

## Potential source apportionment and meteorological conditions involved in airborne I detections in January/February 2017 in Europe

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# Potential source apportionment and meteorological conditions involved in airborne <sup>131</sup>I detections in January/February 2017 in Europe

## Authors

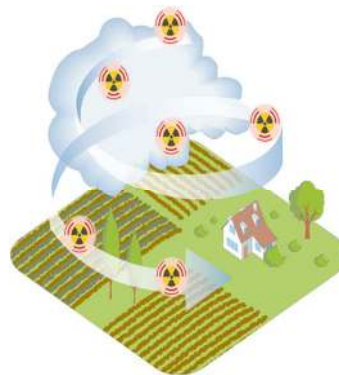
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**ABSTRACT:** Traces of particulate radioactive iodine ( $^{131}\text{I}$ ) were detected in the European atmosphere in January/February 2017. Concentrations of this nuclear fission product were very low, ranging 0.1 to 10  $\mu\text{Bq m}^{-3}$  except at one location in western Russia where they reached up to several  $\text{mBq m}^{-3}$ . Detections have been reported continuously over an 8-week period by about thirty monitoring stations. We examine possible emission source apportionments and rank them considering their expected contribution in terms of orders of magnitude from typical routine releases: radiopharmaceutical production units > sewage sludge incinerators > nuclear power plants > spontaneous fission of uranium in soil. Inverse modeling simulations indicate that the widespread detections of  $^{131}\text{I}$  resulted from the combination of multiple source releases. Among them, those from radiopharmaceutical production units remain the most likely. One of them is located in Western Russia and its estimated source term complies with authorized limits. Other existing sources related to  $^{131}\text{I}$  use (medical purposes or sewage sludge incineration) can explain detections on a rather local scale. As an enhancing factor, the prevailing wintertime meteorological situations marked by strong temperature inversions led to poor dispersion conditions that resulted in higher concentrations exceeding usual detection limits in use within the informal Ring of Five (Ro5) monitoring network.



47

## 48 INTRODUCTION

49 Radioactive  $^{131}\text{I}$  is an important fission product that is produced with high yield by nuclear fission of  
50  $^{235}\text{U}$  (2.9%) or  $^{239}\text{Pu}$  (3.8%). The half-life of  $^{131}\text{I}$  is short enough (8.02 days) to cause high specific  
51 activities, but long enough to allow long-distance (> 3,500 km) atmospheric dispersion when released  
52 into the atmosphere. Elementary iodine ( $\text{I}_2$ ) is a solid under normal conditions of temperature and  
53 pressure and slowly sublimates even at room temperature leading to the formation of gaseous  $\text{I}_2$ . Due  
54 to its volatility and reactivity with many compounds (aerosol, ozone, volatile organic compounds, etc.)  
55 iodine in the ambient atmosphere is distributed in variable proportions between gaseous and  
56 particulate species. The ratio between the various released chemical forms may change along the route  
57 of air masses and from daytime to nighttime as a result of photo-dissociation. Even if the release  
58 occurs only in gaseous form, iodine in air can transform to less volatile aerosol-bound species or  
59 attach to existing particles.<sup>1, 2</sup> All those properties make  $^{131}\text{I}$  one of the most important radionuclides  
60 for the monitoring of airborne radioactive releases.

61 The topic of this study is the early 2017 detection of airborne  $^{131}\text{I}$  at trace levels in Europe that made  
62 headlines in media and that spread rapidly on social networks<sup>3</sup> after the publication of a single map

63 gathering all detections until the first week of February.<sup>4</sup> Here, we report detailed time trend and  
64 geographical evolution of airborne <sup>131</sup>I concentrations and evaluate several potential sources based on  
65 authorized releases or real releases when available. Several significant emissions of <sup>131</sup>I from  
66 radiopharmaceutical production units have already been reported in the literature during the last  
67 decade leading to a similar widespread detection of <sup>131</sup>I at trace levels on the European scale.<sup>5-10</sup> Other  
68 local sources such as sewage plant incinerators, nuclear power plants, as well as possible ubiquitous  
69 and natural production of <sup>131</sup>I from the spontaneous fission of uranium in soils, have also been  
70 examined. Finally, we investigated the meteorological conditions as a “detection enhancing factor”  
71 using airborne naturally occurring radionuclides (namely <sup>7</sup>Be, <sup>22</sup>Na, <sup>210</sup>Pb, and <sup>40</sup>K), an anthropogenic  
72 radionuclide (<sup>137</sup>Cs), and particulate matter with an aerodynamic diameter <10 μm (PM<sub>10</sub>) values as  
73 indicators of atmospheric dispersion by comparing their values during the <sup>131</sup>I detection weeks with  
74 their usual background levels.

75 Most reported data come from atmospheric monitoring stations gathered within the “Ring of Five”  
76 (Ro5) network which is an informal collaboration platform on laboratory level.<sup>11</sup> According to our  
77 experience based on internal Ro5 reports, trace detections of unexpected radionuclides occur on a  
78 more frequent basis than what one might anticipate. On average, there is a handful of unexpected or  
79 unusual radionuclides detected at trace-level each year on a European-wide scale and a dozen on a  
80 local scale. When only one station detects unexpected radionuclides or enhanced activity  
81 concentrations at trace levels, it is a general practice within the Ro5, not to spread the information  
82 outside the country because it may be representative of only a local activity with an isolated impact.  
83 However, in some cases and even for a single detection, it may be worth sharing this information with  
84 the Ro5 community, especially when there is no apparent reason for an anticipated abnormality at this  
85 location, for example when faraway from any nuclear or industrial facility. This was the case in  
86 January 2017, when <sup>131</sup>I was first detected in the Norway’s northeasternmost corner. Most detections  
87 reported by Ro5 members proved to be very close to the minimum detectable activity (MDA), in the  
88 μBq m<sup>-3</sup> range. Given the great public concern regarding releases of anthropogenic radionuclides, the  
89 policy of non-publicizing very localized trace concentrations in the air is normally carried out as long  
90 as levels are of no health concern. However, on few occasions, radionuclide detections simultaneously

91 occur at several monitoring stations, and even if at innocuous levels, these detections should become  
92 public.<sup>1, 12</sup> In the late fall of 2011, the attention of the Ro5 members was triggered by the detection of  
93 <sup>131</sup>I in the European atmosphere as well as in animal thyroid biomonitors after its release from a  
94 radiopharmaceutical facility in Hungary.<sup>13, 14</sup> Other multiple detection events of particulate <sup>131</sup>I at trace  
95 levels were also reported within the Ro5, especially in Scandinavia, in January/February 2012, March  
96 2015, October 2016, and recently in January until beginning of March 2018.<sup>15</sup>

97

## 98 **Materials and Methods**

99 The monitoring stations gathered within the Ro5 network operate according to national regulations  
100 and are usually equipped with high-volume air samplers having an air throughput rate of several  
101 hundred m<sup>3</sup> h<sup>-1</sup> and up to 1,500 m<sup>3</sup> h<sup>-1</sup>. Aerosols are usually sampled on a weekly basis (traditionally  
102 from Monday to Monday). Iodine-131 counting is performed by gamma spectrometry, using low-level  
103 High-Purity Germanium (HPGe) detectors and identifying <sup>131</sup>I by its 364 keV line in the gamma  
104 spectrum. Recent spectrometry improvements (new shields, Compton-suppression systems, higher  
105 detection efficiency, etc.) made it possible to lower the MDA for particulate <sup>131</sup>I down to 0.5 μBq m<sup>-3</sup>,  
106 and even 0.1 μBq m<sup>-3</sup> when using very high-volume samplers. This concentration corresponds to the  
107 staggering amount of 1 atom of <sup>131</sup>I per 10 m<sup>3</sup> of air. Such MDA currently in use within the Ro5  
108 network, allow quantifying concentration levels, which had yet remained undetectable only five years  
109 ago.

110

## 111 **Results and Discussion**

### 112 *Sequence of airborne <sup>131</sup>I detections in Europe.*

113 The first <sup>131</sup>I detections were reported for Week 2 (January 9-16, 2017) by the Norwegian Radiation  
114 Protection Authority at the northernmost European aerosol sampling location in Norway (Svanvik); by  
115 the Finnish Radiation and Nuclear Safety Authority in the northern part of Finland (Rovaniemi) and  
116 by the Central Laboratory for Radiological Protection on the same week in Poland (Warsaw). Over the  
117 entire detection episode (Week 2 to Week 8), 38 particulate <sup>131</sup>I detections above MDA were reported  
118 from 26 locations in Europe (Fig. 1). One unique detection of gaseous <sup>131</sup>I (10.6 ± 3.4 μBq m<sup>-3</sup>) was

119 reported on Week 4 by the laboratory of the “Centro Regionale Radioprotezione” (Arpa Lombardia) in  
120 Milan (Italy) thanks to a high-volume gaseous sampler operating at about  $80 \text{ m}^3 \text{ h}^{-1}$ . Indeed, detection  
121 limits for gaseous  $^{131}\text{I}$  are usually at least 100 times higher than for the aerosol fraction, mainly  
122 because most commercially available charcoal cartridges do not allow for such a high throughput rates  
123 of air (typically between 3 and  $10 \text{ m}^3 \text{ h}^{-1}$ ) like for the aerosol samplers. Unfortunately, the aerosol-  
124 bound  $^{131}\text{I}$  fraction at that location was not detected above a detection limit of  $5 \text{ }\mu\text{Bq m}^{-3}$ , thus  
125 preventing the determination of the gas-to-particle ratio. However, inasmuch as a comparison can be  
126 made with the situations in Europe after the Chernobyl and Fukushima accidents, we assume a similar  
127 gas-to-aerosol ratio between 3 and 5.<sup>1</sup> For the present event, this assumption leads to a particulate  
128 iodine concentration of about 2 to  $3 \text{ }\mu\text{Bq m}^{-3}$ , which would be below the detection limits of the  
129 measurement laboratory in question.

130 All in all, weekly  $^{131}\text{I}$  aerosol concentrations remained in the  $\mu\text{Bq m}^{-3}$  range, except in Hungary  
131 (Budapest) where the level peaked momentarily at  $9.8 \text{ }\mu\text{Bq m}^{-3}$ , and in western Russia where daily  
132 particulate  $^{131}\text{I}$  levels rose up to several tens  $\mu\text{Bq m}^{-3}$  in Dubna and up to 3 orders of magnitude higher  
133 ( $\text{mBq m}^{-3}$  range) in Obninsk (at about 100 km southwest of the Moscow city center), thus indicating a  
134 high concentration gradient in this region.<sup>16</sup> The Dubna station belongs to the International Monitoring  
135 System (IMS) supporting the Comprehensive nuclear Test Ban Treaty Organization (CTBTO).<sup>17, 18</sup>  
136 Due to restrictions implemented by confidentiality agreements, we cannot provide the raw data for the  
137 IMS station. The Obninsk station is managed by the “Typhoon” Research and Production Association  
138 (Typhoon RPA) in connection with the Russian Federal Service for Hydrometeorology and  
139 Environmental Monitoring (Roshydromet).

140 Apart from these two Russian locations, no significant concentration gradient could be observed over  
141 the rest of Europe, which would have helped pointing out the main source. Even when considering that  
142 the gaseous fraction was omitted and that its contribution would have led up to a five-time higher  
143 concentration,  $^{131}\text{I}$  total concentrations would have been so low that they raised no health concern.  
144 They corresponded to 1/1,000 or even 1/10,000 of those observed in 2011 when the diluted

145 Fukushima-labeled air masses were distributed over Europe, which themselves were of no health  
146 concern.<sup>1</sup>

147

#### 148 *Potential sources of airborne iodine*

149 The main feature of this event is the wide spreading of <sup>131</sup>I on the European scale. Even if the  
150 concentrations were very low, they were all in the same range as if they resulted from the dispersion  
151 from a more or less remote source emission or from a combination of several emissions from different  
152 locations. Even if it may have been confined within annual authorization limits, it can also be  
153 considered that the plume rooted in a significant release of radionuclides. For instance, the European-  
154 wide <sup>131</sup>I detection event in fall 2011 resulted from a release of 342 GBq from the Institute of Isotopes  
155 near Budapest (Hungary) and led to somewhat similar, even if slightly higher, <sup>131</sup>I concentrations (0.5  
156 - 68 μBq m<sup>-3</sup> outside Hungary).<sup>8, 14</sup> Source apportionment and ranking in terms of contribution to the  
157 observed values are detailed hereafter.

158

159 Figure 1

160

#### 161 ***Hypothesis for the origin of airborne <sup>131</sup>I: Research reactors or radiopharmaceutical facilities***

162 In nuclear medicine, most diagnostic procedures rely on <sup>99m</sup>Tc whose precursor, <sup>99</sup>Mo, is produced  
163 using low or highly enriched uranium targets.<sup>19</sup> Iodine-131 can be considered as a by-product of the  
164 <sup>99</sup>Mo production.<sup>20</sup> Table 1 summarizes the major <sup>99</sup>Mo, and thus <sup>131</sup>I, suppliers in Europe and Western  
165 Russia. All these facilities have yearly iodine release authorizations ranging between one GBq and up  
166 to several hundred GBq. To cope with an increasing demand for radiopharmaceuticals (see following  
167 hypothesis), there is an increasing effort since several years to increase radiopharmaceutical  
168 production capabilities. For instance in 2015, the <sup>131</sup>I release (850 GBq) from the L. Ya. Karpov  
169 Institute of Physical Chemistry (Obninsk branch) and hereafter named “Karpov Institute”, reached and  
170 exceeded its yearly authorization limit (780 GBq) to fulfil the needs and this limit was about 5 times  
171 higher than that in 2014.<sup>21</sup> Many other facilities are producing medical radionuclides in Russia<sup>22</sup> but



172 regarding the frequent detections in Obninsk, they are closely related ( $r = 0.9$ ) to the release of  
 173 radionuclides from the Karpov Institute in Obninsk.<sup>16,23</sup>

174

175 *Table 1. Main <sup>99</sup>Mo/<sup>131</sup>I producers in Europe and Western Russia.*

Country (Town)	Company or Institute	Maximum yearly authorized <sup>131</sup> I release to the atmosphere (GBq)	Reference
Poland (Otwock- Świerk)	Nuclear Research Radioisotope Centre Polatom	0.1 in 2016	24
Netherlands (Petten)	Mallinckrodt Medical B.V	0.3 in 2017	25
France (Saclay)	UPRA (Cis-BIO international)	0.6 in 2013	26
Belgium (Fleurus)	Institut national des RadioEléments (IRE)	41.8 in 2011	10
Russia (Obninsk)	L. Ya. Karpov Institute of Physical Chemistry (NIFKhI)	780 in 2015	21
Hungary (Budapest)	Institute of Isotopes Ltd (IoI)	1600 in 2011	27

176

177 Apart from the main radiopharmaceuticals producers, there is an increasing number of secondary  
 178 industries that transform concentrated <sup>131</sup>I solutions into diluted ones (injectable or for oral  
 179 administration) as well as into <sup>131</sup>I capsules, which may also represent a potential source of <sup>131</sup>I  
 180 emission. Despite significant amounts of <sup>131</sup>I released within authorized limits, airborne detections of  
 181 <sup>131</sup>I coming from major radiopharmaceutical production units remain rare and unusual. Only close  
 182 monitoring stations may report regular detections at trace concentrations that exhibit negligible doses  
 183 upon inhalation for the local population.<sup>23</sup> Several significant <sup>131</sup>I emissions from radiopharmaceutical  
 184 production units have already been reported in the literature as a result of incidents during the last  
 185 decade, leading to a similar widespread detection of <sup>131</sup>I at trace levels (tenths to tens of  $\mu\text{Bq m}^{-3}$ ) on  
 186 the European scale.<sup>8,9,14,27</sup> In August 2008, an incident release of ca. 48 GBq of gaseous molecular <sup>131</sup>I  
 187 occurred at the Institute of RadioElements (IRE) in Belgium.<sup>8,9</sup> This amount was released at once and  
 188 corresponded to the yearly <sup>131</sup>I release authorization. The incident was rated 3 on the International  
 189 Nuclear and radiological Event Scale (INES). Unfortunately, this event is not documented with regard  
 190 to measurements in the atmosphere. Since this incident, the releases have been reduced by about a  
 191 factor of 10 (IRE Staff, personal communication). The second event concerns the release of 342 GBq  
 192 of <sup>131</sup>I from the Institute of Isotopes Ltd., Hungary in the time interval between September 8 and

7

193 November 16, 2011.<sup>14,27</sup> This incident release was rated 1 on the INES scale. It led to detections in the  
194 range of a few to several tens of  $\mu\text{Bq m}^{-3}$  in Europe, not only by Ro5 stations but also by aerosol  
195 stations of the International Monitoring Systems (IMS). These stations belong to the Comprehensive  
196 Nuclear-Test-Ban Treaty Organization (CTBTO) and are instrumental for the monitoring of the planet  
197 for clandestine nuclear explosions. During 2008 (with 55 of 80 stations being operational), 6.0 % of all  
198 detections within the IMS network were attributed to  $^{131}\text{I}$ . For example, the IMS station RN61 in  
199 Dubna, Russia, has observed “regular emissions from nearby medical or nuclear facilities” in the past.<sup>6</sup>

200 From the box-and-whisker plot shown in Fig. 1, it is also noteworthy mentioning that the “highest”  
201 particulate  $^{131}\text{I}$  concentrations reported in Europe during the studied event came from monitoring  
202 stations located in the vicinity of radiopharmaceutical production units. This was the case at several  
203 locations: in Warsaw (on Week 2) which is ca. 30 km from the Polatom research Centre; in Budapest  
204 (Weeks 4 and 7) which is 9 km away from the Institute of Isotopes and in Obninsk where the Karpov  
205 Institute is located, with  $2.4 \text{ mBq m}^{-3}$  from January, 13 to 16; and  $2.1 \text{ mBq m}^{-3}$  from February, 3 to 6.<sup>28,</sup>  
206 <sup>29</sup>

207 ***Hypothesis for the origin of airborne  $^{131}\text{I}$ : Nuclear medicine hospitals and sewage sludge***  
208 ***incineration plants***

209 Some aerosol sampling stations that detected small traces of  $^{131}\text{I}$  are located in the vicinity of a nuclear  
210 medicine hospital or a sewage sludge plant that collects their waste waters. This was the case in  
211 Freiburg (Germany), Krakow (Poland) or in České Budějovice (Czech Republic). Some national  
212 monitoring bodies or local institutes in charge of the surveillance of the atmosphere were not surprised  
213 to detect minute amounts of  $^{131}\text{I}$  on their filters because such detections are known to occur  
214 sporadically, especially in winter. Involvement of sewage sludge incinerators on the regular airborne  
215  $^{131}\text{I}$  detection has already been pointed out at short distance.<sup>30</sup>

216 Iodine-131 is a frequently used radionuclide in nuclear medicine with therapeutic (rather than  
217 diagnostic) applications. The administered activities vary with the purpose of the treatment.  
218 Hyperthyroidism is treated with lower activities ( $0.2 - 0.5 \text{ GBq } ^{131}\text{I}$  per treatment) than thyroid cancer  
219 ( $1.8 - 9.2 \text{ GBq}$  per treatment).<sup>6</sup> In 1996, some 180,000 patients were treated on a worldwide basis for  
220 hyperthyroidism and another 20,000 for thyroid cancer.<sup>6</sup> The projected requirement for  $^{131}\text{I}$  in the USA

221 for thyroid cancer treatment in 2009 is 170 TBq of  $^{131}\text{I}$ .<sup>6</sup> The widespread use and growing demands  
222 for imaging organ function and disease states or treatment are directly attributable to the development  
223 and availability of a vast range of specific radiopharmaceuticals.<sup>22, 31</sup> Application rates of  $^{131}\text{I}$  in other  
224 countries in recent years illustrate the extent of its use: Argentina: 300 TBq  $\text{y}^{-1}$ ; Bangladesh: 5 TBq  $\text{y}^{-1}$ ;  
225 Chile: 15,000 patients per year requiring 15 TBq  $\text{y}^{-1}$ ; India: 60 TBq  $\text{y}^{-1}$ ; Thailand: requiring 15 TBq in  
226 2008; Austria: 1 TBq in 2008; Australia: approx. 200 patients per year requiring 4 - 6 GBq per  
227 patient.<sup>6</sup> A fraction of the  $^{131}\text{I}$  activity administered to the patient is excreted into the sewer and  
228 transported to water purification plants.<sup>32, 33</sup> This process takes place even when patients are  
229 hospitalized and their excrements (at least urines) are collected for decay, as it is almost unavoidable  
230 that the patients still carry a residual activity of multiple MBq at the time of release from the hospital.  
231 In France, urines are stored for decay in dedicated tanks but faeces are drained out without significant  
232 decay because of the nosocomial hazard that has to be avoided within hospitals. German regulations  
233 allow patient release from hospitals below a residual  $^{131}\text{I}$  body activity of 250 MBq.<sup>34</sup> Radioactive  
234 waste releases from hospitals are subject to authorizations. For instance, in Germany  $^{131}\text{I}$   
235 concentrations in airborne releases are limited to 0.5 Bq  $\text{m}^{-3}$ , and in liquid effluents to 5 Bq  $\text{L}^{-1}$ .<sup>35</sup>  
236 However, except some local or national initiatives such as the one conducted in Germany in the  
237 framework of the Precautionary Radiation Protection Act, there is no European Union (EU) regulation  
238 with regard to release of radionuclides from sewage sludge incinerators, as they are not considered as  
239 radiation controlled areas. Exemption levels for large amounts of solid materials are comparatively  
240 high, e.g.  $10^4$  Bq  $\text{kg}^{-1}$  of  $^{131}\text{I}$  according to the new European radiation protection legislation.<sup>36</sup>

241 Only a fraction of the  $^{131}\text{I}$  reaching the sewage plant is retained in sewage sludge, since radioiodine  
242 remains mainly in aqueous solution.<sup>37</sup> Nevertheless, routine monitoring results for Germany reveal  
243 median  $^{131}\text{I}$  concentration in digested sludge  $C_{\text{I-131}(\text{sludge})}$  of 30 Bq  $\text{kg}^{-1}$  (dry mass).<sup>38, 39</sup> The solid phase  
244 from the purification of waste water is known for its high content of several undesired ingredients such  
245 as heavy metals, pesticides, dioxins, viruses and bacteria,<sup>39</sup> but the content of traces of radioactive  
246 materials from medical applications is rarely considered with regard to potential release, and except  
247 for dose assessment for workers exposed inside the sewage treatment plant. The presence of poisons  
248 reduces the applicability of this material as fertilizers, so in recent years the combustion of sewage

249 sludge has been promoted. The EU recommends the use of sewage sludge as an alternative fuel for  
250 energy production by combustion. This energy can be used on-site to optimize the energetic yield of  
251 the incinerator, for sludge drying or for heating of the premises. It also avoids or minimizes the costly  
252 transportation and storage outside the plant. Total annual production of dry sewage sludge in the 27  
253 countries of the EU currently reaches almost  $10^7$  t, of which a variable percentage is incinerated.<sup>40</sup> For  
254 example, in Poland, only 3 % of the 160,000 t of dry sludge that could have been burnt were actually  
255 incinerated in 2009,<sup>41</sup> but this rapidly increased to 8.7 % in 2011, with aims to increase the proportion  
256 to 30 % by 2020.<sup>42</sup> In Germany, the percentage of sludge being annually incinerated has increased by  
257 20 % from 1999 to 2001 corresponding to  $\sim 0.46 \times 10^6$  t, to about 60 % of the totally produced sludge  
258 amount in 2013 and 2014, and corresponding a sewage sludge sum up  $m_{(\text{sludge per year})}$  of  $10^6$  t (dry  
259 mass).<sup>42,43</sup> High combustion percentages of sewage sludge have been reported also for Belgium,  
260 Austria, France, the highest being for Switzerland (97 %).<sup>42</sup>

261 In all large sewage plants, the sludge designated for incineration is burnt almost immediately to avoid  
262 the necessity of interim storage space (Bremen WWTP administration, personal communication).  
263 Therefore, no significant decay of  $^{131}\text{I}$  needs to be considered and contributes to the release of  $^{131}\text{I}$  into  
264 the atmosphere despite the filtration system efficiency. In order to reduce odor emissions, sewage  
265 sludge mono-incineration facilities operate at temperatures between 850 °C and 950 °C.<sup>45</sup> This  
266 temperature range is far above the boiling points of methyl iodine (42.4 °C) and molecular iodine  
267 (184.4 °C) leading to the two main gaseous species considered with respect to  $^{131}\text{I}$  and its radiological  
268 impact. Modern incineration plants filter out dust particles from the off-gases and apply scrubbing to  
269 the vapors. IRSN has performed measurements (not published yet) of  $^{131}\text{I}$  concentration both in  
270 wastewaters entering a sewage treatment plant and released by its sewage sludge incinerator. The  
271 average concentration in the liquid waste ( $0.4 \text{ Bq L}^{-1}$ ) at an average flow rate ( $2,500 \text{ m}^3 \text{ h}^{-1}$ ) results in  
272  $10^6 \text{ Bq h}^{-1}$ . Measurements made on samples taken in the flue gas of the sewage sludge incinerator after  
273 the fumes had passed through an electro-filter and a charcoal trap, lead to a release of  $7 \times 10^3 \text{ Bq h}^{-1}$   
274 corresponding to  $\sim 1\%$  of the amount of  $^{131}\text{I}$  entering the plant. For the need of establishing orders of  
275 magnitude an emission factor ( $e_f$ ) ranging 1 to 10 % and corresponding to the capture efficiency of gas

276 and fly ash filtering, can be considered. The total  $^{131}\text{I}$  release to the lower atmosphere by sewage  
 277 sludge incineration for recent years can be calculated to be:

$$278 \quad E_{\text{I-131}(\text{year})} = C_{\text{I-131}(\text{sludge})} \cdot m_{(\text{sludge per year})} \cdot e_f \quad (1)$$

279 With the previously mentioned values for Germany ( $C_{\text{I-131}(\text{sludge})} = 30 \text{ Bq kg}^{-1}$ ,  $m_{(\text{sludge per year})} = 10^6$   
 280 tonnes and  $e_f = 1$  to 10%) this results in a yearly release of 0.3 to 3 GBq of  $^{131}\text{I}$  to the lower  
 281 atmosphere, or 0.8 to 8 MBq on a daily basis.

282 The residence time of the released radioiodine will be lower than its radiological lifetime of  $\tau = 11.57$   
 283 d, due to deposition processes. As an estimate, we use the residence time, which can be extracted from  
 284 the records of Fukushima-originating  $^{131}\text{I}$  concentration in the ground level air, obtained in Germany in  
 285 2011. Records from the German Bundesamt für Strahlenschutz (BfS)<sup>45</sup> show an almost exponential  
 286 decrease over several weeks, from which an effective residence time  $\tau_{\text{eff}}$  between 4 and 6 days can be  
 287 extracted. A slightly lower value (3 to 5 d) was found over the North Pacific Ocean for the iodine  
 288 emissions from the Chernobyl accident.<sup>2</sup> Taken an average value of 5 days, the total  $^{131}\text{I}$  activity in the  
 289 atmosphere over Germany is then given by

$$290 \quad A_{\text{tot, I-131}} = E_{\text{I-131}(\text{day})} \cdot \tau_{\text{eff}} \quad (2)$$

291 Using  $\tau_{\text{eff}} = 5 \text{ d}$ , we get  $A_{\text{tot, I-131}} = 4$  to 40 MBq.

292 Assuming a height of the atmospheric mixing layer of  $h_m = 1,000 \text{ m}$ , the average  $^{131}\text{I}$  activity  
 293 concentration in air over Germany can be obtained by

$$294 \quad a_{\text{I-131}} = A_{\text{tot, I-131}} / (F_{\text{Germany}} \cdot h_m) \quad (3)$$

295 which, with  $F_{\text{Germany}} = 3.6 \times 10^{11} \text{ m}^2$ , results in  $a_{\text{I-131}} = 0.01$  to  $0.1 \text{ } \mu\text{Bq m}^{-3}$ .

296

### 297 ***Hypothesis for the origin of airborne $^{131}\text{I}$ : Nuclear Power Plants***

298 Nuclear power plants have  $^{131}\text{I}$  authorization for their release to the atmosphere. Filter and gas trap  
 299 efficiency are regularly tested to verify that the equipment complies with the requirement of a  
 300 minimum scrubbing efficiency (upstream [ $^{131}\text{I}$ ] / downstream [ $^{131}\text{I}$ ] > 1000) for the gaseous fraction.  
 301 The French situation (58 PWR reactors with individual electrical power ranging 900 to 1,450 MWe)  
 302 corresponds to the highest possible expected airborne concentrations and has thus been taken as a  
 303 maximum source contribution. The sum of the reported  $^{131}\text{I}$  releases to the atmosphere from all French

304 NPPs during 2013 leads to 0.3 GBq y<sup>-1</sup>. Note that airborne monitoring performed in the environment  
 305 close to these NPPs in normal operation has never revealed <sup>131</sup>I concentration above detection limits  
 306 and such releases are supposed to occur mainly during the filter efficiency tests performed with CH<sub>3</sub>-  
 307 <sup>131</sup>I, but have been considered here like a chronic release. Taking a one-day release corresponds to 0.8  
 308 MBq and assuming an effective <sup>131</sup>I residence time  $\tau_{\text{eff}}$  of 5 days as for equation (2) leads to  $A_{\text{tot, I-131}} = 4$  MBq. Assuming a height of the mixing layer of  $h_m = 1000$  m, the average total <sup>131</sup>I activity  
 309 concentration in air over France can be obtained by

$$311 \quad a_{\text{I-131}} = A_{\text{tot, I-131}} / (F_{\text{France}} \cdot h_m) \quad (4)$$

312 which, with  $F_{\text{France}}$  the metropolitan area =  $5.5 \times 10^{11}$  m<sup>2</sup> results in  $a_{\text{I-131}} = 0.007$   $\mu\text{Bq m}^{-3}$ . Proper  
 313 comparison with the observations performed in January/February 2017 requires considering a particle-  
 314 to-gas <sup>131</sup>I ratio of about 30%, based on previous works<sup>1</sup> and leads to 0.0022  $\mu\text{Bq m}^{-3}$  for the expected  
 315 <sup>131</sup>I particulate fraction, which is out of reach for the most sensitive gamma-ray detection systems.

316

### 317 ***Hypothesis for the origin of airborne <sup>131</sup>I: Spontaneous fission-derived <sup>131</sup>I from natural <sup>238</sup>U***

318 There is plenty of literature dedicated to emanation of geogas from soil,<sup>47</sup> including radon (<sup>222</sup>Rn).<sup>48</sup>  
 319 Trace amounts of volatile fission products can also be produced by spontaneous fission of naturally  
 320 occurring heavy nuclides. This has already been demonstrated for <sup>129</sup>I, however, it has never been  
 321 evaluated for <sup>131</sup>I, so far. Of all relevant nuclides, <sup>238</sup>U has the highest spontaneous fission rate with 5.5  
 322  $\times 10^{-5}$  % of the entity of disintegrations.<sup>49</sup> The average concentration of U in the upper continental crust  
 323 is 2.8 mg kg<sup>-1</sup>, the vast majority of which is <sup>238</sup>U (99.27 %).<sup>50</sup>

324 The *in situ* production of some fission products by spontaneous fission of <sup>238</sup>U within the rock has  
 325 been estimated and compared with <sup>129</sup>I measured in the Stripa granite, located about 170 km WNW of  
 326 Stockholm, and characterized by an unusual high uranium content (44 ppm) of more than ten times the  
 327 average value for granite.<sup>50, 51</sup> However, there is no available estimation reporting the subsurface  
 328 production of <sup>131</sup>I and its potential emission to the atmosphere. Assumptions about an underground  
 329 source of <sup>131</sup>I are deduced by analogy with <sup>129</sup>I or <sup>222</sup>Rn produced by fission and decay of <sup>238</sup>U,  
 330 respectively before their exhalation (Table 2).

331 In the following, the fission product release estimation is carried out for rock representative of the  
 332 earth's upper crust, characterized by a rather low average  $^{238}\text{U}$  activity of  $30 \text{ Bq kg}^{-1}$ . For such  
 333 concentrations, the induced fission of  $^{235}\text{U}$  has been neglected. The *in situ* production of  $^{131}\text{I}$  by the  
 334 spontaneous fission of  $^{238}\text{U}$  is estimated using equation (5) initially derived for  $^{129}\text{I}$  from <sup>46</sup> which gives  
 335 the equilibrium number of  $^{131}\text{I}$  atoms in  $1 \text{ cm}^3$  of rock, ( $^{131}\text{N}_e$ ):

$$336 \quad ^{131}\text{N}_e = \frac{(^{238}\text{N} \cdot ^{238}\lambda_{sf} \cdot ^{131}\text{Y}_{sf})}{^{131}\lambda} \quad (5)$$

337 where  $^{238}\text{N}$  is the number of  $^{238}\text{U}$  atoms per  $\text{cm}^3$ ,  $^{238}\lambda_{sf}$  and  $^{131}\lambda$  are the spontaneous decay constants of  
 338  $^{238}\text{U}$  and  $^{131}\text{I}$ , respectively and  $^{131}\text{Y}_{sf}$  is the spontaneous fission yield of  $^{131}\text{I}$ .

339 Using the same equation for  $^{129}\text{I}$  leads to a concentration of activity in the rock of  $1.29 \times 10^{-5} \text{ Bq m}^{-3}$   
 340 which is lower than the estimation for Stripa granite ( $2.34 \times 10^{-4} \text{ Bq m}^{-3}$ ) (Table 2).<sup>52</sup> This difference  
 341 results from the higher  $^{238}\text{U}$  activity in the granite ( $550 \text{ Bq kg}^{-1}$ ) compared with the average upper  
 342 continental crust value of  $30 \text{ Bq kg}^{-1}$ .<sup>51</sup> Furthermore, the comparison of the activity of both *in situ*  
 343 produced iodine isotopes shows that the production of  $^{131}\text{I}$  is larger than that of  $^{129}\text{I}$  by about a factor of  
 344 20, due to a higher production rate for  $^{131}\text{I}$ . Thereafter, the loss of iodine from minerals is mainly due  
 345 to the recoil of radionuclides. The high activity level recorded in the Stripa granite water is accounted  
 346 by the diffusion of iodine in the rock porosity and the dissolution in the underground water.<sup>52</sup> Even so,  
 347 these levels are less than 1 % of the estimated concentrations due to *in situ* production and total  
 348 diffusive loss from the granite to the water.<sup>51</sup> Similar reasoning can be applied for other primordial  
 349 radionuclides such as  $^{232}\text{Th}$  (spontaneous fission) and  $^{235}\text{U}$  (induced fission). At a first estimate, the  
 350 consideration of thorium leads to the same order of magnitude as for uranium.

351 The transfer of fissiogenic iodine from subsoil to the atmosphere can be simplified as follows: once  
 352 produced in the rock, volatile iodine diffuses to the soil surface. During the diffusion stage, volatile  
 353 iodine will encounter soil organic matter that will considerably reduce the emission to the atmosphere.  
 354 Iodine sorption on natural organic matter remains complex.<sup>53</sup> However, in case of anoxic conditions  
 355 (e.g. peat bog or water saturated area), poorly sorbed iodine species and volatile iodine species driven  
 356 by microbial processes can be emitted to the atmosphere.<sup>54</sup> In the case of  $^{131}\text{I}$ , its rather short half-life

357 will also compete with the diffusion duration and will limit the activity that will effectively be emitted  
358 to the atmosphere.

359 The  $^{131}\text{I}$  activity measured in the atmosphere soon after underground fission nuclear tests shows that  
360 less than 1 % of  $^{131}\text{I}$  reaches the surface.<sup>55</sup> Further isotopic data from this study shows that there is no  
361 significant fractionation between the xenon isotopes and the precursors, namely iodine isotopes,  
362 suggesting that iodine could behave like a noble gas during diffusion to the surface. Following this  
363 assumption, we use the ratio between activity of  $^{222}\text{Rn}$  in the rock ( $7.8 \times 10^4 \text{ Bq m}^{-3}$ ) and the radon  
364 activity typically recorded in the troposphere ( $10\text{-}30 \text{ Bq m}^{-3}$ ) to derive the maximum activity of  $^{131}\text{I}$  in  
365 the atmosphere, ranging between  $3.6 \times 10^{-8}$  and  $1.1 \times 10^{-7} \text{ Bq m}^{-3}$ , respectively (Table 2). However,  
366 considering the strong sorption of iodine into soils and interaction with organic matter, about 1 % only  
367 of radioiodine can reach the surface.<sup>56</sup> This leads finally to  $^{131}\text{I}$  activity in the troposphere of ca.  $10^{-9}$   
368  $\text{Bq m}^{-3}$ . This is far below the MDA of the highest performing monitoring stations.

369

370 *Table 2: Calculation of the in situ production of  $^{131}\text{I}$  and  $^{129}\text{I}$  by spontaneous fission of  $^{238}\text{U}$  in the rock*  
371 *and estimation of the activity in the troposphere. In situ production of  $^{222}\text{Rn}$  from  $^{238}\text{U}$  decay is also*  
372 *calculated, assuming secular equilibrium among U-daughters.*

		$^{222}\text{Rn}$	$^{131}\text{I}$	$^{129}\text{I}$
Process of <i>in situ</i> production		decay of $^{238}\text{U}$	spontaneous fission of $^{238}\text{U}$	spontaneous fission of $^{238}\text{U}$
calculation parameters	Production half-life (y)	$4.47 \times 10^9$	$8.20 \times 10^{15}$ *	$8.20 \times 10^{15}$ *
	Production yield (%)	100	0.65**	0.03**
	Decay period of produced radionuclide (y)	$1.04 \times 10^{-2}$	$2.20 \times 10^{-2}$	$1.57 \times 10^7$
<i>In situ</i> activity ( $\text{Bq m}^{-3}$ )		$7.8 \times 10^4$	$2.8 \times 10^{-4}$	$1.3 \times 10^{-5}$
calculation results	<i>In situ</i> activity ( $\text{Bq m}^{-3}$ )	$7.8 \times 10^4$	$2.8 \times 10^{-4}$	$1.3 \times 10^{-5}$
	Tropospheric concentration ( $\text{Bq m}^{-3}$ )	10-30	$1.1 \times 10^{-9}$	$5.4 \times 10^{-11}$

373 \* ref.<sup>57</sup>

374 \*\* refs.<sup>58</sup> and <sup>59</sup>

375

### 376 Source apportionment

377 The above estimates for the various  $^{131}\text{I}$  source hypotheses are ranked in Fig. 2. It is clear that, given  
378 the current MDA, any contributions of  $^{131}\text{I}$  either from spontaneous fission of uranium or thorium in  
379 soil or from nuclear power plants remain undetectable by the high-volume sampler and low-level



380 detection equipment of the Ro5 members. The situation is similar for the release source from sewage  
381 sludge incinerators on a global scale but this explanation for detection on a local scale cannot be ruled  
382 out. The most likely release source may concern radiopharmaceutical production units or handling  
383 sites. The release of 342 GBq of  $^{131}\text{I}$  that took place in Hungary in fall 2011 resulted in a detectable  
384 activity range of particulate  $^{131}\text{I}$  about 5 to 10 times higher than the one observed in January/February  
385 2017. Assuming the same dispersion coefficient for both events leads to a rough estimate of the source  
386 term that may have been released in January 2017 of about one tenth, thus about 35 to 70 GBq (Fig.  
387 2). Note that this amount typically matches a monthly average  $^{131}\text{I}$  authorized release of 65 GBq  
388 derived from the yearly authorization ( $780 \text{ GBq y}^{-1} / 12 \text{ months}$ ) from the Karpov Institute alone.

389

390 Figure 2

391

### 392 ***Inverse modeling and attempted localization of the origin of the release***

393 Considering the likelihood of radiopharmaceutical unit involvement, inverse modeling techniques  
394 have been applied by mixing field observations and atmospheric dispersion models. The goal was to  
395 pinpoint the  $^{131}\text{I}$  source and estimate the amount released in the environment. IRSN has developed a  
396 tool to assess an accident release into the environment when the location of the source is  
397 known.<sup>60,61,62,63</sup> The approach consists in the resolution of the inverse problem associated with the  
398 source-receptor relationship:<sup>64</sup>

399

$$\mu = H\sigma + \epsilon \quad (6)$$

400 where  $\mu$  contains air concentrations measurements;  $H$  is the source-receptor matrix calculated using  
401 atmospheric transport models,  $\epsilon$  is a vector that represents errors in the system (model, instrument, and  
402 representativeness errors), and  $\sigma$  is the unknown temporal evolution of the release rate (source term).

403 After all, the aim is to assess the source term  $\sigma$  such as the error  $\epsilon$  is minimal. Considering simplified  
404 assumptions as in,<sup>61</sup> the resolution of the inverse problem (1) involves to minimize the following cost

405 function:  $J(\sigma) = \|\mu - H\sigma\|^2 + \lambda^2\|\sigma\|^2$

406 Subject to  $\sigma \geq 0$

407 The first term of  $J(\sigma)$  measures differences between observed and simulated concentrations while the  
 408 second one is a regularization term, which allows avoiding unreliable solutions. The  $\lambda$  parameter is a  
 409 scalar, which determines the magnitude of the source term fluctuations ( $\sigma$ ). The method is applied  
 410 assuming that each  $^{131}\text{I}$  producer identified in Table 1 is a possible source location. For each producer  
 411  $k$ , ( $1 \leq k \leq 6$ ), the following cost function  $J_k(\sigma_k)$  related to  $k$  is minimized using a quasi-Newton  
 412 algorithm:

$$413 \quad J_k(\sigma_k) = \|\mu - H_k \sigma_k\|^2 + \lambda^2 \|\sigma_k\|^2 \quad (7)$$

414

415 Subject to  $\sigma_k \geq 0$

416 Then, we determine the producer  $k^*$  such as:

$$417 \quad k^* = \arg \min_{1 \leq k \leq 6} \{J_k(\sigma_k)\} \quad (8)$$

418 This approach leads to identify the producer  $k^*$  which is the most able to reproduce the observations  $\mu$   
 419 and which can be therefore considered as the most reliable source location.

420 Atmospheric dispersion simulations have been carried out with the Eulerian ldx model<sup>65</sup> developed by  
 421 IRSN. Meteorological fields are provided by the ARPEGE model developed by Météo-France. The  
 422 spatial resolution of the data is  $0.5^\circ \times 0.5^\circ$  with 3-hour time resolution. The source-receptor matrix  $H$   
 423 is computed using ldx under the approach proposed by ref.<sup>66</sup>. Dimensions of the computational domain  
 424 are [10W, 70E], [35N, 75N] and cover the whole  $^{131}\text{I}$  detection area. The release height is taken for the  
 425 first level of the model between 0 and 40 m. For each  $^{131}\text{I}$  producer  $k$ , daily release rates have been  
 426 assessed from January 5 to February 20 (i.e. 46 days) by minimizing the cost function  $J_k(\sigma_k)$ . It means  
 427 that the number of unknowns related to the producer  $k$  is equal to 46. In total, 196 observations were  
 428 considered in the inversion process apart from the values obtained in Obninsk due to their closeness to  
 429 the Karpov institute (i.e., within the same mesh cell as the source location). Even when a station did  
 430 not report  $^{131}\text{I}$  trace in the atmosphere, it contributed to constraint the inverse problem. Several values  
 431 of  $\lambda$  in the range of  $[10^{-8}, 10^{-5}]$  have been selected.

432 Table 3 gives the values of relative error reduction of the cost function  $J_k(\sigma_k)$  obtained after  
 433 minimization and the total amount of  $^{131}\text{I}$  released between January 5 and February 20. The relative  
 434 error reduction of  $J_k(\sigma_k)$  is given by:

435

$$436 \quad E_r = \frac{(J_k(\sigma_k))_{init} - (J_k(\sigma_k))_{end}}{(J_k(\sigma_k))_{init}} \times 100 \quad (9)$$

437

438 where  $(J_k(\sigma_k))_{init}$  is the initial value of the cost function  $J_k(\sigma_k)$  and  $(J_k(\sigma_k))_{end}$  the minimum value  
 439 of  $J_k(\sigma_k)$  obtained after minimization.

440 The maximum value of  $E_r$  is obtained when the Karpov Institute in Obninsk is assumed to be the  
 441 source location. In that case, the amount of  $^{131}\text{I}$  released estimated is ranging between 65 and 106 GBq,  
 442 which is also consistent with the estimate given previously in the source apportionment section.  
 443 However, these results have to be interpreted cautiously due to the inherent characteristics of the cost  
 444 function  $J_k(\sigma_k)$ , which attributes more weight on the high values of concentrations than on the low  
 445 values. Apart from the values in Obninsk, the highest concentration levels have been reported from  
 446 Dubna with several tens of  $\mu\text{Bq per m}^3$ . This station, only 200 km away from Obninsk, thus has a  
 447 more significant weight than other, farther stations. Apart from that, the results on the source term are  
 448 sensitive to the values of  $\lambda$  due to the rather small number of observations used in the inversion  
 449 process. Therefore, the results given in Table 3 exemplify that the Karpov Institute (Obninsk branch)  
 450 is the  $^{131}\text{I}$  producer, since this location provides a better reproduction of the higher observed values  
 451 reported in Western Russia. However, the values of  $J_k(\sigma_k)$  obtained after minimization do not allow to  
 452 rule on the ability to reproduce the lower levels of  $^{131}\text{I}$  concentrations measured in Western and Central  
 453 Europe.

454 *Table 3. Relative error reduction of  $J_k(\sigma_k)$  after minimization and  $^{131}\text{I}$  total amount estimated for*  
 455 *each  $^{131}\text{I}$  producer  $k$  according to the value of  $\lambda$ .*

$^{131}\text{I}$ producer (location)	Released activity (GBq)	Relative error reduction ( $E_r$ )
Cis-Bio international (Saclay)	6 - 96	0 - 1%
Mallinckrodt Medical (Petten)	14 - 230	0 - 1%
Institut des RadioEléments (Fleurus)	16 - 134	0 - 1%
Polatom (Otwock-Świerk)	44 - 73	2 - 6%

Institute of Isotopes (Budapest)	41 - 256	3 - 4%
Karpov Institute, NIFKhl (Obninsk)	4 2- 78	98 - 99%

456 To validate the relevance of the Karpov Institute as the likely emission location, forward simulations  
 457 were performed in a second stage with Idx using the source term assessed from the Karpov Institute,  
 458 with  $\lambda = 10^{-7}$ . The comparison between simulated and observed concentrations demonstrates that the  
 459 Karpov Institute is likely the origin of  $^{131}\text{I}$  detections in Western Russia and Scandinavia in January  
 460 and also in Western Russia, Poland, Germany and France during the first two weeks of February.  
 461 Figure 3 illustrates the satisfactory agreement between the simulated and observed concentrations in  
 462 these geographical areas. In addition, a video animation showing the hourly progress of the plume  
 463 dispersion from Obninsk between January 5 and February 20, and the weekly field measurements is  
 464 provided in the Supporting Information (SI).

465 However, the simulations are not able to reproduce the  $^{131}\text{I}$  detections in January, located in Western  
 466 Europe (France, Italy, and Germany), Hungary and Poland. Therefore, the hypothesis of concomitant  
 467 releases from different facilities is a serious possibility. Indeed, the fairly high detections reported in  
 468 Budapest ( $10 \mu\text{Bq m}^{-3}$  between 23 and 30 January) may indicate that a release from the nearby  
 469 Institute of Isotopes could have contributed. Forward simulations with Idx show that a release from  
 470 Budapest could explain observations in Spain, France, Germany, and Czech Republic between January  
 471 16 and January 23. More advanced inverse modeling methods, based on Bayesian techniques, which  
 472 do not require any knowledge of potential source locations would be a worthy addition to the present  
 473 simulation results.<sup>8, 65, 68, 69</sup>

474  
 475 Figure 3  
 476

477

#### 478 ***Role of meteorological conditions***

479 The change in the detection locations mainly obeyed the prevailing wind directions over Europe in  
 480 January and February 2017. The meteorological conditions had a strong impact on the detectability of  
 481 the concentrations. The beginning of the year in Europe was especially characterized by poor  
 482 atmospheric dispersion conditions, i.e. high-pressure fields and strong temperature gradient inversions

483 (hereafter called “temperature inversion”) in the lower atmospheric layers, prone to increase any  
484 airborne pollutant concentration. This was the case during Weeks 3, 4, 6, and 7. Valuable information  
485 on the mixing or dispersion conditions prevailing in January 2017 over Western Europe has been  
486 derived from PM<sub>10</sub> increases. Maps of peak values provided by the PREV’AIR modeling system<sup>70</sup>  
487 show how it affects the air quality (Fig. 4) contributing to the atmospheric pollution event.

488 Figure 4  
489

490 Typical PM<sub>10</sub> values are ranging from 10 to 20 μg m<sup>-3</sup>. Simultaneous PM<sub>10</sub> and <sup>137</sup>Cs peak values  
491 indicate that both airborne compounds were similarly affected by meteorological conditions as  
492 exemplified by three French locations (Fig. 5). Atmospheric detections of <sup>137</sup>Cs are typically  
493 associated with the re-suspension of contaminated soil particles.

Figure 5

494  
495 Increased atmospheric activity concentrations have been also reported for other radionuclides during  
496 the <sup>131</sup>I episode, including naturally occurring <sup>40</sup>K and <sup>210</sup>Pb or anthropogenic <sup>137</sup>Cs that are detected on  
497 a routine basis by the organizations involved in the Ro5 network. Airborne concentrations peaked at  
498 their maximum levels since several months or even years. Compared with the last-6-year average  
499 values, they were higher by a factor of 1.8 for <sup>40</sup>K, 2.7 for <sup>210</sup>Pb and up to 5.2 for <sup>137</sup>Cs during the <sup>131</sup>I  
500 episode observed at the French sampling locations. No comparable increasing concentrations have  
501 been reported for <sup>7</sup>Be (factor 0.9) or <sup>22</sup>Na (factor 0.7) both of which are produced in the lower  
502 stratosphere and upper troposphere. This discrepancy points at an atmospheric process, which affects  
503 mostly radionuclides in the lower troposphere, such as <sup>137</sup>Cs or <sup>131</sup>I. Such enhancing factor becomes  
504 explainable by the temperature gradient inversion layer that could develop and last for several days,  
505 due to the radiative cooling of the atmosphere as well as low winds or lack of wind. We took an  
506 average ratio of 3.2 as representative of the average increase ratio observed at French locations for  
507 <sup>137</sup>Cs, <sup>40</sup>K, and <sup>210</sup>Pb during the time when <sup>131</sup>I was detected. Note that this ratio was almost identical  
508 for the PM<sub>10</sub> increase. We called this ratio the “detection enhancing factor” that also affects particulate  
509 <sup>131</sup>I. Dividing <sup>131</sup>I concentrations by this factor leads to hypothetical <sup>131</sup>I activity concentration that

510 would have prevailed without such enhancing meteorological conditions. In such scenario, most of the  
511 observed  $^{131}\text{I}$  concentrations would have fallen below usual detection limits. This is sufficient to  
512 explain why the Ro5 network does not measure  $^{131}\text{I}$  on a regular basis, even if it can be hypothesized  
513 that  $^{131}\text{I}$  will become present more or less routinely at ultra-traces levels as the result of sporadic  
514 release events or routine releases from the booming radiopharmaceutical industry.

515 Detection of harmless traces of airborne  $^{131}\text{I}$  in January/February 2017 in Europe resulted from a  
516 combination of multiple sources and poor atmospheric dispersion conditions. The main releases  
517 capable to imprint trace concentrations on a wide scale belong to the radiopharmaceutical industry that  
518 remains the most probable source in the present case study. Based on field observations and inverse  
519 dispersion modeling, the Obninsk region appears as the most likely involved emission area, but with a  
520 release amount consistent with usual authorized limits for the Karpov Institute. This makes this  
521 unusual  $^{131}\text{I}$  episode over Europe the first case of a regular/routine release of a radionuclide resulting in  
522 continental detections by members of the Ro5 monitoring network. Other releases from  
523 radiopharmaceutical production units located in Hungary and to a lesser extent in Poland may explain  
524 the concentrations observed on Week 2 and Week 4, respectively on a local to regional scale. Apart  
525 from radiopharmaceutical release sources, there were multiple secondary additional emissions (e.g.  
526 from nuclear medicine hospitals or sewage sludge incinerators) that were too weak to significantly  
527 contribute to European-wide detections when considering them alone, but that may have exhibited a  
528 dominant contribution on a local scale. In any case, nuclear power plant-derived releases of  $^{131}\text{I}$  appear  
529 too weak to be detected even by monitoring stations located in their vicinity. All of those detections  
530 would probably have gone unnoticed without unusually disadvantageous weather conditions marked  
531 by strong inversions of the vertical temperature gradient in the lower atmospheric layers. As a result,  
532 they led to concentrations exceeding the usual detection limits in use within the Ro5 network. While  
533 likely to be dominant, the lack of reports of the gaseous fraction of  $^{131}\text{I}$  also denotes the demand to  
534 improve both its detection limits and the number of samplers for gaseous radioiodine. Finally this  
535 event also confirms the capability of the organizations involved in the Ro5 informal network, and in  
536 charge of monitoring airborne radioactivity to detect unexpected radionuclides or unusual

537 concentrations, down to the 0.1 – 1  $\mu\text{Bq m}^{-3}$  range, thus ensuring a highly sensitive monitoring even at  
538 concentration levels of no concern for human health.

### 539 **Associated content**

540 Supporting Information: Video animation of the plume dispersion from Obninsk.

541

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544

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### 555 **Disclaimer**

556 The views expressed in this study are those of the authors and do not necessarily reflect the views of  
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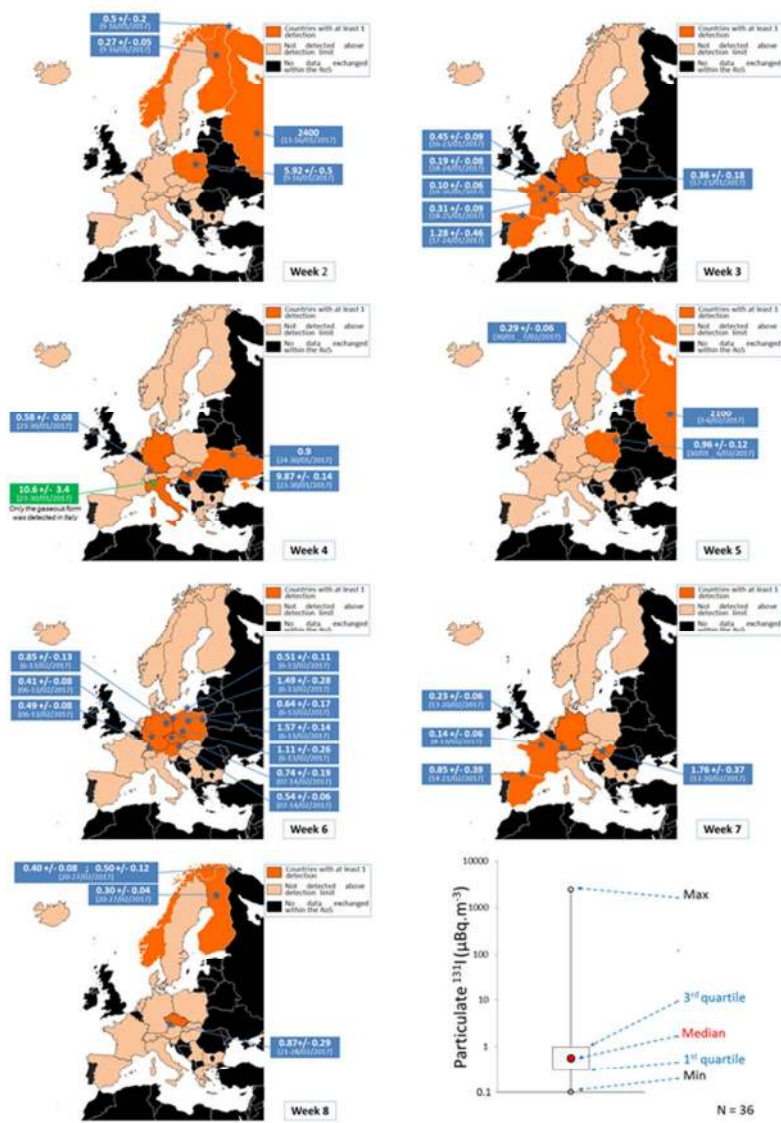


Figure 1. Weekly changes of particulate  $^{131}\text{I}$  ( $\mu\text{Bq m}^{-3}$ ) in January / February 2017 and box-and-whisker plot of the 36 weekly values. CTBTO data are not presented since based on daily samples.

190x254mm (96 x 96 DPI)

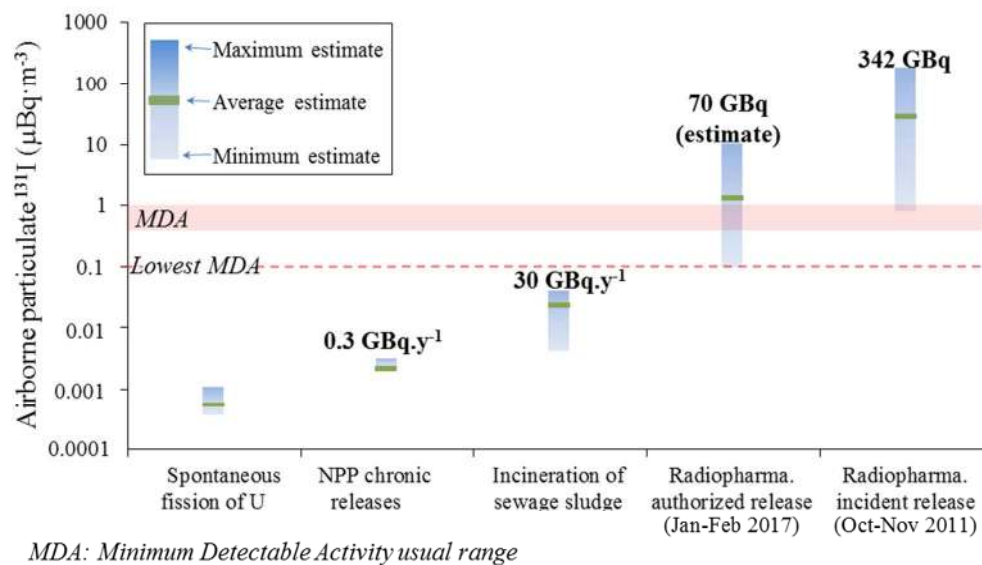


Figure 2. Source apportionment of particulate <sup>131</sup>I in the air over Europe in January/February 2017 based on source term estimates.

254x190mm (96 x 96 DPI)

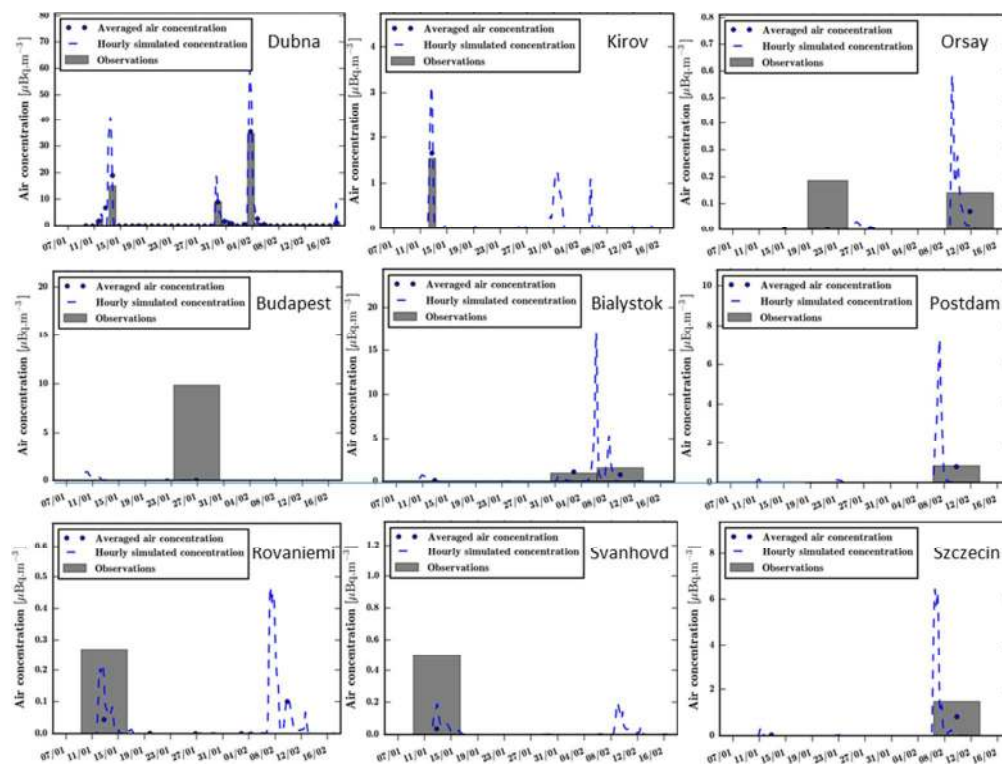


Figure 3. Observed  $^{131}\text{I}$  activity concentrations (grey blocks), and simulated activity concentrations derived from the source term assessed for the Karpov Institute. Blue dashes are hourly simulated concentrations and blue circles are centered on the respective time interval measurements.

254x190mm (96 x 96 DPI)

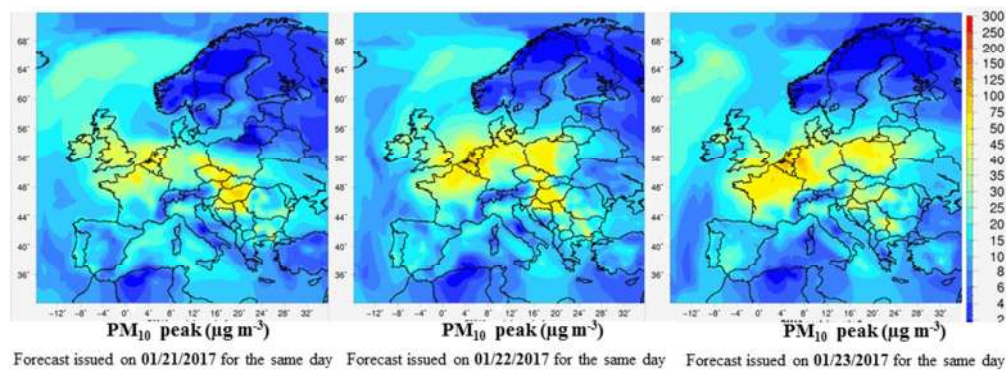


Figure 4. PM<sub>10</sub> peak value forecasts ( $\mu\text{g m}^{-3}$ ) over Europe, 21-23 January 2017. The maps are representative for large-scale phenomena; they cannot reproduce local aspects of air pollution.

254x190mm (96 x 96 DPI)

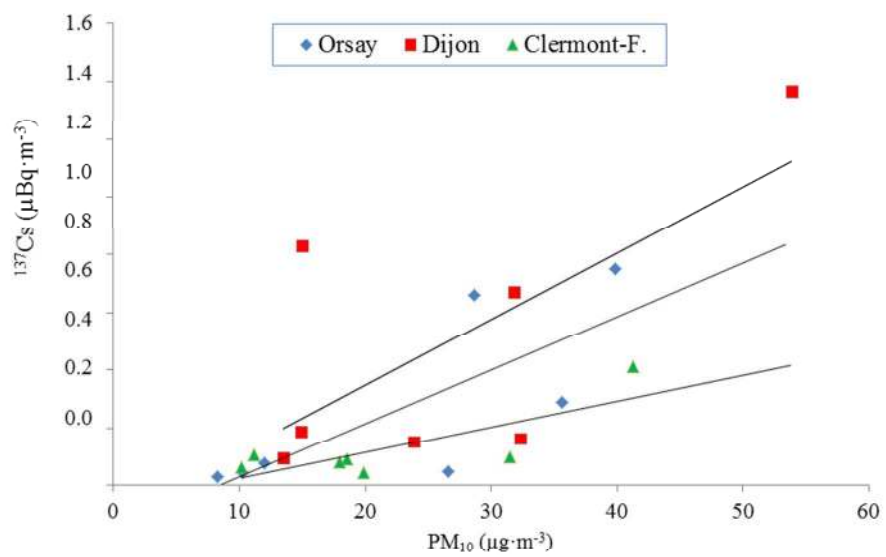


Figure 5. Example of weekly PM<sub>10</sub> and <sup>137</sup>Cs at French sampling locations having <sup>131</sup>I detections in January and February 2017.

254x190mm (96 x 96 DPI)





254x190mm (96 x 96 DPI)