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Europe-wide atmospheric radionuclide dispersion by unprecedented wildfires in the Chernobyl Exclusion Zone, April 2020

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1 Abstract

2 From early April 2020, wildfires raged in the highly contaminated areas around the Chernobyl nuclear 3 power plant (CNPP), Ukraine. For about four weeks, the fires spread around and into the Chernobyl 4 exclusion zone (CEZ) and came within a few kilometres of both the CNPP and radioactive waste storage 5 facilities. Wildfires occurred on several occasions throughout the month of April. They were extinguished, 6 but weather conditions and the spread of fires by airborne embers and smoldering fires led to new fires 7 starting at different locations of the CEZ. The forest fires were only completely under control at the 8 beginning of May, thanks to the tireless and incessant work of the firefighters and a period of sustained 9 precipitation. In total, 0.7-1.2 TBq ¹³⁷Cs were released into the atmosphere. Smoke plumes partly spread 10 south and west and contributed to the detection of airborne ¹³⁷Cs over the Ukrainian territory and as far away 11 as Western Europe. The increase in airborne ¹³⁷Cs ranged from several hundred µBq·m⁻³ in northern Ukraine to trace levels of a few μ Bq·m⁻³ or even within the usual background level in other European countries. 12 13 Dispersion modeling determined the plume arrival time and was helpful in the assessment of the possible 14 increase in airborne 137 Cs concentrations in Europe. Detections of airborne 90 Sr (emission estimate 345 – 15 612 GBq) and Pu (up to 75 GBq, mostly ²⁴¹Pu) were reported from the CEZ. Americium-241 represented 16 only 1.4% of the total source term corresponding to the studied anthropogenic radionuclides but would have

17 contributed up to 80% of the inhalation dose.

18 Synopsis

Wildfires in highly radioactive environment can re-emit radionuclides into the atmosphere. Such emissionspresent a potential health risk for firefighters.

21 Introduction

22 As a result of global fallout from atmospheric nuclear explosions and the Chernobyl nuclear accident, the 23 Eurasian boreal forest represents one of the greatest stocks of long-lived anthropogenic radionuclides in the terrestrial environment, primarily ¹³⁷Cs ($T_{1/2} = 30.1$ yr.).¹ Since large wildfires in 1992, the fire hazard in the 24 Chernobyl area has been viewed with serious concern.² These fire events are capable of emitting 25 26 radionuclides (RN) into the atmosphere and can redistribute part of the already deposited RN.³ These RN are found in topsoil lavers, forest litter, and in the biomass. Emission of natural RN such as ²¹⁰Po are also 27 28 known to occur from wildfire events.^{4, 5} Wildfires in heavily contaminated areas generate radioactive smoke 29 particles and thus an additional radiation exposure through inhalation or ingestion of contaminated foodstuff, following RN re-deposition.⁶ The consequences of wildfires in the highly contaminated area 30 31 around the CNPP (parts of northern Ukraine, southern Belarus and the western part of the Russian 32 Federation) as well as emission factors or resuspension factors have already been investigated at a local

level.⁷⁻¹² Evidence for long-range transport of RN from fires on an international scale is more recent.^{1, 13} 33 Considerable efforts have been made by Ukraine, Belarus, and the Russian Federation to limit the 34 35 consequences of fires in contaminated areas.^{14, 15} However, despite preventative measures (e.g., controled 36 fires, fire-breaks, access trails, limitation of fuel quantities in some areas, minimization of human presence) 37 intended to limit both ignition and the spread of wildfires, they occur on a yearly basis in the Chernobyl area^{16, 17} and affect wildlife.¹⁸ The Chernobyl ecosystem has regularly suffered from major wildfires notably 38 in 1992, 1999, 2000, 2002-2004, 2006, 2010, 2015, 2016, and 2018^{15, 19} with major impacts on the 39 vegetation cover.²⁰ For a brief historic review of wildfires in contaminated areas, see the Supporting 40 41 Information (SI).

42

43 Herein, we investigate the devastating April 2020 wildfires, which lasted for about four weeks in the 44 Ukrainian part of the contaminated areas around the CNPP. The detailed geographic analysis and timeline 45 is provided in the SI. The fire situation in the CEZ and bordering areas was characterized by a combination 46 of numerous ignitions and subsequent spread of fires. Their magnitude varied according to different 47 parameters: 1) biomass type, vegetation density, and location accessibility (forest, meadow, peatland, and 48 marshland); 2) meteorological parameters (wind speed, wind direction, precipitation frequency and 49 amount). These multiple factors hindered firefighting, despite the mobilization of nearly 400 firefighters 50 and 90 specialized aerial and terrestrial vehicles (two AN-32P airplanes, one Mi-8 water-bombing 51 helicopter, heavy engineering equipment, and seven additional road construction machines of the Armed 52 Forces of Ukraine). The first three weeks of April saw the development of particularly large and numerous 53 fires. Two main fire areas were identified during this period: in the Polisske district and in the Kopachi-54 Chistogalovka-CNPP cooling pond (<12 km from CNPP). Daily information about burned areas including 55 vegetation cover, contamination density, and radionuclide emissions were then published by the Ukrainian Hydrometeorological Institute (UHMI).²¹ According to the UHMI, 870 km² were burned in total, including 56 57 65 km² in proximity to the CNPP and 20 km² on the left bank of the Pripyat river.

58 Because of a period of easterly and southerly winds, slight increases in the airborne ¹³⁷Cs were observed at 59 a few western European locations while most of the airborne ¹³⁷Cs concentrations remained within the 60 μ Bq·m⁻³ range on a weekly sampling basis, which corresponds to the usual background level. Such western 61 detections are somewhat rare since the general air mass circulation is usually easterly. A similar situation 62 had already occurred from the end of August to early September 2002 with a slight increase in airborne ¹³⁷Cs concentrations in the western European atmosphere.²² The follow-up of the April 2020 wildfire 63 situation was scrutinized on a daily basis by the UHMI,²¹ and regularly commented by the SCK-CEN^{23, 24} 64 and the IRSN.²⁵⁻²⁸ In total about 1,100 ¹³⁷Cs results were gathered throughout Europe including already 65 66 published data from the International Monitoring System (IMS) in support to the Comprehensive NuclearTest-Ban Treaty Organization.^{23, 24} This collection, which also includes a hundred values for ⁹⁰Sr, ²³⁸Pu and ²³⁹⁺²⁴⁰Pu from Ukraine, corresponds to the most comprehensive available dataset related to that event (see Tables S6 and S7, SI). The purpose of this study is to investigate both the RNs source terms and the additional exposure to RNs for firefighters and inhabitants of Kiev. Part of these assessments are based on numerous measurements performed during this event and dispersion calculation. RNs that were not determined have been estimated based on RN ratios typical of the Chernobyl accident and ²¹⁰Po results obtained during other wildfires.

74

75 Background

76 Starting on April 26, 1986 and for a period of ten days, the Chernobyl accident released harmful quantities 77 of radionuclides of I, Cs, Te, Sr, Pu, and others (see Table S1, SI). Some regions of Belarus, Ukraine and the Russian Federation were seriously affected by the radioactive fallout from the CNPP accident.²⁹ About 78 79 6 million ha of forest including 2.5 million ha in Ukraine were heavily contaminated. In the most 80 contaminated regions following the accident the dominant forests were young or middle-aged pine and pine-81 hardwood stands, with a high fire risk.⁸ The highest radionuclide deposition density occurred in the area 82 surrounding the CNPP, in the so-called Chernobyl Exclusion Zone (CEZ) in Ukraine and in the Polesie 83 State Radioecological Reserve (PER) in Belarus. The CEZ, initially about 30 km in radius around the CNPP 84 was subsequently enlarged to an oblong area of 2,600 km² with a 439 km circumference. It is located approximately 100 km north of Kiev (see Figure S1, SI). The CEZ is mostly covered by forest where 85 86 radionuclides are distributed between soil, forest litter, and vegetation. Between 57 and 79% of the total ¹³⁷Cs contamination is stored within the upper soil layer (0- 2 cm).³⁰ Only a few percent of the ¹³⁷Cs 87 inventory is contained in the living biomass, where 137 Cs behaves like potassium, its chemical analogue. 88 89 During fire events in forested areas, the main source of radioactive aerosols is the burning forest litter. In comparison, the trees affected by the fire emit minor amounts of ¹³⁷Cs and ⁹⁰Sr and only trace amounts of 90 Pu isotopes and ¹⁴⁴Ce.^{31, 32} More information about radionuclide apportionment in the terrestrial ecosystem 91 92 and fire impact is provided in the SI.

Present day contamination is the result of radionuclides released during the accident with a medium or long radioactive half-life [137 Cs ($T_{1/2} = 30.07 \text{ yr.}$), 90 Sr ($T_{1/2} = 29.14 \text{ yr.}$), 238 Pu ($T_{1/2} = 87.76 \text{ yr.}$), 239 Pu ($T_{1/2} =$ 24.13 10³ yr.), 240 Pu ($T_{1/2} = 6.57 \times 10^3 \text{ yr.}$), 241 Pu ($T_{1/2} = 14.35 \text{ yr.}$) and those arising as decay products of them: 241 Am ($T_{1/2} = 432 \text{ yr.}$, 237 Np ($T_{1/2} = 2.144 \times 10^6 \text{ yr.}$)]. Conversely to 137 Cs which spread and deposited all over Europe, 90 Sr, and Pu isotopes as nuclear fuel debris were mainly deposited in Ukraine and Belarus.³³ Cesium-137 is the preferred RN used in forecasting the long-term radiological consequences after an accidental release, because of its radiotoxicity, bioaccumulation, comparatively long half-life andstraightforward measurement procedure.

101 Once forests become contaminated with radiocesium, any further significant redistribution is limited. 102 Processes of small scale redistribution include resuspension, fire and erosion/runoff.³ According to the 103 International Atomic Energy Agency (IAEA), none of these processes are likely to result in any significant 104 migration of radiocesium beyond the location of initial deposition.³⁴ The question of radionuclide 105 redistribution may be a matter of interest for nearest adjacent areas that would have not been initially 106 contaminated or with a much lower contamination of their ecosystem. This gives interest for radionuclide 107 redistribution on a local scale and detection of trans-border fire plumes. Depending on the wildfire intensity, 108 smoke plumes may reach atmospheric layers up to several kilometres above ground level and travel for 109 thousands of kilometers¹, or be maintained in the atmosphere up to about 20 days.³⁵ Previous studies have 110 already brought some reassuring general information.³⁶ Over already contaminated areas the additional 111 contamination due to redistribution of artificial radionuclides by a fire remains low, about 1% of the previous inventory.³² Thanks to atmospheric dispersion and buoyancy effects, a sharp decrease in the airborne 112 113 concentration can be expected with distance (a dozen fold less at a distance of 100 m and thousands fold less at a distance of several kilometres from the fire line.^{32, 37} Regarding the redistribution or loss of RNs 114 115 from a burned area, it has been suggested that a wildfire outbreak might export at least 40% and up to 90% of the ¹³⁷Cs inventory.⁷⁻¹² In other words, this would mean that a fire could virtually "clean" an area from 116 117 ¹³⁷Cs. Current and historical research does not in any way support this proposition.

118 In several studies, it has been tried to determine the proportion of radionuclides that can be emitted into the 119 atmosphere from a zone with a given contamination density or emitted from burning material during a fire, 120 either through a resuspension factor K (m^{-1}) or an emission factor E_f (%). A compilation of both parameters 121 can be found in the SI. The UHMI calculated estimates of radionuclide emission into the atmosphere daily. 122 Their estimates relied on detailed satellite observations of the extent of the fires and combined the specific 123 contamination density of the burned areas, the vegetation type, and the specific radionuclide distribution in 124 the ecosystem. For the April 2020 fire event, the UHMI used a ¹³⁷Cs emission factor of 5% because of the 125 exceptional intensity of the crown fires. As of April 20, a total of 690 GBq of ¹³⁷Cs would have been released 126 into the atmosphere.²¹ The ¹³⁷Cs source term was later re-evaluated including emission from April 20 to 30, 127 leading to a release between 600 and 860 GBq for the CEZ, as well as 60 to 85 GBq for the Zhytomyr 128 region.³⁰ Additionally, 13.5 GBg of ⁹⁰Sr and 0.059 GBg of Pu isotopes were estimated by the UHMI to have 129 been released during fires. Using a similar approach (satellite observations of fire spots, burned areas, emission factors), Evangeliou & Eckhardt estimated that 341 GBq of ¹³⁷Cs, 51 GBq of ⁹⁰Sr, 2 GBq of ²³⁸Pu, 130 0.033 GBg of ²³⁹Pu, 0.066 GBg of ²⁴⁰Pu and 0.504 GBg of ²⁴¹Am were released between April 1 – 22, 131 132 2020.38

134 **Plume detection in Ukraine**

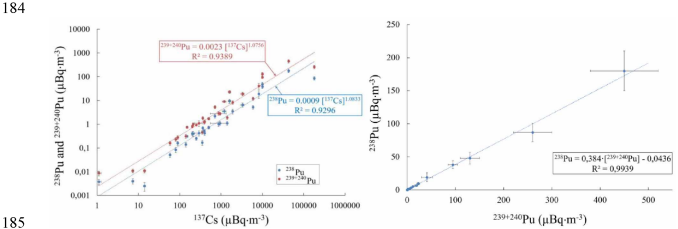
135 Plume detection was revealed by a 1,000 to 10,000-fold ¹³⁷Cs increase (i.e. up to a fraction of a Bq·m⁻³ in the CEZ and up to a fraction of a mBq·m⁻³ in the Kiev area) as compared to the usual 137 Cs average 136 137 background levels of about 3.5 mBg m^{-3} in the CEZ and 6 μ Bg m^{-3} in Kiev.³² Levels in excess of critical 138 threshold concentrations (i.e. a reference level of 0.21 mBq·m⁻³ for ¹³⁷Cs in Ukraine) were observed in 139 proximity to fire lines, owing to wildfire magnitude. High airborne contamination values were observed due 140 to the proximity of aerosol samplers with the fire lines (i.e. 42 mBq·m⁻³ on April 12 at the Korogodske forestry "square 11" station). An even higher ¹³⁷Cs value of 180 mBq·m⁻³ (0.18 Bq·m⁻³) was momentarily 141 reported on April 13 at the CNPP "Ukrenergomontazh - Open SwitchGear-750 point", at about 500 m from 142 143 the damaged reactor. Such orders of magnitude were similar to those reported during previous wildfire 144 outbreaks: 250 mBq·m⁻³ at the end of June, 2015 near the abandoned village of Polis'ke, 150 mBq·m⁻³ on 145 July 29, 2016 in the "Red forest", 25 mBq·m⁻³ on June 29, 2017 in the CEZ, too.^{39, 40} During the April 2020 146 fires, the UHMI estimated that the additional ground surface ¹³⁷Cs deposition in the CEZ peaked at about 147 $65 \text{ Bq} \cdot \text{m}^{-2}$ where the surface contamination was already in the MBq $\cdot \text{m}^{-2}$ to the tens of MBq $\cdot \text{m}^{-2}$ range (pers. 148 comm., O. Voitsekhovych, UHMI).

149 Strontium-90 was also detected in up to 20 locations, all of them in the highly contaminated area (see Table S7, SI). Based on about 40 pairs of ¹³⁷Cs and ⁹⁰Sr measurements, the ⁹⁰Sr /¹³⁷Cs activity ratio exhibited such 150 151 a significant variation (0.04 - 6.67) that it could not reasonably be represented by an aggregated statistic 152 (average value of 0.91). Hereafter, the highest ratio (6.67) was considered as an outlier and hence not taken 153 into account as representative of the whole fire situation. It corresponded to the pair of ¹³⁷Cs and ⁹⁰Sr values 154 that were by far the highest (as reported on April 13 from the OSG-750 station with 137 Cs=180 mBg·m⁻³ and 155 90 Sr=1,200 mBq·m⁻³) and because these concentrations were obtained over a very short period of time (~half an hour). 80% of the ⁹⁰Sr /¹³⁷Cs ratios were less than or equal to 1. Inversely, 20% of airborne ⁹⁰Sr 156 157 concentrations were higher than ¹³⁷Cs. This suggests that ⁹⁰Sr and ¹³⁷Cs biomass contamination levels and magnitude of the fires were much more likely to be of significance in the observed ⁹⁰Sr /¹³⁷Cs ratio. 158 159 Measurements performed using aerosol impactors (Yoschenko et al.) showed in general that ⁹⁰Sr bound to 160 coarse particles with an activity median aerodynamic diameter (AMAD) of > 25 μ m, while ¹³⁷Cs 161 predominantly bound to the finer aerosol fraction.³² Therefore, it is likely that the largest part of ⁹⁰Sr 162 emissions remained airborne for only 1 - 2 km. The distance between mobile aerosol sampling units and 163 the fire line thus participated to the variability of the 90 Sr / 137 Cs ratio. 164 Contrarily to 90 Sr which behaves like calcium (a factor of main influence on the plant physiology), 137 Cs is

- Contrarily to "Sr which behaves like calcium (a factor of main influence on the plant physiology),
- 165 much less bioaccumulated in wood .¹⁸ Recently, Holiaka et al.⁴¹ have reported a rather constant average ⁹⁰Sr

166 137 Cs ratio of about 2.5 in wood disks of Scots Pine stems sampled approximately 5 km north of the CNPP. 167 This value is clearly different from that found on average in aerosols (0.76, range 0.04 - 3.1) during the April wildfires. However, when looking into detail to the radial distribution of ¹³⁷Cs and ⁹⁰Sr in the wood 168 169 disks the authors concluded that, due to the year-by-year root uptake increase, a newly formed annual ring 170 receives a bigger amount of radiocesium than a ring formed in the previous years. In addition, ¹³⁷Cs is also translocated into these rings from older sapwood.⁴¹ Converselv, the radial distribution of ⁹⁰Sr shows 171 decreasing concentrations from heartwood to sapwood, i.e. decreasing ⁹⁰Sr concentrations from inner to 172 173 outer parts.⁴¹ The lower ⁹⁰Sr /¹³⁷Cs ratio observed in aerosols could thus be consistent with a partial combustion of aged tree trunks which remain charred after the fire goes through while the outer parts of the 174 175 tree (peripheral annual rings with a lower ⁹⁰Sr /¹³⁷Cs ratio) are burned.

177 In addition to ¹³⁷Cs and ⁹⁰Sr, about thirty measurements for both airborne ²³⁸Pu and ²³⁹⁺²⁴⁰Pu were reported 178 from the CEZ (see Table S7, SI). Maximum values for hourly measurements reached 180 µBq·m⁻³ for ²³⁸Pu 179 and 450 µBq·m⁻³ for ²³⁹⁺²⁴⁰Pu on April 12 at the Korohodske forestry monitoring station. The average 238 Pu/ $^{239+240}$ Pu ratio found was 0.40, which is slightly lower than the value of 0.48 - 0.50 reported during the 180 Chernobyl accident in 1986 or 0.47 found by Kashparov et al. ⁴²⁻⁴⁴ for the residual contamination of the 181 environment in 2000. The ²³⁸Pu/¹³⁷Cs, ²³⁹⁺²⁴⁰Pu/¹³⁷Cs and ²³⁸Pu/²³⁹⁺²⁴⁰Pu relationships shown in Figure 1 are 182 183 those that will be used hereafter in source term assessments.



185

Figure 1: (Left) Airborne ²³⁸Pu and ²³⁹⁺²⁴⁰Pu vs. ¹³⁷Cs concentrations in the CEZ, Ukraine, April 2020. 186 (Right) Airborne ²³⁸Pu concentration vs. ²³⁹⁺²⁴⁰Pu concentration in aerosols sampled in the CEZ, April 2020. 187 188

189 Except close to the fire spreading in the CEZ where the gamma dose equivalent rate was in the 0.5 -190 $30 \,\mu$ Sv·h⁻¹ range, the highest airborne concentrations measured elsewhere in Ukraine were not high enough 191 to significantly increase the ambient gamma dose equivalent rate. Daily average airborne ¹³⁷Cs 192 concentrations measured by the UHMI in Kiev reached 290 µBq·m⁻³ from April 8 to 9 and up to 700 µBq·m⁻³ 193 from April 10 to 11. The maximum observed average value during each daily sampling period remained far

below the National Radiation Safety Standards of Ukraine⁴⁵ establishing the allowed concentration of ¹³⁷Cs

195 to 800 mBq·m⁻³. Such concentration increases were concomitant with the smoke plume arrival in Kiev and

- have to be compared with the yearly average local background concentration of 6 μ Bq·m⁻³ (usual range 3 –
- 197 8 μBq·m⁻³).
- 198

199 Plume detection outside Ukraine

200 All the positive airborne ¹³⁷Cs measurements outside Ukraine during April 2020 were in the µBq·m⁻³ range, 201 i.e. within or just above the typical background ¹³⁷Cs level usually observed in the springtime (see Table S6, 202 SI). The reason for this background and the low-level airborne 137 Cs persistence is examined in the 203 Supporting Information. Considering the airborne ¹³⁷Cs background routinely observed and resulting from 204 resuspension on a local scale, it was difficult to assert if the measured concentrations in april 2020 included 205 a tiny and remote fire contribution. For instance, a weekly average value of $5.67 \pm 0.58 \,\mu\text{Bg}\cdot\text{m}^{-3}$ was 206 observed at Seehausen in north-eastern Germany (52.891 N; 11.729 E). This value is significantly higher 207 than the airborne level in the westernmost Europe. However, it remains in the usual range of variability as 208 a result of a comparatively more significant local Chernobyl deposition in 1986. In addition, Seehausen is 209 also known for ¹³⁷Cs resuspension during periods of dry meteorological conditions (Pers. Comm. A. 210 Dalheimer, DWD). Moreover, dispersion calculation did not reveal a noticeable transportation of airborne 211 137 Cs to Germany. It means that this location was not affected by the fire plume. In Norway, positive 137 Cs 212 detections were observed at the two northernmost sampling locations (Viksjofjell, 69.62 N; 30.72 E) and 213 Svanhovd (69.45 N ; 30.04 E) with very low ¹³⁷Cs concentrations ($< 1 \mu Bq \cdot m^{-3}$). However, because of a persistent snow cover prone to prevent ¹³⁷Cs resuspension from soil both on a local and regional scales, it 214 215 can be stated that airborne ¹³⁷Cs were mostly related to forest fires in Ukraine, as suggested by the 216 coincidence between simulation of plume arrival and sampling dates with higher than usual airborne ¹³⁷Cs 217 concentration (see Plume dispersion analysis section). Contrarily to what was first expected, in Eastern 218 Europe and relatively close to Ukraine, the fire plume, although more concentrated than in Western Europe, 219 did not ensue a significant ¹³⁷Cs increase above the usual local or regional ¹³⁷Cs background level which is 220 usually also higher than in Western Europe. At some Eastern Europe locations, the weekly average ¹³⁷Cs 221 activity may have hidden the peak value that would have been found if the sampling period had coincided 222 with the sampling duration. In such case, the atmospheric dispersion modeling is the only way to retrieve 223 this information (see *Plume dispersion analysis* section). For instance, in Budapest, it is estimated that the smoke plume arrived between April 5 and 6 and remained until April 9. A ¹³⁷Cs peak value of about 224 225 $25 \mu Bq \cdot m^{-3}$ was assessed. This peak cannot be foresee based on the weekly-average sampling value (12.8 μ Bq·m⁻³). In Poland, the estimated ¹³⁷Cs peak value was about 50 μ Bq·m⁻³ while the corresponding 226

227 weekly average value was at most about 6 μ Bq·m⁻³ which is not much higher than the usual ¹³⁷Cs background 228 level ranging from less than 1 μ Bq·m⁻³ to a few μ Bq·m⁻³.

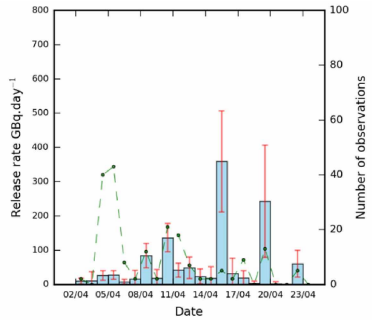
In France, the maximum weekly ¹³⁷Cs activity level reached $1.31 \pm 0.24 \mu$ Bg·m⁻³ from April 6 to 14, 2020 229 230 and was observed in the southeastern corner. It confirms model forecasts indicating that the highest value 231 occurred in that region during this period. The weekly average ¹³⁷Cs value measured during the fire event 232 was thus 4 to 8 fold higher than the ¹³⁷Cs background level in SE of France. The IRSN estimated that the 233 average ¹³⁷Cs level added by fires in Ukraine during the presence of the air mass in France was at most 2 234 μ Bq m⁻³ which is of no health concern for the public. For that purpose, the average regional background 235 137 Cs level (0.15 to 0.30 μ Bq·m⁻³) determined in a period ranging from March 15 to May 15 over the past 5 236 vears was removed for the remaining sampling period. Finally, the amount of 137 Cs corresponding to the 237 presence of the smoke plume was divided by the air volume filtered during its estimated presence. The 238 concentration of airborne ¹³⁷Cs measured in France in April 2020 had not been observed since 2002 (from 239 the end of August to the end of September 2002) when airborne 137 Cs concentrations increased up to 240 $3.5 \,\mu\text{Bq}\cdot\text{m}^{-3}$. This increase also resulted from wildfires raging in the Chernobyl area when for short periods 241 the usual prevailing westerly wind was not in place. Winds were in fact easterly during Week 36 and Week 242 38, 2002, leading to twin spikes in the concentration of airborne ¹³⁷Cs in France, Germany, Czech Republic, 243 and Austria. Outside the Chernobyl area, the highest airborne ¹³⁷Cs activity level was measured in Vilnius, 244 Lithuania with up to 196 µBq·m⁻³ during Week 36, 2002.¹³ At that time, there were also a lot of fires in the vicinity of Vilnius,⁴⁶ which were responsible for an increase in PM_{10} up to 370 µg·m⁻³. Due to a higher 245 246 water-soluble ¹³⁷Cs percentage, Lujaniene et al. (2006) concluded that these particulate matters were transported to Lithuania from forest fires occurring in Ukraine and Belarus.¹³ Rising ¹³⁷Cs activity 247 248 concentrations from smoke are due to both the enhancement of the airborne dust load, acting as ¹³⁷Cs carrier, 249 but also to the fact that smoke is significantly rich in ¹³⁷Cs.³¹

250 We took the opportunity of this event to check if other radionuclides emitted into the atmosphere by 251 wildfires might be used, in combination with ¹³⁷Cs, as a tool to attest the contribution of the wildfire plume 252 far from the Chernobyl area. Strontium-90, plutonium isotopes, and ²⁴¹Am might be candidates. However, 253 their tedious radiochemistry complicates their determination and their much lower expected airborne 254 concentrations as compared to ¹³⁷Cs usually requires to gather together several weekly filters to exceed the 255 detection limit in the composite sample. These RN are thus difficult to quantify above detection limits on a 256 weekly aerosol-sampling basis. Looking for a more convenient-to-measure radionuclide whose activity ratio with ¹³⁷Cs might be relevant for wildfire plume detection at long distances, we also examined the airborne 257 258 ⁴⁰K concentrations (See SI).

259

260 Source term assessment methodology

261 The UHMI performed RN source terms assessment based on environmental parameters such as satellite 262 observations of burned areas, biomass density, and contamination density maps. Another way to assess 263 radionuclide amounts emitted from the burned areas is to apply inverse modeling techniques combining 264 atmospheric transport model and observed airborne concentrations. Such methodology was implemented 265 using the comprehensive airborne ¹³⁷Cs dataset acquired on the European scale (see Table S6, SI) to estimate 266 first the amounts of ¹³⁷Cs emitted into the atmosphere between April 2 and April 24, 2020. The method is 267 described in the SI. Twenty-two different daily releases were estimated by inverse modeling between April 268 2 and 24, 2020. Although satellite images after April 24 indicate a persistent residual fire, releases were 269 assumed insignificant after that date. In addition, the available measurements after April 24 are clearly not 270 sufficient for an inverse modeling estimation of the source term. Our inverse modelling is based on a 271 variational approach which consists in the minimization of a least-squares cost function assuming log-272 normal observations errors and wihout considering any additional background term (see SI). Performing 273 Monte-Carlo simulations, 15,000 different source terms were computed in order to take into account 274 uncertainties resulting both from dispersion modeling, meteorological fields and representation errors of observations. For instance, this analysis indicates that the estimated ¹³⁷Cs source term for the entire fire 275 276 period lies between 700 and 1,200 GBq. This range reflects the set of all the above mentioned uncertainties. 277 From April 2 to 15, our results indicate daily ¹³⁷Cs emissions ranging from 10 to 138 GBq (Figure 2). For 278 this period, the source term estimation proves to be fairly robust since the sensitivity to observation 279 perturbations is weak. The large number of observed values considered in the inverse modeling process 280 reasonably explains the standard deviation on daily release rates remaining low. From April 15, the 281 magnitude of the daily release rates varied significantly. The maximum daily release rate reached 362 GBq 282 on April 16 and 241 GBq on April 20 thus corresponding to the highest releases estimated during the entire 283 fire period. However, the standard deviations calculated for April 16 and 19 are large. This increases the 284 source term margin of error and leads to a higher level of uncertainty.



Date Figure 2: April 2 to 24 ¹³⁷Cs average daily release rates reconstructed by Monte Carlo analysis using n= 15,000 samples (blue rectangles) and the associated standard deviations (orange bars). The green dashed line represents the number of observations used for each daily release assessed by inverse modeling.

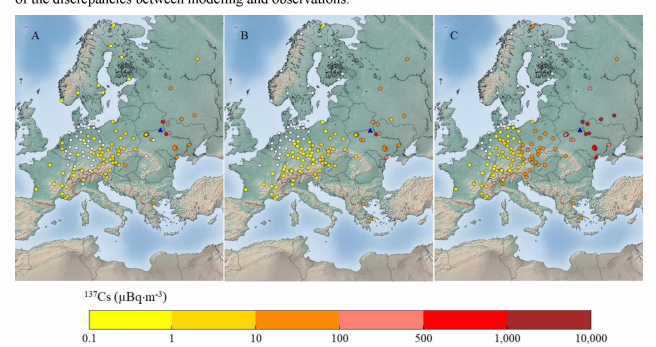
290 **Plume dispersion analysis**

291 The video of the plume dispersion simulation is available in the SI. An average source term was deduced 292 from the Monte Carlo analysis as an input parameter. The smoke plume first went in the direction of the 293 Russian Federation between April 2 and 3. From April 3 to 5, it moved to the south of Ukraine (including 294 the Kiev region). From April 5 to 7, the plume continued to move towards both the west and south. It reached 295 Romania, Hungary, the Czech Republic, eastern Poland, Austria and Slovenia during this period. The 296 simulated hourly ¹³⁷Cs concentrations are about 10 - 50 µBq·m⁻³, sometimes higher in eastern Romania and much lower to the west. From April 7, according to the model, there were very low ¹³⁷Cs concentrations 297 298 (just above 1 μ Bq·m⁻³) in southern Germany, France and Italy. These levels are close to the detection limits 299 (DL). The model provides an explanation as to why the vast majority of sampling stations located in this 300 geographical area did not report any concentration higher than DL. Furthermore, even in the case of a 301 measurement > DL, as for instance in Austria and Poland, these measurements did not vary from the usual 302 seasonal values. Modeling still tends to emphasize a contribution in ¹³⁷Cs concentrations from the fire, 303 especially in the Czech Republic, Austria, Italy, Slovenia and France. Between April 8 and 10, a new plume 304 reached the south of Ukraine again. The plume extended to Greece as of April 11. The hourly-simulated 305 concentration reached up to $100 \mu Bq \cdot m^{-3}$ in the Thessaloniki area.

This estimate is fully consistent with measurements performed in Thessaloniki between April 11 and 13 with an average concentration of 25.5 μ Bq·m⁻³ and a residence time of 3.5 ± 0.2 d.⁴⁷

308 The wind direction changed again between April 12 and 13. Consequently, the wind blew towards Russia. 309 This change in direction corresponded to the most significant release period when the hourly simulated 310 concentrations exceeded 100 uBq·m⁻³ at the IMS Russian station in Dubna. The plume then moved northwest, to the northernmost part of Norway. The simulation shows hourly values up to $30 \mu Bq \cdot m^{-3}$ over 311 312 this area on April 14. The plume passed through this region relatively quickly, in about a few hours. This 313 explains why the weekly sampling carried out in Svanhovd showed little variation compared to weekly 314 averages with a concentration of only $0.5 \,\mu\text{Bq}\cdot\text{m}^{-3}$. This value is likely the result of fires being controlled. 315 During this period, very low weekly average ¹³⁷Cs concentrations were reported in Western Europe (about 316 $1 \mu Bq m^{-3}$). From April 15 to 19, the wind mainly blew east and southeast. Several slightly contaminated 317 air masses mainly affected the Kiev region and the more southern regions of Ukraine. On April 21, the 318 plume reached Greece again. The simulation consistently matched measurements carried out in Thessaloniki 319 $(9.6 \pm 0.8 \mu \text{Bq} \cdot \text{m}^{-3})$ on April 21 with an estimated residence time of $11 \pm 3 \text{ d}^{.47}$

55% of the simulated ¹³⁷Cs concentrations were within a factor of 2 compared to the observed concentrations 320 321 (see Table S5, SI). This score provides significant validation of the reconstructed source term (see SI). 322 Maximum simulated and observed ¹³⁷Cs concentrations are compared from April 2 to 24 (Figure 3). The 323 observed and simulated ¹³⁷Cs maximum concentrations (maps A and B) are very similar. The maximum 324 simulated concentrations in Ukraine, Belarus and the Russian Federation are the highest and are consistent 325 with the maximum concentration levels reported in these countries. Further west, the correlation between 326 simulated and observed concentrations is also satisfactory, although the model tends to underestimate the 327 maximum observed concentrations in Austria. In any case, the maximum concentrations measured in this 328 area are low and the usual ¹³⁷Cs background level would have to be rigorously taken into account, if known, 329 in order to make a proper comparison between observed concentrations and simulated concentrations added 330 by the plumes. As a result of the agreement between simulations and observations, the hourly maximum simulated ¹³⁷Cs concentration was derived at each sampling location (Figure 3, map C). Due to the spatial 331 332 resolution of the long-range dispersion model, values obtained at less than 50 km from the CNPP were not 333 taken into account because of the significant associated uncertainties. The maximum hourly concentrations 334 simulated in Eastern Ukraine, the western part of the Russian Federation and southern Belarus were above 335 $1 \text{ mBq} \cdot \text{m}^{-3}$ (Figure 3, map C). Simulated concentrations then gradually decreased to the west. They were 336 above 10 µBq·m⁻³ over a central band stretching from northern Norway to southern Greece. Further west, 337 concentrations were even lower but still above 1 µBq·m⁻³ in southern France, Switzerland, western Austria 338 and Germany. This confirms that the ¹³⁷Cs concentrations measured in these countries were partly due to 339 the influence of the remote fire despite values being very close to observed seasonal measurements. However, in Belgium and in the Netherlands, unusual ¹³⁷Cs concentrations (5.3 and 1.59 μ Bq·m⁻³, respectively) were locally reported, that both our dispersion simulation and that of SCK-CEN cannot reproduce. It has been suggested that a local/regional simultaneous resuspension event might be the reason of the discrepancies between modeling and observations.

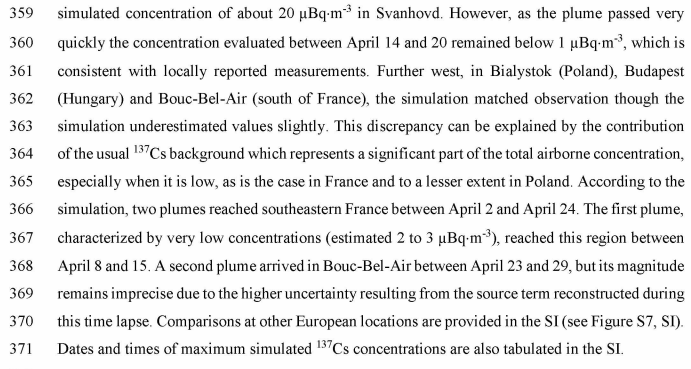


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Figure 3: ¹³⁷Cs airborne concentration maps (μ Bq·m⁻³) over the course of the April, 2020 wildfire event in Ukraine. (A) Maximum observed concentrations; (B) Maximum simulated concentrations; (C) Maximum values based on hourly simulated concentrations (i.e. maximum time-resolved peak concentrations). The same color scale applies to all maps.

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350 According to Figure 4, the plume passed through Ukraine several times. At the Baryshivka station, 351 hourly simulated concentrations exceeded 1 mBq·m⁻³ between April 17 and 20. In Greece, fire 352 plumes reached the Thessaloniki area (northern Greece) on three occasions. The most significant episode around April 13 in this area was characterized by hourly simulated concentrations up to 353 354 100 μ Bq·m⁻³. The average ¹³⁷Cs estimated concentration over the air sampling period (20 to 30 μ Bq·m⁻³) remains consistent with measurement taken in Thessaloniki (25.5 μ Bq·m⁻³). Further 355 south, in Athens, no observable 137 Cs activity concentration (> 3 µBq·m⁻³) was determined on three 356 consecutive filters sampled from April 3 - 15. The geographical extent of the plume was very large 357 358 as it reached as far as the northernmost part of Norway for a few hours on April 15 with a peak



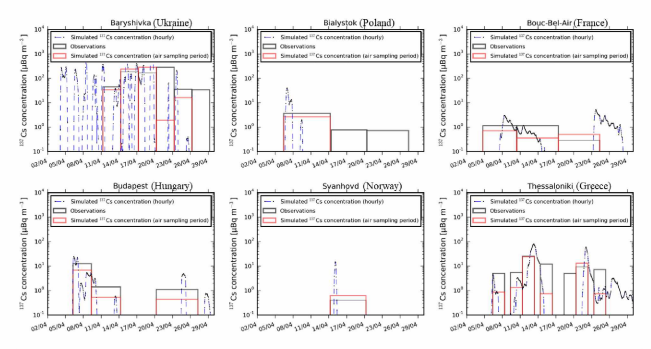




Figure 4: Example of comparisons between simulated and observed airborne ¹³⁷Cs concentrations at several European locations. The grey line shows the observed concentrations for each sampling period and the red line the simulated concentrations. The blue dashed line represents the simulated hourly concentrations.

378

379 ⁹⁰Sr, ²³⁸Pu, ²³⁹⁺²⁴⁰Pu emissions

380 Because of the facility researchers have measuring 137 Cs at trace levels using γ -spectrometry, most 381 investigations and reported data during this event focused on airborne ¹³⁷Cs as a tracer of biomass burnings, 382 both on a nation-wide scale (i.e. in Ukraine) and on a continental scale. However, owing to their respective 383 dose coefficients, there is much concern about the emission by wildfires of radionuclides such as ⁹⁰Sr, Pu 384 isotopes, and ²⁴¹Am. Strontium-90 as well as ²³⁸Pu and ²³⁹⁺²⁴⁰Pu were measured only inside the CEZ but not 385 on a wider scale. This prevented the use of the inverse methodology described for ¹³⁷Cs to assess: 1) the 386 corresponding source terms 2) resulting airborne concentrations elsewhere in Ukraine and in the rest of 387 Europe and 3) internal exposure. In France, all the filters taken at sampling stations that were assumed to be 388 in the path of the fire plume according to the dispersion analysis, in addition to a slightly higher observed 389 137 Cs concentration than normal, were gathered in a composite sample representing 365,775 m³. The 390 following analyses were performed on this composite sample: α-spectrometry (²³⁸Pu, ²³⁹⁺²⁴⁰Pu, ²⁴¹Am), ICP-391 MS (²³⁹Pu and ²⁴⁰Pu separately), proportional couting (⁹⁰Sr). Results for ²³⁸Pu ($0.12 \pm 0.07 \text{ nBq} \cdot \text{m}^{-3}$) and 392 $^{239+240}$ Pu (3.43 ± 1.09 nBq·m⁻³) were in the usual background ranges reported in France over the last decade $(0.03 - 0.42 \text{ nBq} \cdot \text{m}^{-3} \text{ for } ^{238}\text{Pu} \text{ and } 0.10 - 2.85 \text{ nBq} \cdot \text{m}^{-3} \text{ for } ^{239+240}\text{Pu})$. Stontium-90 remained below a decision 393 394 threshold of 62 nBq·m⁻³. As previously mentioned (see section *Plume detection in Ukraine*), emitted ⁹⁰Sr 395 bound in general to the coarse aerosol fraction and is found to travel only short distances from a fire. 396 Conversely, Yoschenko et al. demonstrated that transuranic elements which are in general bound to the fine 397 aerosol fraction, are prone to travel much greater distances from the emission point.³² Because of a high (60%) relative uncertainty associated to the ²³⁸Pu result, the ²³⁸Pu/²³⁹⁺²⁴⁰Pu activity ratio observed in France 398 399 was not appropriate to discriminate a possible Chernobyl signature. However, the ²⁴⁰Pu/²³⁹Pu mass ratio 400 (0.25 ± 0.07) was slightly higher than the average value of 0.176 ± 0.03 usually observed in France during 401 the last decade and which is typical of global fallout. Given the 0.41-0.42 typical signature of the Chernobyl 402 fallout, this intermediate value could correspond to the added contribution following the smoke plume 403 arrival in France in April.

To cope with the lack of large scale Sr and Pu results during this event we used the generally accepted Chernobyl-ratios between the above-mentioned radionuclides and airborne ¹³⁷Cs, and assumed these ratios to be representative of the emissions. Given the internuclide relationships derived from previous measured concentrations in the CEZ⁴⁸ and taking into account the previously estimated ¹³⁷Cs source term range (700 -1,200 GBq), we considered the uncertainty associated with the relationships established between ⁹⁰Sr and ¹³⁷Cs on one hand and between ^{238, 239+240}Pu and ¹³⁷Cs on the other hand (Table 1).

41 ITable 1: Estimations of Chernobyl-labelled radionuclides emissions (in GBq) during the April 2020 wildfires in Ukraine. Values for ¹³⁷Cs, ⁹⁰Sr, ²³⁸Pu and 412³⁹⁺²⁴⁰Pu are derived from field measurements. Those for ²³⁹Pu, ²⁴⁰Pu, ²⁴¹Pu and ²⁴¹Am are deduced from typical activity or isotopic ratios characteristic 4136 the Chernobyl accident. The grand total source term (¹³⁷Cs + ⁹⁰Sr + ²³⁸Pu + ²³⁹Pu + ²⁴⁰Pu + ²⁴¹Pu + ²⁴¹Pu + ²⁴¹Am) has been evaluated from values in bold characters.

, in the second s		This study						Other studies				
			R ² coef.	Estimated	Rounded	Ratio to		Protsak	Tabachnyi	Talerko		Evangeliou
		Relationships used		emission	average	¹³⁷ Cs	total ST	et al. 21	et al. 30	et al. 49	et al. 50	et al. 38
Radionuclide				range (GBq)	(GBq)	(%)	(%)	(GBq)	(GBq)	(GBq)	(GBq)	(GBq)
¹³⁷ Cs	(M)			700 – 1,200	950	100	62.2	690	660 - 945	574	220 -1,810	341
⁹⁰ Sr	(M)	${}^{90}\text{Sr} = 0.3313 \; [{}^{137}\text{Cs}]{}^{1.061 \pm 0.030}$	0.8318	346 - 613	480	50.4	31.5	13.5	n.d.	n.d.	n.d.	51
²³⁸ Pu	(M)	${}^{238}\mathbf{Pu} = 0.0009 \; [{}^{137}\mathbf{Cs}]{}^{1.0833}$	0.9296	1.1 – 1.9	1.5	0.16	0.1	n.d.	n.d.	n.d.	n.d.	2
²³⁹⁺²⁴⁰ Pu	(M)	$^{239+240}$ Pu = 0.0023 [137 Cs] $^{1.0756}$	0.9389	2.6 - 4.7	3.7	0.39	0.2	n.d.	n.d.	n.d.	n.d.	0.099
²³⁸ Pu+ ²³⁹⁺²⁴⁰ Pu	(M)	^{238,239+240} Pu = 0.0032 [¹³⁷ Cs] ^{1.0771}	0.9375	3.7 - 6.7	5.2	0.55	0.3	0.059	n.d.	n.d.	n.d.	2.1
²³⁹ Pu	(D)	239 Pu/ $^{239+240}$ Pu = 0.403 (*)		1.1 - 1.9	1.5	0.16	0.1	n.d.	n.d.	n.d.	n.d.	0.033
²⁴⁰ Pu	(D)	240 Pu / 239 Pu (activity) = 1.496 (**)		1.6 - 2.9	2.2	0.23	0.1	n.d.	n.d.	n.d.	n.d.	0.066
²⁴¹ Pu (2020)	(D)	241 Pu/ $^{239+240}$ Pu (activity) = 19.1		50.5 - 90.1	70.3	7.4	4.6	n.d.	n.d.	n.d.	n.d.	0.000
²⁴¹ Pu (2020)	(D)	241 Pu/ 239 Pu (activity) = 39.75		42.3 - 75.6	59.0	6.2	3.9	n.d.	n.d.	n.d.	n.d.	n.d.
²⁴¹ Am (2020)	(D)	241 Am / 241 Pu = 0.328		13.9 - 29.6	21.7	2.3	1.4	n.d.	n.d.	n.d.	n.d.	0.504
¹³⁷ Cs+ ⁹⁰ Sr+ΣPu+ ²⁴¹ Am				1,114 – 1,939	1,530		100					

414 (M): measured, (D): deduced, ST: Source Term, n.d.: not determined

415 (*) Values taken from ref.⁴⁸ for the organic layer of the surface soil (litter and humus).

416 (**) derived from the low-uncertainty 240 Pu / 239 Pu (isotopic) ratio 0.408 ± 0.003 obtained by Muramatsu et al.⁴⁸ in the Chernobyl environment.

417

Our ¹³⁷Cs source term estimation is fully consistent with that proposed by Ukrainian researchers: 690 GBq 418 as of April 20 or 660 to 945 GBg until the end of the blaze event.^{21, 30} Theses estimates are based on a very 419 420 different method combining land cover and vegetation features, radionuclide distribution in the ecosystem, 421 biomass burning emission factors, and fire satellite detections. Evangeliou & Eckhardt indicated that their 422 model underestimated measurements by about 70%, which means that in their study the average modeled 423 concentration was almost half of the average measured concentration.³⁸ Based on observed airborne 424 concentrations, our estimated average ⁹⁰Sr emission (480 GBq) is much higher than those estimated by Protsak et al. (13.5 GBq) and Evangeliou & Eckhardt (51 GBq). These two ⁹⁰Sr assessments combined 425 426 estimations of the residual ⁹⁰Sr biomass contamination, satellite detections and used a ⁹⁰Sr emission factor 427 of 0.2% (i.e. 25-fold lower than for ¹³⁷Cs, based on their research). Our ²³⁸Pu estimate is also consistent with that of Evangeliou & Eckhardt.³⁸ However, the ²³⁸Pu/²³⁹⁺²⁴⁰Pu activity ratio (~20) indicated by these 428 429 authors³⁸ represents a serious discrepancy and is inconsistent with the characteristic Chernobyl Pu isotopic 430 signature of 0.48 or with the value of 0.47 typical of the residual contamination observed in 2000 in Ukraine.^{42, 44} In our study, the relationship obtained between 238 Pu and $^{239+240}$ Pu can be expressed by 238 Pu = 431 $0.384[^{239+240}Pu]-0.0436$ (R² = 0.9939). The slope of the $^{238}Pu/^{239+240}Pu$ adjustement (0.384) fits the one 432 433 characteristic of the Chernobyl accident (0.4 - 0.5).

434

435 ²³⁹Pu and ²⁴⁰Pu emissions

436 Distinct determinations of ²³⁹Pu and ²⁴⁰Pu were not reported. Both Pu isotopes behave similarly in the 437 environment and have a much longer half-life as compared with the period of time that went by since the 438 accident, thus we can neglect their differential decay. We can assume that the ²⁴⁰Pu/²³⁹Pu ratio at the emission to be the same as in soil or in the biomass. Muramatsu et al.⁴⁸ indicated a relative consistency of 439 440 the mass (or atom) ratio (average 0.408 ± 0.003 , range 0.386 - 0.412) regardless of the ²³⁹⁺²⁴⁰Pu 441 concentrations range in surface soil samples (i.e. organic rich lavers). In some moss and soil samples of 442 Chernobyl, Jakopic et al.⁵¹ reported mass ratios of 0.3624 ± 0.0011 and 0.4140 ± 0.0035 , respectively. In forest soil, a mass ratio ranging from 0.186 to 0.348 was found.⁵² We retain an average ²⁴⁰Pu/²³⁹Pu mass 443 ratio of 0.41 which once expressed in ²⁴⁰Pu/²³⁹Pu activity ratio corresponds to a value of 1.50, close to the 444 445 characteristic activity ratio (1.40) that can be deduced from the Chernobyl releases (see Table S1, SI). Plutonium-239 can also be expressed with reference to $^{239+240}$ Pu with a typical activity ratios (239 Pu/ $^{239+240}$ Pu) 446 of 0.403.⁴⁸ Based on our ²³⁹⁺²⁴⁰Pu emission estimate (2.6 – 4.7 GBq) we derived a ²³⁹Pu emission between 447 448 1.1 and 1.9 GBg and a ²⁴⁰Pu emission between 1.6 and 2.9 GBg.

449

450 ²⁴¹Pu and ²⁴¹Am emissions

Unfortunately, for the sake of study, no ²⁴¹Pu or ²⁴¹Am determination was reported for that event. Plutonium-451 452 241 releases during the Chernobyl accident ($\sim 6 PBq$) was the largest contributor to the total plutonium released amount (see Table S1, SI). Despite its rather short half-life ($T_{1/2} = 14.4$ yr.), ²⁴¹Pu still denotes a 453 454 significant reservoir of environmental radioactivity. However, as a pure β emitter (maximum energy of only 455 20.8 keV) ²⁴¹Pu represents a lesser radiological risk compared with other α -emitting plutonium isotopes 456 (except via its decay product ²⁴¹Am). Since the Chernobyl accident, only 19.1% of the ²⁴¹Pu amount released in 1986 is still present in the environment. It is possible to assess the ²⁴¹Pu emission by the April 2020 457 458 wildfires using the original 241 Pu/ $^{239+240}$ Pu activity ratio and applying a simple 241 Pu decay. This ratio was 459 estimated for May, 1986 in a range between 70 and 100.43, 53-57 or even in a range of 82 to 120 in upper parts of lichens, 98 in air filter (May 1, 1986), 95 in grass (May 1, 1986).⁵⁶ Considering a global average ratio of 460 100 at the time of the accident and neglecting ²³⁹Pu and ²⁴⁰Pu decays as of April 2020, the ²⁴¹Pu/²³⁹⁺²⁴⁰Pu 461 462 activity ratio would have been about 19.1 during the wildfire event. Given the previously estimated $^{239+240}$ Pu emission (2.6 – 4.7 GBq) results in a ²⁴¹Pu emission between 50.5 and 90.1 GBq. Another estimation of the 463 ²⁴¹Pu emission can be performed using the initial ²⁴¹Pu/²³⁹Pu mass ratio of 0.123 ± 0.007 (ref.⁵⁵) for May 464 465 1986 and confirmed by the value of 0.0384 ± 0.0022 determined for 2009.⁵¹ Once expressed in activity ratio 466 and ²⁴¹Pu-decay corrected to date, the ²⁴¹Pu/²³⁹Pu activity ratio is 39.75. Considering the above estimated 467 ²³⁹Pu emission range (1.1 - 1.9 GBg) leads to a ²⁴¹Pu emission ranging 42.3 - 75.6 GBg, consistent with the above-mentioned estimation based on the ${}^{241}Pu/{}^{239+240}Pu$ ratio (50.5 – 90.1 GBq). 468

469 Most radiological concern comes from ²⁴¹Am. This nuclide results from ²⁴¹Pu decay and is characterized by 470 a much longer half-life ($T_{1/2} = 432.7$ yr.), a much higher radiotoxicity (as an α - γ emitter), and a higher 471 environmental mobility than its parent.⁵³ As a result, the in-growth of ²⁴¹Am from ²⁴¹Pu decay exhibits an increasing radiation risk with time. By 2058, the ²⁴¹Am activity will exceed that of plutonium isotopes by 472 473 about a factor of two.⁵⁸ In addition, its inhalation effective dose coefficient is about 245 fold higher than for 474 ²⁴¹Pu. A deeper insight into the differential soil-to-plant transfer of both Pu and Am is necessary to assess the ²⁴¹Am emission from that of ²⁴¹Pu. For both radionuclides, the downward migration in soil is expected 475 476 in the same range $(0.1 - 0.5 \text{ cm} \cdot \text{yr}^{-1})$. However, a 1.5 fold higher migration velocity has been reported for 477 ²⁴¹Am as compared with Pu in the Chernobyl-contaminated environment of Belarus characterized by sandy 478 soil of various types (soddy podzolic sand, loamy sand and peat bog).⁵⁸ Despite a high variability induced by both plant species and soil structure and composition, both concentration factors (Cf_{Am} and Cf_{Pu}) between 479 480 soil and plant (root uptake path) remain much lower (2 to 3 order of magnitude less) compared to ¹³⁷Cs and even more so to 90 Sr. ${}^{54, 58, 59}$ As a rule of thumb, the 241 Am soil-to-plant concentration factor (Cf_{Am}) exceeds 481 that for Pu.^{58, 60} Sokolik et al.⁵⁸ proposed an average Cf_{Am} to Cf_{Pu} ratio of 2.2 (1 – 6 range) for meadow 482 483 grasses. Other determinations indicating a higher Cf_{Am} to Cf_{Pu} ratio were reported: about a factor of 2 in 484 crops⁶¹ and 2.3 (1.6 - 3.3 range) in bilberry and lingonberry plants.⁵⁹ Concentration factors from forest litter 485 to spruce and pine needles, to pine root and stem, to spruce bark, to fern, alder and heather have been 486 summarized in Mietelski et al.⁵⁴ with a similar average of 2.6 (0.3 - 5.7 range). Based on the above-mentioned 487 values and as a conservative approach, we considered hereafter a differential soil-to-plant transfer between ²⁴¹Am and ²⁴¹Pu (Cf_{Am}/Cf_{Pu}) of 2.4. In the same period of time (34 yr.) since the Chernobyl accident, the 488 ingrown ²⁴¹Am represents 2.6% of the ²⁴¹Pu initially released. We can neglect the ingrowth of ²⁴¹Am stored 489 490 in the burnable biomass because 90% of the Pu is stored in the litter with a degradation rate of about a few 491 years.³² As of April 2020 the theoretical activity ratio ²⁴¹Am/²⁴¹Pu from Chernobyl was thus about 13%. Multiplying this ratio by the above-mentioned Cf_{Am}/Cf_{Pu} ratio of 2.4 leads to a ratio of 0.328 between the 492 493 amounts of ²⁴¹Am and remaining ²⁴¹Pu emitted into the atmosphere in April 2020. To cope with the lack of 494 experimental determination the same biomass burning emission factor (i.e. 1%) can safely be considered 495 for both RN since Pu and Am have a much higher boiling point as compared with the maximum expected 496 fire temperature. In such a case, their emission during wildfires is assumed to be mainly limited through the 497 uplift of ashes and not through the gas phase.^{5, 62} The higher ²⁴¹Am release potential thus likely arises from its higher upstream soil-to-plant transfer. The slightly higher ²⁴¹Am release potential than for Pu is validated 498 499 by the experimental determination of the resuspension factor by wildfires (K) (see Tables S2 and S3, SI).^{62,} ⁶³ In short, given the overall range estimated for the ²⁴¹Pu emission (42.3 - 90.1 GBq) results in an ²⁴¹Am 500 501 emission estimation between 13.9 GBq (42.3 GBq x 0.328) and 29.6 GBq (90.1 GBq x 0.328).

502 The particular case of storage facility risks in the vicinity of the CNPP are detailed in the information notes 503 published by the IRSN.²⁵⁻²⁸ The Ukrainian authorities indicated that the spent fuel storage facility N1 (which 504 contains spent fuel from the Chernobyl NPP decommissioned in 2000) was located in the immediate area 505 of the NPP, thus safe from the fires. The spent fuel storage facility N2 located in the CEZ contains empty 506 tanks made of huge reinforced concrete structures surrounded by a fence. The authorities also said that the 507 Pidlisne storage facility, made of fireproof reinforced concrete structures and used to store nuclear waste, 508 was also located in a place at low risk to fires. As a precaution, the forest was cut down around the storage 509 facility to avoid the threat of fire, and the distance from green spaces was more than 100 meters.⁶⁴ In addition 510 to waste managed in the industrial zone around the sarcophagus and engineered disposal sites, non-511 engineered near-surface trench dumps were used as waste repositories. In the early 2000's, ²⁴¹Pu was responsible for approximately 20% of the overall activity, while ²⁴¹Am and ²³⁸Pu were responsible for 45% 512 513 and 15% of the total α -activity, respectively.⁶⁵ Considering the numerous waste disposal sites that contain 514 large amounts of radionuclides and that are in the immediate proximity to the CNPP, it is likely that their 515 radionuclide content was spared from fire as previously mentioned.⁶⁶ Indeed, their soil cover naturally protects them. However, vegetation can grow in this soil, uptake radionuclides, and as a result, become 516 highly contaminated.⁶⁷ After the trenches were covered with soil, some of them were also planted with small 517 518 pine trees.⁶⁵ During the course of the trees' growth, partial uptake of the radionuclides buried in the trenches 519 occurs. These radionuclides are then distributed between the trees and the topsoil through litterfall. Waste 520 trenches are also periodically flooded by groundwater. As a result of a solution pH in the 4.8 - 5.1 range, 521 radionuclide dissolution of fuel particles and migration of the corresponding radionuclides can be more effective in their proximity.⁶⁸ Thirv et al.⁶⁷ estimated that the maximum ⁹⁰Sr transfer will be reached 40 522 523 years after planting with 7% of the total ⁹⁰Sr content in the trench being transferred to the trees, and 12% to 524 the forest litter. According to the transfer calculation conducted by Thirv et al.⁶⁷ on waste buried (trenches), 525 the transfer of ⁹⁰Sr from soil to tree and litter has been estimated to be at most 7% and 12%, respectively 526 thus 19% for the burnable biomass. The ⁹⁰Sr transfer is thus 216 fold more efficient than that of ¹³⁷Cs,⁶⁷ resulting in a maximum ¹³⁷Cs transfer of only 0.09%. Recent work by Kashparov et al. confirms that the 527 activity of the mobile form of ⁹⁰Sr in the trench has presently reached its maximum value.⁶⁹ This results 528 529 from the decrease of the amount of remaining nuclear fuel particles not yet dissolved in topsoil and the 530 reduction of the ⁹⁰Sr soil-to-plant transfer due to its radioactive decay. Regarding Pu, it can be emphasized 531 that, unlike ¹³⁷Cs and ⁹⁰Sr, the transfer of plutonium to plants is extremely low; this element is therefore 532 present at only trace amounts in forest organic matter and remains fixed in the mineral soil fraction 533 contaminated in 1986. Yoschenko et al. estimated that its transfer to trees plus litter was 3.5 fold lower than for ¹³⁷Cs.³² Using this ratio we can estimate a Pu concentration factor from soil to plant + litter burnable 534 535 biomass (Cf_{Pu}) of 0.026%. Taken this factor and considering the previously mentioned ratio Cf_{Am} / Cf_{Pu} of 536 2.4 results in a transfer factor for 241 Am (Cf_{Am}) of 0.0624%. In short, considering the biomass emission 537 factor of 4% (for ¹³⁷Cs and ⁹⁰Sr) and 1% (for Pu isotopes and ²⁴¹Am), the proportion of radionuclides emitted 538 into the atmosphere by the wildfires as compared to their trench content can be estimated at 0.036‰, 7.6‰, 539 0.0026‰ for ¹³⁷Cs, ⁹⁰Sr and Pu isotopes, respectively. The volume of waste in the replanted trenches is not precisely known. The trenches are scattered over a total area of 450 ha in the CEZ (around 10⁶ m³).⁶⁵ A 540 541 more refined estimate of the potential radionuclide emissions could be performed based on the buried waste 542 amount and existing biomass on their surface. However, such specific estimates require further investigation 543 which is beyond the scope of this study.

544 **Dose assessment**

A dose assessment was performed on two categories of people: firefighters who took part in firefighting in the CEZ and inhabitants of Kiev at about 100 km from the CEZ. Firefighters have already been identified at risk in the event of a forest/bush fire over the dumps (storage trenches), as their occupational dose may exceed the constraint of $0.3 \text{ mSv} \cdot \text{yr}^{-1.65}$ Their inhalation dose assessment was performed taking deliberately a conservative approach with the followings assumptions: 1) the firefighters had no respiratory protection (as part of Personal Protective Equipment, PPE) such as a self-contained breathing apparatus (SCBA) or a facemask, 2) all aerosol sizes resulting from wildfires were belonging to the respirable fraction (<10 μ m), 552 3) a total working time in the exclusion zone of a hundred hours for each of them (10 days with 10 working hours per day), 4) a breathing rate of $3 \text{ m}^3 \cdot \text{h}^{-1}$ corresponding to a very heavy exercise.⁷⁰ The airborne 553 concentrations considered were based on field measurements where ¹³⁷Cs, ⁹⁰Sr, ²³⁸Pu and ²³⁹⁺²⁴⁰Pu were 554 simultaneously measured (see Table S7, SI). Detailed estimations for ²³⁹Pu, ²⁴⁰Pu, ²⁴¹Pu and ²⁴¹Am (both not 555 measured) were derived from measured ¹³⁷Cs concencentrations and Pu isotopic ratio according to the 556 557 relationships mentioned in Table 1. Two possibilities were considered for the measured values: maximum 558 observed or average values. The first assumption, leading to airborne concentrations of 1 $Bq \cdot m^{-3}$ (rounded values) for both ¹³⁷Cs and ⁹⁰Sr, and 1 mBq·m⁻³ for ²³⁸Pu, is very pessimistic since the highest measured 559 560 concentrations at similar orders of magnitude were only reported during short periods of time (i.e. peak 561 values over half an hour). Thus the consideration of such constant concentrations over a hundred-hour 562 exposure is likely excessive and must be considered as an upper limit that could not be exceeded. This led to the following calculated airborne concentrations: 1 mBq·m⁻³ for ²³⁹Pu, 1.5 mBq·m⁻³ for ²⁴⁰Pu, 44 mBq·m⁻³ 563 for ²⁴¹Pu and 12 mBq·m⁻³ for ²⁴¹Am. Given the above-mentioned considerations and the effective dose 564 565 coefficients for workers (Table 2), the corresponding committed effective dose over an integration time of 566 50 years by inhalation of radioactive smoke at such RN concentrations would have reached 170 microSievert 567 (μ Sv) in total of which 80% comes from ²⁴¹Am (Table 3). A much more likely dose assessment even if may 568 be an underestimation can be established based on spatially averaged concentrations (Table 3). In this case, 569 the resulting dose induced by inhalation of all artificial RN considered in this study over a period of 570 100 hours would be $1.3 \,\mu$ Sv. These estimated doses are much lower than the internationally accepted maximum dose for the public from external sources (1 mSv·yr⁻¹). 571

572

573 Table 2: Effective dose coefficients (Sv·Bq⁻¹) of RN of interest for the general public (adult) and for

574 workers during wildfires.⁷¹

Radionuclide \rightarrow	137 Cs	⁹⁰ Sr	²³⁸ Pu, ²³⁹ Pu, ²⁴⁰ Pu	²⁴¹ Pu	²⁴¹ Am
Absorption type \rightarrow	F	М	S	S	Μ
Dose coefficient (Sv Bq ⁻¹) Adult of the Public (*)	4.6 10-9	3.6 10-8	1.6 10-5	1.7 10-7	4.2 10-5
Dose coefficient (Sv Bq ⁻¹) Workers (*)	4.8 10-9	3.6 10-8	1.5 10-5	1.6 10-7	3.9 10-5

575 (*) aerosol diameter of 1 μ m.

576 Notes: Absorption types: F = Fast, M = Moderate, S = Slow. For ⁹⁰Sr, the type M corresponding to fuel 577 fragments or when unspecified forms is recommended for the general public ⁷² and for workers ⁷³ even if 578 there is no recommendation in ⁷¹. For Pu isotopes a S-type solubility has been considered as a result of the 579 Pu oxide forms released during the Chernobyl accident.

580

581 Despite rather high airborne ¹³⁷Cs concentrations measured near fire lines, the low ¹³⁷Cs dose coefficient, as

582 compared with that of ⁹⁰Sr or Pu isotopes, minimizes its dose impact (only 1% of the total inhalation dose,

Table 3). It is important to point out that the 137 Cs dose coefficient is 10 fold less significant than that of 90 Sr

584	(Table 2). Since airborne 90 Sr concentrations were only 0.8 fold lower than those of 137 Cs on average, 90 Sr
585	resulted in a greater average dose impact. As α -emitters, ²³⁸ Pu, ²³⁹ Pu and ²⁴⁰ Pu do not cause any significant
586	external exposure. However, owing to their high radiotoxicity, the contribution of transuranic elements by
587	inhalation to the exposure of firefighters is not something to ignore. Airborne Pu concentrations in the
588	proximity of fire lines were about 250 fold lower than ¹³⁷ Cs concentrations on average. But the Pu effective
589	dose coefficients are about 3,500 fold higher than that of ¹³⁷ Cs. These significant differences in dose
590	coefficients make ^{238,239,240} Pu isotopes definitively more significant dose contributors. This is almost the
591	same for ²⁴¹ Am which emits mostly high-energy α -particles in addition to low-energy γ -rays (E = 60 keV,
592	13%). Eventually, the highest inhalation dose coefficient among the studied RN belongs to 241 Am which has
593	a dose coefficient of about 250 fold higher than that of ²⁴¹ Pu. In any case, the estimated effective dose for
594	firefighters (0.013 μ Sv·h ⁻¹) as a result of inhalation of those radionuclides remained lower than or similar to
595	the external exposure to radiation from the highly contaminated environment of the CEZ which is most
596	often between 0.1 and 1 μ Sv·h ⁻¹ , and with maximum values about 10 μ Sv·h ⁻¹ . ^{6, 32} However, they may have
597	been significantly reduced (1 to 2 orders of magnitude) if protective equipment has been actually used. ⁶

601

Table 3: Estimated effective doses (in µSv) by inhalation of artificial RN at a breathing rate of 3 m³·h⁻¹ and 600 during 100 hours, for firefighters in the CEZ during the April 2020 wildfires, for two scenarios: (A) observed peak concentrations, (B) spatially averaged concentrations.

Scenario	Radionuclide	¹³⁷ Cs	⁹⁰ Sr	²³⁸ Pu	²³⁹ Pu	²⁴⁰ Pu	²⁴¹ Pu	²⁴¹ Am	Total
A	Maximum airborne concentration (mBq·m ⁻³)	1000	1000	1	1	1.5	44	12	
	Bq inhaled after 100 h	300	300	0.3	0.3	0.4	13.2	3.6	
	Dose (µSv)	1.44	10.8	4.5	4.2	6.3	2.1	141	170
В	Average airborne concentration (mBq·m ⁻³)	10	10	0.01	0.007	0.01	0.3	0.09	
	Bq inhaled after 100 h	3	3	0.003	0.002	0.003	0.09	0.027	
	Dose (µSv)	0.014	0.108	0.045	0.029	0.042	0.014	1.05	1.3
	Contribution to the total inhalation dose	1.1%	8.3%	3.5%	2.2% 8,9%	3.2%	1.1%	80.6%	

602

603 The inhalation dose rate (0.013 µSv·h⁻¹) would have also remained about 8 fold lower than the average ambient dose rate of 0.1 µSv·h⁻¹ from the natural background.⁶ When considering maximum airborne 604 concentrations, the dose rate estimation $(1.7 \,\mu \text{Sv} \cdot \text{h}^{-1})$ by inhalation of artificial RN for firefighters is similar 605

606 to the order of magnitude of the external exposure dose rate from the highly contaminated environment (1 607 $-10 \,\mu \text{Sv}\cdot\text{h}^{-1}$). It would have been 10 to 20 fold higher than the external ambient dose rate from natural 608 background radiation of 0.1 μ Sv·h⁻¹ (range 0.07 - 0.23 μ Sv·h⁻¹) and the internal dose of 0.18 μ Sv·h⁻¹ (range $0.05 - 1.3 \mu \text{Sv} \cdot \text{h}^{-1}$), respectively.^{6, 63} Our estimates are consistent with those from previous studies,^{6, 62, 63} 609 610 which also indicate the predominant contribution of transuranic elements in the internal inhalation 611 exposure.⁶ These studies also point out that the dose received by firefighters because of smoke inhalation 612 (internal dose) was only about 1% of the dose induced by ground shine and the effective external dose would 613 have exceeded the expected internal dose for firefighters even without protective equipment.⁶ The dose of 614 external radiation from the smoke (cloud shine) in case of fire was not taken into account as it is negligible 615 $(10^4 - 10^5)$ fold lower) as compared to the external dose (ground shine) from the contaminated environment.³²

616 Chernobyl-labeled radionuclides aside, naturally occurring radionuclides with a high dose coefficient and 617 that are prone to emission during a wildfire have to be considered. This is typically the case for, among others, 210 Po (T_{1/2} = 138 d.) as a progeny of the relatively long-lived 210 Pb (T_{1/2} = 22 yr.) and which 618 619 accumulates in the biomass through foliar uptake. Polonium-210 has an effective dose coefficient of 3.3 620 10^{-6} Sv/Bg and 3.0 10^{-6} Sv/Bg for an adult of the public and for a worker, respectively, and given a type M 621 solubility corresponding to chloride, hydroxide, volatilized Po and all unspecified Po forms. Polonium is 622 among the radioactive elements with a low fusion point (about 254 °C for elemental Po under 1 atm). The 623 volatilization points of common polonium compounds are about 390 °C thus much lower as compared to mean wildfire temperatures (> 500 °C with maximum of 1,000 -1,200 °C).⁵ As a result, ²¹⁰Po is easily 624 emitted during a fire. Carvalho et al. suggested that, as a result of combustion, a percentage of the ²¹⁰Po 625 626 initially in the biomass becomes concentrated in flying ash particles which corresponds to the refractory 627 remaining fraction after organic and water losses, while another percentage of the ²¹⁰Po forms gaseous ions 628 after volatilization which are likely to be captured by electrostatic forces onto smaller aerosol particles (< 629 $0.5 \,\mu\text{m}$), due to their higher surface/mass (or volume) ratio.⁵ Because of its high dose coefficient these 630 processes are a further reason to take ²¹⁰Po into account when estimating internal exposure during a wildfire. Carvalho et al. measured a maximum airborne ²¹⁰Po concentration of 70 mBq·m⁻³ in the proximity of a fire 631 632 line in Portugal in the summer 2012.⁵ This concentration is as much as 1,000 fold higher as compared with the airborne ²¹⁰Po background level of about a dozen μ Bq·m⁻³ in the Northern hemisphere.⁷⁴ Since ²¹⁰Po was 633 not measured in April 2020, it is difficult to assert that such concentration would have been reached in the 634 635 CEZ even though wildfire conditions can be assumed to be similar. In order to adapt to the absence of ²¹⁰Po 636 measurement we propose to use a wider range of possible airborne ²¹⁰Po concentrations encompassing the concentration found in Portugal, i.e. 1, 10 and 100 mBq·m⁻³, in order to provide the order of magnitude of 637 the inhalation dose assessment. We can estimate a maximum breathing rate at 3 m³·h⁻¹ for a firefighter 638

working 10 hours a day (as established by Kashparov et al., 2015) during 10 days.⁶ Based on a ²¹⁰Po concentration of 10 mBq·m⁻³, the inhalation dose would increase by 9 μ Sv. This would represent 5% of the inhalation dose from radionuclides originating from Chernobyl (¹³⁷Cs + ⁹⁰Sr + Σ Pu + Am) when considering maximum airborne concentrations. When considering average airborne concentrations this would result in a 7 fold higher dose.

644 With the exception of the area in the immediate proximity to the fire, concentrations rapidly decreased and 645 did not present any concern for public health. At greater distances from the blaze, as in Kiev, the airborne 646 ¹³⁷Cs measured concentrations, or calculated concentrations for the other RN, remained between 1,000 and 647 10,000 fold lower on average than those in the CEZ and did not present any risk for the population, even 648 when considering ingestion of foodstuffs subject to radionuclide deposition. To compute airborne RN 649 concentrations for an inhabitant of Kiev, we used the previously calculated RN source terms (Table 1) as an 650 input parameter in the Eulerian ldx dispersion / deposition model developed by the IRSN (see Source term 651 assessment methodology). Subsequent RN deposition was computed assuming a dry deposition velocity of 652 $0.2 \text{ cm} \cdot \text{s}^{-1}$. Based on actual meteorological data, the effective dose induced both by inhalation (respiratory 653 rate of 22.18 m³·d⁻¹) and ingestion of foodstuffs following RN deposition was determined to be 150 nSv for 654 an adult (100 nSv for inhalation from April 1 to 22, 2020 and 50 nSv for ingestion over a period of 1 year after deposition). The detailed inhalation dose per radionuclide is as follows: ¹³⁷Cs 0.1 nSv, ⁹⁰Sr 1 nSv, ²³⁸Pu 655 5 nSv, ²³⁹Pu 5 nSv, ²⁴⁰Pu 10 nSv, ²⁴¹Pu 5 nSv (ΣPu 25 nSv) and ²⁴¹Am 75 nSv. For the ingestion dose 656 657 calculation, the main agricultural products that were considered include vegetables in season (April), dairy 658 products and meat. The daily consumption of 500 g of leafy vegetables is considered as the most penalizing scenario. As confirmed by Talerko et al.,⁴⁹ the dose induced by exposure to the cloud shine (immersion) 659 660 was negligible as compared with internal exposure. The total exposure was also negligible compared to the 661 annual public exposure limit of 1 mSv according to Ukrainian Radiation Safety Standards for the general public as an added effective dose⁴⁵ or when compared to the average annual global exposure of 2.4 mSv 662 induced by natural background radiation.²⁹ 663

Elsewhere in Europe, doses were even lower as airborne concentrations were much lower. At some distant locations the contribution of the fire plume to the ¹³⁷Cs airborne concentration was estimated to be between 2 and 8-fold at most the usual ¹³⁷Cs trace-level concentration. Assuming consistent soil contamination as a routine source of ¹³⁷Cs background emission (through soil particle resuspension) the use of the airborne ¹³⁷Cs/⁴⁰K ratio has also proven to be helpful in the determination of fire plume contribution. However, the proper use of this tool requires the knowledge of a local baseline value to distinguish any additional remote ¹³⁷Cs input.

In anticipation of future wildfires in the Chernobyl area, the detailed study of the re-emission into the atmosphere of Pu isotopes released during the Chernobyl accident or arising as their decay products (²⁴¹Am,

²³⁷Np), in addition to ⁹⁰Sr, ¹³⁷Cs, ^{243,244}Cm and naturally occurring radionuclides such as ²¹⁰Po during 673 674 wildfires is recommended for a more comprehensive estimate of the internal exposure by inhalation for 675 firefighters and for the population. It is also essential to point out that our knowledge of respective amounts 676 of radionuclides emitted both in gaseous phase according to their volatilization point and as aerosol particles 677 (flying ashes) is insufficient and requires further study. In the future, if forests are not thinned, exposure 678 risks from forest fire emissions are expected to increase due to the accumulation of debris, litter and standing 679 dead trees and because of early and lengthy droughts in the framework of climate change. 680 Lengthy wildfire outbreaks are a challenge for inverse modeling computation when they both vary in

681 location and magnitude. Such outbreaks are also an opportunity to strengthen international collaboration 682 between radioprotection organizations and demonstrate the benefits of rapid information sharing, which is 683 the main goal of the informal Ring of Five (Ro5) European monitoring network.

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691 692

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707 Supporting information

- 708 The Supporting information contains the complete airborne radionuclide concentration dataset,
- satellite images of fires spots and a video of the smoke plume dispersion over Europe. It also
- 710 contains 1) a review of historic wildfires in contaminated ecosystems, 2) knowledge about
- radionuclide emission by fires in forested and non-forested lands, 3) information about long-lasting
- persistence of airborne 137 Cs at trace-levels in Europe, 4) geographic analysis and timeline of the
- April 2020 wilfire event in Ukraine, 5) information about the poor air quality observed in Kiev on
- April 16, 2020, 6) radionuclide apportionment in the terrestrial ecosystem, 7) information about
- data collection 8) information about the use of airborne 40 K and 137 Cs/ 40 K ratio for the identification
- of a fire plume contribution, and 9) information about the methodology used for the source term
- assessment.
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719 **Competing interests**

The authors declare that they have no conflicts of interest.

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