455 Figure A.1. shows example of set-up for the one sampling cylinder method. Connect the set-up to the
456 sampling port of the container as for a direct connection. Fill the cylinder once and wait 1 min to allow
457 the conditioning of the inside wall, purge, evacuate and fill it again The use of self-sealing fittings
458 facilitates all aspects of sample handling.

459 When purging, a collecting bag or similar equipment should be used in order to avoid any release of
460 SF₆ to the atmosphere. A few litres of SF₆ can then be passed through the cylinder to condition its
461 internal walls. After analysis, SF₆ samples should be reclaimed.

462 **A.1.3.2 Two sampling cylinders method**



463
464 **Figure A.2. – Two sampling cylinders method set-up**

465 Figure A.2. shows an example of set-up for the two sampling cylinders method. Two (2) cylinders are
466 connected in parallel with tubing and distribution block to the sampling port of the container as for a
467 direct connection. Each cylinder is equipped with a manometer in order to confirm that the cylinders
468 are under vacuum before the sampling. Open the valve of the waste cylinder in order to purge the
469 tubing and distribution block. Pressure in the sampling module and waste cylinder will increase
470 according to the pressure inside the sampled equipment. Close the valve of the waste cylinder when

471 pressure is stabilized. Open the valve of the sampling cylinder. Close the valve when the pressure is
472 stabilized.

473 No evacuation and collection bag or similar equipment is necessary on-site. The cylinders have to be
474 prepared according to the procedure describe in paragraph A.1.2. The tubing, manometer and valve
475 connected to the cylinders has to be properly designed and assembled with care in order to keep the
476 cylinders under vacuum for a long period until their utilization. After analysis, SF₆ samples and waste
477 materials should be reclaimed.

478 **A.2 On-site analysis**

479 Tubes that detect water vapour, HF, SO₂ and mineral oil are available from several manufacturers.
480 The use of such tubes is no longer recommended for doing analysis in SF₆. This tubes originally used
481 for measuring concentrations of several components in air to estimate harmful concentrations of
482 mostly toxic products in a defined volume of air. Other detection systems like SO₂ sensors or multi-
483 detector apparatus are more precise and made for the use in SF₆ systems.

484 **A.2.1 SF₆ concentration meter**

485 With this device, the SF₆ concentration is determined by a density meter which measures the speed of
486 sound in a gas sample. This type of meter is usually calibrated for measuring pure SF₆ percentage in
487 air with a precision of about ±1 %; however, this accuracy may be affected by the presence of other
488 gases (for example CF₄, decomposition product).

489 In all cases the manufacturer's instructions shall be followed.

490 **A.2.2 Hygrometers**

491 The following types of apparatus are available:

492 **Chilled mirror hygrometer**

493 A chilled mirror hygrometer determines the dew point by measuring the temperature at which
494 condensation occurs on a mirror. The temperature at which the mirror is just fogged is called the dew
495 point. Instructions in order to convert the dew point of water vapour to mass concentration.

496 **Electronic hygrometer**

497 An electronic hygrometer is a sensor which measures the partial pressure of water vapour in a gas
498 mixture, generally by a change in electric capacitance as the gas passes through a semi-permeable
499 layer. The dew point of the water is univocally set by its partial pressure.

500 **Electrochemical sensor hygrometer**

501 The electrochemical hygrometer determines the humidity by measuring the steady current needed to
502 electrolyze the water in the gas flow.

503 **Caution:** Measurements of SF₆ with HF content can damage the equipment. In all cases, the
504 manufacturer's instructions shall be followed.

505 **A.3 Laboratory analysis**

506 **A.3.1 Gas chromatography**

507 Chromatography is used to analyse SF₆ and its mixtures in order to identify and quantify the
508 composition of the gas and its contaminants.

509 Chromatography is a separation technique used for chemical substances (homogeneous liquid or
510 gaseous mixture) which is based on differences behavior between a current mobile phase (ex: gas)
511 and a stationary phase (an analytical column).

Each different compound requires a different time to traverse the column which is related to its chemical interaction with the latter. The result is a chromatogram, which is a time series of peaks corresponding to the components of the sample. For peak detection different detector types are available, for multi-purpose measuring and stability as needed for SF₆ analysis the basic TCD is recommended but other detectors (e.g. discharge ionisation detector (DID) or electron capture detector (ECD) etc.) are available and can be used too. (Figure A.3.).

A.3.1.1 Gas chromatography with detector systems with linear accuracy over minimum 3 decades, here as an example the thermal conductivity detector (TCD)

Air, CF₄, CO₂, SF₆ and SOF₂ are readily analysed. However, the quantitative measurement of SO₂F₂ and SO₂ may be difficult. The analysis of HF is not possible, as it is not detectable with this chromatographic technique.

Typical accuracy for the quantification of CF₄, N₂, and air are given in IEC 60376. The accuracy of the recommended method as described in the IEC 60376 Standard can be improved as follows:

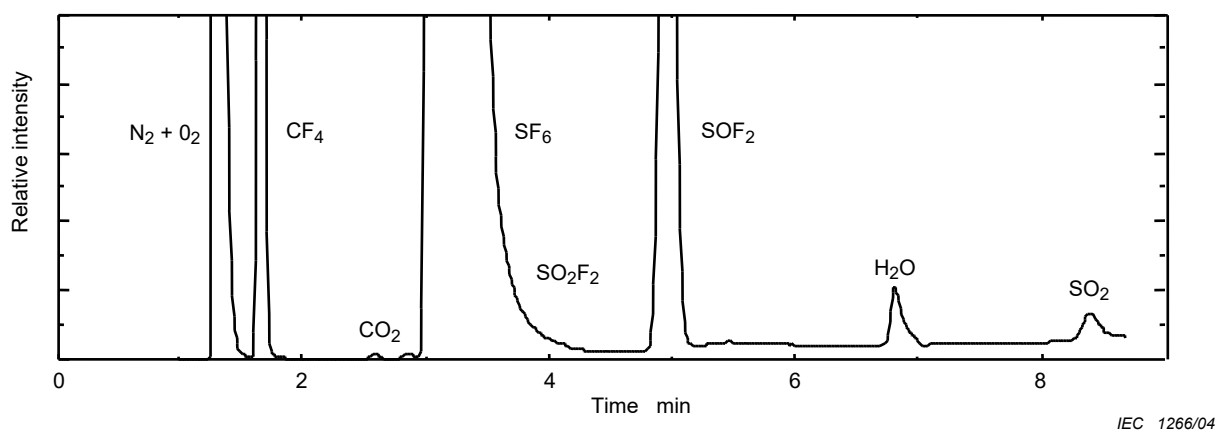
- An automatic injection unit, using a sampling loop (0,1 ml to 1 ml) and a multi-port switching valve, is preferred to the manual injection with a glass syringe. The chromatograph should be equipped with an inlet port to which the sampling bottle is attached;
- An improved separation of trace impurities is obtained with a temperature gradient (typically 60 °C to 180 °C), rather than by isothermal. In this case, a peak corresponding to the water vapour present in the gas sample can be seen in the chromatogram. However, moisture content can only be determined semi-quantitatively in this case;
- It is important to note that this chromatographic method will produce adequate quantitative measurements if, and only if, the chromatograph systems are well conditioned. This means that the instrumentation should be kept running and used routinely. Otherwise, prior to use, it will need a thorough preconditioning by injecting suitable gas mixtures containing several hundreds of ppm_v of SO₂ and/or SOF₂ and/or HF in air (or SF₆) followed by a series of calibration runs. Indeed, there is no other known means of detecting and quantifying low concentrations of decomposition products with the confidence specified in this standard.

A.3.1.1.1 Representative chromatograph set-up

If the following condition is used, reliable results are expected with the following parameters:

- Carrier gas:
helium or hydrogen (10-25 ml/min, a slightly better resolution can be obtained with H₂); flow rate should be optimized for the type of column used.
- Analytical column:
stainless steel tube packed by porous polymers type Porapak Q (80/100 mesh), 3-4 m × 3 mm;
wide-bore capillary (Poraplot type), 20-30 m, 0,53 mm;
new GS GasPro (J&W), 60 m, 0,32 mm.
- Oven programme:
initial temperature: 60-80 °C;
final temperature: 120-180 °C;
heating rate: 10-20 °C/min.

A chromatogram of SF₆ impurities and decomposition by-products is shown in Figure A.3.



IEC 1266/04

Figure A.3 – Example of a gas chromatogram in one print out for showing the different possible by-products after decomposition. (For determination of a compound concentration the substance peak needs to have ideal form for performing integration) (analysis performed with a Porapak Q column)

A.3.1.2 Example of Gas chromatography with mass spectrometer detector (MS)

This chromatographic technique allows to check the quality of pure SF_6 gas or of SF_6 mixture (N_2 or CF_4). It also allows identification and quantification of the following contaminants: CF_4 , SO_2 , N_2 , O_2 , COS , SO_2F_2 and SOF_2 .

A.3.1.2.1 Representative chromatograph set-up

If the following condition is used, reliable results are expected with the following parameters:

- Carrier gas: Helium (flow rate: 2.5 mL/min (40 cm/sec) flow rate should be optimized for the type of column used.
- Analytical column: Agilent Select Low Sulfur # CP8575 , 60 m, 0.32 mm
- Oven programme:
 - Initial temperature: 30°C
 - Final temperature: 105°C
 - Heating rate: 25°C/min
- Interface GC /MS temperature: 250°C
- Detector
 - Source temperature: 230°C
 - Quadrupole temperature : 150°C

A typical chromatogram of SF_6 impurities and decomposition by-products is shown in Figure A.4.

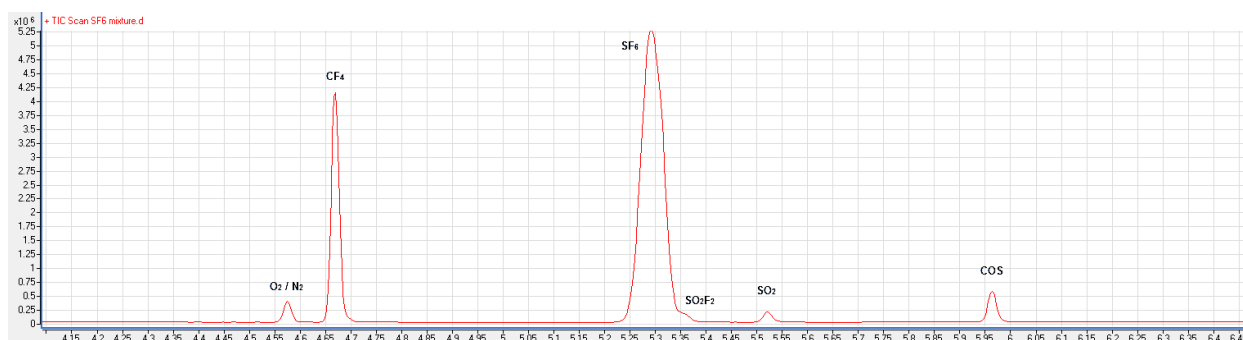


Figure A.4. – Typical GCMS chromatogram of decomposed SF₆

(Courtesy of Hydro-Québec)

A.3.1.2.2 Example for the determination of oil by Gas chromatography with flame ionization detector (FID)

The oil content in SF₆ can be measured with the following method, which is applicable to the analysis of C10 to C50 petroleum hydrocarbons. The oil may be obtained from a filter placed in a transfer line, or directly from a container, by washing with cyclohexane. With a high temperature flame ionization detector, the minimum detection limit is about 4 µg oil per 1 ml of cyclohexane. The minimum detection limit of oil in SF₆ will depend on the mass of SF₆ that passed through the filter or was in the container and the volume of cyclohexane used to extract the oil.

The following conditions represent one example that will achieve the desired results:

- Analytical column: Stainless steel megabore column, 5 m in length × 0,53 mm inner diameter, with a 0,15 µm film thickness.
- Carrier and make-up gas: Ultra-high purity helium at 15 ml/min each, for a total flow rate to the flame ionization detector of 30 ml/min.
- Flame ionization detector gas: Ultra-high purity hydrogen at 30 ml/min, and air at 300 ml/min.

The chromatogram is analysed by peak grouping to measure the area of the C10-C24 and the C24-C50 petroleum hydrocarbon ranges. These two ranges are calibrated with n-hexadecane (n-C16) and n-hexatriacontane (n-C36), respectively.

A.3.1.3 Ion chromatography

Ion chromatography is based on ion exchange columns. It can be used for the analysis of the following decomposition products: SO₂, SOF₂, SO₂F₂, SF₄, HF. Carbonate eluent has very low background yield and excellent sensitivity. This method has been validated for a variety of environmental matrices using the ION PAC AS4A. Further details can be found in US EPA catalogue [1].

A.3.2 Infrared spectroscopy

A.3.2.1 Principle

A beam of infrared light directed through a material sample to a light sensitive detector is attenuated. The ratio of the transmitted light to the incident as a function of wavelength forms the infrared absorption spectrum of the material sample.

The infrared absorption spectrum of a gas sample will exhibit peaks at the absorption wavelength of the gas. The size, shape and location of peaks in a spectrum can be used to identify the presence and quantity of a gas sample.

Most of the impurities of SF₆ can be identified by infrared absorption except oxygen and nitrogen; diatomic gases and monatomic gases (e.g. argon) do not have a significant infrared absorption. The presence of some impurities may be obscured by the spectrum of SF₆. The spectral region at frequencies less than 580 cm⁻¹ is clear of any SF₆ interference.

A.3.2.2 Fourier transform infrared spectrometer (FTIR)

FTIR spectrometers provide high resolution with a fast response. The whole infrared spectrum is measured many times and averaged to reduce the effect of noise. Spectra shall be obtained at a resolution sufficient to resolve the absorption bands to identify and quantify the sample components.

A.3.2.3 Absorption cell

The gas sample is introduced into an absorption cell within the spectrometer for analysis. The cell is preferably made of stainless steel to minimize the reaction with the impurity, HF, of the sample. The cell shall be fitted with KBr windows for infrared transmission down to, at least, 500 cm⁻¹. The optical path length of the cell shall be at least 10 cm. Path lengths of several metres are obtained by path-folding mirrors within the cell to minimize the total volume of the cell. The optimum path length depends on the signal-to-noise ratio of the spectrometer at the absorption frequencies of interest and on the minimum detection limit required.

A.3.2.4 Analysis

The method of spectral analysis is based on the linear relationship between absorption and concentration. The linearity domain depends upon the equipment used and the absorptivity.

A.3.2.5 Absorption frequencies of gases

The principal frequencies of absorption peaks for SF₆ and the contaminants of SF₆ are shown in Table A.2.

Table A.2 – Peak absorption of SF₆ and contaminants

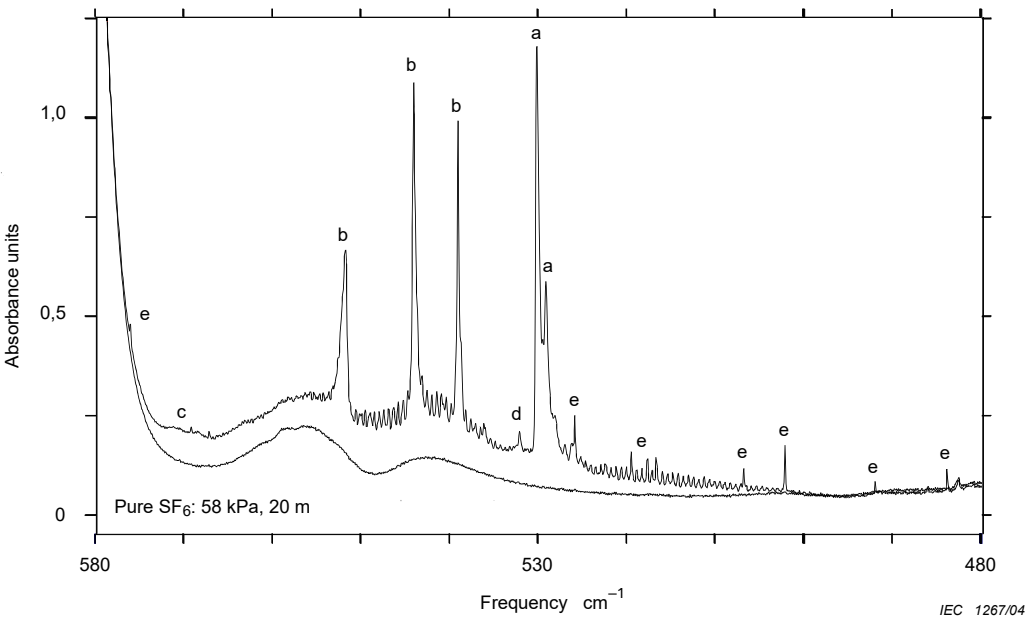
Gas	Maxima of absorption cm ⁻¹	Peak absorptivities x 10 ⁻⁶ (kPa x ml/l x m) ⁻¹
SO ₂	491, 494, 497, 500, 503, 506	~ 1,1 each
SOF ₂	530, 808	12, 46
SO ₂ F ₂	539, 544, 552	21, 25, 15
SOF ₄	570, 752	4, 8
SF ₄	532, 730	9, 80
CF ₄	1 283, 2 186	550; 2.2
HF	3 644, 3 693	Lines too narrow to quantify
Mineral oil	2 930	Not available
SF ₆	610, 860, 950, 1260, 1560 (broad bands)	Not given

639

The presence of water vapour can be observed but is hard to be quantified by FTIR. A complete method for mineral oil is found in IEC 60376.

The peak absorptivity's in Table A.2 are provided as a guide to select and/or calibrate the conditions of pressure and path length for the spectra of the samples. Note that the presence of SF₆ does modify the appearance of the spectra, broadening and in some cases shifting the peaks. Reference spectra of contaminants shall be obtained in conditions similar to the conditions for samples.

646 A typical spectrogram of SF₆ and impurities recorded in the above conditions is shown in Figure A.5.



647
648 a: SOF₂ c: SOF₄ e: H₂O
649 b: SO₂F₂ d: SF₄
650

Figure A.5 – IR spectrum of contaminated SF₆

Annex B

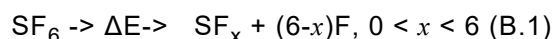
By-products of SF₆ and its mixtures

B.1 Decomposition of SF₆ and its mixtures

B.1.1 General

For SF₆ mixtures, the usual SF₆ by-products and specific mixture by-products, such as Nitrogen Oxide(s) and Nitrogen Fluoride(s) for SF₆/N₂ and Fluorocarbon(s) for SF₆/CF₄, will be produced. The quantities will depend on the mixture composition, contaminants and energy introduced. For typical SF₆ mixtures, the gas decomposition rates are expected not to exceed those for undiluted SF₆.

When arcing occurs in SF₆ due either to normal switching operations or fault clearances, or in the unlikely event of an internal arcing fault, different by-products are generated simultaneously in varying quantities [2], and [3]. When the SF₆ molecule is stressed by temperature, radiation or electrical discharge and separation of fluorine atoms occurs, a number of radicals, ions, or neutral molecules are produced, depending on the type of excitation and the energy input, according to:



When the input of energy ΔE ceases, most of the atoms recombine to form SF₆, whilst others combine with different substances in the system to form a variety of stable end products. Such substances include in particular oxygen and water and also materials used in the construction of the equipment.

These by-products are considered here in relation to the energy delivered to the SF₆.

B.1.2 The behaviour of SF₆ in an electric arc

Heavy current arcing occurs normally during circuit-breaker switching and fault clearance operations, and abnormally during an internal arc fault.

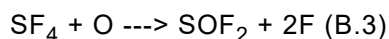
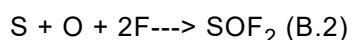
From temperatures of 500 °C, SF₆ begins to decompose into its constituent parts, with the degree of decomposition being directly proportional to the quantity of energy converted. In the dissociation process defined in reaction (B.1), the SF₆ molecules are broken down into sulphur and fluorine atoms at about 3 000 °C.

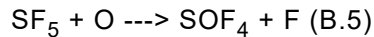
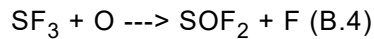
The large quantity of heat adsorbed during this process is dissipated away from the arc zone by radiation and convection. Below a temperature of about 1 000 °C, the atoms recombine or react with other substances, such as vaporised electrode metal, the vessel wall, plastics or impurities. Gaseous and solid by-products can arise, including metal fluorides and sulphur fluorides, of which the most important are CuF₂, AlF₃, WF₆, CF₄ and SF₄.

These products, generally known as primary by-products, are formed during or shortly after a equal discharge in the less-than-one-second range. Dust-like deposits which may appear on the surfaces of insulators during normal operation have no detrimental effect on their dielectric performance.

Some of the by-products are chemically stable; others are very unstable, particularly in presence of water.

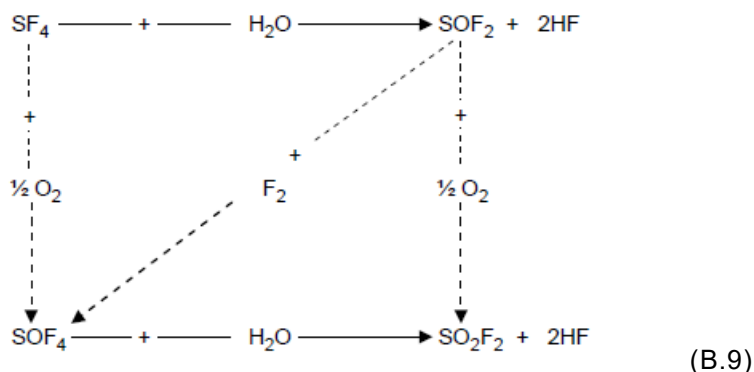
In presence of oxygen, by-products can arise as follows:





The oxygen involved in the reactions (B.2), (B.3), (B.4), and (B.5) may remain as result of the evacuation process or may be released by the electrode materials during arcing. SOF_2 is the major by-product.

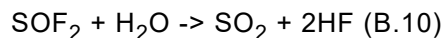
In the presence of moisture, the following reactions occur:



The dashed lines in reaction (B.9) indicate reactions that occur only to a limited degree during SF_6 decomposition in an arc. Whilst the reactions of (B.9) imply the formation of significant quantities of HF, this product has not been reported in large quantities following power arcing in electric power equipment. This is probably because by-product formation is mainly according to (B.2), (B.3) and (B.4). It is also possible that HF formed by the reactions of (B.9) further reacts with metal vapours to produce metal fluorides.

SF_4 is produced in significant quantities but hydrolyses rapidly (B.9) in the presence of moisture.

In the continued presence of moisture, further hydrolysis occurs, leading to:



Other reactions have been reported in the literature; additional by-products of arcing such as S_2F_{10} may be formed. However, the quantity of S_2F_{10} formed under arcing conditions is extremely low, because SF_5 radicals, produced at high temperatures, form S_2F_{10} only when cooled very rapidly, a condition not likely to apply in the arc [4].

B.1.3 SF_6 decomposition with low current discharges

Whilst operating voltage is applied to equipment containing SF_6 , the possibility of low current discharges such as corona, sparking and partial discharges cannot be ruled out. However, the concentrations of by-products resulting from such discharges are likely to be very low. When SF_6 is decomposed in spark discharges, the reactions shown in dashed lines in (B.9) will predominate [4].

717 **B.1.4 Catalytic decomposition of SF₆ (high-temperature behaviour)**

718 SF₆ can be heated to 500 °C in quartz vessels without decomposing. Up to temperatures of about
719 150 °C, all commonly used materials, such as metals, glass, ceramics, rubber and polyester resin are
720 fully resistant to SF₆. It is only at temperatures higher than 200 °C that some metals begin to have a
721 decomposing effect on the gas, but in the case of the metals and alloys normally used, this effect is
722 not observed to any marked degree until the temperature range of 400 °C to 600 °C is reached [3].

723 As the maximum operating temperatures inside electric power equipment in the absence of arcing are
724 far below these values, no SF₆ decomposition of this kind is to be expected during operation.

725 **B.2 Corrosion behaviour of SF₆ and its by-products**

726 SF₆ is a completely non-reactive gas. There is no possibility therefore that corrosion will be caused
727 directly by the SF₆ itself. However, the primary and secondary by-products, in presence of moisture,
728 may form corrosive electrolytes which may lead to damage some of the material used inside the
729 equipment.

730 The metals commonly used, such as aluminum, steel, copper and brass, are hardly attacked at all, but
731 materials such as glass, porcelain, insulation paper and the like are more vulnerable to damage,
732 depending upon the concentration of the corrosive substances concerned. Other insulating materials,
733 such as epoxy resin, polyester, polyethylene, polymethyleneoxide, PTFE, PVC, etc., are not
734 significantly affected. It is important that measures be taken in the design to take account of the
735 corrosive properties of the by-products. Corrosion can be prevented by the thorough exclusion of
736 moisture and by using suitable materials.

737 **B.3 Measures for the removal of by-products**

738 Moisture and by-products inside equipment in service can be reduced to acceptable levels by
739 adsorption. Materials such as alumina, molecular sieves or mixtures thereof are suitable for this
740 purpose. They adsorb the gaseous reactive by-products very effectively and practically irreversibly,
741 and at the same time ensure that the gas maintains a low dew point [5]. Disposal of by-products
742 removed from equipment is straightforward. The acid components (sulphuric acid and hydrogen
743 fluoride) are degraded by means of alkaline compounds. Most of the solid reaction products are not
744 soluble in water, or can be dissolved only with difficulty, but certain metal fluorides can react with
745 water to form hydrofluoric acid. It is necessary therefore to treat the solid reaction products with
746 calcium hydroxide (lime), for example, to neutralise the acid components. The resultant sludge can
747 then be disposed of according to local regulations.

748 **B.4 Physiological characteristics of by-products**

749 By-products can cause irritation of the skin, eyes and mucous membranes, such as in the respiratory
750 tract, and in high concentrations can cause pulmonary oedema, given sufficient time of exposure. SF₆
751 containing by-products have an unpleasant pungent smell that is itself associated with an irritant
752 effect. The olfactory thresholds, especially for SOF₂, SO₂ and HF, are of the same order as the TLVs.
753 Because of these characteristics, even small quantities of gaseous by-products may give rise to
754 unmistakable warning indications within a matter of seconds, before any risk of poisoning can arise
755 [2].

756

Annex C (informative)

Procedure for evaluating the potential effects on health from by-products of SF₆ and its mixtures

C.1 General

This annex suggests procedures for the evaluation of the risks to health due to by-products of SF₆ and its mixtures released into the local atmosphere.

Within the decomposition products generated in mixtures, SF₆ by-products generally are predominant in terms of quantity and toxicity. Safety procedures related to the presence of the usual SF₆ by-products must therefore apply in applications with SF₆ mixtures.

During normal service SF₆ and its mixtures remains inside the electric power equipment and the gaseous by-products formed are neutralised by molecular sieves as well as by natural recombination processes. SF₆ and its mixtures can become present in the atmosphere because of leakage or if a gas filled compartment fails to contain the gas, e.g. internal arc fault. It is necessary to differentiate clearly between leakage conditions and internal arc fault situations leading to a sudden release of SF₆, when evaluating health risk.

In case of leakage it is necessary to consider the effects of long-term exposure to the gaseous by-products of SF₆. The concentrations of these by-products in the air should remain low enough to present no threat to unprotected personnel during a normal working period of e.g. 8 h.

In case of a sudden release of SF₆ due to an internal arc fault, the emergency procedure required by the local regulation may imply a momentary exposure. By-product concentrations of higher levels than would be tolerable during e.g. 8 h can be tolerated if the exposure time is considerably reduced. Clearly in this case account should be taken of all possible sources of toxic emissions and this requires detailed knowledge of all of the by-products formed. In this respect a full treatment should consider contributions from metal vapour, burnt plastics, cable insulation, paint, etc., on an equal footing to those attributable to SF₆.

The following clauses give the procedures for calculating the risks associated with the presence of SF₆ by-products in the atmosphere due to leakage and to internal arc fault together with the principles adopted for performing those calculations.

C.2 Formation and health effects of SF₆ by-products

C.2.1 Formation of SF₆ by-products

During high power arcing in SF₆ the arc core reaches temperatures of the order of 10 000 K. At these temperatures, the molecules of the gas are completely broken down into their parent atoms, sulphur and fluorine, as reported in B.1.2. Any impurities present, such as air or moisture, are dissociated in a similar manner. The result of this is a localised region containing only single atoms of: S, F, H, N, O and diverse ions. The heating of the electrodes and the insulating parts by the arc adds vapours of Cu, W, C and Al to this atmosphere.

After arc extinction or in regions where cooling commences, these atoms start to bind together again and reform mainly SF₆. However, chemical reactions take place with the impurities present and in particular with moisture and oxygen, giving rise to the so-called arc by-products, as reported in Annex B. The quantities formed are directly related to the power injected into the gas volume.

In case of power arcing, the most frequently encountered gaseous by-products such as SOF_2 , SO_2 , HF and also CF_4 , SF_4 and SO_2F_2 [5], WF_6 [2], COF_2 [6] and solid by-products such as CuF_2 [2], AlF_3 [2]. In case of low energy electrical discharges, S_2F_{10} is also formed in extremely small quantities [7], [8] and [9].

804 **C.2.2 Effects of SF_6 by-products on health**

805 **C.2.2.1 General**

If SF_6 , which has been subjected to arcing or to low-energy discharges, is exhausted into the work place then the potential health risk will depend on the concentration of each by-product in the air, and hence on the volume of the room containing the equipment. Toxicity estimations should take into account the concentration of each by-product in relation to the permissible concentrations for the appropriate exposure time. Equation (C.1) should be adopted to calculate the total risk associated to the by-products.

$$812 \quad \text{Risk}_{\text{tot}} = \sum_i \frac{\text{Concentration (by - product)}_i}{\text{Threshold (by - product)}_i} \leq 1 \quad (\text{C.1})$$

813 **C.2.2.2 Health effects of arc decomposed SF_6**

814 For multi-component mixtures, toxicologists define three general cases:

- 815 • each component acts in a different manner, or on different target organs; the effects are hence not
816 cumulative and each component is separately treated;
- 817 • the components act in a similar manner on the same target organs; their effect is thus cumulative
818 and calculations will take this into account;
- 819 • one component largely outweighs the contribution of the others; the overall toxic effect should be
820 estimated by studying the concentration of this component alone.

821 A survey of the majority of the work carried out over the past decades on high power arc
822 decomposition of SF_6 [5], [10] concludes that the total health risk to personnel, due to arc
823 decomposed SF_6 , is predominantly dependent on the SOF_2 concentrations generated.

824 Hydrolysis of SOF_2 may occur in the presence of significant concentrations of moisture, producing
825 SO_2 and HF, as reported in B.1.2. For the time being, no OEL (Occupational Exposure Limit) has been
826 defined for SOF_2 . For this reason, it is assumed that hydrolysis always takes place giving HF and
827 SO_2 . The potential effect on health is then evaluated on those gaseous by-products (see Table C.1).

828 **C.2.2.3 Exposure duration and dilution in air**

829 The two key notions essential to any evaluation of risk to health due to toxic substances are:

- 830 • concentration within the surrounding volume;
- 831 • time duration of exposure, leading to the selection of the appropriate exposure limit.

832 When a leakage situation is calculated, the OEL (Occupational Exposure Limit) concentration, defined
833 as TWA (Time Weighted Average over an 8 h per day, 40 h per week exposure limit), should be used.

834 Under abnormal conditions e.g. internal arc fault, personnel immediately leave the room of the electric
835 power equipment and the exposure is hence momentary. Under those conditions, concentrations
836 defined as C (Ceiling exposure limit, values never to be exceeded), should be employed. When the C
837 value is not defined, the STEL (Short Term Exposure Limit) could be adopted. The STEL refers to an
838 average exposure of 15 minutes that should not be exceeded during the 8 hours working time.

The occupational exposure limits (OELs) defined by the American Conference of Governmental Industrial Hygienists (ACGIH) [11] for SO₂, HF, and S₂F₁₀ are given in Table C.1.

Table C.1 – OELs for SO₂, HF, and S₂F₁₀

OEL	SO ₂	HF	S ₂ F ₁₀
TWA (μl/l)	2	0,5	Not defined
STEL (μl/l)	5	Not defined	Not defined
C (μl/l)	Not defined	2	0,01

NOTE So far, no OEL has been defined for SOF₂

C.2.3 Quantitative estimation of gaseous by-products

C.2.3.1 General

The quantity of gaseous by-products formed during an electrical discharge is generally referred to the energy in joule dissipated in the gas. This is called as production rate, r .

Experimental results are generally quoted in mol/J but another unit widely adopted is l/kJ. The conversion is based on the fact that 1 mol of ideal gas occupies 24,37 l, at 20 °C and at 100 kPa, and is given in equation (C.2).

$$1 \text{ mol/J} = 24.37 \times 10^3 \text{ l/kJ} \quad (\text{H.2})$$

The estimation of the production rates of SOF₂ due to arcing and S₂F₁₀ due to low energy discharge, both sparking and partial discharges in the gas, is given in the next clauses C.2.3.2, C.2.3.3 and C.2.3.4., respectively.

C.2.3.2 Estimation of the SOF₂ production rates due to arcing

The SOF₂ production rate is experimentally determined and depends on the electrode material used and the type of discharge considered. Exothermic reactions, which occur with aluminium electrodes, seem to enhance the production rate.

The values used here have been averaged over the range of presently available data found in the literature [5] and [10].

Table C.2 – SOF₂ production rate

Electrode material	SOF ₂ production rate (r)	
	mol/J	l/kJ
Cu, Fe, WCu	150×10^{-9}	$3,7 \times 10^{-3}$
Al	600×10^{-9}	15×10^{-3}
NOTE Aluminium electrodes are assumed only for GIS bus-bar situations.		

C.2.3.3 Estimation of the S₂F₁₀ production rate due to sparking

Manoeuvring disconnectors produces sparking, which is considered as a low energy discharge. Average parameters are: 1 kV arc voltage drop, 0.25 A capacitive current and 1 s time duration, resulting in 0.25 kJ per each sparking event. However, individual sparks may reach up to 3 kA peak current for only a few tens of microseconds.

868 In the absence of published data, the production rate of $0,05 \times 10^{-9}$ mol/J (or $1,22 \times 10^{-6}$ l/kJ) is
869 chosen for S_2F_{10} due to sparking, which reflects a situation at the lower end of the spark discharge
870 range but more than 2 000 times greater than for arcs. This has been confirmed as being realistic by
871 measurements on a real disconnector under highly accelerated operation conditions [8].

872 **C.2.3.4 Estimation of the S_2F_{10} production rate due to partial discharges**

873 Single components of high-voltage switchgear and controlgear or sub-assemblies in which they are
874 contained shall not exceed the maximum permissible partial discharge level of $q = 5$ pC (see sub
875 clause 6.2.9.102 of IEC 62271-203).

876 At rated voltage U_r , the energy dissipated during each single partial discharge event is therefore:

$$877 \quad E = q \times \frac{U_r}{\sqrt{3}} \quad (C.3)$$

878 which always falls into the micro to nano-joule energy range.

879 The S_2F_{10} production rate due to partial discharges at power frequency is not available in the
880 literature. A value of $0,2 \times 10^{-9}$ mol/J (or $4,88 \times 10^{-6}$ l/kJ) is obtained by extrapolating the experimental
881 results for sparking in the very low energy region.

882 **C.2.4 Procedures for health risk evaluation**

883 The procedures for the evaluation of the potential effects on health of gaseous by-products are based
884 on the following assumptions:

- 885 • The equipment room is completely closed and ventilation is inoperative during the period of
886 interest;
- 887 • The adsorbers fitted into the electric power equipment do not reduce the amount of gaseous
888 by-products during the period of interest;
- 889 • The gas emitted uniformly mixes with the air in the room containing the electric power equipment
890 in a short time with respect to the working day or exposure duration.

891 Figure C.1 describes the typical procedure to follow for the evaluation of the potential effects on
892 health of the by-products formed during arcing (either power interruption or capacitive switching) in
893 SF_6 . It is applicable to both leakage and internal arc fault cases.

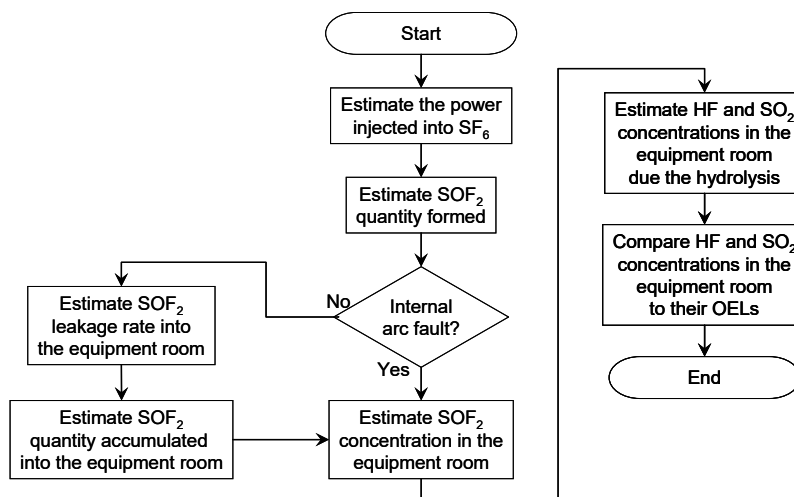


Figure C.1 – Procedure for the evaluation of the potential effects on health due to arcing

The power injected during one single event into the electric power equipment should be evaluated. Depending on the circumstances, e.g. power arcing, capacitive arcing, corona discharge, this could be thought as arc voltage drop times arc current times arc duration or as voltage drop times charge transferred. The cumulated power is then evaluated multiplying the power of the single event by the number of events taking place during the reference period.

The quantity of SOF_2 formed within the electric power equipment is the power itself times the SOF_2 production rate. The SOF_2 production rate is a non-linear function of the power, the quantity of SF_6 , the time duration and the electrode material. Numerical values are determined by experiments (see C.2.3.2.).

In case of internal arc fault, the bursting disk bursts and the whole SOF_2 quantity is suddenly released into the switchgear room, which is the worst case scenario.

In case of leakage, only a small portion of the SOF_2 quantity formed into the electric power equipment transfers to the switchgear room and accumulates over a certain period of time. Under this condition, if $V_{\text{SOF}_2, \text{equipment}}$ is the SOF_2 quantity formed inside the electric power equipment in litres and $F_{p, \text{rel}}$ is the leakage rate of the electric power equipment in % p.a., the SOF_2 quantity accumulated into the switchgear room during one day is:

$$V_{\text{SOF}_2, \text{room}} = \frac{V_{\text{SOF}_2, \text{equipment}} * F_{p, \text{rel}}}{100 * 365} \quad (\text{C.4})$$

The SOF_2 concentration in the equipment room is the SOF_2 quantity into the equipment room divided by the volume of the switchgear room. For outdoor installations, the same evaluation should be performed considering the volume of a virtual equipment room.

Due to hydrolysis in the equipment room, the HF concentration in $\mu\text{l/l}$ is twice the SOF_2 concentration in $\mu\text{l/l}$ and the SO_2 concentration in $\mu\text{l/l}$ is equal to the SOF_2 concentration in $\mu\text{l/l}$.

The potential effect on health of the by-products is then evaluated comparing the HF and the SO_2 concentrations in the equipment room to the OELs (Occupational Exposure Limits). The total risk R_{tot} should be defined as:

$$R_{\text{tot}} = \frac{\text{Concentration}(\text{SO}_2)}{\text{TLV}(\text{SO}_2)} + \frac{\text{Concentration}(\text{HF})}{\text{TLV}(\text{HF})} \leq 1 \quad (\text{C.5})$$

A similar procedure can be adopted for the evaluation of the potential effects on health of S_2F_{10} due to low energy electrical discharges. In this case, no hydrolysis takes place. The procedure is described in Figure C.2.

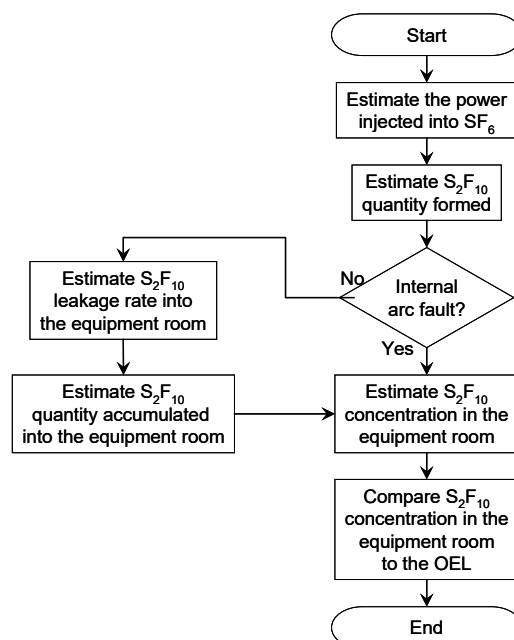


Figure C.2 – Procedure for the evaluation of the potential effects on health due to low energy discharges

C.3 Conclusions

Calculations, based on the state-of-the-art, show that, for leakage situations, there is no risk to health due to exposure to by-products. The by-products, formed by arcing and by low-energy discharges, released due to leakage from SF_6 -filled electric power equipment, reach negligible concentrations in the workplace atmosphere. Therefore, there is no cause for concern and no need for precautions other than the normal ventilation practices for low-lying areas. This is still valid even in the case of abnormal leakage situations (a leakage rate for example 2 orders of magnitude higher than the normal rate).

However, in the unlikely event of an internal fault leading to a release of SF_6 , significant concentrations of by-products can occur in an equipment room. In any situation of this sort, basic safety procedures include evacuation rules are designed to ensure that personnel are exposed to exhausted materials for a time as short as possible. Furthermore, forced ventilation and/or venting ensure the concentration levels can be reduced within minutes.

For outdoor installations, the volume of air into which the arc decomposed SF_6 escapes is large if not infinite. Prevailing winds and the high exhaust velocity also speed up dispersion.

It is thus concluded that, as long as basic safety procedures are followed, the risk specifically associated with the use of SF_6 in electric power equipment is minimised.

Annex D
(informative) Cryogenic reclaim of SF₆

D.1 General

This annex describes as example a method used in Australia during the past 15 years to reclaim SF₆ based on a cryogenic technique [12], [13] and [14].

D.2 Applications

The principal use of the cryogenic process is to remove contaminants from used SF₆ stored in cylinders returned to the main process plant from a site where it has been removed from electric power equipment.

The process can be used to:

- Aggregate the remainder of SF₆ from containers that have been used to fill electric power equipment;
- Recover SF₆ from electric power equipment to a pressure of below 100 Pa without the need for a recovery compressor;
- Restore used SF₆, even heavily arced or contaminated by air to the standard of technical grade SF₆;
- Separate SF₆ from a mixture of SF₆ and nitrogen (N₂). All the SF₆ can be recovered and essentially none is released to the atmosphere;
- Obtain CF₄ from an SF₆ / CF₄ mixture.

D.3 Physical Background

The process relies on the variation of the saturated vapour pressure of SF₆ and its typical contaminants with temperature (see Figure D.1).

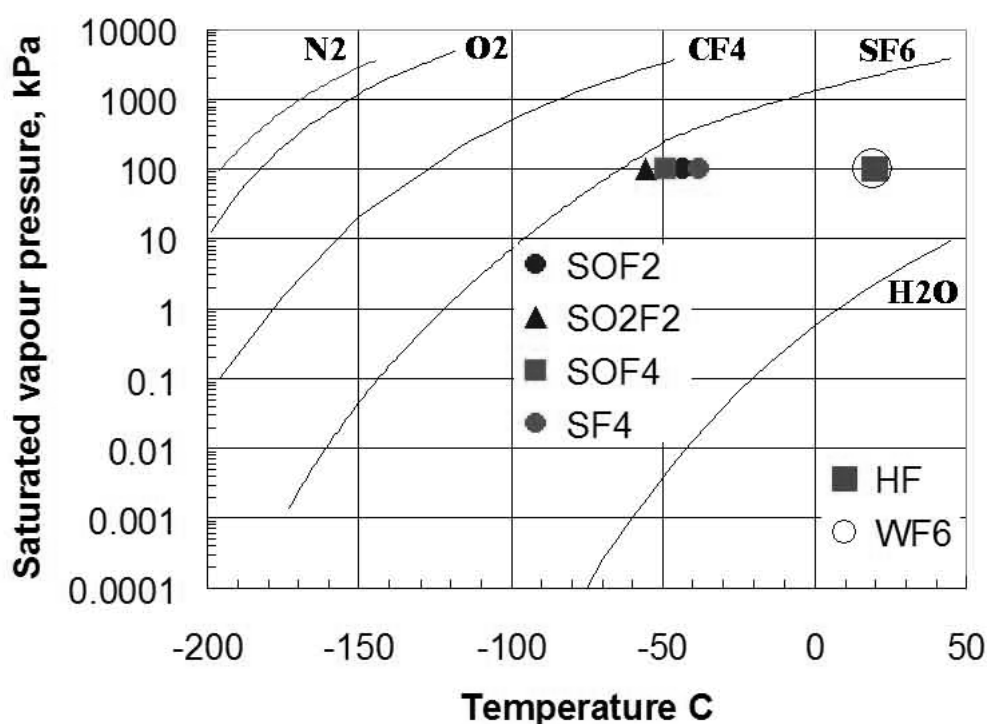


Figure D.1 – Saturated vapour pressure of various gases as a function of temperature

D.4 Cryogenic processes

SF₆ is recovered by means of evacuating and cooling a cylinder called capture cylinder. The pressure differential between the cylinder and the gas-filled compartment or the container containing the SF₆ to be processed causes the SF₆ to be transferred into the cylinder where it will liquefy or solidify due to the low temperatures achieved by liquid nitrogen thereby maintaining a pressure differential until recovery is completed.

The collection process is scalable. A number of cylinders can be connected in parallel to increase collection capacity. Smaller cylinders can be used if the masses of SF₆ involved are small. The speed of mass transfer is mainly limited by the diameter of the connecting pipes and with appropriate sized piping the practical limit given by the speed of sound can be attained.

This process can also be used to separate air from SF₆ as – provided SF₆ is cooled sufficiently – SF₆ will freeze inside the capture cylinder allowing the air to be released or evacuated by a conventional vacuum pump.

The process comprises a second step, where a second cylinder called collection cylinder is cooled by liquid nitrogen and connected to the capture cylinder which is allowed to warm up. The pressure differential between the cylinders causes SF₆ to be transferred into the collection cylinder while the contaminants remain frozen in the capture cylinder. During the SF₆ transfer, the capture cylinder warms up from approx. -150 °C to approx. -45 °C and the pressure in the capture cylinder can be obtained by closing the valve controlling the SF₆ transfer and reading the pressure gauge on top of the capture cylinder. This pressure (see Figure D.1) is representative of the highest temperature portion of SF₆ in the capture cylinder, even though temperature gradients may be present.

The upper temperature limit in the capture cylinder (i.e. approx. -45 °C) is set to limit the concentration (dew point) of moisture transferred to the collection cylinder. This upper temperature

limit must also be low enough to also prevent transfer of reactive gaseous by-products that may be present in the capture cylinder.

The capture cylinder may be partly immersed in a refrigerated bath to allow for a better temperature control.

D.5 Description of a cryogenic reclaimer [13]

A cryogenic reclaimer comprises the following components:

- Gas cylinders (see IEC 62271-4) in addition suitable for cryogenic temperatures);
- Insulated dewars meeting local safety requirements and capable of withstanding -220 °C;
- Vacuum pump (see IEC 62271-4);
- Optional filter, used for reclaim heavily arced gas (see IEC 62271-4);
- Sampling point (see IEC 62271-4)
- Valves, fittings, pressure gauges and gas piping (see IEC 62271-4), in addition suitable for cryogenic temperatures where relevant).
- Liquid nitrogen purchased as needed

The components can be put together in different configurations according to the task at hand.

Figure D.2 is the flow chart of a cryogenic reclaimer suitable for SF₆ recovery on site;

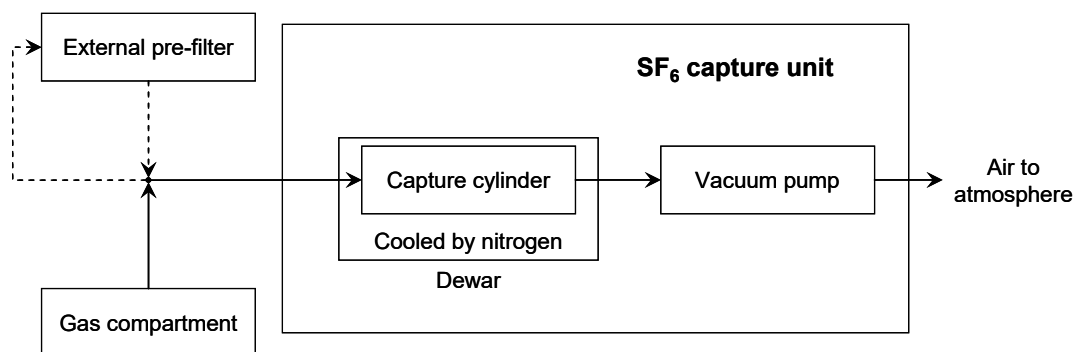


Figure D.2 – Typical cryogenic reclaimer for SF₆ recovery on site

The procedure for operation of the manual process is as follows:

- The whole reclaimer is evacuated with the vacuum pump.
- The capture cylinder is chilled with liquid nitrogen and the vacuum in the cylinder itself is maintained by the freezing of SF₆ which is continuously drawn in.
- The impure gas entering the process can be directed through a filter if it contains a significant amount of solid contaminants as in the case of heavily arced gas.
- Once the gas capture capacity has been reached, the upstream valves are closed and the downstream valves opened.
- The vacuum pump can be used to evacuate and draw off all the gaseous components above the frozen SF₆ i.e. oxygen and nitrogen.
- The downstream valves are closed and the cylinder can be removed, warmed up and transported.

Figure D.3 is the flow chart of a cryogenic reclaimer suitable for removing contaminants.

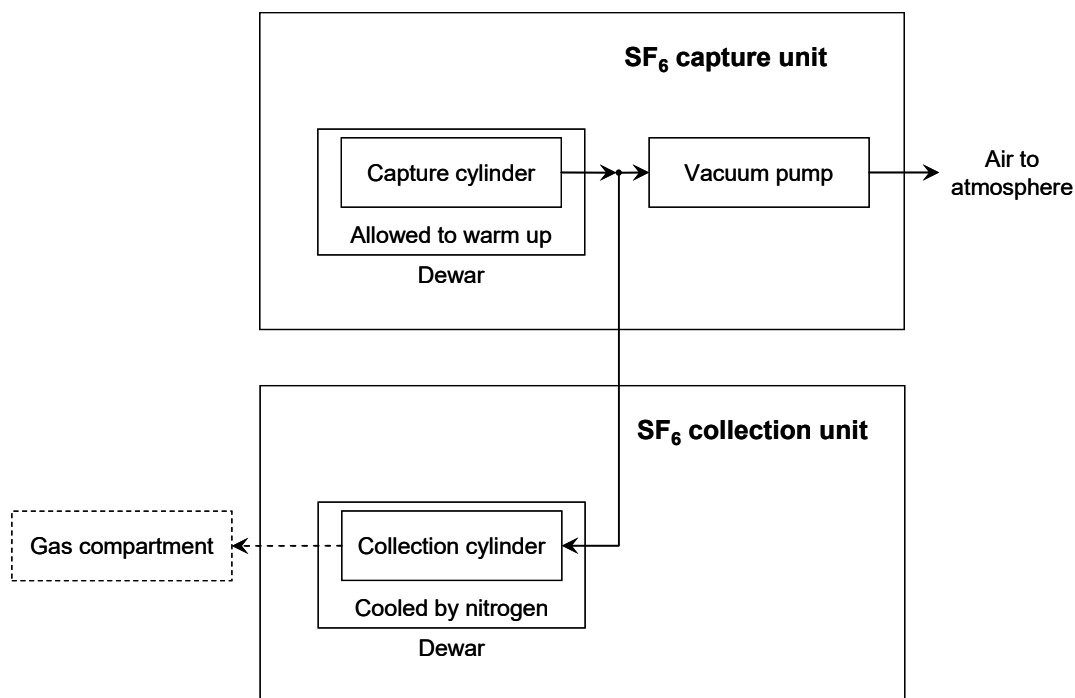


Figure D.3 – Typical cryogenic reclaimer for removing contaminants

The procedure for operation of the manual process is as follows:

- The capture cylinder and the collection cylinder are chilled with liquid nitrogen.
- The vacuum pump can be used to evacuate and draw off all the gaseous components above the frozen SF₆ i.e. oxygen and nitrogen.
- The capture cylinder is then allowed to warm up to the selected upper temperature limit (e.g. -45 °C)
- Pure SF₆ is transferred into the collection cylinder as a consequence of the temperature difference.

The capture cylinder is likely to end up containing solid contaminants from heavily arced SF₆ and requires cleaning after use.

Annex E (informative)

Reclaiming recommendations

E.1 General

On-site reclaiming procedures are based on the absorption of impurities. Operational contamination should already be absorbed with the user's filter unit. Such filters are already an integral part of the company's SF₆ maintenance devices, or are available as separate filter units.

E.2 Filtering requirements

The filters should meet the following requirements:

- they must reliably remove the mentioned contaminants;
- they filters should be of cartridge type for safe and easy disposal;
- input and output should be equipped with the same self-sealing couplings;
- changing filters should not require disassembly of any fittings, tubing, or any other connection to eliminate the possibility of leakage;
- changing the filter is recommended for each purification operation.

E.3 Transport of used SF₆ in gas cylinders and containers by road

See IEC 62271-4.

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