

1 **Response to** Anonymous Referee #1
2 Received and published: 16 May 2017

3 General Comments:

4 =====

5 The manuscript under review presents an inversion analysis using the FLEXPART model
6 of radio-isotopes of Cs and I, emitted after the Chernobyl nuclear power plant accident,
7 using an expanded set of measurements (including deposition) to estimate the source
8 term.

9 The source estimates were evaluated using a Eulerian transport model, and systematic
10 uncertainties are quantified with sensitivity tests.

11 The authors address scientific questions within the scope of ACP, with valid scientific
12 methods and assumptions. Overall, the description of experiments and calculations are
13 sufficiently complete and precise. The language and presentation can be somewhat
14 improved (see specific comments below).

15
16 Specific Comments:
17 =====

18 1. On the choice of re-analysis dataset - isn't ERA-Interim more suitable as a primary
19 database due to the higher resolution and 4D-Var assimilation? Also, see point 6 below
20 on how it was decided which dataset gives better results. Can't the uncertainty be
21 additionally quantified using an independent reanalysis dataset to drive the inverse
22 model (e.g. NCEP)?

23 **Response:** Indeed, we expected that ERA-interim would give better results since it's a
24 newer, highly resolved assimilated dataset. However, it was clearly shown in the
25 manuscript why it was not selected (detailed analysis in [section 4.1](#)).

26 We agree that a better uncertainty could be quantified using an independent dataset. In
27 fact, we tried to do what the reviewer suggested and get FNL and GFS fields from NCEP
28 for 1986, for which we have tested flexpart in the past. However, neither the FNL data
29 (<https://rda.ucar.edu/datasets/ds083.2/>) nor the GFS
30 ([https://www.ncdc.noaa.gov/data-access/model-data/model-datasets/global-forecast-
31 system-gfs](https://www.ncdc.noaa.gov/data-access/model-data/model-datasets/global-forecast-system-gfs)) were available for 1986 for use in flexpart model. Of course, there are other
32 re-analysis datasets from NCEP that cover 1986 meteorology. Though, they do not
33 contain all the necessary variable for a proper run with the flexpart model and they have
34 never been tested before. Therefore, we ended up using 2 different products from
35 ECMWF.

36
37 2. Though there is indeed uncertainty about the gas/particulate fraction of Iodine-131
38 emitted at Chernobyl, it's not well advised to model the radionuclide in the particulate
39 form. Atmospheric measurements have revealed a gaseous/total atmospheric ratio of
40 ~80% (Ring of Five). This could significantly impact atmospheric residence times and
41 deposition patterns. Thus a quantification of the uncertainty impact on the final
42 emission estimate is required.

43 **Response:** The partition of gaseous and particulate iodine has been well studied in the
44 recent accident of Fukushima and, as stated in the Ring of Five measurement network, it
45 is close to 20% in particulate form and 80% as a gas.

46 When performing inverse calculation of iodine, there is no point to state "it is not
47 advised to model iodine as aerosol", because everything depends on the available
48 measurements that will be used to perform the inversion and improve the source-term.
49 In the case of Chernobyl, the existing infrastructure of 1986 resulted in having available

1 observations of the particulate fraction of iodine and, thus, this is the reason that we
2 modeled iodine as an aerosol in flexpart. It would make no sense to model iodine as a
3 gas and perform the inversion, while having aerosol surface activity concentration
4 measurements. This would be out of question.
5 Nevertheless, all the above do not, of course, imply that there was iodine only in
6 particulate form released from Chernobyl in 1986. Of course, there should be gaseous
7 iodine as well, which we cannot calculate with inverse modeling due to the lack of
8 gaseous-phase measurements of iodine.
9

10 3. Section 3.3: When dealing with priors as an ensemble are they simply numerically
11 averaged or do their individual uncertainties used to weight (as is more appropriate)?

12 **Response:** As it is stated in the paper, the prior estimation of the source emissions was
13 calculated by assessing the core inventory of the reactor and estimating what has been
14 consumed from the last time that the fuel was updated (see references within the
15 paper). In this way, they were able to assess what it has been produced in the nuclear
16 reactor and, thus, what it has been released. All these happened in 1986.
17 In all these inventories a constant relative uncertainty of 50% was assigned, which
18 stems from this rough estimation of the source term that was rather arbitrary. This is
19 the reason that we have not weighted the prior ensemble using individual uncertainties;
20 simply because we know absolutely nothing about the daily uncertainties. Therefore, we
21 thought that the best was to simply average prior emissions numerically.
22

23 4. Section 3.4: The i) magnitude, and ii) doubling of deposition over concentration
24 uncertainties seems arbitrary, and should be better motivated.

25 **Response:** We agree with the reviewer's comment and we have further given more
26 information on the use of these uncertainty values (see page 11-Manuscript with track
27 changes enabled).
28

29 5. Section 3.5: Doesn't nudging the dynamical model to ERA-Interim (used in the
30 inversion) limit it's potential as an independent check of the emission inventory
31 improvement?

32 **Response:** We fully agree with this comment on the use of data from ECMWF only (ERA-
33 40 & ERA-Interim). However, the two available datasets from NCEP (GFS & FNL), which
34 have been tested and found to contain all the necessary variables that flexpart needs, in
35 order to run, are limited to more recent years. Please see a more detailed response in
36 the first comment.
37

38 6. Section 4.1: It is not clear if comparison of ERA-40 and ERA-Interim was and selection
39 of the "proper" using RMSE was done selectively for certain "previously inaccurate"
40 regions only. Perhaps the text can be reworked to be made clearer?

41 **Response:** We appreciate the reviewer for pointing us to section 4.1, because, indeed, it
42 was not clear. We have tried to rework on it in order to make it clearer now (see page
43 13).
44

45 Technical corrections:

46 =====

47 Abstract:

48

49 1.2 The present paper -> This paper

1 **Response:** Corrected.
2
3 l.6 the real magnitude -> the magnitude
4 **Response:** Corrected.
5
6 l.10 Remove "because radioactivity is a sensitive topic for the public and attracts a lot of
7 attention"
8 **Response:** We prefer to leave this argument present in the manuscript. This argument
9 is true, as after the accident the official authorities stepped back in releasing
10 measurements in public, in order to minimize the public concern of radioactivity and
11 avoid terrifying the population.
12
13 l.17 give such kind of -> provide available
14 **Response:** Corrected.
15
16 l.25 Please rephrase sentence "The results were of the present inversion were
17 confirmed using an independent Eulerian model, for which deposition patterns were also
18 improved when using the estimated posterior releases" to make meaning more clear
19 **Response:** Corrected.
20
21 Section 2:
22
23 p.4 l.33 good estimations -> estimates
24 **Response:** Corrected.
25
26 p.4 l.17 largest -> bulk of the
27 **Response:** Corrected.
28
29 p.4 l.31 supposedly -> reportedly
30 **Response:** Corrected.
31
32 Section 3:
33
34 p.5 l.20 bomb -> weapon
35 **Response:** Corrected.
36
37 Section 4:
38
39 Rename section 4.1 to "Meteorological datasets results comparison"
40 **Response:** We prefer to keep the title of the section as is, because it describes exactly
41 what we did; we selected the proper meteorological dataset to proceed with the
42 inversions. A title like "Meteorological datasets results comparison" seems less clear in
43 our opinion.
44
45 p. 14 l.12: May 3rd, 4th and 5th -> May 3-5 (and similarly elsewhere...)
46 **Response:** Corrected.
47
48 p. 14 l.14: Chapter 2?
49 **Response:** Corrected (section).

1
2 p.15 l.8: less -> lower
3 **Response:** Corrected.

4
5 p.16 l.12: is the main advantage of -> improvement by
6 **Response:** Corrected.

7
8 p.18 l.23-27: No need to list all countries (and country codes) here. Please remove and
9 reference for brevity.
10 **Response:** Corrected.

11
12 p.19 l.7-9: Rephrase sentence: "Unfortunately, considering that 131I has a lifetime of
13 only about 8 days, it was impossible to gather any observations of 131I deposition over
14 Europe" -> "observations of 131I deposition over Europe are unavailable, due to the
15 short half-life"
16 **Response:** Corrected.

17
18 Figure 1: The uncertainty bands of the different priors are unintelligible. This Fig is not
19 necessary and can be removed. Priors per radionuclide can be broken into individual
20 figures and provided in a Supplement.

21 **Response:** We had a long discussion about the presentation of our emissions. The
22 majority of the coauthors believes that the prior emissions must be presented in the
23 paper (and not in the Supplements). In this way, the readers would be able to see right
24 away how different the posterior emissions are, compared to the prior ones; thus, how
25 drastic the change in the posterior emissions is, compared to the prior ones, using not
26 only concentrations, but also deposition observations for the inversions.
27 If there is no problem with the length and the number of Figures in the manuscript, we
28 like to keep the Figure in the paper.

29
30 Figure 5: Why are only 3 emission heights levels plotted, and not all calculated?

31 **Response:** The source-receptor relationship (see section 3.2) was calculated in 3-hourly
32 intervals (8 per day) for 12 days of release (26 Apr 1986 at 00:00 to 8 May 1986 at
33 00:00) and at 6 layers (0 to 3000 m separated every 500 m). This gives $8 \times 12 \times 6 = 576$
34 unknowns that the inversion needs to solve. If we separate between more vertical levels,
35 then the unknowns increase dramatically and cannot be easily treated using our
36 available computational resources. Furthermore, so far, all the previously published
37 assessments for the source term from the Chernobyl accident reported a maximum
38 altitude of emission of about 2.2 to 2.5 km (see relevant references in the manuscript),
39 based on eyewitnesses. Therefore, we thought it is challenging to define 1 additional
40 layer, in case the observations drive the inversion into releases at higher altitudes. From
41 what we saw in the obtained results, indeed emissions at 2.5-3 km occurred, although
42 they were only 4%-13% of the total emissions (Table 1). Hence we expect emissions to
43 be rather insignificant at higher altitudes.

44

1 Anonymous Referee #2

2 Received and published: 17 May 2017

3 This paper appears to be the next in a series by this research group to recreate the
4 events associated with the Chernobyl disaster in an attempt to better understand the
5 emissions produced and how they were dispersed and deposited across the land- scape.
6 A previous paper (or two) in the series attempted to reassemble the original deposition
7 dataset used to generate the maps published by De Cort et al in 1998, and these data are
8 now being used via an inversion process to model (predict) the source term (actual
9 amounts) of radioactive materials that were released during the disaster. I was shocked
10 to read that only a very small percentage of the original data was dis- coverable by this
11 research team which reaffirms the growing philosophy of requiring empirical data to at
12 least be archived and accessible under some defined conditions.

13 C1

14 This latest effort represents a highly innovative and creative approach to refine esti-
15 mates of the source terms for the Chernobyl accident. Perhaps I am easily impressed,
16 but I found this exercise to be nothing short of brilliant. Of particular note is the re- vised
17 injection height profile suggested by the current modeling effort which may have
18 broader relevance to other contamination systems that may be influenced by fire (e.g.
19 Fukushima).

20 Of course, what is equally interesting from this exercise is the fact that there is no
21 substitute for good direct measurements – even the best models are approximations and
22 usually miss many important features of the phenomenon under study. The physicist
23 might suggest that the addition of other variables/factors might solve this problem but
24 this may not be possible under most real-world situations. This exploration is as
25 valuable as a revealer of what is not predictable as it is as a predictor.

26 Overall, this is a very interesting and well-written paper that makes a substantial and
27 original contribution to this literature. Studies of this sort are increasingly valuable as
28 we face greater threats of nuclear incidents and other environmental hazards in the
29 coming years.

30 **Response:** We acknowledge the reviewer for his kind comments. Indeed, I was always
31 fascinated by the Chernobyl disaster and its consequences since May 1986 that was
32 young and listened all these terrifying statements about this “transparent nightmare” as
33 they called in the news 30 years ago. My interest became much more intense as a
34 scientist later, when I found out that there were >500,000 measurements all over
35 Europe that somebody does not want to share for reasons that nobody explains!!! The
36 funniest thing is that I officially contacted both the IAEA (official organization for
37 accidental release issues) and the JRC (official organization for the measurements taken
38 after the accident) and they both responded that these measurements have been lost. No
39 one from these institutes has no idea about them blaming the Russians that they hide
40 them. It is suspicious that nobody from the JRC has kept a copy of the data, since the JRC
41 researchers processed all these 500k measurements to create the deposition map of
42 Cesium after the accident, which was presented in the ATLAS (see relevant reference in
43 the manuscript).

44 So, since 2003, I started gathering measurements in an excel file. I feel really satisfied
45 that I can now exploit all these measurements and define a more detailed source-term
46 from the Chernobyl accident. In this direction, I have made these measurements
47 publically available, exactly as the resulting posterior releases from the present paper,
48 so anyone can perform any kind of analyses using these measurements and the resulting

- 1 release ratios. Last but not least, whenever I find new measurements, I update the
- 2 dataset.
- 3

1 **Inverse modelling of the Chernobyl source term using**
2 **atmospheric concentration and deposition measurements**

3
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6
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15
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17

1 **Abstract**

2 | This paper describes the results of an inverse modelling study for the determination of
3 the source term of the radionuclides ¹³⁴Cs, ¹³⁷Cs and ¹³¹I released after the Chernobyl
4 accident. The accident occurred on 26 April 1986 in the Former Soviet Union and released
5 about 10¹⁹ Bq of radioactive materials that were transported as far away as the USA and
6 | Japan. Thereafter, several attempts to assess the magnitude of the emissions were made that
7 were based on the knowledge of the core inventory and the levels of the spent fuel. More
8 recently, when modelling tools were further developed, inverse modelling techniques were
9 applied to the Chernobyl case for source term quantification. However, because radioactivity
10 is a sensitive topic for the public and attracts a lot of attention, high quality measurements,
11 that are essential for inverse modelling, were not made available except for a few sparse
12 activity concentration measurements far from the source and far from the main direction of
13 the radioactive fallout.

14 For the first time, we apply Bayesian inversion of the Chernobyl source term using not
15 only activity concentrations, but also deposition measurements from the most recent public
16 dataset. These observations refer to a data rescue attempt that started more than 10 years ago,
17 | with a final goal to provide available measurements into anyone interested. As regards to our
18 inverse modelling results, emissions of ¹³⁴Cs were estimated to be 80 PBq or 30–50% higher
19 than what was previously published. From the released amount of ¹³⁴Cs, about 70 PBq were
20 deposited all over Europe. Similar to ¹³⁴Cs, emissions of ¹³⁷Cs were estimated as 86 PBq, in
21 the same order with previously reported results. Finally, ¹³¹I emissions of 1365 PBq were
22 found, which are about 10% less than the prior total releases.

23 The inversion pushes the injection heights of the three radionuclides to higher altitudes
24 (up to about 3 km) than previously assumed (≈2.2 km) in order to better match both
25 | concentration and deposition observations over Europe. The results of the present inversion
26 were confirmed using an independent Eulerian model, for which deposition patterns were also
27 improved when using the estimated posterior releases. Although the independent model tends
28 to underestimate deposition in countries that are not in the main direction of the plume, it
29 reproduces country levels of deposition very efficiently. The results were also tested for
30 robustness against different set-ups of the inversion through sensitivity runs. The source term
31 data from this study are made publically available.

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1 Introduction

2 About 30 years ago, on April 26th 1986, the worst nuclear accident in human history
3 took place in the nuclear power plant (NPP) complex “V. I. Lenin” of the Former Soviet
4 Union (FSU), near the city of Pripyat and in proximity to the administrative border of Ukraine
5 with Belarus. The disaster began during a systems test at reactor four. There was a sudden and
6 unexpected power surge, and when an emergency shutdown was attempted, a much larger
7 spike in power output occurred, which led to a reactor vessel rupture and a series of steam
8 explosions. These events exposed the graphite moderator of the reactor to air, causing it to
9 ignite (Burakov et al., 1996; Medvedev, 1990). The resulting fire sent a plume of highly
10 radioactive fallout into the atmosphere, which dispersed over an extensive geographical area.
11 Around 10 EBq (10^{19} Bq) of fission products were released, of which the largest fraction were
12 noble gases (De Cort et al., 1998). The most severe contamination occurred in FSU countries
13 (Ukraine, Belarus and Russia) (Izrael et al., 1990, 1996). From 1986 to 2000, 350 to 400
14 thousand people were evacuated and resettled from the most severely contaminated areas of
15 Belarus, Russia, and Ukraine (Cardis et al., 1996; Fairlie and Sumner, 2006).

16 Shortly after the accident became known to the public, concerns were put forward about
17 the extent of radioactive pollution and the exposure to radiation of the European population.
18 Many countries in Europe (e.g., Hamilton et al., 1986; Kauppinen et al., 1986) and outside
19 Europe, such as the USA (e.g., Bondieth and Brantley, 1986), Taiwan (e.g., Chung and Lo,
20 1986) and Japan (e.g., Aoyama et al., 1987; Nishizaw et al., 1986) started reporting unusually
21 high levels of radioactivity in environmental media. After the accident, the REM
22 (Radioactivity Environmental Monitoring) programme was established with the aim to
23 improve procedures for the collection, evaluation and harmonization of environmental
24 radioactivity concentrations and the modelling of the migration of radioactivity in the
25 environment for routine and emergency conditions (JRC, 2016). In parallel, several research
26 groups worldwide started independently reporting observations of surface atmospheric
27 concentrations and deposition taken for research purposes. A direct outcome of the REM
28 project was the "Atlas of caesium deposition on Europe after the Chernobyl accident"
29 (hereafter: "Atlas") based on 500 thousand measurements all over Europe, 60% of which had
30 been collected in the FSU.

31 2 Earlier estimates of the source term and purpose of the paper

32 Early studies published just after the accident (e.g., IAEA, 1992; SCUAE, 1986) gave
33 estimates of the total emitted activity and made first assessments of the temporal release

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1 profiles, although not all of them agreed on the total emitted amount or the vertical position of
2 the releases, i.e., the heights to which radioactive material was vented by the explosions and
3 fires. However, it was clearly emphasized that the releases might have probably reached the
4 free troposphere, due to the pronounced steam explosions and the following thermal
5 explosion. First estimates were published by a USSR report (SCUAE, 1986) estimating that
6 1.76 EBq of ^{131}I , 85 PBq of ^{137}Cs , and 54 PBq of ^{134}Cs were released, Abagyan et al. (1986)
7 reported releases of 38.7 PBq of ^{137}Cs and 482 PBq of ^{131}I . Persson et al. (1987), as well as
8 Albergel et al. (1988), reported a similar source term as in SCUAE (1986) but with different
9 injection altitudes. Later on, Devell et al. (1995) and De Cort et al. (1998) published more
10 refined estimations of the release history. Official results on the source term were published
11 almost ten years after the accident in the 1995 OECD report (Waight et al., 1995), which
12 estimated total releases for ^{131}I , ^{137}Cs , and ^{134}Cs that were the same as in SCUAE (1986).
13 More recently, Brandt et al. (2002) used these official emissions estimates and found
14 excellent agreement between modelling results and observations for surface concentrations.
15 Finally, Davoine and Bocquet (2007) reported releases of 1.82 EBq of ^{131}I , 136 PBq of ^{137}Cs ,
16 and 35 PBq of ^{134}Cs , respectively.

17 The bulk of the releases lasted for about ten days, while later releases were several
18 orders of magnitude lower (De Cort et al., 1998). The first three days correspond to the initial
19 explosions (steam and thermal explosions) characterized by ejections of fuel fragments. The
20 next four days weaker releases occurred due to the fire extinguishing attempts of the
21 firefighters. The last three days the emissions rose up again due to the fuel fire and the core
22 melt-down. The altitude at which emissions were injected into the atmosphere was even more
23 difficult to estimate due to the numerous parameters that have to be taken into consideration
24 (mechanical factors characterizing the explosions, generated heat, local meteorological
25 factors, local scavenging conditions, boundary layer diurnal cycles, etc.). Albergel et al.
26 (1988) and Gudiksen et al. (1989) reported that the first release must have reached 2000 m or
27 more. A similar profile of the Chernobyl emissions was proposed by Lange et al. (1988) and
28 Hass et al. (1990).

29 The goal of this paper is to reconstruct and assess the source term based on inverse
30 modelling techniques. We focus on the temporal variations and the altitude of the releases.
31 Although reportedly 500 thousand deposition measurements were used to create the Atlas
32 map, only five thousand deposition measurements were made available to the public in the
33 REM database, and very few of these data referred to the FSU countries, where the highest
34 contamination occurred. Therefore, inverse modelling studies for the quantification of the

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1 | source, term of Chernobyl were mainly based on atmospheric concentrations only (e.g.
2 Davoine and Bocquet, 2007). For the first time, we perform inverse modelling calculations
3 using an extended dataset of deposition observations for ^{134}Cs , ^{137}Cs and ^{131}I (Evangelidou et
4 al., 2016) together with surface atmospheric activity concentrations. The dataset that we used
5 consists of three thousand observation for ^{134}Cs and eleven thousand observations for ^{137}Cs ,
6 60% of which were made in the FSU countries. The data originate from the public REM
7 database of the Joint Research Centre, enriched with measurements from Ukraine, Belarus
8 and Russia and a few other countries. All of these data were used for creating the original
9 Atlas map, but they were not included in the public REM database and were thus inaccessible.
10 These data have been recovered in a recent data rescue effort (see Evangelidou et al., 2016).
11 All simulations regarding the inversion were performed using FLEXPART version 10
12 combined with a Bayesian inversion algorithm (see next section).

13 3 Methodology

14 3.1 Experimental set-up

15 We used the Lagrangian particle dispersion model FLEXPART version 10 (Stohl et al.,
16 1998, 2005) to simulate transport and deposition of radionuclides. This model was originally
17 developed for calculating the dispersion of radioactive material from nuclear emergencies, but
18 since then it has been used for many other applications as well. Nuclear emergency
19 applications include simulations of the transport of radioactive materials from NPPs and other
20 facilities (Andreev et al., 1998; Wotawa et al., 2010) or from nuclear weapon tests (Becker et
21 al., 2010). The model has a detailed description of particle dispersion in the boundary layer
22 and a convection scheme to describe particle transport in clouds (Forster et al., 2007).

23 Due to the fact that the Chernobyl accident took place 30 years ago, when
24 meteorological models were much simpler than nowadays, the quality of the operational
25 meteorological analyses at that time was low compared to current operational data. For this
26 reason, to drive FLEXPART we used ERA-40 (Uppala et al., 2005), which is an European
27 Centre for Medium-range Weather Forecast (ECMWF) re-analysis (using 3-dimensional
28 variational data assimilation (3D-Var)) of the global atmosphere and surface conditions for 45
29 years (1957–2002) at a 125 km resolution. Furthermore, we used ERA-Interim (Dee et al.,
30 2011), which is a global atmospheric reanalysis from 1979, continuously updated in real time.
31 This system includes a 4-dimensional variational analysis (4D-Var) with a 12-hour analysis

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1 window. The spatial resolution of the data set is approximately 80 km on 60 vertical levels
2 from the surface up to 0.1 hPa.

3 We discretized the emissions from Chernobyl into 576 distinct pulses (six vertical
4 layers \times 96 3-h intervals between 00:00 UTC on 26 April and 00:00 UTC on 8 May) and ran
5 the dispersion model forward in time for each one of the 576 emission array elements. Each
6 one of these simulations quantified the sensitivity of downwind atmospheric activity
7 concentrations and depositions to the emissions in a single time-height emission array
8 element. 300,000 particles per release were used for each simulation, giving a total of about
9 172.8 million particles. To assess the impact of a given release scenario, we also used the
10 model in the same set-up but using time- and altitude-varying emissions instead of pulse
11 emissions. Three aerosol tracers (for ^{134}Cs , ^{137}Cs and ^{131}I) subject to wet and dry deposition
12 were used. While cesium is almost entirely attached onto particle surfaces, iodine can be
13 present in the atmosphere as molecular I_2 , as organic iodide, or as iodide salts. While I_2 is a
14 gas, iodide salts are aerosols. In which form iodine is released to the environment from a
15 nuclear facility depends on its operating conditions (Simondi-Teisseire et al., 2013).
16 Furthermore, iodine chemistry in the atmosphere is complex and can involve, for instance,
17 chemical transformation of the different compounds and particle formation (Saiz-Lopez et al.,
18 2012). Without further information, it is impossible to accurately model the atmospheric
19 processes related to the radioiodine release from Chernobyl. Therefore, we chose a simple
20 approach for our modelling, namely assuming that all released ^{131}I was in particulate form.
21 Radioactive decay was not included in the model simulations, since all radionuclide
22 observations and also the a priori emission data were decay-corrected to the time of the
23 accident for the purpose of the inverse modelling.

24 The simulations accounted for wet and dry deposition, assuming a particle density of
25 2500 kg m^{-3} and four different fractions of each radionuclide with aerodynamic mean
26 diameters of 0.4, 1.2, 1.8 and $5.0 \mu\text{m}$ and logarithmic standard deviations of 1.35, 1.25, 1.20
27 and 1.35, respectively. The four different size bins (0.4, 1.2, 1.8 and $5.0 \mu\text{m}$) received 15, 30,
28 40, and 15% of the emitted mass following Malá et al. (2013). The wet deposition scheme
29 considers below-cloud and in-cloud scavenging separately based on cloud liquid water and
30 cloud ice content, precipitation rate and cloud depth from ECMWF, as described in (Grythe et
31 al., 2017).

1 3.2 Inverse modelling

2 We used the inversion algorithm previously used to calculate the source term of ^{133}Xe
3 and ^{137}Cs in the recent accident in Fukushima NPP (Japan) in 2011 (Stohl et al., 2012), the
4 emissions of greenhouse gases (Stohl et al., 2008), and volcanic sulfur dioxide and ash
5 emissions (Kristiansen et al., 2010; Stohl et al., 2011). The algorithm is based on original
6 work by Seibert (2000), incorporates different types of observation data and can be based on
7 forward or backward calculations with FLEXPART. A full description of the algorithm has
8 been given elsewhere (Seibert et al., 2011). The inversion setup is almost identical to that
9 described by Stohl et al. (2012) for determining the Fukushima emissions as a function of
10 time and altitude.

11 We determine radionuclide emissions as a function of time for 96 3-hourly intervals
12 between 00:00 UTC on 26 April and 00:00 UTC on 8 May. While basically all published
13 estimates (e.g., De Cort et al., 1998) suggest that the emissions after 5 May were about six
14 orders of magnitude lower than before, we included also 6 and 7 May in our inversion, to
15 verify this. The inversion was also done for six vertical levels (0–0.5 km, 0.5–1.0 km, 1.0–
16 1.5 km, 1.5–2.0 km, 2.0–2.5 km, 2.5–3.0 km), yielding a total of $n = 576$ unknowns (i.e.,
17 emission values) denoted as vector x . For each one of the n unknowns, a unit amount of
18 radionuclide was emitted in FLEXPART and the model results (surface concentrations or
19 deposition values) were matched (i.e., ensuring spatiotemporal co-location) with m
20 radionuclide observations put into a vector y_0 . Modelled values y corresponding to the
21 observations can be calculated as:

$$22 \quad y = M \cdot x \quad (\text{Eq. 1})$$

23 where M is the $m \times n$ matrix of source-receptor relationships calculated with FLEXPART. As
24 the problem is ill-conditioned, with the measurement data not giving a strong constraint on all
25 elements of the source vector, additional a priori information on the emissions is necessary to
26 stabilize the solution. Including the a priori (prior) source vector x^a , Eq. 1 becomes:

$$27 \quad M \cdot (x - x^a) \approx y^o - M \cdot x^a \quad (\text{Eq. 2})$$

28 and as an abbreviation:

$$29 \quad M \cdot \bar{x} \approx \bar{y} \quad (\text{Eq. 3})$$

30 Considering standard deviations of the errors while assuming the errors to be uncorrelated,
31 the cost function is:

$$32 \quad J = (M \cdot \bar{x} - \bar{y})^T \cdot \text{diag}(\sigma_o^{-2}) \cdot (M \cdot \bar{x} - \bar{y}) + \bar{x}^T \cdot \text{diag}(\sigma_x^{-2}) \cdot \bar{x} + (D\bar{x})^T \cdot \text{diag}(\epsilon) \cdot D\bar{x}$$

33 (Eq. 4)

1 The first term on the right hand side of Eq. (4) measures the model–observation misfit,
2 the second term is the deviation from the a priori values, and the third term measures the
3 deviation of the temporal emission profile from smoothness. Vector σ_o is the standard errors
4 of the observations, and vector σ_x the standard errors of the a priori values. The operator
5 $diag(a)$ yields a diagonal matrix with the elements of a in the diagonal. D is a matrix with
6 elements equal to -2 or 1 , giving a discrete representation of the second derivative, and ϵ is a
7 regularization parameter determining the weight of the smoothness constraint compared to the
8 other two terms.

9 Eq. 4 implies normally distributed, uncorrelated errors, a condition that is not generally
10 fulfilled. To quantify the model errors, we used an ensemble of FLEXPART simulations
11 using two meteorological datasets (ERA–40 and ERA–Interim) for all three radionuclides of
12 interest. The inversion algorithm reads the source–receptor relationships calculated in
13 FLEXPART simultaneously in order to evaluate a range of prior modeled concentrations and
14 deposition densities. Observation errors may be correlated with neighbouring values, and
15 deviations from the prior sources are likely to be asymmetric, with overestimation being more
16 likely than underestimation as zero is a natural bound. The justification for using this
17 approach is that the problem becomes much easier to solve, detailed error statistics are
18 unknown anyway, and experience shows that reasonable results can be obtained. Negative
19 emission values can occur in this set-up but were removed in an iterative procedure by
20 binding them more strongly to the positive a priori values (i.e., by reducing the uncertainty of
21 these emission elements).

22 For ^{134}Cs and ^{137}Cs , we have used measurements of both atmospheric activity
23 concentrations as well as deposition to constrain the source term (see section 3.4), despite the
24 additional uncertainties contained in the modelled deposition values, primarily related to
25 errors in precipitation information and the scavenging formulation (Gudiksen et al., 1989).
26 However, if the measurements are sparse, all available data should be used, even if not all
27 data can be modelled with the same accuracy. The limitations on performing an inversion
28 using deposited activity measurements were highlighted by Gudiksen et al. (1989). These
29 limitations are associated with the uncertainties of precipitation in the meteorological datasets
30 and of the scavenging schemes used in models, as well as to the unknown mass of ^{137}Cs
31 deposited over Europe, as a result of nuclear weapon tests in the past. However, 30 years after
32 the accident, the mass of ^{137}Cs attributed to the nuclear weapon tests has been well–
33 documented for Europe, it has been reported to be up to 3.5 kBq m^{-2} (De Cort et al., 1998)
34 and has been removed from the observation datasets. In addition, meteorological data have

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1 been improved tremendously with the generation of reanalysis fields (e.g., from ECMWF,
2 Dee et al., 2011; Uppala et al., 2005), which are more accurate and have better spatial
3 resolution compared to operational data available at the time of the accident. The latter in
4 conjunction to the more sophisticated and realistic scavenging schemes used currently in
5 models (e.g., Grythe et al., 2017) support also more accurate simulations of the atmospheric
6 dispersion of radioactive material.

7 In the present case, model and measurement error were combined into the observation
8 error $\sigma_o = \sqrt{\sigma_{meas}^2 + \sigma_{mod}^2}$, where σ_{meas} is the measurement error and σ_{mod} the model error.
9 While the inversion method formally propagates stochastic errors in the input data into an a
10 posteriori emission error, the overall error is determined also by partly systematic other errors,
11 which are difficult to quantify. One possible such error source is systematic errors in
12 simulating the deposition process, leading to biases in atmospheric aerosol lifetime. In that
13 respect, it is beneficial to use both atmospheric concentration and deposition measurements,
14 as errors in modelling the deposition process will affect atmospheric concentrations and
15 deposition values (at least partly) in the opposite way (i.e., overestimating deposition will lead
16 to underestimates of atmospheric concentrations). Thus, combining these two types of data
17 will partly lead to error compensation in inverse modelling.

18 3.3 Prior emissions of ^{134}Cs , ^{137}Cs and ^{131}I

19 [Figure 1](#) shows the time profiles of the released quantities of ^{131}I , ^{134}Cs and ^{137}Cs
20 published in different studies of the Chernobyl accident (Abagyan et al., 1986; Brandt et al.,
21 2002; Izrael et al., 1990; Persson et al., 1987; Talerko, 2005a, 2005b). These estimates were
22 used as an ensemble of different alternative a priori source vectors in our inversion. It should
23 be noted that only source terms published with sufficient temporal and emission height
24 information were considered. In Brandt et al. (2002), total released amounts of ^{134}Cs , ^{137}Cs
25 and ^{131}I , were 54, 85 PBq and 1.76 EBq, respectively, and the highest altitude of the release
26 was 2.2 km on April 26th, gradually decreasing during the following days. For this first
27 release (Prior 1, [Figure 1](#)), we assumed that each particle was injected exactly at each specific
28 altitude without giving any range in the altitude. For the second one (Prior 2), the same mass
29 as in Brandt et al. (2002) was released, but it was equally distributed within the corresponding
30 height layer used for the inverse modelling ([Figure 1](#)). For instance, instead of injecting the
31 released mass of ^{137}Cs at exactly 2.2 km, we injected it between 2.0 km to 2.5 km.

32 The next source profile (Prior 3) was from Persson et al. (1987), who reported the same
33 release amount but reported a release height that reached 2.5 km during the first day

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1 (compared to 2.2 km in Brandt et al. (2002)) and 1.0 km in the following release days
2 (compared to 0.4 km in Brandt et al. (2002)). Izrael et al. (1990) reported emission amounts
3 for ^{137}Cs and ^{131}I only and found that 73 and 483 PBq were released, respectively, at heights
4 of up to 1.5 km during the first two days, at 0.5 km during the third and fourth day, and again
5 up to 1500 m in the following (Prior 4). In one of the first assessments of the source term
6 (Prior 5), Abagyan et al. (1986) reported lower releases than the other studies for ^{137}Cs (39
7 PBq), while ^{131}I releases (482 PBq) were more comparable. The vertical profile of the release
8 was the same as in Izrael et al. (1990). Finally, the last release (Prior 6) was adopted from
9 Talerko (2005a, 2005b), who reported that 73 PBq of ^{137}Cs and 976 PBq of ^{131}I were released
10 at the same heights as Abagyan et al. (1986) and Izrael et al. (1990) had suggested before.

11 To define our a priori emissions (vector x^a) and their uncertainties (vector σ_x in eq. 4),
12 we have used the aforementioned published releases as an ensemble to calculate the daily
13 average emissions of ^{134}Cs , ^{137}Cs and ^{131}I and the respective standard deviations (Figure 2).
14 Accordingly, 54 ± 9 PBq of ^{134}Cs , 74 ± 15 PBq of ^{137}Cs and 1510 ± 395 PBq of ^{131}I were emitted
15 in total during the 10-day period of the releases. As expected, the most uncertain releases
16 occurred during the two first days of the accident, when a dual explosion took place, and
17 during the last two days, when the fuel was ignited. These events were accounted for quite
18 differently in the previously published estimates (see Figure 1).

19 All previous studies suggested that emissions ended abruptly on 5 May, with later
20 emissions being lower by six orders of magnitude (De Cort et al., 1998). For our inversion,
21 we extended the potential emission period by two days, to identify potential late emissions.
22 For this, we used prior emissions of 5 TBq d^{-1} for ^{134}Cs , 10 TBq d^{-1} for ^{137}Cs and 100 TBq d^{-1}
23 for ^{131}I on 6 and 7 May (i.e., about three orders of magnitude smaller than on 5 May)
24 associated with an uncertainty of 0.5 PBq d^{-1} , 1 PBq d^{-1} and 10 PBq d^{-1} , respectively (see
25 Figure 2). Uncertainties of the last two daily emissions were left quite high, in order to allow
26 inversion to calculate potential posterior releases that are much higher than the reported six
27 orders of magnitude lesser levels.

28 3.4 Surface activity concentration and deposition observations

29 Measurements of surface activity concentrations and deposition densities from all over
30 Europe were adopted from Evangeliou et al. (2016). The database consists of surface air
31 activity concentration measurements (in Bq m^{-3}) of ^{134}Cs (1,927 observations), ^{137}Cs (1,601)
32 and ^{131}I (2,041) and deposition density observations (in kBq m^{-2}) of ^{134}Cs (2,966) and ^{137}Cs
33 (11,334) as shown in Figure 3. Of the 11,334 deposition observations for ^{137}Cs , 4,077 were

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1 adopted from the public REM dataset, and the remainder were made available from Talerko
2 (2005a, 2005b) and Kashparov et al. (2003). The data of ¹³⁷Cs deposition over the FSU
3 countries were collected using the standardized method adopted previously in the former
4 USSR (Tsaturov et al., 1996). The samples were collected within national framework
5 programmes for the determination of radioactive deposition in settlements; they were
6 included into the database used for the creation of the Atlas map, but not in the public REM
7 database. Air concentrations in areas closer to the vicinity of the plant were determined using
8 airborne gamma spectrometers mounted on aircraft or helicopters capable of flying at low
9 altitudes (25–100 m) during the initial period after the accident. In countries where
10 concentrations were lower, surface air was sucked through filters for a long time (e.g., hours
11 to days depending on the relevant detection limits and the air concentrations) using high-
12 volume samplers. Then, the filters were measured with gamma spectrometry.

13 As regards to the relative measurement errors (it is combined with model error to give
14 observation error, section 3.2), the experience gathered from Fukushima was used (Stohl et
15 al., 2012). The use of deposition observations in the inversion involves additional uncertainty
16 compared to surface concentrations, due to the unknown mass of each long-lived radionuclide
17 that was deposited previously in the area (e.g., from nuclear weapon tests) and due to the
18 uncertainty of precipitation that differs from different meteorological datasets (Gudiksen et
19 al., 1989). For these reasons, the relative measurement errors were chosen to be double (60%)
20 for deposition densities compared to the concentration values (30%). This, together with the
21 often higher model error, gives deposition values less weight in the inversions in order to
22 account for the aforementioned associated uncertainties. Activity concentrations used in the
23 present inversion were selected from areas with coordinates 10°–20°E and 40°–60°N
24 excluding measurements from Budapest (Hungary), Göttingen (Germany) and Prague
25 (Czechia). All the measurements outside this domain together with the excluded ones were
26 used for validation. Similar to concentrations, deposition measurements from another domain
27 (10°–40°E and 40°–60°N) were used in the inversion due to the different density of
28 observations, whereas the rest were used for validation.

29 3.5 The Eulerian Chemistry – Transport Model (CTM) LMDz-OR-INCA

30 In order to assess the improvement of the emissions achieved by the inversion, we used
31 the LMDz-OR-INCA global Chemistry – Transport Model (CTM) to simulate prior and
32 posterior emissions of ¹³⁷Cs. The model is totally different from FLEXPART and couples the
33 LMDz (Laboratoire de Météorologie Dynamique) General Circulation Model (GCM)

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1 (Hourdin et al., 2006) and the INCA (INteraction with Chemistry and Aerosols) model
2 (Folberth et al., 2006) (Hauglustaine et al., 2004). The atmospheric model was furthermore
3 coupled to the land surface model ORCHIDEE (ORganizing Carbon and Hydrology In
4 Dynamic Ecosystems) dynamical vegetation model (Krinner et al., 2005). In the present
5 configuration, the model consists of 19 hybrid vertical levels extending to the stratosphere,
6 and a horizontal resolution of $2.5^{\circ} \times 1.3^{\circ}$ (144 grid-cells in longitude, 142 in latitude).
7 However, the GCM offers the possibility to zoom over specific regions by stretching the grid
8 with the same number of grid-boxes. In the present study, a zoom over Europe (10°W – 60°E ,
9 20°N – 80°N) was applied achieving a maximum horizontal resolution of 0.45 degrees in
10 longitude and 0.51 degrees in latitude. A more detailed description and an extended
11 evaluation of the GCM can be found in Hourdin et al. (2006). The large-scale advection of
12 tracers was calculated based on a monotonic finite-volume second-order scheme (Hourdin
13 and Armengaud, 1999). Deep convection was parameterized according to the scheme of
14 Emanuel (1991). The turbulent mixing in the planetary boundary layer (PBL) was based on a
15 local second-order closure formalism.

16 The model simulates the distribution of natural (e.g., sea-salt and dust) and
17 anthropogenic aerosols (sulfates, black carbon, radionuclides). It keeps track of both the
18 number and the mass of aerosols using a modal approach to treat the size distribution, which
19 is described by a superposition of 5 log-normal modes (Schulz, 2007), each with a fixed
20 spread. The aerosols are treated in three particle modes, sub-micronic (diameter $< 1 \mu\text{m}$)
21 corresponding to the accumulation mode, micronic (diameter $1\text{--}10 \mu\text{m}$) corresponding to
22 coarse particles, and super-micronic or super coarse particles (diameter $> 10 \mu\text{m}$). In the
23 present study, four different particle diameters (0.4, 1.2, 1.8 and $5.0 \mu\text{m}$) were assumed for
24 each of the radionuclides (one in sub-micronic mode and three belonging in the micronic
25 mode) using the prior and posterior emissions, exactly as in the runs with FLEXPART.
26 LMDz-OR-INCA accounts for emissions, transport (resolved and sub-grid scale), and
27 scavenging (dry deposition and washout) of chemical species and aerosols interactively in the
28 GCM.

29 Each simulation using LMDz-OR-INCA lasted nine months (April to December 1986).
30 Using the present experimental set-up and considering that the lifetime of ^{137}Cs in the model
31 is around seven days (Evangelidou et al., 2013), the atmospheric burden of ^{137}Cs in Europe
32 nine months after the accident is almost zero and everything has been deposited. For this
33 study, the model ran in a nudged mode using 6-hourly ERA Interim Re-analysis data (Dee et
34 al., 2011) with a relaxation time of 10 days (Hourdin and Issartel, 2000).

1 4 Results

2 4.1 Selection of the proper meteorological dataset

3 In order to select the meteorological input dataset that is more suitable for simulating
4 the dispersion of the Chernobyl radioactive cloud, we simulated the accident with
5 FLEXPART using the prior source term (Figure 2) and the two available re-analysis datasets
6 (ERA-40 and ERA-Interim). Figure 4 shows the relative difference (%) in deposition (i.e.,
7 $(ERA_{40} - ERA_{Interim})/ERA_{Interim}$) over Europe averaged for the studied radionuclides
8 (^{134}Cs , ^{137}Cs and ^{131}I). Moreover, the simulated deposition of ^{134}Cs , ^{137}Cs and ^{131}I over Europe
9 using both meteorological datasets for the prior emissions can be seen in Figure S 1. The main
10 problem of most model simulations of the accident has been a failure to reproduce
11 concentrations and deposition in remote areas (e.g., Brandt et al., 2002; Evangeliou et al.,
12 2013; Hass et al., 1990; Hatano et al., 1998), where measurements have revealed quite
13 significant contamination (De Cort et al., 1998; Evangeliou et al., 2016). A characteristic
14 example is the Scandinavian countries, Austria and Germany, where measurements have
15 shown deposition densities of ^{134}Cs and ^{137}Cs above 10 kBq m^{-2} (Figure 3). In these regions,
16 simulations using the ERA-Interim data failed to deposit such large quantities, whereas using
17 the ERA-40 dataset led to a more realistic deposited mass (Figure S 1).

18 This was confirmed by the root mean square error (RMSE), which is an absolute
19 measure of fit of a variable to observations and it is interpreted as the standard deviation of
20 the unexplained variance; hence it is in the same units as the response variable. RMSE values
21 spatially were estimated for areas that deposition of radionuclides has shown large
22 discrepancies from observations. The RMSE values averaged for all Scandinavian countries
23 were estimated to be 47 kBq m^{-2} for ^{134}Cs and 36 kBq m^{-2} for ^{137}Cs using the ERA-Interim
24 dataset and only 36 and 27 kBq m^{-2} (for ^{134}Cs and ^{137}Cs) using the ERA-40 fields. In
25 Germany, RMSEs for ^{134}Cs and ^{137}Cs were 49 and 43 kBq m^{-2} using the ERA-Interim and 41
26 and 32 kBq m^{-2} using the ERA-40 fields, whereas in Austria, they decreased from 48 and 40
27 kBq m^{-2} to 44 and 35 kBq m^{-2} , respectively. A different representation of deposition was also
28 achieved for ^{131}I , although there are not enough measurements to clearly decide which data
29 set gave better results (Figure 3).

30 According to Evangeliou et al. (2016), the total deposition of ^{137}Cs in Europe was 75
31 PBq, based on approximately 12 thousand measurements (shown also in Figure 3) that were
32 interpolated onto a regular grid. We calculated that 71 PBq of ^{137}Cs were deposited over
33 Europe using the prior release (Figure 2) and ERA-40 fields. On the contrary, deposition of

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1 ^{137}Cs using ERA–Interim was much lower (56 PBq). The same deposition pattern was found
2 for ^{131}I and ^{134}Cs , with deposited amounts to be 35% higher when using the ERA–40 re-
3 analysis dataset. The largest relative increase in deposition was estimated in Scandinavia,
4 where models have struggled to reproduce deposition, in Belarus and in different parts of
5 Russia. While it is somewhat surprising that ERA-40 allowed more realistic simulations than
6 the more modern ERA-Interim dataset, we therefore selected the ERA–40 data as our
7 reference dataset for the inversion. The simulations performed with the ERA–Interim dataset
8 were used as ensemble members in the inversion to quantify the model uncertainties.

9 4.2 Posterior emissions of ^{134}Cs , ^{137}Cs and ^{131}I

10 In this section, the results of the inversion using the prior source term shown in [Figure 2](#),
11 are discussed. According to our inversion, 80 ± 5 PBq of ^{134}Cs were released in total, with the
12 highest emissions occurring on April 26th and 28th. Then the releases declined substantially
13 but increased again on May 3–5, due to the fuel fire and the core melt–down ([Figure 5](#)). This
14 was consistent with what was previously reported for the accident (see [section 2](#) and
15 references therein). We estimated that about 70 PBq of ^{134}Cs were deposited all over Europe.
16 Unfortunately, there exists no direct calculation of the total deposition of ^{134}Cs over Europe
17 based on measurements, due to the relatively short-lived nature of this radionuclide and thus
18 lack of data. However, considering that the isotopic ratio $^{134}\text{Cs}/^{137}\text{Cs}$ for the Chernobyl
19 accident was reported as 0.6 (Arvela et al., 1990) and about 75 PBq of ^{137}Cs were deposited
20 all over Europe according to measurements (Evangelidou et al., 2016), our ^{134}Cs source term
21 might be a slight overestimate. With respect to the emission altitudes of ^{134}Cs averaged for the
22 12–day period, 37% were released below 0.5 km (against 71% in the prior), 5% at 0.5–1.0 km
23 (against 4%), 10% at 1.0–1.5 km (against 14%), 16% at 1.5–2.0 km (against 9%), 19% at 2.0–
24 2.5 km (against 2%) and 13% at 2.5–3.0 km (nothing was released above 2.5 km in the prior
25 source term) ([Table 1](#)). Our optimised inversion lifted 47% of the releases above 1.5 km, in
26 contrast to only 11% in the prior source term.

27 Like the prior emissions, the posterior emissions of ^{137}Cs were high at the beginning of
28 the accident, due to the initial explosions, then decreased until they rose up again due to fuel
29 melt–down ([Figure 5](#)). Although our total posterior emissions are nearly the same as the prior
30 emissions (86 against 74 PBq), posterior simulations resulted in less efficient deposition at
31 close distances and more deposition over remote regions (see next section). The main
32 difference in the source terms is a much higher release during the first day of the accident (29
33 PBq against 19 PBq in the prior emissions). Furthermore, the releases on the first day

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1 occurred at much higher altitudes: 1.2 PBq were released at altitudes up to 0.5 km, 0.5 PBq
2 between 0.5 and 1.0 km, 10.3 PBq between 1.0 and 1.5 km, 9 PBq at 1.5–2.0 km, 5 PBq at
3 2.0–2.5 km and 3 PBq at 2.5–3.0 km. The corresponding values in the prior source term were
4 0, 0.3, 9.5, 7.5, 2 and 0 PBq. Thus, our inversion emits 28% of the releases of the first day
5 above 2.0 km and 10% above 2.5 km, in contrast to only 9% and 0% in the prior emissions,
6 respectively. For the whole 12-day period, 21% of the posterior emissions were released
7 above 2 km, compared to only 2% of the prior emissions (see [Table 1](#)).

8 Finally, the posterior emissions of ^{131}I were estimated as 1365 PBq in total, about 10%
9 lower than the prior total releases. The temporal pattern of the posterior releases remained
10 almost exactly as in the prior emissions (high emissions on April 26th, then a decrease
11 followed by a slight increase towards the end of the 12-day period) ([Figure 5](#)). The most
12 notable difference was again related to the altitude of the injection. We estimate that 70% of
13 the mass emitted was injected between below 1 km, 21% between 1 and 2 km and the rest
14 (9%) above 2 km. The vertical profile of the prior releases was 76% at 0–1 km, 19% at 1–2
15 km and 5% at 2–3 km.

16 Overall, we found that the inversion shifted the emissions to higher altitudes compared
17 with the prior estimates in order to better match observations. Specifically, 13% of the total
18 emitted mass of ^{134}Cs , 10% of ^{137}Cs and 4% of ^{131}I were injected above 2.5 km, where no
19 prior emissions occurred ([Table 1](#)). It seems likely that higher emission altitudes lead to
20 reductions of the efficiency of dry and possibly also the wet deposition. As a consequence of
21 this, increased atmospheric burdens, transport over longer distances and enhanced deposition
22 in areas located far from the source can be expected. Another major change was that the
23 inversion increased the emissions of ^{134}Cs and ^{137}Cs emissions on the first day by factors of
24 2.8 and 1.5, respectively.

25 4.3 Deposition over Europe using the optimised emissions

26 The pronounced elevation of the posterior emissions of ^{134}Cs resulted in a higher
27 deposition in remote areas compared to the simulation using prior emissions. More
28 specifically, an indistinguishable increase of 5% was estimated in Scandinavia, mostly in
29 Finland (north of Tampere) ([Figure 6](#)). Another spatial increase in deposition was observed in
30 the Alpine environments of Austria and Switzerland, where deposition was almost doubled
31 ([Figure 6](#)). Finally, in the FSU countries of Belarus and Russia deposition also increased by
32 20% and 64%, respectively, whereas the same amount as in the simulation using the prior
33 emissions was found in Ukraine but shifted slightly to the east ([Figure 6](#)). Country-by-country

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1 comparison of deposition of ^{134}Cs was not performed due to the lack of available
2 measurements of ^{134}Cs over Europe.

3 The optimised emissions of ^{137}Cs resulted in a more accurate deposition over Europe
4 compared to the published deposition maps (De Cort et al., 1998; Evangeliou et al., 2016)
5 (Figure 6). For instance, only trace amounts were deposited in the Baltic countries (Estonia,
6 Latvia, Lithuania) using the optimised fluxes in contrast to the prior source term. Decreased
7 deposition compared to when using prior emissions was also observed in Eastern Europe
8 (Poland, Romania, Czechia) or in the Balkan countries (Bulgaria, Former Yugoslavia,
9 Greece), and it is also seen in the Atlas (De Cort et al., 1998). On the contrary, about 30%
10 higher deposition was observed in remote regions of Europe such as in Norway, Sweden and
11 Finland, where measurements presented both in the Atlas and in Evangeliou et al. (2016)
12 reveal ^{137}Cs values of more than 40 kBq m^{-2} . This improvement by the presented posterior
13 fluxes of ^{137}Cs mainly resulted in much higher deposited quantities in areas where to date
14 most of the models have failed to reproduce the high observed deposition values (Brandt et
15 al., 2002; Evangeliou et al., 2013; Hass et al., 1990b; Hatano et al., 1998). In addition, our
16 results capture well the southeastern part of the Black Sea, where observations have not been
17 included in the Atlas, but independent measurements have proved that deposition of ^{137}Cs
18 exceeded 40 kBq m^{-2} there (Köse et al., 1994; Varinlioğlu et al., 1994). The latter is also
19 captured well when using the prior releases. The only discord with our optimised fluxes is the
20 existence of additional deposition in Northwestern Russia, which is not seen in the Atlas.
21 However, since ground-based measurements from this area are lacking, it remains unclear by
22 which measurement data the Atlas results are actually supported.

23 Due to the short-lived nature of ^{131}I , few deposition measurements exist over Europe
24 and it is not possible to compare our deposition maps with observations. However, despite the
25 slightly lower posterior emissions (by 10%), no difference in deposition patterns can be
26 observed comparing to the prior emissions (Figure 6). Nevertheless, absolute numbers show
27 that deposition is slightly lower over Scandinavia, as well as also in regions of Central Europe
28 (e.g. Austria, southern Germany, Poland). The only way to validate these findings for ^{131}I is
29 comparison with atmospheric activity concentrations reported by various groups in Europe
30 (see next section).

31 4.4 Validation of the inversion results against observations

32 It was mentioned in section 3.4 that a fraction of the measurements was excluded from
33 the inversion. These data were used here for investigating the improvement obtained with the

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1 posterior source term compared to the prior source term. Comparison of simulated surface
2 activity concentrations and deposition values of ^{134}Cs , ^{137}Cs and ^{131}I using the prior and the
3 optimized (posterior) source terms are shown in [Figure 7](#). Furthermore, time-series of activity
4 concentrations that were excluded from the inversion (see section 3.4) were compared with
5 simulated concentrations obtained using the prior and posterior emissions ([Figure 8](#)) for
6 remote stations (Athens (Greece), Glasgow (United Kingdom) and Umea (Sweden)) and
7 stations located closer to the Chernobyl NPP (Budapest (Hungary), Göttingen (Germany) and
8 Prague (Czechia)).

9 The comparison of simulated and measured activity concentrations of ^{134}Cs using the
10 prior and posterior fluxes showed generally low correlation coefficients ($R^2 \leq 0.4$) but small
11 improvements when using posterior emissions (R^2 increased from 0.2 to 0.4). Furthermore,
12 the fraction of modelled values, which are within a factor of 10 from the measurements
13 increased from 63% to 75%. This is also shown in the example time-series for Budapest
14 (Hungary), Göttingen (Germany) and Umea (Sweden), where the posterior concentration
15 levels are closer to the observations in the beginning of the accident, but there was a drastic
16 decrease afterwards ([Figure 8](#)).

17 For the ^{134}Cs deposition data, the simulation using the posterior emissions increased the
18 fraction of the data that are within a factor of 10 from the measurements (65%, compared to
19 58% using the prior emissions) ([Figure 7](#)). RMSEs were improved in Scandinavia (31 kBq m^{-2}
20 using the posterior releases compared to 36 kBq m^{-2} using the prior releases), as well as
21 along the borders of Ukraine, Belarus and Russia, where a large portion was deposited.
22 However, observations from these regions were included in the inversion and thus RMSEs
23 cannot be estimated for independent measurements. The largest deviations close to the NPP
24 were observed in the eastern part of Belarus (near Gomel), where measurements showed high
25 deposition of radionuclides (see Atlas). However, deposition observations of ^{134}Cs from this
26 particular area were limited ([Figure 3](#)) and had a limited impact on the inversion.

27 For ^{137}Cs surface activity concentrations, there was a drastic improvement in agreement
28 with independent data when using the posterior instead of the prior emissions ([Figure 7](#)). The
29 fraction of modelled data that were within a factor of 10 of the measured values increased
30 from 18% using the prior emissions to 84% using the posterior emissions ([Figure 7](#)). This is
31 also apparent in the example of prior and posterior modelled and measured time-series
32 concentrations shown in [Figure 8](#) for Athens (Greece). Although the modelled concentrations
33 were already in the right order of magnitude using the prior emissions, the inversion improved
34 the agreement further, especially after May 15th. During this period, surface concentrations of

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1 ^{137}Cs using the prior releases were several orders of magnitude lower than in the observations
2 (Figure 8).

3 Similar to concentrations, deposition densities of ^{137}Cs using the posterior emissions
4 showed better results than using the prior ones (Figure 7). As for the concentrations, the
5 posterior deposition values were generally increased, which is in better agreement with the
6 observations. Specifically, the inversion increased the fraction of the modelled values that are
7 within a factor 10 of the observations from only 54% using the prior emissions to 72%, when
8 the posterior source term was used. RMSEs decreased from 27 to 19 kBq m⁻² in Scandinavia,
9 from 32 to 29 kBq m⁻² in Germany and from 35 to 27 kBq m⁻² in Austria confirming this
10 better representation of deposition. However, near the NPP our results show poor agreement
11 with the Atlas map estimating a generally lower deposition in Belarus and Ukraine and
12 relatively higher deposition values in the Russian territory close to the borders with Ukraine
13 and Belarus. This is probably the result of injecting posterior emissions at higher altitudes,
14 which causes slower deposition of ^{137}Cs to nearby areas and enhances deposition over remote
15 regions.

16 As the comparison of modelled grid-cell values with point observations is always
17 problematic, we have also calculated the total modelled deposition in all European countries,
18 We compare these values to the country totals from Evangeliou et al. (2016) and the Atlas
19 (De Cort et al., 1998). Notice that for calculating the country totals, measurement data were
20 used that were also ingested by the inversion. The results are shown in Figure 9, for the
21 simulations using the prior and the posterior emissions. Deposition of ^{137}Cs over Europe is
22 already captured very well using the prior emissions with high correlation coefficients
23 ($R^2 \sim 0.9$). However, it is obvious that using the posterior fluxes, the deposition values of
24 ^{137}Cs are closer to the identity line for both observation datasets, while high correlations are
25 maintained (Figure 9). High deposition in the countries of the FSU is also captured quite well,
26 whereas deposition in Western Europe is slightly underestimated (e.g., in Belgium, Denmark,
27 Ireland, Luxembourg and Netherlands).

28 Finally, the releases of ^{131}I were estimated to be 1365 ± 185 PBq, which is about 11%
29 lower than in the prior emissions (1510 ± 395 PBq). Comparing with independent
30 observations, modelled ^{131}I concentrations over Europe showed a slight improvement with
31 68% of the data within a factor of 10 from the observations in the posterior emissions,
32 compared to 62% with the prior emissions. Unfortunately, observations of ^{131}I deposition over
33 Europe were unavailable due to the short half-life of ^{131}I .

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Deleted: , (specifically Austria (AT), Belarus (BY), Belgium (BE), Croatia (HR), Czech Republic (CZ), Denmark (DK), Estonia (EE), Finland (FI), France (FR), Germany (DE), Greece (GR), Hungary (HU), Ireland (IE), Italy (IT), Latvia (LV), Lithuania (LT), Luxembourg (LU), Moldavia (MD), Netherlands (NL), Norway (NO), Poland (PL), Rumania (RU), Russia (RU, European part), Slovak Republic (SK), Slovenia (SL), Spain (ES), Sweden (SE), Switzerland (CH), Ukraine (UA) and United Kingdom (GB))

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1 Another point worth highlighting is whether the model was able to correctly simulate
2 the arrival times of the radioactive fallout. An example may be seen in [Figure 8](#) for the six
3 different stations with independent data (Athens, Glasgow, Umea, Budapest, Göttingen and
4 Prague), for which the time–series of concentrations have been plotted. It is obvious that the
5 model was able to predict the arrival times to the measurement stations quite accurately. More
6 specifically, it captured arrival times with a delay of up to 1 day, at maximum, in Southern
7 Europe (Greece) and in Western (United Kingdom), Central (Germany) and Eastern Europe
8 (Czechia and Hungary). In Northern Europe (Umea, Sweden), although the model captured
9 the arrival time of the plume quite well, it failed to capture the right levels of the modelled
10 concentrations that were several orders of magnitude lower.

11 Finally, the model did not reproduce well the duration of the plume passage, with
12 typically a too rapid concentration decrease after the peak concentrations were reached. This
13 is probably attributed to potential remobilisation of the deposited radionuclides and has been
14 also confirmed both for Chernobyl (Garger et al., 1997, 1998; Nicholson, 1989; Rosner and
15 Winkler, 2001) and Fukushima (Steinhauser et al., 2015; Stohl et al., 2012; Yamauchi, 2012).
16 It has been found that after the first passage of the plume and the atmospheric removal of the
17 transported radionuclides, radioactivity can be resuspended by the prevailing winds causing a
18 secondary contamination. This is likely the reason that all three radionuclides were detected
19 continuously in the measurements ([Figure 8](#)) after the initial event, even when the air is not
20 even coming from Chernobyl. The remobilisation is also a problem for the inversion, which
21 attempts to attribute the measured activity concentrations to direct releases from NPP.
22 However, given that measured concentrations during such remobilisation events are several
23 orders of magnitude smaller than during the initial plume passage, this is not a severe
24 problem.

25 5 Discussion

26 5.1 Further validation of the posterior emissions using a Eulerian Chemistry – 27 Transport Model (CTM)

28 We have used the LMDz-OR-INCA model to simulate the accident of Chernobyl
29 independently of FLEXPART using both the prior and posterior emissions ([Figure 10](#)). The
30 simulated surface activity concentrations and deposition densities of ^{137}Cs are compared with
31 the most recently updated measurement dataset (Evangelidou et al., 2016). [Figure 10](#) shows
32 that a much larger amount of ^{137}Cs was deposited over Europe using the obtained posterior

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1 emissions. In total numbers, 75 PBq out of 86 PBq (or 87% of the total released amount) were
2 deposited over Europe using the optimised emissions and 63 PBq out of 74 PBq (or 85% of
3 the total released amount) using the prior emissions out. The posterior number agrees very
4 well with the 77 PBq total deposition of ¹³⁷Cs over Europe reported in the Atlas (De Cort et
5 al., 1998). It is also consistent with the estimated total deposition of ¹³⁷Cs over Europe of
6 75 PBq based on the measurement dataset presented in Evangeliou et al. (2016). It is
7 furthermore consistent with the deposited amount calculated using FLEXPART,
8 approximately 80 PBq.

9 The improvement when using posterior emissions can also be seen in the direct
10 comparison of simulated concentrations and deposition densities with measurements (Figure
11 10, lower panel). It seems that the release of ¹³⁷Cs at higher altitudes in the posterior
12 emissions resulted in much smaller wet and dry deposition in areas close to Chernobyl and
13 more long-range transport of the radioactive fallout. This is translated in higher surface
14 activity concentrations and deposition in remote regions of Europe. Accordingly, with the
15 posterior emissions 85% of the modelled concentration values (in contrast to 47% using the
16 prior emissions) are within a factor of 10 from measurements.

17 Comparison of simulated deposition with measurements did not show a large
18 improvement using the prior and optimized emissions of ¹³⁷Cs despite the pronounced better
19 representation of deposition over Europe. This is due to the fact that most of the
20 measurements were collected close to the Chernobyl NPP and, therefore, hundreds of
21 observations can be located within a single grid-cell of LMDz-OR-INCA. Nevertheless,
22 RMSEs decreased from 35 to 22 kBq m⁻² in Scandinavia, from 48 to 45 kBq m⁻² in Germany
23 and from 45 to 31 kBq m⁻² in Austria. To better assess the resulting deposition, we calculated
24 again modelled country totals of ¹³⁷Cs deposition using both the prior and the posterior
25 releases and compared them with the respective values from the Atlas and Evangeliou et al.
26 (2016) (Figure 11). In general, even with posterior emissions the model still underestimates
27 deposition in countries that are not within the main direction of the fallout, such as Belgium,
28 Netherlands, Spain, France, Great Britain, Ireland and Italy. However, it manages to
29 reproduce levels of contamination in Ukraine, Belarus and Russia, in Scandinavia (except for
30 Norway that is still underestimated), in Central Europe (Poland, Germany and Austria), as
31 well as around the Baltic countries. Almost all values were less than an order of magnitude
32 lower than the observations maintaining high correlation coefficients for both datasets
33 ($R^2 > 0.8$).

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1 5.2 Uncertainty analysis

2 While we propagate uncertainties in the inversion, it can be argued that true posterior
3 uncertainties may be quite different from what we obtain. One reason is that even the prior
4 uncertainties are not well characterized; another reason is that the inversion assumes that all
5 data are independent and normally distributed. Furthermore, all measurements were taken
6 about 31 years ago, from several different groups all over Europe that used various different
7 techniques to determine radionuclide levels in soil or atmospheric aerosol; this induces an
8 uncertainty that cannot be easily defined. Inversion uncertainty also depends on the
9 uncertainty of the model, which is a function of the way it treats atmospheric transport and
10 removal, both of which depend both on the meteorological input data as well as model
11 parameterizations.

12 To better characterise the true uncertainty of our results and examine how robust our
13 inversion is to different set-ups, we have performed numerous sensitivity tests. In each of
14 them, we tuned different parameters of the inversion. More specifically, we have performed
15 inversions (a) using six different prior source terms, (b) using three different injection profiles
16 in the prior emissions, (c) using two different meteorological datasets (ECMWF ERA-40 and
17 ERA-Interim), (d) including only deposition observations or (e) only activity concentrations,
18 and (f) including only observations (both concentrations and deposition densities) from areas
19 close to the NPP (28°E–32°E, 48°N–52°N). For each of the sensitivities, the standard
20 deviations of the daily posterior emissions were calculated for the whole period, which are
21 plotted as step function in [Figure S 2](#) (TBq s⁻¹). The results are averaged for the 12-day period
22 in [Table 2](#) for ¹³⁷Cs only, assuming that in relative terms they would be similar for ¹³⁴Cs and
23 ¹³¹I, as – except for different decay corrections – they are treated in the same way within the
24 model.

25 When six different prior source terms were used in the inversion (sensitivity test a), the
26 total posterior emissions changed by only 10% ([Table 2](#)). This shows that the posterior
27 emissions are robust against changes in prior emissions and the general pattern of high
28 releases in the beginning and in the end of the 12-day period is well maintained in all six
29 cases ([Figure S 2](#)). The largest differences in the posterior source term occurred for the first
30 two time steps of April 26th and for the later days of the releases. The first is attributed to the
31 large differences of the prior emissions during April 26th, which in some cases reached up to
32 70%.

33 For the very first days of the accident the events that led to the releases of radionuclides
34 are well-known. Two explosions were witnessed immediately after the accident and the

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1 altitude of the injection was assessed pretty well. This is apparent from already published
2 results from model simulations elsewhere (see Brandt et al., 2002; Evangeliou et al., 2013,
3 and references therein). Small variations of the emission altitude (sensitivity test b) affect the
4 inversion rather insignificantly, changing posterior emissions by only 8.5% (Table 2). Much
5 larger differences, 55%, were obtained when switching between different meteorological
6 datasets (ERA-40 to ERA-Interim) (test c, Table 2). This is expected, as the precipitation
7 fields in the ECMWF re-analysis are quite different from those in ERA-40, causing
8 substantial differences in ^{137}Cs deposition (see Figure 4).

9 Other tests explored the sensitivity to using different subsets of measurements. For
10 instance, when only concentration measurements were used in the inversion (test e) including
11 a relative uncertainty in the measurements of 30% (see section 3.4), emissions changed by
12 only 22%. When only deposition observations were used (associated with a relative
13 uncertainty of 60%, test d), posterior emissions were 67% higher than in our reference case.
14 When the inversion was applied using the closest deposition and activity concentration
15 observations (28°E–32°E and 48°N–52°N, test f), the obtained posterior emissions were
16 doubled (Table 2). A likely reason for this deviation can be the aerosol lifetime in
17 FLEXPART (see Grythe et al., 2017). 70% of the emitted mass of each of the three
18 radionuclides was in the sub-micronic and micronic mode. For particles in this range, dry
19 deposition in FLEXPART is slow and also the below-cloud removal is not very efficient close
20 to the source (Grythe et al., 2017). This probably led to an underestimate of deposition near
21 the source, leading to emission increases when the higher observed deposition data are used.
22 Lack of enough deposition was pronounced near the NPP (see Figure 6) and, hence, when
23 using measurements from this small domain (28°E–32°E and 48°N–52°N), the inversion is
24 forced towards higher releases.

25 6 Conclusions

26 We present a detailed inversion analysis of the most important radionuclides (^{134}Cs ,
27 ^{137}Cs and ^{131}I) released after the worst nuclear accident in human history, which occurred on
28 26 April 1986 in the FSU (nowadays, near the borders of Ukraine with Belarus and Russia).
29 For the first time, in addition to atmospheric activity concentration observations, we have also
30 included deposition measurements adopted from a recently compiled dataset, to determine the
31 Chernobyl source term. To constrain the inversion, we have used an ensemble of six different
32 previously published source terms that include different injection altitudes, different total

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1 emitted mass and temporal variation of emissions in order to derive a prior source term and its
2 associated uncertainty.

3 To drive the dispersion model, we used ECMWF re-analysis data. In tests we found
4 that the model produced more realistic radionuclide deposition patterns with the ERA-40 re-
5 analysis dataset than with the ERA-Interim, especially in areas located far from the source
6 (e.g., in Scandinavia, Southeastern France and Scotland), thus ERA-40 was used as a
7 reference for the inverse modelling. We calculated that 71 PBq of ^{137}Cs were deposited over
8 Europe using the prior release (Figure 2) and ERA-40 fields, achieving 35% more deposition,
9 at maximum, than using the ERA-Interim re-analysis dataset.

10 Regarding the posterior emissions of ^{134}Cs , about 80 PBq were released in total with the
11 same temporal pattern as in the prior source term, although these emissions are 32% higher
12 than in Brandt et al. (2002), SCUAE (1986) and Waight et al. (1995) and 55% higher than
13 those reported from Davoine and Bocquet (2007). From the released amount of ^{134}Cs , about
14 70 PBq were deposited all over Europe. Using as approximation for the deposited quantity,
15 we estimated that these emissions might be slightly overestimated.

16 The posterior emissions of ^{137}Cs were high initially (due to the two explosions), then
17 decreased and rose up again during the first days of May (due to the fuel melt-down). The
18 total emissions of ^{137}Cs were estimated to be 86 PBq (against 74 PBq in the prior). Their
19 magnitude is comparable to the emissions reported previously in Brandt et al. (2002), Izrael et
20 al. (1990), Talerko (2005a) and Waight et al. (1995), and significantly lower than those
21 reported by (Davoine and Bocquet, 2007) (136 PBq).

22 Finally, the posterior emissions of ^{131}I were estimated as 1365 PBq or about 10% less
23 than the prior total releases. This is 16% lower than the emissions reported in Brandt et al.
24 (2002) and Davoine and Bocquet (2007), but almost 3 times higher than those reported in
25 Izrael et al. (1990) and Persson et al. (1987) and about 35% higher than those in Talerko
26 (2005b).

27 The most important conclusion for the optimised emissions of all three radionuclides
28 included in this study is the characteristic tendency of the inversion to inject released amounts
29 at higher altitudes. About 47% of the released ^{134}Cs were injected above 1.5 km, in contrast to
30 only 11% in the prior source term. For ^{137}Cs , the portion that was injected above 1.5 km
31 altitude was 26%, relative to only 12% in the prior source term. The differences in prior and
32 posterior emission profiles were smaller for ^{131}I (17% above 1.5 km in posterior emissions,
33 14% in prior ones), probably due to the limited amount of available observations over Europe.

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1 The posterior emissions of ^{137}Cs were assessed independently using a Eulerian
2 Chemistry Transport Model (LMDz-OR-INCA) to simulate transport and deposition. We
3 calculated that 87% (or 75 PBq) of ^{137}Cs posterior releases were deposited over Europe with
4 LMDz-OR-INCA versus 85% (63 PBq) using the prior releases of ^{137}Cs (in prior and
5 posterior fluxes, the total released amount were 74 and 86 PBq, respectively). This deposited
6 amount in Europe is similar to the reported one in the Atlas (77 PBq) and identical to the most
7 recently published estimation that used different data but the same methodology as in the
8 Atlas (75 PBq). The model tends to underestimate deposition in countries that are not in the
9 main direction of the fallout, but it manages to reproduce contamination levels in most
10 countries with correlation coefficients above 0.8.

11 Overall, the results of our inversion for the radionuclides ^{134}Cs , ^{137}Cs and ^{131}I released
12 after the Chernobyl accident were very robust against different set-ups of the inversion. From
13 all sensitivity tests performed here, the maximum variation in the posterior emissions resulted
14 when using measurements from a domain that includes only the highest deposition regions
15 (28°E – 32°E and 48°N – 52°N). The relatively inefficient modeled deposition near the NPP
16 together with relatively high amounts of observed deposition increased the posterior
17 emissions substantially. The source terms obtained in this study are available as an electronic
18 supplement to this publication.

19

20 **Data availability.** All data used for the present publication can be obtained from the
21 corresponding author upon request.

22

23 **Competing interests.** The authors declare that they have no conflict of interest.

24

25 **Author Contributions.** N. Evangeliou designed and performed the experiments and wrote the
26 paper. T. Hamburger designed the experiments and provided corrections on the inversion
27 algorithm. Y. Balkanski wrote part of the paper. A. Cozic provided updates in the LMDz-OR-
28 INCA model. A. Stohl designed and supervised the study and wrote parts of the paper.

29

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4 **References**

- 5 Abagyan, A. A., Ilyin, L. A., Izrael, Y. A., Legasov, V. A. and Petrov, V. E.: The information
6 on the Chernobyl accident and its consequences prepared for IAEA. *Atomic Energy* 3 (5), *At.*
7 *Energy*, 3(4), doi:10.1007/BF01122262, 1986.
- 8 Albergel, A., Martin, D., Strauss, B. and Gros, J. M.: The chernobyl accident: Modelling of
9 dispersion over europe of the radioactive plume and comparison with air activity
10 measurements, *Atmos. Environ.*, 22(11), 2431–2444, doi:10.1016/0004-6981(88)90475-1,
11 1988.
- 12 Andreev, I., Hittenberger, M., Hofer, P., Kromp-Kolb, H., Kromp, W., Seibert, P. and
13 Wotawa, G.: Risks due to beyond design base accidents of nuclear power plants in Europe -
14 The methodology of riskmap, *J. Hazard. Mater.*, 61(1–3), 257–262, doi:10.1016/S0304-
15 3894(98)00130-7, 1998.
- 16 Aoyama, M., Hirose, K. and Sugimura, Y.: Deposition of gamma-emitting nuclides in Japan
17 after the reactor-IV accident at Chernobyl', *J. Radioanal. Nucl. Chem. Artic.*, 116(2), 291–
18 306, doi:10.1007/BF02035773, 1987.
- 19 Arvela, H., Markkanen, M. and Lemmela, H.: Mobile Survey of environmental gamma
20 radiation and fallout level in Finland after Chernobyl accident., *Radiat. Prot. Dosimetry*, 32,
21 177–184, doi:10.1093/oxfordjournals.rpd.a080734, 1990.
- 22 Becker, A., Wotawa, G., Ringbom, A. and Saey, P. R. J.: Backtracking of noble gas
23 measurements taken in the aftermath of the announced october 2006 event in North Korea by
24 means of pts methods in nuclear source estimation and reconstruction, *Pure Appl. Geophys.*,
25 167(4–5), 581–599, doi:10.1007/s00024-009-0025-0, 2010.
- 26 Bondiett, E. A. and Brantley, J. N.: Characteristics of Chernobyl radioactivity in Tennessee,
27 *Nature*, 322(6077), 313–314, doi:10.1038/322313b0, 1986.
- 28 Brandt, J., Christensen, J. H. and Frohn, L. M.: Modelling transport and deposition of caesium
29 and iodine from the Chernobyl accident using the DREAM model, *Atmos. Chem. Phys.*
30 *Discuss.*, 2(3), 825–874, doi:10.5194/acpd-2-825-2002, 2002.
- 31 Burakov, B. E., Strykanova, E. E. and Anderson, E. B.: Secondary Uranium Minerals on the
32 Surface of Chernobyl "Lava," *MRS Proc.*, 465, doi:10.1557/PROC-465-1309, 1996.
- 33 Cardis, E. L., Anspaugh, L., Ivanov, V. K., Likhtarev, I., Prisyazhniuk, A., Mabichi, K. and
34 Okeanov, A. E.: Estimated long term health effects of the Chernobyl accident, in *International*
35 *Conference on one decade after Chernobyl: summing up the consequences of the accident*, pp.
36 241–279, Vienna. [online] Available from: <http://www.osti.gov/scitech/servlets/purl/381695>,
37 1996.
- 38 Chung, C. and Lo, J. G.: Radioactive I-131 over Taiwan after the Chernobyl accident, *J.*
39 *Radioanal. Nucl. Chem.*, 105(6), 325–333, doi:10.1007/BF02166343, 1986.
- 40 De Cort, M., Dubois, G., Fridman, S. D., Germenchuk, M., G., Izrael, Y. A., Janssens, A.,
41 Jones, A. R., Kelly, G., N., Kvasnikova, E., V., Matveenko, I., I., Nazarov, I., N., Pokumeiko,
42 Y., M., Sitak, V. A., Stukin, E., D., Tabachny, L., Y., Tsaturov, Y. S. and Avdyushin, S., I.:
43 Atlas of caesium deposition on Europe after the Chernobyl accident, EU - Office for Official
44 Publications of the European Communities, Luxembourg., 1998.
- 45 Davoine, X. and Bocquet, M.: Inverse modelling-based reconstruction of the Chernobyl
46 source term available for long-range transport, *Atmos. Chem. Phys.*, 7(1), 1–43,
47 doi:10.5194/acpd-7-1-2007, 2007.
- 48 Dee, D. P., Uppala, S. M., Simmons, A. J., Berrisford, P., Poli, P., Kobayashi, S., Andrae, U.,

1 Balmaseda, M. A., Balsamo, G., Bauer, P., Bechtold, P., Beljaars, A. C. M., van de Berg, L.,
2 Bidlot, J., Bormann, N., Delsol, C., Dragani, R., Fuentes, M., Geer, A. J., Haimberger, L.,
3 Healy, S. B., Hersbach, H., H??lm, E. V., Isaksen, I., K??llberg, P., K??hler, M., Matricardi,
4 M., McNally, A. P., Monge-Sanz, B. M., Morcrette, J. J., Park, B. K., Peubey, C., de Rosnay,
5 P., Tavolato, C., Th??paut, J. N. and Vitart, F.: The ERA-Interim reanalysis: Configuration
6 and performance of the data assimilation system, *Q. J. R. Meteorol. Soc.*, 137(656), 553–597,
7 doi:10.1002/qj.828, 2011.

8 Devell, L., Guntay, S. and Powers, D. A.: The Chernobyl reactor accident source term,
9 Organisation for Economic Co-operation and Development, Nuclear Energy Agency, Paris.,
10 1995.

11 Emanuel, K. A.: A Scheme for Representing Cumulus Convection in Large-Scale Models, *J.*
12 *Atmos. Sci.*, 48(21), 2313–2329, doi:10.1175/1520-
13 0469(1991)048<2313:ASFRCC>2.0.CO;2, 1991.

14 Evangeliou, N., Balkanski, Y., Cozic, A. and Møller, A. P.: Simulations of the transport and
15 deposition of 137Cs over Europe after the Chernobyl Nuclear Power Plant accident: Influence
16 of varying emission-altitude and model horizontal and vertical resolution, *Atmos. Chem.*
17 *Phys.*, 13(14), 7183–7193, doi:10.5194/acp-13-7183-2013, 2013.

18 Evangeliou, N., Hamburger, T., Talerko, N., Zibtsev, S., Bondar, Y., Stohl, A., Balkanski, Y.,
19 Mousseau, T. A. and Møller, A. P.: Reconstructing the Chernobyl Nuclear Power Plant
20 (CNPP) accident 30 years after. A unique database of air concentration and deposition
21 measurements over Europe, *Environ. Pollut.*, (August), doi:10.1016/j.envpol.2016.05.030,
22 2016.

23 Fairlie, I. and Sumner, D.: The other report on Chernobyl (TORCH), Berlin, Brussels, Kiev.
24 [online] Available from: <http://www.chernobylreport.org/torch.pdf>, 2006.

25 Folberth, G. A., Hauglustaine, D. A., Lathière, J. and Brocheton, F.: Interactive chemistry in
26 the Laboratoire de Météorologie Dynamique general circulation model: model description and
27 impact analysis of biogenic hydrocarbons on tropospheric chemistry, *Atmos. Chem. Phys.*,
28 6(8), 2273–2319, doi:10.5194/acp-6-2273-2006, 2006.

29 Forster, C., Stohl, A. and Seibert, P.: Parameterization of convective transport in a Lagrangian
30 particle dispersion model and its evaluation, *J. Appl. Meteorol. Climatol.*, 46(4), 403–422,
31 doi:10.1175/JAM2470.1, 2007.

32 Garger, E. K., Hoffman, F. O. and Thiessen, K. M.: Uncertainty of the long-term
33 resuspension factor, *Atmos. Environ.*, 31(11), 1647–1656, doi:10.1016/S1352-
34 2310(96)00345-7, 1997.

35 Garger, E. K., Kashpur, V., Paretzke, H. G. and Tschiersch, J.: Measurement of resuspended
36 aerosol in the Chernobyl area: Part II. Size distribution of radioactive particles, *Radiat.*
37 *Environ. Biophys.*, 36(4), 275–283, doi:10.1007/s004110050082, 1998.

38 Grythe, H., Kristiansen, N. I., Groot Zwaafink, C. D., Eckhardt, S., Ström, J., Tunved, P.,
39 Krejci, R. and Stohl, A.: A new aerosol wet removal scheme for the Lagrangian particle
40 model FLEXPARTv10, *Geosci. Model Dev.*, 10, 1447–1466, doi:10.5194/gmd-10-1447-
41 2017, 2017.

42 Gudiksen, H. P., Harvey, T. F. and Lange, R.: Chernobyl source term, atmospheric
43 dispersion, and dose estimation, *Health Phys.*, 57(5), 697–706, doi:10.1097/00004032-
44 198911000-00001, 1989.

45 Hamilton, E. I., Zou, B. and Clifton, R. J.: The Chernobyl accident - Radionuclide fallout in
46 S.W.England_, *Sci. Total Environ.*, 57, 231–251, 1986.

47 Hass, H., Memmesheimer, M., Geiß, H., Jakobs, H. J., Laube, M. and Ebel, A.: Simulation of
48 the chernobyl radioactive cloud over Europe using the eurad model, *Atmos. Environ. Part A,*
49 *Gen. Top.*, 24(3), 673–692, doi:10.1016/0960-1686(90)90022-F, 1990a.

50 Hass, H., Memmesheimer, M., Geiß, H., Jakobs, H. J., Laube, M. and Ebel, A.: Simulation of

1 the chernobyl radioactive cloud over Europe using the eurad model, *Atmos. Environ. Part A.*
2 *Gen. Top.*, 24(3), 673–692, doi:10.1016/0960-1686(90)90022-F, 1990b.
3 Hatano, Y., Hatano, N., Amano, H., Ueno, T., Sukhoruchkin, A. K. and Kazakov, S. V.:
4 Aerosol migration near Chernobyl: Long-term data and modeling, *Atmos. Environ.*, 32(14–
5 15), 2587–2594, doi:10.1016/S1352-2310(97)00511-6, 1998.
6 Hauglustaine, D. A., Hourdin, F., Jourdain, L., Filiberti, M.-A., Walters, S., Lamarque, J.-F.
7 and Holland, E. A.: Interactive chemistry in the Laboratoire de Meteorologie Dynamique
8 general circulation model: Description and background tropospheric chemistry evaluation, *J.*
9 *Geophys. Res.*, 109(D04314), doi:10.1029/2003JD003957, 2004.
10 Hourdin, F. and Armengaud, A.: The Use of Finite-Volume Methods for Atmospheric
11 Advection of Trace Species. Part I: Test of Various Formulations in a General Circulation
12 Model, *Mon. Weather Rev.*, 127(5), 822–837, doi:10.1175/1520-
13 0493(1999)127<0822:TUOFVM>2.0.CO;2, 1999.
14 Hourdin, F. and Issartel, J. P.: Sub-surface nuclear tests monitoring through the CTBT xenon
15 network, *Geophys. Res. Lett.*, 27(15), 2245–2248, doi:10.1029/1999GL010909, 2000.
16 Hourdin, F., Musat, I., Bony, S., Braconnot, P., Codron, F., Dufresne, J. L., Fairhead, L.,
17 Filiberti, M. A., Friedlingstein, P., Grandpeix, J. Y., Krinner, G., LeVan, P., Li, Z. X. and
18 Lott, F.: The LMDZ4 general circulation model: Climate performance and sensitivity to
19 parametrized physics with emphasis on tropical convection, *Clim. Dyn.*, 27(7–8), 787–813,
20 doi:10.1007/s00382-006-0158-0, 2006.
21 IAEA: The Chernobyl accident: Updating of INSAG-1. A report by the International Nuclear
22 Safety Advisory Group, Safety Ser., International Atomic Energy Agency (IAEA), Vienna.
23 [online] Available from: [http://www-](http://www-pub.iaea.org/MTCD/publications/PDF/Pub913e_web.pdf)
24 [pub.iaea.org/MTCD/publications/PDF/Pub913e_web.pdf](http://www-pub.iaea.org/MTCD/publications/PDF/Pub913e_web.pdf), 1992.
25 Izrael, Y. A., Vakulovsky, S. M., Vetrov, V. A., Petrov, V. N., Rovinsky, F. Y. and Stukin, E.
26 D.: Chernobyl: Radioactive Contamination of the Environment, Gidrometeoizdat, 1990.
27 Izrael, Y. A., De Cort, M., Jones, A. R., Nazarov, I. M., Fridman, S. D., Kvasnikova, E. V.,
28 Stukin, E. D., Kelly, G. N., Matveenko, I. I., Pokumeiko, Y. M., Tabatchnyi, L. Y. and
29 Tsaturon, Y.: The atlas of caesium-137 contamination of Europe after the Chernobyl
30 accident., in *The radiological consequences of the Chernobyl accident*, edited by A.
31 Karaoglou, G. Desmet, G. N. Kelly, and H. G. Menzel, pp. 1–10, European Commission,
32 EUR 16544 EN., 1996.
33 JRC: Radioactivity Environmental Monitoring (REM), Eur. Union – Jt. Res. Cent. [online]
34 Available from: <http://rem.jrc.ec.europa.eu/RemWeb/Index.aspx#> (Accessed 22 November
35 2016), 2016.
36 Kashparov, V. A., Lundin, S. M., Zvarych, S. I., Yoshchenko, V. I., Levchuk, S. E.,
37 Khomutinin, Y. V., Maloshtan, I. M. and Protsak, V. P.: Territory contamination with the
38 radionuclides representing the fuel component of Chernobyl fallout, *Sci. Total Environ.*,
39 317(1–3), 105–119, doi:10.1016/S0048-9697(03)00336-X, 2003.
40 Kauppinen, E. I., Hillamo, R. E., Aaltonen, S. H. and Sinkko, K. T. S.: Radioactivity size
41 distributions of ambient aerosols in Helsinki, Finland, during May 1986 after Chernobyl
42 accident: preliminary report, *Environ. Sci. Technol.*, 20(12), 1257–1259,
43 doi:10.1021/es00154a011, 1986.
44 Köse, A., Topcuoğlu, S., Varinlioğlu, A., Kopya, A. I., Azar, A., Uzun, O. and Karal, H.: The
45 levels of cesium radionuclides in lichens in the eastern black sea area of turkey, *Toxicol.*
46 *Environ. Chem.*, 45(3–4), 221–224, doi:10.1080/02772249409358087, 1994.
47 Krinner, G., Viovy, N., de Noblet-Ducoudré, N., Ogée, J., Polcher, J., Friedlingstein, P.,
48 Ciais, P., Sitch, S. and Prentice, I. C.: A dynamic global vegetation model for studies of the
49 coupled atmosphere-biosphere system, *Global Biogeochem. Cycles*, 19(1), n/a--n/a,
50 doi:10.1029/2003GB002199, 2005.

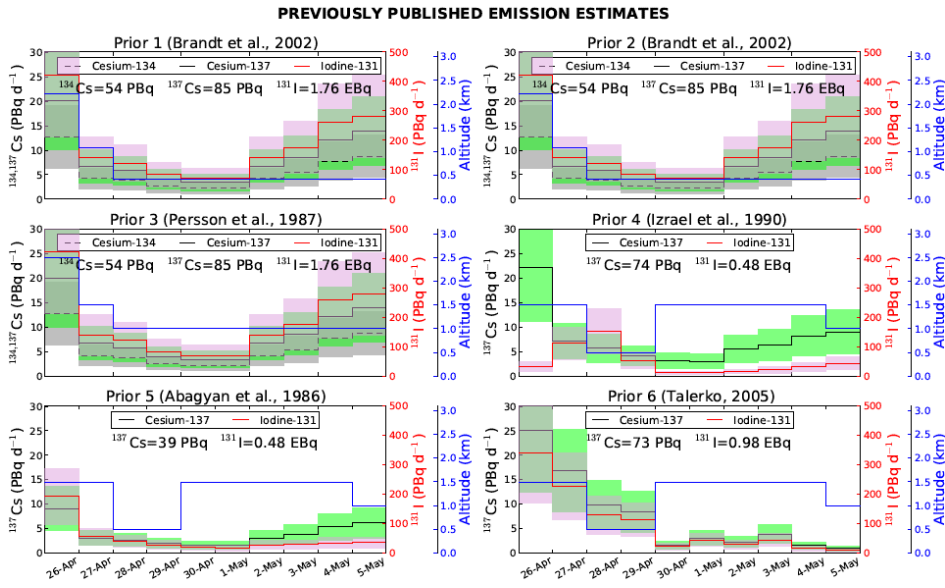
1 Kristiansen, N. I., Stohl, A., Prata, A. J., Richter, A., Eckhardt, S., Seibert, P., Hoffmann, A.,
2 Ritter, C., Bitar, L., Duck, T. J. and Stebel, K.: Remote sensing and inverse transport
3 modeling of the Kasatochi eruption sulfur dioxide cloud, *J. Geophys. Res. Atmos.*, 115(21),
4 1–18, doi:10.1029/2009JD013286, 2010.
5 Lange, R., Dickerson, M. H. and Gudiksen, P. H.: Dose Estimates from the Chernobyl
6 Accident, *Nucl. Technol.*, 82(3), 311–323, 1988.
7 Malá, H., Rulík, P., Bečková, V., Mihalík, J. and Slezáková, M.: Particle size distribution of
8 radioactive aerosols after the Fukushima and the Chernobyl accidents, *J. Environ. Radioact.*,
9 126, 92–98, doi:10.1016/j.jenvrad.2013.07.016, 2013.
10 Medvedev, Z. A.: The legacy of Chernobyl, New York : W.W. Norton, 1990. [online]
11 Available from: <https://search.library.wisc.edu/catalog/999619878702121>, 1990.
12 Nicholson, K. W.: The deposition, resuspension and weathering of Chernobyl derived
13 material in the UK, *Nature*, 9(2), 113–119, doi:10.1088/0952-4746/9/2/004, 1989.
14 Nishizaw, K., Takata, K., Hamada, N., Ogata, Y., Kojima, S., Yamashit, O., Ohshima, M. and
15 Kayama, Y.: I-131 in milk and rain Chernobyl, *Nature*, 342(6095), 308,
16 doi:10.1038/324308a0, 1986.
17 Persson, C., Rodhe, H. and De Geer, L.-E.: The Chernobyl accident – A meteorological
18 analysis of how radionuclides reached and were deposited in Sweden, *Ambio*, 16(1), 20–31,
19 1987.
20 Rosner, G. and Winkler, R.: Long-term variation (1986-1998) of post-Chernobyl ⁹⁰Sr,
21 ¹³⁷Cs, ²³⁸Pu and ^{239,240}Pu concentrations in air, depositions to ground, resuspension
22 factors and resuspension rates in south Germany, *Sci. Total Environ.*, 273(1–3), 11–25,
23 doi:10.1016/S0048-9697(00)00716-6, 2001.
24 Saiz-Lopez, A., Plane, J. M. C., Baker, A. R., Carpenter, L. J., von Glasow, R., Gómez
25 Martín, J. C., McFiggans, G. and Saunders, R. W.: Atmospheric Chemistry of Iodine, *Chem.*
26 *Rev.*, 112(3), 1773–1804, doi:10.1021/cr200029u, 2012.
27 Schulz, M.: Constraining model estimates of the aerosol Radiative Forcing, *Université Pierre*
28 *et Marie Curie, Paris VI*, 2007.
29 SCUAE: The accident at the Chernobyl Nuclear Power Plant and its consequences, USSR
30 State Committee on The Utilization of Atomic Energy, Parts I and II, Information compiled
31 for the IAEA experts' Meeting, Vienna, 25-29 August., 1986.
32 Seibert, P.: Inverse modelling of sulfur emissions in Europe based on trajectories, in *Inverse*
33 *Methods in Global Biogeochemical Cycles*, edited by P. Kasibhatla, M. Heimann, P. Rayner,
34 N. Mahowald, R. G. Prinn, and D. E. Hartley, pp. 147–154, American Geophysical Union,
35 *Geophysical Monograph 114*, Washington, DC., 2000.
36 Seibert, P., Kristiansen, N. I., Richter, A., Eckhardt, S., Prata, A. J. and Stohl, A.:
37 Uncertainties in the inverse modelling of sulphur dioxide eruption profiles, *Geomatics, Nat.*
38 *Hazards Risk*, 2(3), 201–216, doi:10.1080/19475705.2011.590533, 2011.
39 Simondi-Teisseire, B., Girault, N., Payot, F. and Clément, B.: Iodine behaviour in the
40 containment in Phœbus FP tests, *Ann. Nucl. Energy*, 61, 157–169,
41 doi:10.1016/j.anucene.2013.02.039, 2013.
42 Steinhäuser, G., Niisoe, T., Harada, K. H., Shozugawa, K., Schneider, S., Synal, H. A.,
43 Walther, C., Christl, M., Nanba, K., Ishikawa, H. and Koizumi, A.: Post-Accident Sporadic
44 Releases of Airborne Radionuclides from the Fukushima Daiichi Nuclear Power Plant Site,
45 *Environ. Sci. Technol.*, 49(24), 14028–14035, doi:10.1021/acs.est.5b03155, 2015.
46 Stohl, A., Hittenberger, M. and Wotawa, G.: Validation of the lagrangian particle dispersion
47 model FLEXPART against large-scale tracer experiment data, *Atmos. Environ.*, 32(24),
48 4245–4264, doi:10.1016/S1352-2310(98)00184-8, 1998.
49 Stohl, A., Forster, C., Frank, A., Seibert, P. and Wotawa, G.: Technical note: The Lagrangian
50 particle dispersion model FLEXPART version 6.2, *Atmos. Chem. Phys.*, 5(9), 2461–2474,

1 doi:10.5194/acp-5-2461-2005, 2005.
2 Stohl, A., Seibert, P., Arduini, J., Eckhardt, S., Fraser, P., Grealley, B. R., Maione, M.,
3 O'Doherty, S., Prinn, R. G., Reimann, S., Saito, T., Schmidbauer, N., Simmonds, P. G.,
4 Vollmer, M. K., Weiss, R. F. and Yokouchi, Y.: A new analytical inversion method for
5 determining regional and global emissions of greenhouse gases: sensitivity studies and
6 application to halocarbons, *Atmos. Chem. Phys. Discuss.*, 8(6), 19063–19121,
7 doi:10.5194/acpd-8-19063-2008, 2008.
8 Stohl, A., Prata, A. J., Eckhardt, S., Clarisse, L., Durant, A., Henne, S., Kristiansen, N. I.,
9 Minikin, A., Schumann, U., Seibert, P., Stebel, K., Thomas, H. E., Thorsteinsson, T., Tørseth,
10 K. and Weinzierl, B.: Determination of time-and height-resolved volcanic ash emissions and
11 their use for quantitative ash dispersion modeling: The 2010 Eyjafjallajökull eruption, *Atmos.*
12 *Chem. Phys.*, 11(9), 4333–4351, doi:10.5194/acp-11-4333-2011, 2011.
13 Stohl, A., Seibert, P., Wotawa, G., Arnold, D., Burkhart, J. F., Eckhardt, S., Tapia, C.,
14 Vargas, A. and Yasunari, T. J.: Xenon-133 and caesium-137 releases into the atmosphere
15 from the Fukushima Dai-ichi nuclear power plant: Determination of the source term,
16 atmospheric dispersion, and deposition, *Atmos. Chem. Phys.*, 12(5), 2313–2343,
17 doi:10.5194/acp-12-2313-2012, 2012.
18 Talerko, N.: Mesoscale modelling of radioactive contamination formation in Ukraine caused
19 by the Chernobyl accident, *J. Environ. Radioact.*, 78(3), 311–329,
20 doi:10.1016/j.jenvrad.2004.04.008, 2005a.
21 Talerko, N.: Reconstruction of 131I radioactive contamination in Ukraine caused by the
22 Chernobyl accident using atmospheric transport modelling, *J. Environ. Radioact.*, 84(3), 343–
23 362, doi:10.1016/j.jenvrad.2005.04.005, 2005b.
24 Tsaturov, Y. S., Cort, M. De, Dubois, G., Tabachnyi, L. Y., Matvienko, I. I. and
25 Guermenchuk, M. G.: The Need for Standardisation in the Analysis, Sampling and
26 Measurement of deposited Radionuclides, edited by A. Karaoglou, G. Desmet, G. N. Kelly,
27 and H. G. Menzel, pp. 425–433, European Commission, Brussels. [online] Available from:
28 http://www.iaea.org/inis/collection/NCLCollectionStore/_Public/31/056/31056873.pdf, 1996.
29 Uppala, S. M., Kallberg, P. W., Simmons, A. J., Andrae, U., Bechtold, V. D., Fiorino, M.,
30 Gibson, J. K., Haseler, J., Hernandez, A., Kelly, G. A., Li, X., Onogi, K., Saarinen, S., Sokka,
31 N., Allan, R. P., Andersson, E., Arpe, K., Balmaseda, M. A., Beljaars, A. C. M., Van De
32 Berg, L., Bidlot, J., Bormann, N., Caires, S., Chevallier, F., Dethof, A., Dragosavac, M.,
33 Fisher, M., Fuentes, M., Hagemann, S., Holm, E., Hoskins, B. J., Isaksen, I., Janssen, P. A.
34 E. M., Jenne, R., McNally, A. P., Mahfouf, J. F., Morcrette, J. J., Rayner, N. A., Saunders, R.
35 W., Simon, P., Sterl, A., Trenberth, K. E., Untch, A., Vasiljevic, D., Viterbo, P. and Woollen,
36 J.: The ERA-40 re-analysis, *Q. J. R. Meteorol. Soc.*, 131(612), 2961–3012,
37 doi:10.1256/qj.04.176, 2005.
38 Varinlioglu, A., Topcuoglu, S., Köse, A., Kopya, A. I., Uzun, O., Azar, A. and Karal, H.:
39 Levels of cesium radionuclides in mosses in the eastern Black Sea area of Turkey, *J.*
40 *Radioanal. Nucl. Chem.*, 187(6), 435–440, doi:10.1007/BF02165773, 1994.
41 Waight, P., Metivier, H., Jacob, P., Soulchkevitch, G., Viktorsson, C., Bennett, B., Hance, R.,
42 Yumazawa, S., Kusumi, S., Bouville, A., Sinnaeve, J., Ilari, O. and Lazo, E.: Chernobyl ten
43 years on. Radiological and health impact. an assessment by the NEA Committee on the
44 Radiation Protection and Public Health, OECD Nuclear Agency., 1995.
45 Wotawa, G., Becker, A., Kalinowski, M., Saey, P., Tuma, M. and Zähringer, M.:
46 Computation and analysis of the global distribution of the radi xenon isotope ¹³³Xe based on
47 emissions from nuclear power plants and radioisotope production facilities and its relevance
48 for the verification of the nuclear-test-ban treaty, *Pure Appl. Geophys.*, 167(4–5), 541–557,
49 doi:10.1007/s00024-009-0033-0, 2010.
50 Yamauchi: Secondary wind transport of radioactive materials after the Fukushima accident,

1 Earth, Planets Sp., 64(1), e1–e4, doi:10.5047/eps.2012.01.002, 2012.
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1 **FIGURES AND LEGENDS**

2

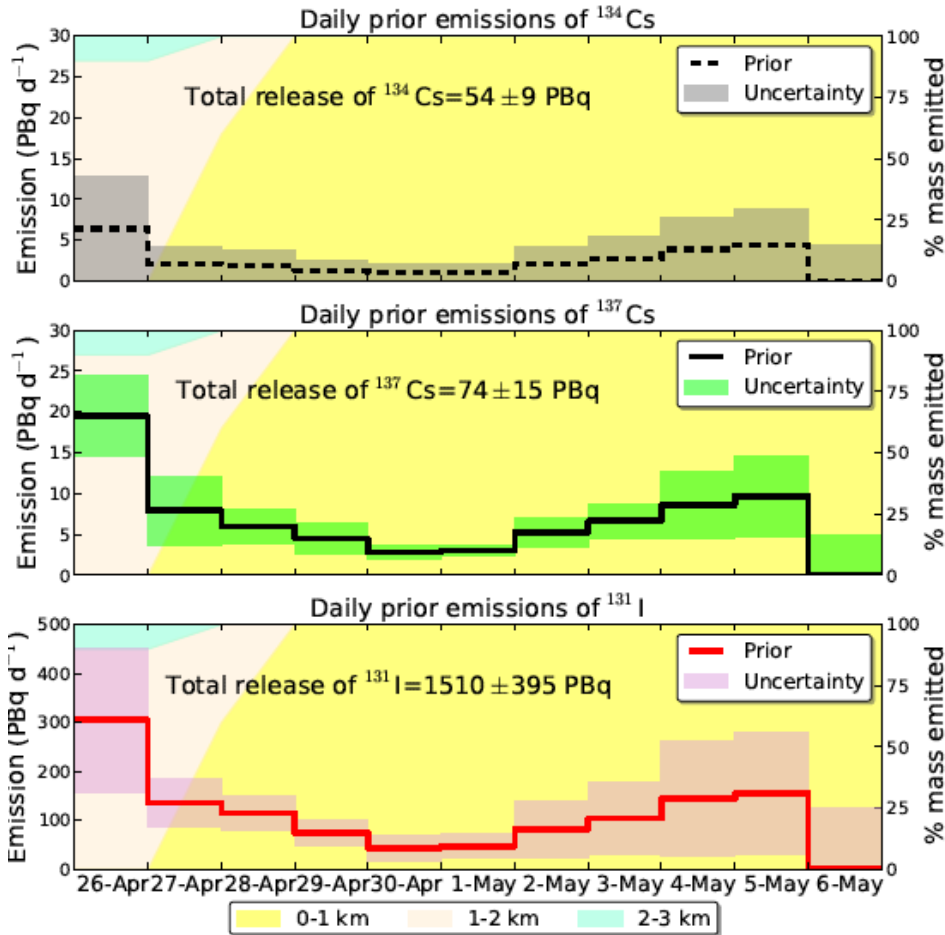


3

4 **Figure 1.** Six different profiles of source releases for ^{134}Cs (black dashed line), ^{137}Cs (black
 5 line) and ^{131}I (red line), published after the Chernobyl accident. These emissions were used to
 6 calculate the a priori (prior) source information and the relative uncertainty of the inversion.
 7 Blue line indicates the maximum altitude of the emissions.

8

PRIOR SOURCE EMISSIONS



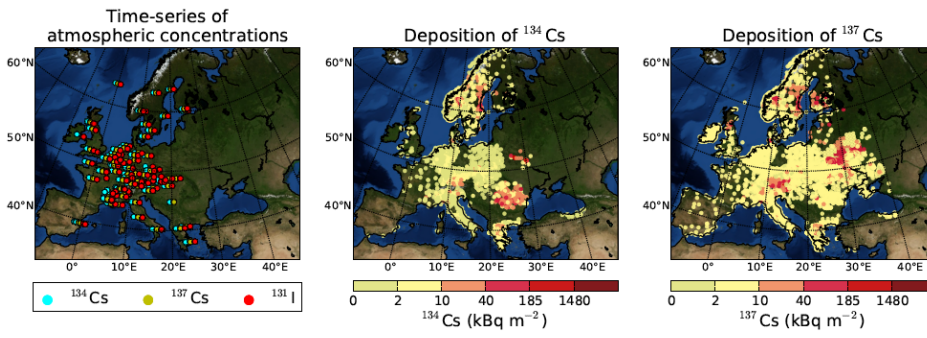
1
 2 **Figure 2.** Calculated prior source term and uncertainty for ¹³⁴Cs (black dashed line), ¹³⁷Cs
 3 (black line) and ¹³¹I (red line) from 26 April to 7 May 1986. Note that emissions are plotted
 4 only until 6 May for simplicity, as on 6 and 7 May 1986 they were reported to be zero. The
 5 prior releases were calculated as the average and standard deviation of the six previously
 6 published source terms (Prior 1–6) shown in [Figure 1](#). On the right axis the vertical
 7 distribution of the emissions at altitudes 0–1 km (yellow), 1–2 km (beige) and 2–3 km
 8 (turquoise) is plotted as shaded background colours.

9

Nikolaos Evangeliou 16/6/2017 17:54

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OBSERVATIONS USED IN THE INVERSION



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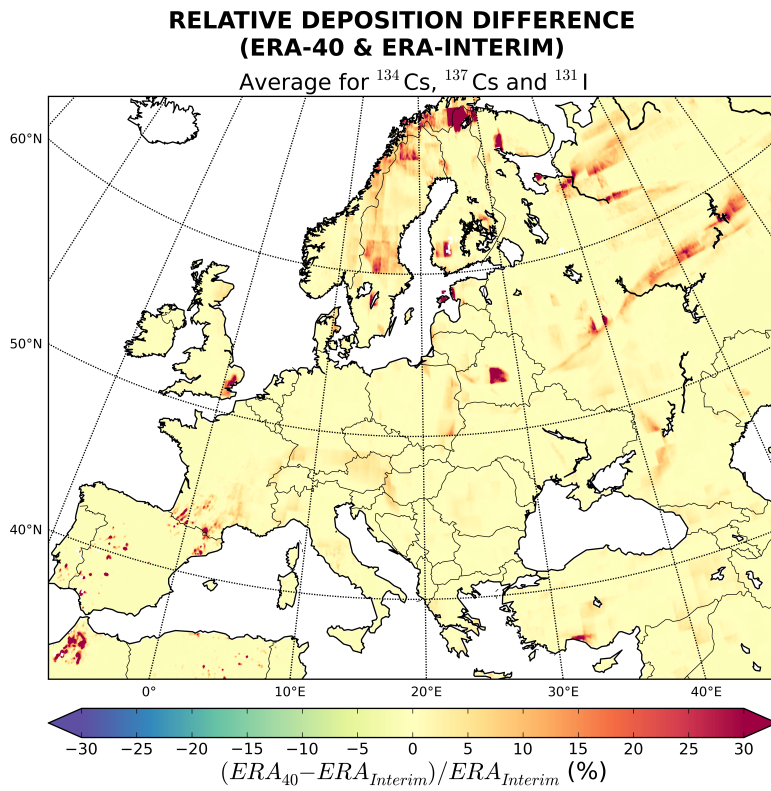
2 **Figure 3.** Locations of atmospheric activity concentration measurements of ^{134}Cs , ^{137}Cs and
3 ^{131}I and deposition locations and levels of ^{134}Cs and ^{137}Cs over Europe adopted from
4 Evangeliou et al. (2016).

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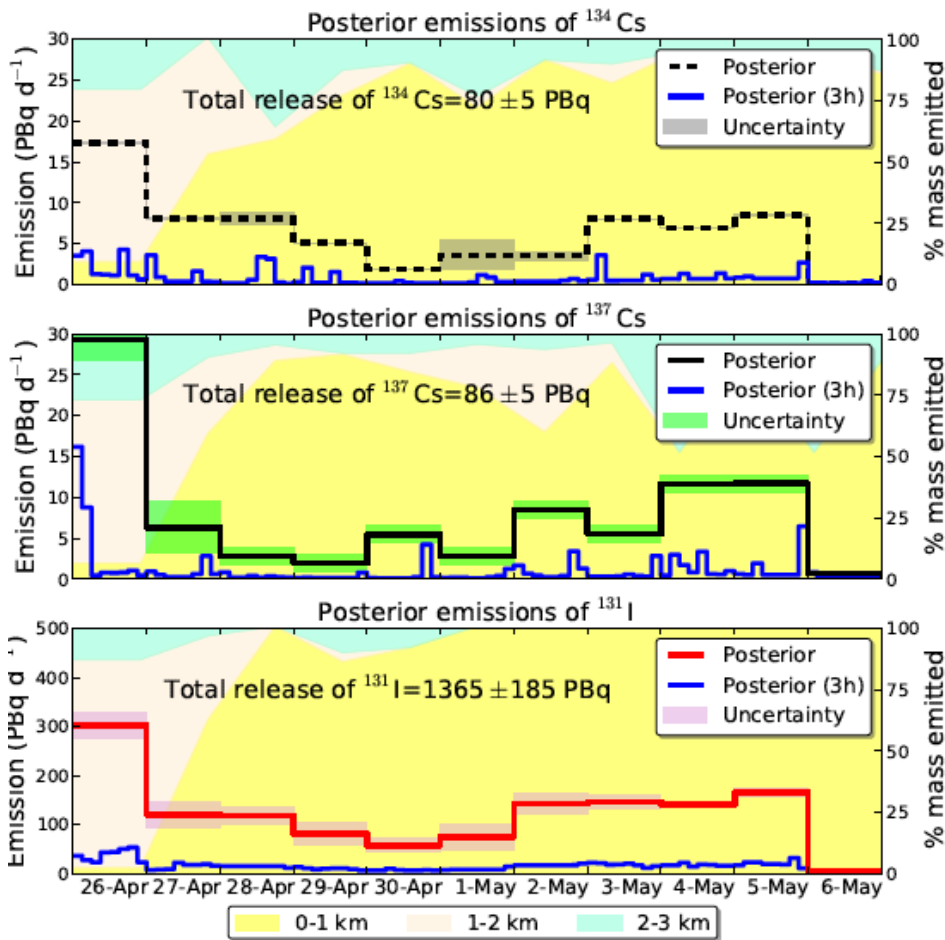
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1
 2 **Figure 4.** Percentage (%) deposition difference between the ERA-40 and ERA-Interim data
 3 sets, i.e., $(ERA_{40} - ERA_{Interim}) / ERA_{Interim}$ in FLEXPART. The relative difference is an
 4 average for the radionuclides ^{134}Cs , ^{137}Cs and ^{131}I .
 5

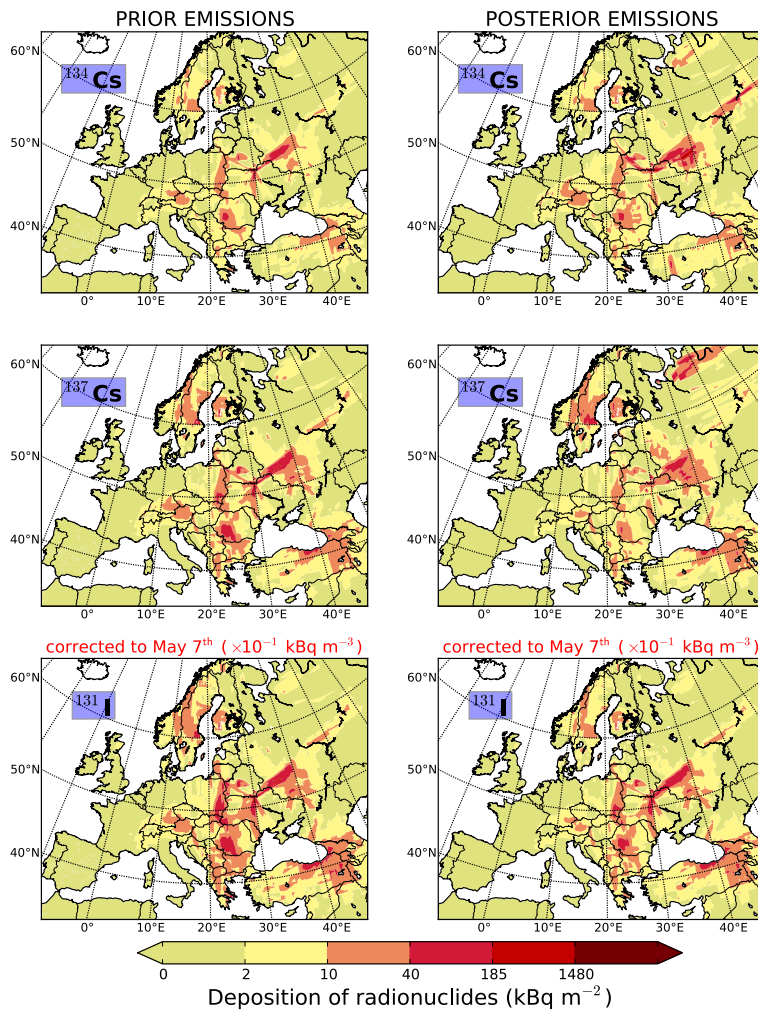
POSTERIOR SOURCE EMISSIONS



1
 2 **Figure 5.** Posterior emissions of ^{134}Cs , ^{137}Cs and ^{131}I against uncertainty plotted daily (similar
 3 to prior ones), as well as per time-step (3 h) from 26 April to 7 May 1986. Note that
 4 emissions are plotted only until 6 May for simplicity, as they were close to zero during 7 May
 5 1986. On the right axis the vertical distribution of the emissions at altitudes 0–1 km (yellow),
 6 1–2 km (beige) and 2–3 km (turquoise) is plotted as shaded background colours.

7

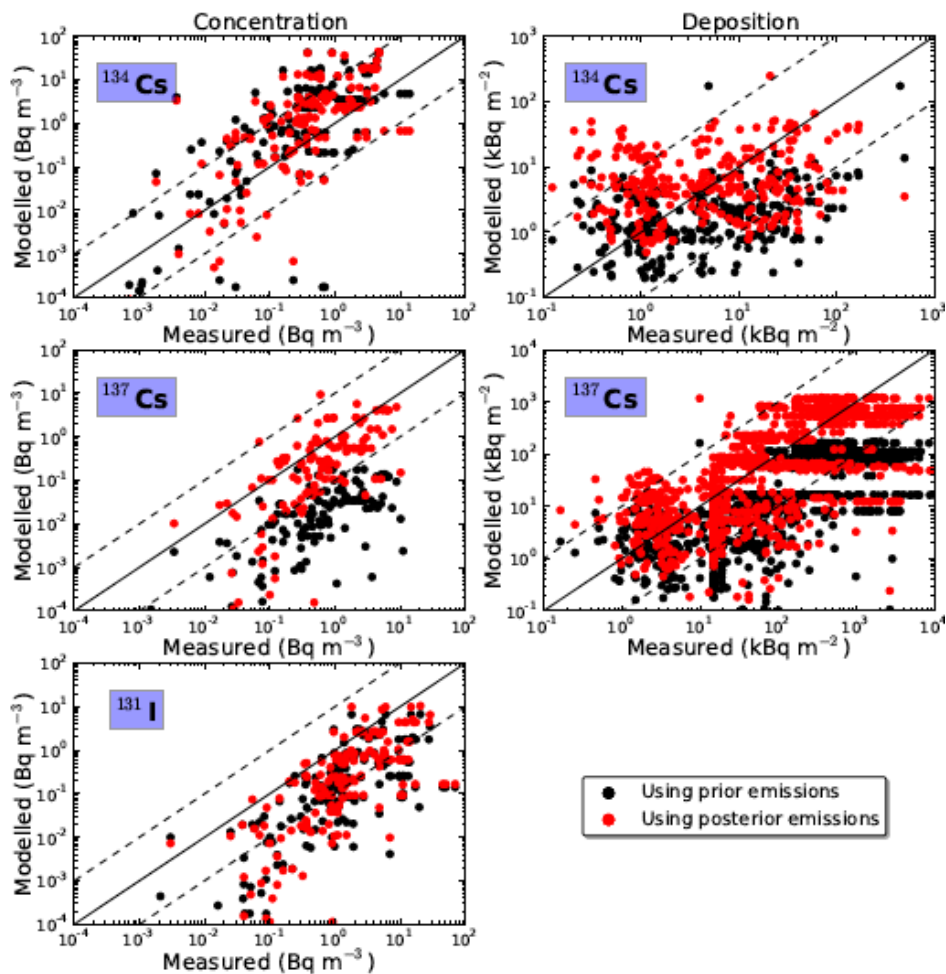
**TOTAL DEPOSITION BEFORE & AFTER
THE INVERSION**



1
2 **Figure 6.** Cumulative deposition of ^{134}Cs , ^{137}Cs and ^{131}I using prior (left column) and
3 posterior emissions (right column). Note that deposition of ^{131}I was corrected for radioactive
4 decay to the end date of the releases (May 7th). Considering that emissions of ^{131}I were about
5 20 times higher than those of ^{134}Cs and ^{137}Cs , total cumulative deposition of ^{131}I was scaled
6 by a factor 0.1 in order to be able to use the same colour scale as for the other radionuclides.

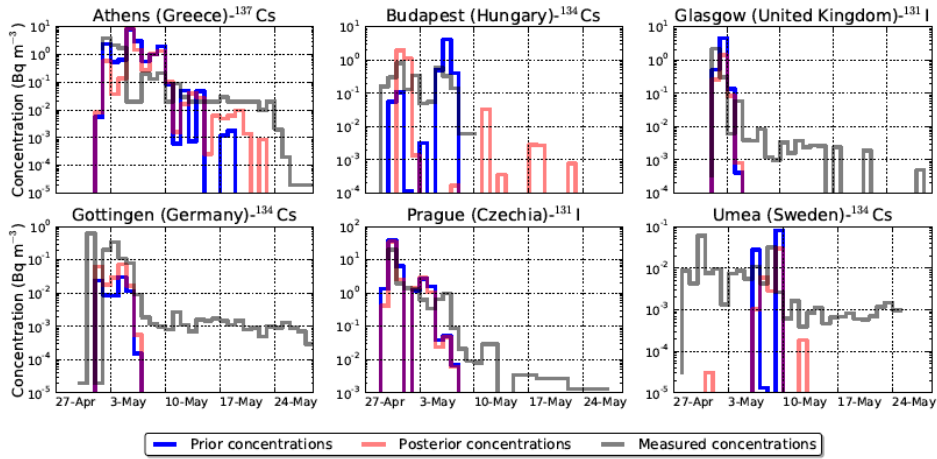
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VALIDATION USING OBSERVATIONS EXCLUDED FROM THE INVERSION



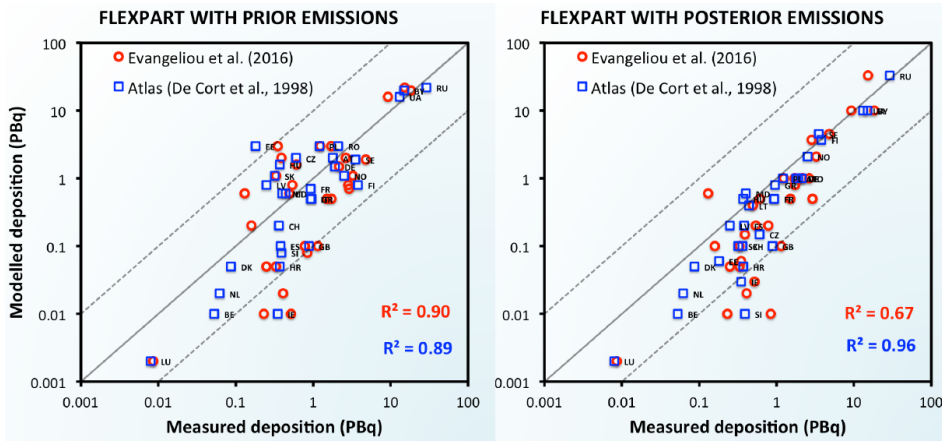
1
 2 **Figure 7.** Left column: Comparison of modelled concentrations with observations excluded
 3 from the inversions for ^{134}Cs ($N_{134} = 318$), ^{137}Cs ($N_{137} = 232$) and ^{131}I ($N_{131} = 318$).
 4 Right column: Comparison of modelled deposition densities with observations excluded from
 5 the inversions for ^{134}Cs ($N_{134} = 273$) and ^{137}Cs ($N_{137} = 1115$).
 6

TIME-SERIES OF ATMOSPHERIC ACTIVITY CONCENTRATIONS



1
 2 **Figure 8.** Time-series of measured (grey) and simulated prior (blue) and posterior (red)
 3 concentrations of ¹³⁴Cs, ¹³⁷Cs and ¹³¹I for the stations Athens (Greece), Budapest (Hungary),
 4 Glasgow (United Kingdom), Göttingen (Germany), Prague (Czechia) and Umea (Sweden).

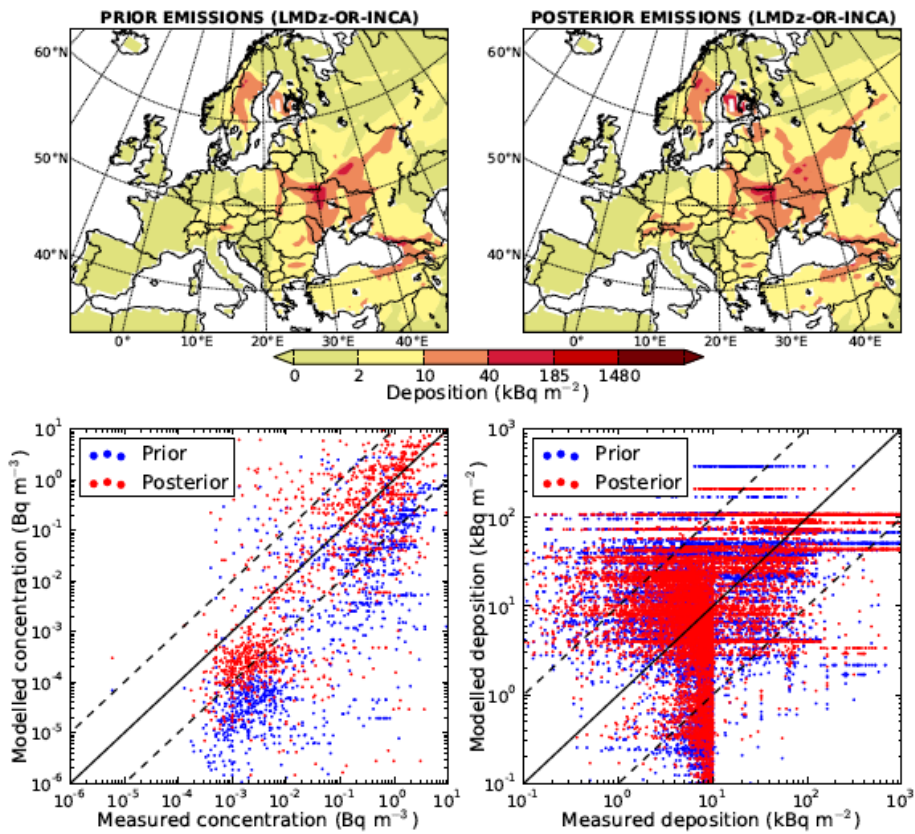
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1
 2 **Figure 9.** Country-by-country total cumulative deposition of ^{137}Cs simulated with
 3 FLEXPART model using the prior and posterior emissions versus the estimated ones from the
 4 Atlas (De Cort et al., 1998) and from Evangeliou et al. (2016). The different countries are
 5 highlighted using their official country codes (Austria (AT), Belarus (BY), Belgium (BE),
 6 Croatia (HR), Czech Republic (CZ), Denmark (DK), Estonia (EE), Finland (FI), France (FR),
 7 Germany (DE), Greece (GR), Hungary (HU), Ireland (IE), Italy (IT), Latvia (LV), Lithuania
 8 (LT), Luxembourg (LU), Moldavia (MD), Netherlands (NL), Norway (NO), Poland (PL),
 9 Rumania (RU), Russia (RU, European part), Slovak Republic (SK), Slovenia (SL), Spain
 10 (ES), Sweden (SE), Switzerland (CH), Ukraine (UA) and United Kingdom (GB)).

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DEPOSITION OF CESIUM-137 OVER EUROPE & COMPARISON WITH OBSERVATIONS



1
 2 **Figure 10.** Deposition of ^{137}Cs using the Eulerian LMDz-OR-INCA Chemistry Transport
 3 Model prior emissions from the ensemble of six a priori releases and the optimised emissions
 4 resulting from our inversion. Comparison of modelled surface concentrations and depositions
 5 of ^{137}Cs with observations from a recently published dataset (Evangelidou et al., 2016).

6

1 **SUPPLEMENTARY FIGURE LEGENDS**

2

3 **Figure S 1.** Deposition of ^{134}Cs , ^{137}Cs and ^{131}I based on the prior emissions used in the
4 present inversion using ERA-40 and ERA-Interim meteorological datasets (Dee et al., 2011;
5 Uppala et al., 2005).

6

7 **Figure S 2.** Sensitivity of the inversion of ^{137}Cs to modification of different parameters.
8 Sensitivity tests accounted for (a) six different prior source terms, (b) three different injection
9 profiles in the prior emissions, (c) two different meteorological datasets (ECMWF ERA-40
10 and ERA-Interim), (d) only deposition observations or (e) only activity concentrations and (f)
11 only observations (both concentrations and deposition densities) from areas close to the NPP
12 (28°E – 32°E , 48°N – 52°N). Uncertainties for each case are plotted as step function showing
13 the range of uncertainty for every time step (TBq s^{-1}).

14

- 1 **Table 1.** Injection altitude (% of total released mass) of prior and posterior emissions of ^{134}Cs ,
 2 ^{137}Cs and ^{131}I averaged over the 12-day period from April 26th 1986 until May 7th 1986.

	0–0.5 km	0.5–1.0 km	1.0–1.5 km	1.5–2.0 km	2.0–2.5 km	2.5–3.0 km
<i>Prior releases</i>						
^{134}Cs	71%	4%	14%	9%	2%	-
^{137}Cs	70%	4%	14%	10%	2%	-
^{131}I	68%	8%	10%	9%	5%	-
<i>Posterior releases</i>						
^{134}Cs	37%	5%	10%	16%	19%	13%
^{137}Cs	37%	23%	14%	5%	11%	10%
^{131}I	38%	32%	13%	8%	5%	4%

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4

1 **Table 2.** Sensitivity of posterior total emissions to (a) different prior emissions (six different previously published assessments), (b) using
 2 different injection altitudes in the prior source term, (c) replacing the ECMWF ERA-40 with the ERA-Interim meteorological data, (d) using
 3 only deposition data or (e) surface activity concentration measurements only and (f) using both deposition and concentration observations close
 4 to the NPP (28°E–32°E, 48°N–52°N), expressed as relative differences to the reference inversion.

	Different prior emissions	Different injection profiles	Different meteorology	Deposition only	Concentration only	28°E–32°E 48°N–52°N
Posterior difference	10%	8.5%	55%	67%	22%	96%

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